```
import pandas as pd
from sklearn.preprocessing import StandardScaler
import matplotlib.pyplot as plt
import numpy as np
from sklearn.impute import SimpleImputer
from sklearn.model_selection import train_test_split, cross_val_score, Stratifie
from sklearn.tree import DecisionTreeClassifier
from sklearn.svm import SVC
from sklearn.pipeline import make_pipeline
from sklearn.metrics import accuracy_score, roc_curve, auc, fbeta_score, classif
import seaborn as sns
```

### Exercise 1 - Data Preparation

1.1 - Load the dataset, inspect feature names and target distribution. Comment on dataset imbalance.

```
In [42]: df = pd.read_csv('data.csv')
         # Finn ut hva som menes med Feature names og target distribution, boka maybe
         # Feature names
         print("Feature names:", list(df.columns))
         # Target distribution
         print(df['diagnosis'].value_counts())
         print("\nFraction:\n", df['diagnosis'].value_counts(normalize=True))
        Feature names: ['id', 'diagnosis', 'radius_mean', 'texture_mean', 'perimeter_mean
        ', 'area_mean', 'smoothness_mean', 'compactness_mean', 'concavity_mean', 'concave
        points_mean', 'symmetry_mean', 'fractal_dimension_mean', 'radius_se', 'texture_se
         , 'perimeter_se', 'area_se', 'smoothness_se', 'compactness_se', 'concavity_se',
        'concave points_se', 'symmetry_se', 'fractal_dimension_se', 'radius_worst', 'text
        ure_worst', 'perimeter_worst', 'area_worst', 'smoothness_worst', 'compactness_wor
        st', 'concavity_worst', 'concave points_worst', 'symmetry_worst', 'fractal_dimens
        ion worst']
        diagnosis
        R
            357
             212
        Name: count, dtype: int64
        Fraction:
         diagnosis
            0.627417
             0.372583
        Name: proportion, dtype: float64
```

The dataset is a bit unbalanced, as it does have more benign than malicious data, but I think the amount of malicious data is enough. If I am unlucky the random splits might turn out with too few malicious but it should be enough.

1.2 - Analyze all features with and without standardization (i.e., zero mean and unit variance). Plot the feature analysis

with and without standardization and decide which version is more suitable.

```
In [43]: # Drop id and diagnosis to get features, these are not numerical
         X = df.drop(columns=["id", "diagnosis"], errors="ignore")
         # Standardize features
         scaler = StandardScaler()
         X_scaled = pd.DataFrame(scaler.fit_transform(X), columns=X.columns, index=X.inde
         # Describe statistics before and after
         print("No Standardization\n", X.describe().T[['mean','std']])
         print("\nStandardization:\n", X_scaled.describe().T[['mean','std']])
         # Plot boxplots for all features
         plt.figure(figsize=(14,6))
         plt.subplot(1,2,1)
         X.boxplot()
         plt.title("Features without standardization")
         plt.xticks(rotation=90)
         plt.subplot(1,2,2)
         X_scaled.boxplot()
         plt.title("Features with standardization")
         plt.xticks(rotation=90)
         plt.tight_layout()
         plt.show()
```

NO Scandar dizaction		
	mean	std
radius_mean	14.127292	3.524049
texture_mean	19.289649	4.301036
perimeter_mean	91.969033	24.298981
area_mean	654.889104	351.914129
smoothness_mean	0.096360	0.014064
compactness_mean	0.104341	0.052813
concavity_mean	0.088799	0.079720
concave points_mean	0.048919	0.038803
symmetry_mean	0.181162	0.027414
fractal_dimension_mean	0.062798	0.007060
radius_se	0.405172	0.277313
texture_se		0.551648
perimeter_se	2.866059	2.021855
area_se	40.337079	45.491006
smoothness_se	0.007041	0.003003
compactness_se	0.025478	0.017908
concavity_se	0.031894	0.030186
concave points_se	0.011796	0.006170
symmetry_se	0.020542	0.008266
<pre>fractal_dimension_se</pre>	0.003795	0.002646
radius_worst	16.269190	4.833242
texture_worst	25.677223	6.146258
perimeter_worst	107.261213	33.602542
area_worst	880.583128	569.356993
smoothness_worst	0.132369	0.022832
compactness_worst	0.254265	0.157336
concavity_worst	0.272188	0.208624
concave points_worst	0.114606	0.065732
symmetry_worst	0.290076	0.061867
<pre>fractal_dimension_worst</pre>	0.083946	0.018061

#### Standardization:

Standaruization.		
	mean	std
radius_mean	-1.373633e-16	1.00088
texture_mean	6.868164e-17	1.00088
perimeter_mean	-1.248757e-16	1.00088
area_mean	-2.185325e-16	1.00088
smoothness_mean	-8.366672e-16	1.00088
compactness_mean	1.873136e-16	1.00088
concavity_mean	4.995028e-17	1.00088
concave points_mean	-4.995028e-17	1.00088
symmetry_mean	1.748260e-16	1.00088
fractal_dimension_mean	4.745277e-16	1.00088
radius_se	2.372638e-16	1.00088
texture_se	-1.123881e-16	1.00088
perimeter_se	-1.123881e-16	1.00088
area_se	-1.311195e-16	1.00088
smoothness_se	-1.529727e-16	1.00088
compactness_se	1.748260e-16	1.00088
concavity_se	1.623384e-16	1.00088
concave points_se	0.000000e+00	1.00088
symmetry_se	8.741299e-17	1.00088
<pre>fractal_dimension_se</pre>	-6.243785e-18	1.00088
radius_worst	-8.241796e-16	1.00088
texture_worst	1.248757e-17	1.00088
perimeter_worst	-3.746271e-16	1.00088
area_worst	0.000000e+00	1.00088
smoothness_worst	-2.372638e-16	1.00088

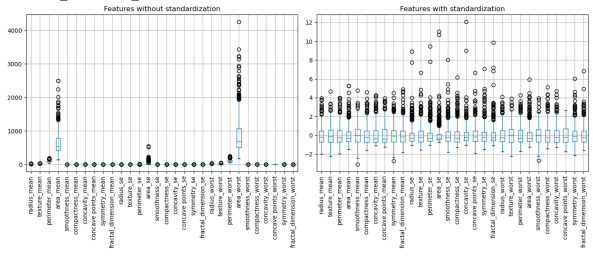
```
      compactness_worst
      -3.371644e-16
      1.00088

      concavity_worst
      7.492542e-17
      1.00088

      concave points_worst
      2.247763e-16
      1.00088

      symmetry_worst
      2.622390e-16
      1.00088

      fractal_dimension_worst
      -5.744282e-16
      1.00088
```



Looking at the plots it seems like it is best to used the standardized one, as the left plot shows that there are huge difference in scaling, where some values are in the thousands, and others are barely over zero. Using the standardized one means the scales are more comparable and therefore not having some values dominate the models. And the esthetics of the right boxplot looks right to use.

## 1.3 - Comment on importance of three way split with respect to hyperparameter search and robustness of any learned model.

The three way split is important in regards to hyperparameters as we dont use the test set to cheat the hyperparameters to fit the test set. For robustness its good because it keeps the test set seperate from the hyperparameter search, and thus we don't risk to fit the hyperparameters to the test data, but rather use the test data to see how the hyperparameters works on new data.

### **Exercise 2 - Decision Trees**

### **Splits**

#### DO NOT TOUCH

(After closer reconsideration, it turn out the random state makes sure the splits are the same, this was made to make sure it was the same, woops. Keeping this now so i dont have to split anymore i guess.)

```
In [44]: # Features and target
X = df.drop(columns=["id", "diagnosis"], errors="ignore")
y = df["diagnosis"].map({"M": 1, "B": 0})
# Split into 60% train, 20% val, 20% test
```

```
X_train, X_temp, y_train, y_temp = train_test_split(
     X, y, test_size=0.4, stratify=y, random_state=42
)
X_val, X_test, y_val, y_test = train_test_split(
     X_temp, y_temp, test_size=0.5, stratify=y_temp, random_state=42
)
print("Train:", X_train.shape)
print("Validation:", X_val.shape)
print("Test:", X_test.shape)
```

Train: (341, 30)
Validation: (114, 30)
Test: (114, 30)

2.1 - Train a Decision Tree classifier using default parameters. Evaluate it on validation sets from original splits (report accuracy mean and std).

```
In [45]: # Only training set
    # Modell with default parameters
    clf = DecisionTreeClassifier(random_state=42)

# 5-fold cross-validation on training set
    cv_scores = cross_val_score(clf, X_train, y_train, cv=5, scoring="accuracy")

print("Cross Validation Accuracy mean:", np.mean(cv_scores))
    print("Cross Validation Accuracy std: ", np.std(cv_scores))

decison_tree_train_result = cv_scores
```

Cross Validation Accuracy mean: 0.9238704177323104 Cross Validation Accuracy std: 0.040627367474774134

```
In [46]: # Validation set
    clf.fit(X_train, y_train)
    val_score = clf.score(X_val, y_val)
    print("Validation Accuracy:", val_score)
```

Validation Accuracy: 0.9385964912280702

Validation set better accuracy than cross validation mean

2.2 - From the trained model, comment on feature importance values and identify the top 3 features from your model.

```
In [47]: # Get important features from the trained model, all features, and sorting them
importances = clf.feature_importances_
feature_names = X.columns
feature_importances = pd.Series(importances, index=feature_names).sort_values(as print("Feature importances:\n", feature_importances)
```

```
Feature importances:
                       0.704881
perimeter_worst
                     0.136016
concave points_worst
texture_worst
                      0.038957
smoothness_worst
area_se
                      0.019922
symmetry_mean
                      0.010456
                      0.009410
radius_mean
radius_worst
                      0.009410
                      0.008365
area_worst
smoothness_se
                      0.001726
                     0.000000
0.000000
area_mean
perimeter_mean
texture_mean
                      0.000000
                      0.000000
perimeter_se
texture_se
                      0.000000
                      0.000000
radius se
fractal_dimension_mean     0.000000
                      0.000000
concavity_mean
concave points_mean 0.000000
smoothness_mean
                      0.000000
                      0.000000
compactness_mean
                      0.000000
symmetry_se
                      0.000000
compactness_se
concavity_se
                      0.000000
                     0.000000
0.000000
0.000000
concave points_se
compactness_worst
concavity_worst
symmetry worst
                      0.000000
fractal_dimension_worst     0.000000
dtype: float64
```

The model places high importance perimieter\_worst, a little important on concave points\_worst. It also places absolutely no importance at all on about 20 features. To me, this feels a little weird but perimeter\_worst might just be a good classifier between malicious and benign. It might also be a case of overfitting when it relies that mch on one value.

2.3 - Vary the max depth parameter (e.g., depth 2-10). Use validation accuracy (mean ± std from cross-validation on the training set) to choose the best depth. Provide performance for each chosen depth (at-least 5 to be reported) and discuss the aspects of overfitting vs. underfitting.

```
In [48]: # Varying max depth from 2 to 10
depths = range(2, 11)

for d in depths:
    clf = DecisionTreeClassifier(max_depth=d, random_state=42)
    cv_scores = cross_val_score(clf, X_train, y_train, cv=5, scoring="accuracy")
    print(f"Max Depth: {d}, CV Accuracy mean: {np.mean(cv_scores):.4f}, std: {np
```

```
Max Depth: 2, CV Accuracy mean: 0.9298, std: 0.0309
Max Depth: 3, CV Accuracy mean: 0.9355, std: 0.0236
Max Depth: 4, CV Accuracy mean: 0.9326, std: 0.0343
Max Depth: 5, CV Accuracy mean: 0.9298, std: 0.0309
Max Depth: 6, CV Accuracy mean: 0.9298, std: 0.0395
Max Depth: 7, CV Accuracy mean: 0.9239, std: 0.0280
Max Depth: 8, CV Accuracy mean: 0.9239, std: 0.0406
Max Depth: 9, CV Accuracy mean: 0.9239, std: 0.0406
Max Depth: 10, CV Accuracy mean: 0.9239, std: 0.0406
```

The accuracy is about the same, allthough Depth=3 does have the highest mean aswell as the lowest std. The variance does increase whilst mean starts to decrease from max depth 6 and onwards, except for std on 7, this might be a sign of overfitting. The accuracy and std is worse on max depth 2 than 3, this could be some underfitting. The first CV training was also worse than depth 2

```
In [49]: # Max depth 3 on validation set
    clf = DecisionTreeClassifier(max_depth=3, random_state=42)
    clf.fit(X_train, y_train)
    val_score_dt = clf.score(X_val, y_val)
    print("Validation Accuracy with max depth 3:", val_score_dt)

best_decision_tree_results = {
        "max_depth": 3,
        "val_acc": val_score_dt
}
```

Validation Accuracy with max depth 3: 0.9385964912280702

Max depth 3 does get higher accuraccy

## 2.4 - Repeat previous exercise with different min samples leaf values. Which setting generalizes best according to the validation set?

Cross Validation Accuracy mean with min_samples_leaf	1: 0.92387 mean
Cross Validation Accuracy std with min_samples_leaf	1: 0.04063 std
Validation Accuracy with min_samples_leaf	1: 0.9386 val
Cross Validation Accuracy mean with min_samples_leaf	2: 0.93261 mean
Cross Validation Accuracy std with min_samples_leaf	2: 0.03151 std
Validation Accuracy with min_samples_leaf	2: 0.9386 val
, <u> </u>	
Cross Validation Accuracy mean with min_samples_leaf	3: 0.93261 mean
Cross Validation Accuracy std with min_samples_leaf	3: 0.02709 std
Validation Accuracy with min_samples_leaf	3: 0.92105 val
	4. 0.02664
Cross Validation Accuracy mean with min_samples_leaf Cross Validation Accuracy std with min_samples_leaf	4: 0.92664 mean 4: 0.02471 std
Validation Accuracy with min_samples_leaf	4: 0.92105 val
variation Accaracy with min_samples_real	4. 0.32103 Vai
Cross Validation Accuracy mean with min_samples_leaf	5: 0.92084 mean
Cross Validation Accuracy std with min_samples_leaf	5: 0.02381 std
Validation Accuracy with min_samples_leaf	5: 0.91228 val
Cross Validation Accuracy mean with min_samples_leaf	6: 0.91799 mean
Cross Validation Accuracy std with min_samples_leaf	6: 0.02989 std
Validation Accuracy with min_samples_leaf	6: 0.91228 val
Cross Validation Accuracy mean with min_samples_leaf	7: 0.91211 mean
Cross Validation Accuracy std with min_samples_leaf	7: 0.02755 std
Validation Accuracy with min_samples_leaf	7: 0.92982 val
Cross Validation Accuracy mean with min_samples_leaf	8: 0.92093 mean
Cross Validation Accuracy std with min_samples_leaf	8: 0.02514 std
Validation Accuracy with min_samples_leaf	8: 0.92982 val
Cross Validation Accuracy mean with min samples leaf	9: 0.92387 mean
Cross Validation Accuracy std with min_samples_leaf	9: 0.02953 std
Validation Accuracy with min_samples_leaf	9: 0.92982 val
, <u> </u>	
Cross Validation Accuracy mean with min_samples_leaf	10: 0.92387 mean
Cross Validation Accuracy std with min_samples_leaf	10: 0.02953 std
Validation Accuracy with min_samples_leaf	10: 0.92982 val
Cross Validation Accuracy mean with min_samples_leaf	11: 0.92387 mean
Cross Validation Accuracy std with min_samples_leaf	11: 0.02803 std
Validation Accuracy with min samples leaf	11: 0.92982 val
Cross Validation Accuracy mean with min_samples_leaf	12: 0.92093 mean
Cross Validation Accuracy std with min_samples_leaf	12: 0.02986 std
Validation Accuracy with min_samples_leaf	12: 0.92982 val
Cross Validation Accuracy mean with min_samples_leaf	13: 0.91211 mean
Cross Validation Accuracy mean with min_samples_leaf	13: 0.91211 mean 13: 0.02421 std
Validation Accuracy with min_samples_leaf	13: 0.92982 val
Cross Validation Accuracy mean with min_samples_leaf	14: 0.90921 mean
Cross Validation Accuracy std with min_samples_leaf	14: 0.02797 std
Validation Accuracy with min_samples_leaf	14: 0.92982 val
Construction As 111 1 2 2 2 2	15. 0.00007
Cross Validation Accuracy mean with min_samples_leaf	15: 0.90627 mean
Cross Validation Accuracy std with min_samples_leaf Validation Accuracy with min_samples_leaf	15: 0.02831 std 15: 0.92982 val
variuation Accuracy with min_samples_lear	TO. M.ATAQT AGT

```
Cross Validation Accuracy mean with min_samples_leaf 16: 0.90332 mean
Cross Validation Accuracy std with min_samples_leaf 16: 0.02835 std
Validation Accuracy with min_samples_leaf
                                                     16: 0.92982 val
Cross Validation Accuracy mean with min_samples_leaf 17: 0.90332 mean
Cross Validation Accuracy std with min_samples leaf
                                                    17: 0.02835 std
Validation Accuracy with min_samples_leaf
                                                     17: 0.92982 val
Cross Validation Accuracy mean with min_samples_leaf
                                                     18: 0.89744 mean
Cross Validation Accuracy std with min_samples_leaf
                                                     18: 0.03188 std
                                                     18: 0.92982 val
Validation Accuracy with min_samples_leaf
Cross Validation Accuracy mean with min_samples_leaf 19: 0.89744 mean
Cross Validation Accuracy std with min_samples_leaf
                                                     19: 0.03188 std
Validation Accuracy with min_samples_leaf
                                                     19: 0.9386 val
Cross Validation Accuracy mean with min_samples_leaf 20: 0.89744 mean
Cross Validation Accuracy std with min_samples_leaf 20: 0.03188 std
Validation Accuracy with min_samples_leaf
                                                     20: 0.9386 val
```

For min\_samples\_leaf 1, 2, 19, 20 gave the same accuracy on the validation set. Low number of leaf samples might have high variance and therefore overfit, whilst the the (longer,deeper?) tree might have to high bias and underfit. Also interesting is the mean from the training splits keep getting worse the more min samples gets added.

These min samples gives the same result as the best max depth setting.

As a summary, decision tree gives the same validation accuracy regardless of settings, be it max depth or min sample leaves or default settings.

### Exercise 3: Support Vector MAchines (SVM)

3.1 - Train a linear SVM (kernel="linear") and evaluate on validation sets (use the original 5 splits that was created in previous exercise). Report accuracy mean, std and plot the ROC for each split.

```
roc_auc = auc(fpr, tpr)
  plt.plot(fpr, tpr, label=f'Fold {i} (AUC = {roc_auc:.2f})')

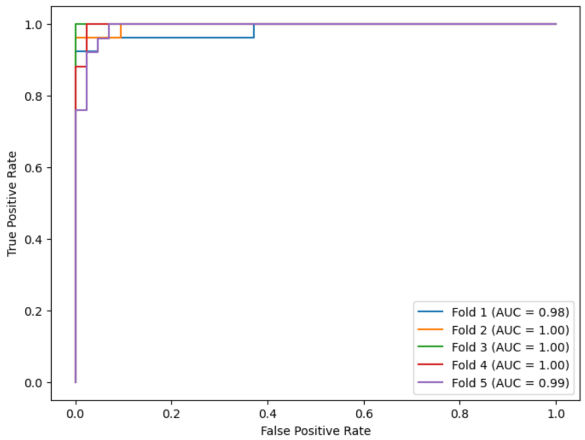
plt.xlabel('False Positive Rate')
plt.ylabel('True Positive Rate')
plt.title('ROC Curve for SVM Classifier')
plt.legend()
plt.show()

svm_clf.fit(X_train, y_train)
val_score_svm_linear = svm_clf.score(X_val, y_val)
print("SVM kernel=linear Validation Accuracy:", np.round(val_score_svm_linear, 5

# Best results for linear is this one, so values saved for comparison
```

SVM Cross Validation Accuracy mean: 0.96483 mean SVM Cross Validation Accuracy std: 0.01492 std

#### ROC Curve for SVM Classifier



SVM kernel=linear Validation Accuracy: 0.96491 val

ROC shows how good the model seperates by class. False positive rate - how many begign being classified as malignant here, false as true True positive rate - Malignant being classified as malignant, true as true

## 3.2 - Train an Radial Basis Function (RBF) kernel SVM. Compare its performance to the linear kernel using validation accuracy. Plot ROC for these models.

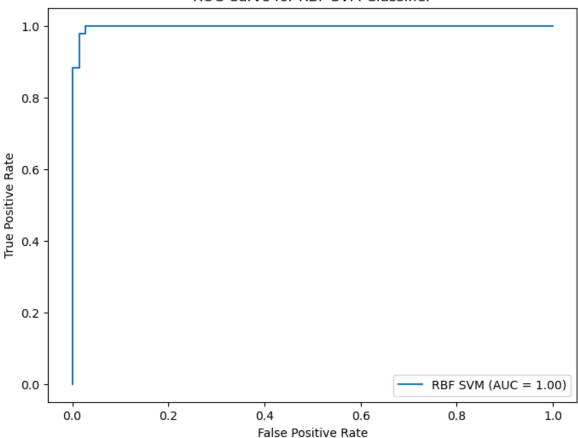
```
In [52]: # Train same splits on RBF model
svm_rbf_clf = make_pipeline(StandardScaler(), SVC(kernel="rbf", gamma="scale", r

rbf_cv = cross_val_score(svm_rbf_clf, X_train, y_train, cv=5, scoring="accuracy")
```

```
print("RBF SVM Cross Validation Accuracy mean:", np.round(np.mean(rbf_cv), 5), "
print("RBF SVM Cross Validation Accuracy std:", np.round(np.std(rbf_cv), 5), "st
# ROC on validation set
y_proba_rbf = svm_rbf_clf.fit(X_train, y_train).predict_proba(X_val)[:, 1]
fpr_rbf, tpr_rbf, _ = roc_curve(y_val, y_proba_rbf)
roc_auc_rbf = auc(fpr_rbf, tpr_rbf)
plt.figure(figsize=(8,6))
plt.plot(fpr_rbf, tpr_rbf, label=f'RBF SVM (AUC = {roc_auc_rbf:.2f})')
plt.xlabel('False Positive Rate')
plt.ylabel('True Positive Rate')
plt.title('ROC Curve for RBF SVM Classifier')
plt.legend()
plt.show()
# Saving best fold results for comparison
best_rbf_fold_idx = int(np.argmax(rbf_cv))
best_rbf_fold_score = rbf_cv[best_rbf_fold_idx]
rbf_results_trainingset = {
    "cv_scores": rbf_cv,
    "cv_mean": rbf_cv.mean(),
    "cv_std": rbf_cv.std(),
    "best_fold": best_rbf_fold_idx + 1,
    "best_fold_acc": best_rbf_fold_score,
    "roc_auc": roc_auc_rbf
}
# Pointy curves cause few data??
```

RBF SVM Cross Validation Accuracy mean: 0.96475 mean RBF SVM Cross Validation Accuracy std: 0.01506 std

#### ROC Curve for RBF SVM Classifier



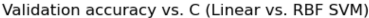
Comparing the validation accuracy between these two models we get:

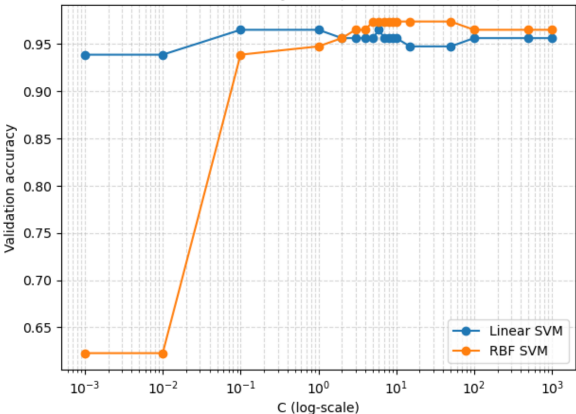
- SVM kernel=linear Validation Accuracy: 0.96491
- RBF SVM Cross Validation Accuracy mean: 0.96475

Currently the SVM with linear kernel is 0.0002 ahead, or 0.02%, which is basically no difference in this dataset, considering the size of the set

## 3.3 - Experiment with different values of C (regularization strength). Use the validation set to select the best C. Report results as a plot of accuracy vs. C

```
In [53]: Cs = [0.001, 0.01, 0.1, 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 50, 100, 500, 1000]
        # Pipelines for linear and RBF
        def make_lin(C): return make_pipeline(StandardScaler(), SVC(kernel="linear", C=C
        def make_rbf(C): return make_pipeline(StandardScaler(), SVC(kernel="rbf",
         lin val acc = []
         rbf_val_acc = []
         # Train on training set, evaluate on validation set for each C
         for C in Cs:
            lin = make_lin(C)
            rbf = make_rbf(C)
            lin.fit(X_train, y_train)
            rbf.fit(X_train, y_train)
            lin_val_acc.append(lin.score(X_val, y_val))
            rbf_val_acc.append(rbf.score(X_val, y_val))
         # Plot accuracy vs C for both models
         plt.figure(figsize=(7,5))
         plt.plot(Cs, lin_val_acc, marker='o', label='Linear SVM')
         plt.plot(Cs, rbf_val_acc, marker='o', label='RBF SVM')
         plt.xscale('log')
         plt.xlabel('C (log-scale)')
         plt.ylabel('Validation accuracy')
         plt.title('Validation accuracy vs. C (Linear vs. RBF SVM)')
         plt.legend()
         plt.grid(True, which='both', linestyle='--', alpha=0.5)
         plt.show()
         # Best C and accuracy for both models
         best_lin_idx = int(np.argmax(lin_val_acc))
         best_rbf_idx = int(np.argmax(rbf_val_acc))
         print(f"Best Linear SVM: C={Cs[best_lin_idx]} | Val acc={lin_val_acc[best_lin_
         # Save for comparions
         best_rbf_idx = int(np.argmax(rbf_val_acc))
         best_rbf_results = {
            "C": Cs[best_rbf_idx],
            "val_acc": rbf_val_acc[best_rbf_idx]
```





Best Linear SVM: C=0.1 | Val acc=0.9649 Best RBF SVM: C=5 | Val acc=0.9737

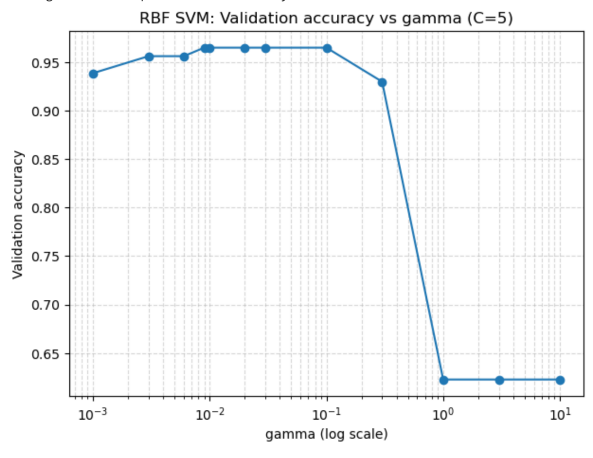
C strength is how much the SVM tries to seperate classes. Low C means it allows more mistakes during training and can give better generalization, also might underfit. High C is more strict when classifying during training, also risks overfitting.

3.4 - Experiment with different  $\gamma$  values for the RBF kernel. Discuss the effect on bias-variance trade-off for all experimented values. Select the best  $\gamma$  using the validation set and report the performance on validation set.

```
print("Gammas:", gammas)
print("Validation accuracies:", [round(a, 4) for a in val_accs])
print(f"Best gamma: {best_gamma} | Validation accuracy: {best_val_acc:.4f}")

plt.figure(figsize=(7,5))
plt.plot(gammas, val_accs, marker='o')
plt.xscale('log')
plt.xscale('log')
plt.xlabel('gamma (log scale)')
plt.ylabel('Validation accuracy')
plt.title(f'RBF SVM: Validation accuracy vs gamma (C={C_fixed})')
plt.grid(True, which='both', linestyle='--', alpha=0.5)
plt.show()
```

Gammas: [0.001, 0.003, 0.006, 0.009, 0.01, 0.02, 0.03, 0.1, 0.3, 1, 3, 10] Validation accuracies: [0.9386, 0.9561, 0.9561, 0.9649, 0.9649, 0.9649, 0.9649, 0.6228, 0.6228] Best gamma: 0.009 | Validation accuracy: 0.9649



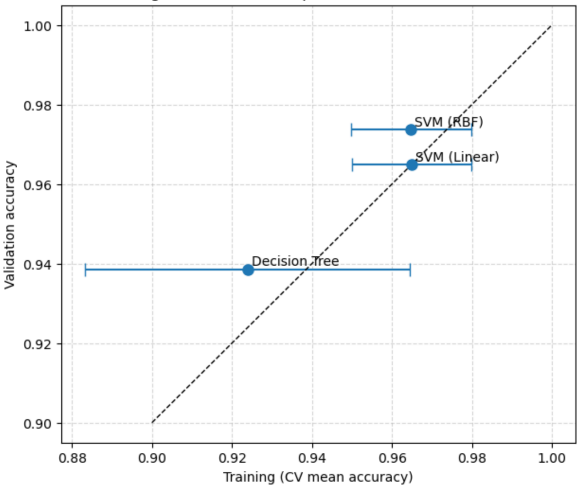
## Exercise - 4: Model Comparison

4.1 - Compare Decision Tree and SVM results from the training set (cross-validation mean ± std) and validation set. Plot the performance comparison plots (e.g., scatter plots, ROC curves). Which model generalizes better?

```
In [55]: # Training result decision tree
  decison_tree_train_result
  # Best decision tree results for comparison
  best_decision_tree_results
# Linear SVM results for comparison
```

```
np.mean(cv_scores_svm_linear)
         np.std(cv_scores_svm_linear)
         val_score_svm_linear
         # Best RBF training set results for comparison
         rbf results trainingset
         # RBF SVM results for comparison
         best_rbf_results
Out[55]: {'C': 5, 'val_acc': 0.9736842105263158}
In [56]: # Summary table for all models, easy reading
         results = pd.DataFrame({
             "Model": ["Decision Tree", "SVM (Linear)", "SVM (RBF)"],
             "CV Mean": [
                 np.mean(decison_tree_train_result),
                 np.mean(cv scores svm linear),
                 rbf_results_trainingset["cv_mean"]
             ],
             "CV Std": [
                 np.std(decison_tree_train_result),
                 np.std(cv scores svm linear),
                 rbf results trainingset["cv std"]
             "Validation Accuracy": [
                 best_decision_tree_results["val_acc"],
                 val_score_svm_linear,
                 best rbf results["val acc"]
             ]
         })
         print(results)
                   Model CV Mean CV Std Validation Accuracy
        0 Decision Tree 0.923870 0.040627
                                                        0.938596
            SVM (Linear) 0.964834 0.014924
        1
                                                         0.964912
               SVM (RBF) 0.964749 0.015056
                                                         0.973684
In [57]: plt.figure(figsize=(7,6))
         plt.errorbar(results["CV Mean"], results["Validation Accuracy"],
                      xerr=results["CV Std"], fmt='o', capsize=5, markersize=8)
         for i, row in results.iterrows():
             plt.text(row["CV Mean"]+0.001, row["Validation Accuracy"]+0.001,
                      row["Model"], fontsize=10)
         # Perfect generalization line
         plt.plot([0.9, 1.0], [0.9, 1.0], 'k--', lw=1)
         plt.xlabel("Training (CV mean accuracy)")
         plt.ylabel("Validation accuracy")
         plt.title("Model generalization comparison: Decision Tree vs SVM")
         plt.grid(True, linestyle="--", alpha=0.5)
         plt.show()
```





The decision tree has a bit lower accuracy for both validation and training than the SVM variants. The decision tree also has higher variance and these characteristics tell me that the decision tree is worse for this task than the SVM variants. To my understanding the SVM RBF variant has a slight tendency to overfits as the point is above the diagoanl. The SVM linear variant is bang on the line but the validation accuracy is a bit lower. Both their variance is about the same. Even though the RBF kernel is not directly on the diagonal, it is close enough to say it is robust enough and generalizes best because of the increased accuracy.

# 4.2 - Discuss the trade-off between usability and accuracy for this dataset. Which model would you recommend for a medical decision-support system, and why? (Hint - Make use of $F\beta$ score analysis)

```
In [58]: # Predictions for each model
y_pred_dt = clf.predict(X_val)  # Decision Tree
y_pred_svm = svm_clf.predict(X_val)  # Linear SVM
y_pred_rbf = svm_rbf_clf.predict(X_val)  # RBF SVM

# F0.5 (6=0.5), F1 (6=1), F2 (6=2) and F3 (6=3) scores
for name, y_pred in [
    ("Decision Tree", y_pred_dt),
    ("SVM (Linear)", y_pred_svm),
    ("SVM (RBF)", y_pred_rbf)
```

Considering this data is from detecting breast cancer, accuracy is very important so we can get all cases and also not misclassify. F1 shows how balanced the model is, between precision and recall. F2 emphazises recall more, meaning its better at picking up false negative, i.e diagnosing someone with cancer as not having it, which is very dangerous. Since the linear scores better for both metric, this seems like the obvious choice.

4.3 - Use GridSearchCV with the training set to tune hyperparameters for both Decision Trees and SVMs. Confirm your final choice with the validation set. Summarize the best settings and provide relevant performance plots

```
cv5 = StratifiedKFold(n_splits=5, shuffle=False)
# Decision Tree with GridSearchCV
dt = DecisionTreeClassifier(random_state=42)
dt_grid = {
    "max_depth": [None, 2, 3, 4, 5, 6, 8, 10],
    "min_samples_leaf": [1, 2, 5, 10, 20],
    "min_samples_split": [2, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 20],
    "criterion": ["gini", "entropy"],
dt_gs = GridSearchCV(dt, dt_grid, scoring="accuracy", cv=cv5, n_jobs=-1, refit=T
dt_gs.fit(X_train, y_train)
# Full table of CV results (sorted)
dt_res = pd.DataFrame(dt_gs.cv_results_)
dt_res = dt_res[["params", "mean_test_score", "std_test_score", "rank_test_score
dt_res = dt_res.sort_values(["rank_test_score", "std_test_score"]).reset_index(d
print("\n=== Decision Tree - all CV results ===")
print(dt res)
# Best summary + validation
dt_cv_mean = dt_gs.best_score_
dt_cv_std = dt_gs.cv_results_["std_test_score"][dt_gs.best_index_]
dt_val_acc = accuracy_score(y_val, dt_gs.predict(X_val))
print("\n=== Decision Tree - best ===")
print("Best params:", dt_gs.best_params_)
print(f"CV accuracy (mean ± std): {dt_cv_mean:.4f} ± {dt_cv_std:.4f}")
print(f"Validation accuracy: { dt val acc:.4f}")
dt_best = {"best_params": dt_gs.best_params_, "cv_mean": dt_cv_mean, "cv_std": d
```

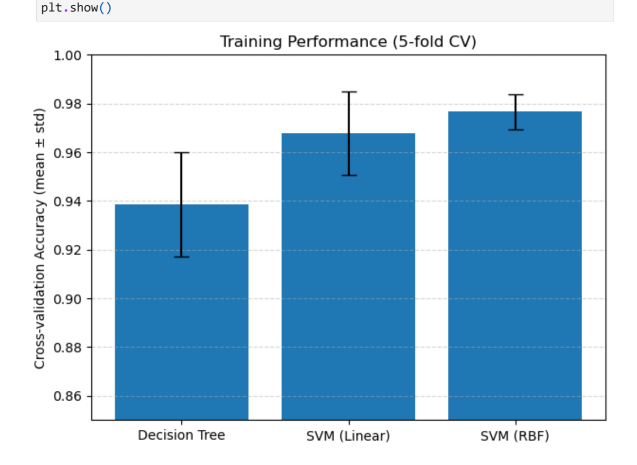
```
=== Decision Tree - all CV results ===
                                                       params mean_test_score \
            {'criterion': 'gini', 'max_depth': 3, 'min_sam...
                                                                      0.938491
       1
            {'criterion': 'gini', 'max_depth': 3, 'min_sam...
                                                                      0.938491
            {'criterion': 'gini', 'max_depth': 3, 'min_sam...
       2
                                                                    0.938491
            {'criterion': 'gini', 'max_depth': 3, 'min_sam...
        3
                                                                     0.938491
            {'criterion': 'gini', 'max_depth': 3, 'min_sam...
       4
                                                                      0.938491
       955 {'criterion': 'gini', 'max_depth': 10, 'min_sa...
                                                                      0.897442
       956 {'criterion': 'gini', 'max_depth': 10, 'min_sa...
                                                                      0.897442
       957 {'criterion': 'gini', 'max_depth': 10, 'min_sa...
                                                                      0.897442
       958 {'criterion': 'gini', 'max_depth': 10, 'min_sa...
                                                                    0.897442
       959 {'criterion': 'gini', 'max_depth': 10, 'min_sa...
                                                                      0.897442
            std_test_score rank_test_score
       0
                  0.021304
        1
                  0.021304
       2
                  0.021304
                                          1
        3
                  0.021304
                  0.021304
                                          1
       4
        . .
                       . . .
                                        . . .
       955
                0.031876
                                        865
       956
                 0.031876
                                       865
       957
                                        865
                  0.031876
       958
                  0.031876
                                        865
       959
                 0.031876
                                        865
        [960 rows x 4 columns]
       === Decision Tree - best ===
       Best params: {'criterion': 'gini', 'max_depth': 3, 'min_samples_leaf': 1, 'min_sa
       mples_split': 6}
       CV accuracy (mean \pm std): 0.9385 \pm 0.0213
       Validation accuracy:
In [60]: # SVM with linear kernel and GridSearchCV
         svm_lin = make_pipeline(StandardScaler(), SVC(kernel="linear", probability=True,
         svm lin grid = {"svc C": [0.001, 0.01, 0.1, 0.5, 0.6, 0.7, 0.8, 0.9, 1, 1.1, 2,
         svm_lin_gs = GridSearchCV(svm_lin, svm_lin_grid, scoring="accuracy", cv=cv5, n_j
         svm_lin_gs.fit(X_train, y_train)
         # Full table of CV results (sorted)
         svm_lin_res = pd.DataFrame(svm_lin_gs.cv_results_)
         svm_lin_res = svm_lin_res[["params", "mean_test_score", "std_test_score", "rank_
         svm_lin_res = svm_lin_res.sort_values(["rank_test_score", "std_test_score"]).res
         print("\n=== SVM (Linear) - all CV results ===")
         print(svm_lin_res)
         # Best summary + validation
         lin_cv_mean = svm_lin_gs.best_score_
         lin_cv_std = svm_lin_gs.cv_results_["std_test_score"][svm_lin_gs.best_index_]
         lin_val_acc = accuracy_score(y_val, svm_lin_gs.predict(X_val))
         print("\n=== SVM (Linear) - best ===")
         print("Best params:", svm_lin_gs.best_params_)
         print(f"CV accuracy (mean ± std): {lin_cv_mean:.4f} ± {lin_cv_std:.4f}")
         print(f"Validation accuracy:
                                          {lin_val_acc:.4f}")
         svm_lin_best = {"best_params": svm_lin_gs.best_params_, "cv_mean": lin_cv_mean,
```

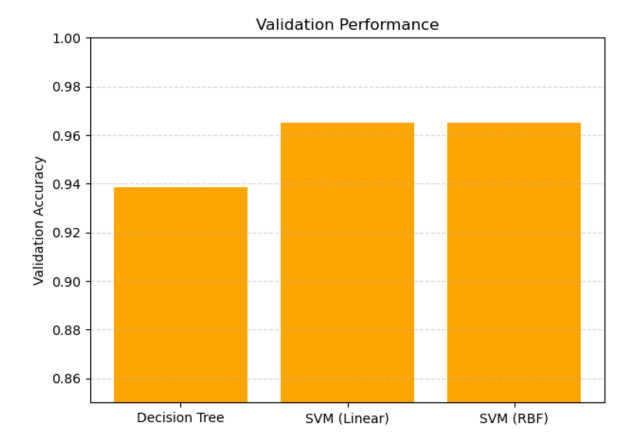
```
=== SVM (Linear) - all CV results ===
                      params mean_test_score std_test_score rank_test_score
        0
             {'svc__C': 0.7}
                                     0.967775
                                                     0.017064
                                                                             1
        1
             {'svc__C': 0.8}
                                     0.967775
                                                     0.017064
                                                                             1
        2
             {'svc__C': 0.9}
                                     0.967775
                                                     0.017064
                                                                             1
        3
             {'svc__C': 0.6}
                                     0.964834
                                                     0.014924
                                                                             4
        4
               {'svc__C': 1}
                                     0.964834
                                                     0.014924
                                                                             4
        5
            {'svc__C': 1.1}
                                                     0.014924
                                     0.964834
               {'svc__C': 2}
                                                                             7
        6
                                     0.964791
                                                     0.023916
        7
               {'svc__C': 5}
                                     0.964791
                                                     0.023916
                                                                            7
            {'svc__C': 0.5}
                                                                            9
        8
                                     0.961893
                                                     0.014949
               {'svc_C': 3}
                                     0.961893
                                                     0.023864
                                                                            9
           {'svc__C': 0.01}
        10
                                     0.961850
                                                     0.007310
                                                                            11
        11
            {'svc__C': 0.1}
                                     0.958951
                                                     0.010974
                                                                            12
        12
               {'svc__C': 7}
                                     0.958909
                                                     0.025341
                                                                            13
            {'svc__C': 100}
       13
                                     0.953026
                                                     0.028569
                                                                            14
             {'svc__C': 10}
        14
                                     0.953026
                                                     0.036541
                                                                            14
        15 {'svc__C': 0.001}
                                     0.932609
                                                     0.021775
                                                                            16
        === SVM (Linear) - best ===
        Best params: {'svc__C': 0.7}
        CV accuracy (mean \pm std): 0.9678 \pm 0.0171
       Validation accuracy:
In [61]: # SVM with RBF kernel and GridSearchCV
         svm_rbf = make_pipeline(StandardScaler(), SVC(kernel="rbf", probability=True, ra
         svm rbf grid = {
             "svc__C":
                           [0.001, 0.01, 0.1, 1, 3, 5, 6, 7, 8, 10, 100],
             "svc_gamma": ["scale", 0.001, 0.003, 0.01, 0.03, 0.1, 0.3, 1, 3, 10],
         }
         svm_rbf_gs = GridSearchCV(svm_rbf, svm_rbf_grid, scoring="accuracy", cv=cv5, n_j
         svm_rbf_gs.fit(X_train, y_train)
         # Full table of CV results (sorted)
         svm rbf res = pd.DataFrame(svm rbf gs.cv results )
         svm_rbf_res = svm_rbf_res[["params", "mean_test_score", "std_test_score", "rank_
         svm rbf res = svm rbf res.sort values(["rank test score", "std test score"]).res
         print("\n=== SVM (RBF) - all CV results ===")
         print(svm_rbf_res)
         # Best summary + validation
         rbf_cv_mean = svm_rbf_gs.best_score_
         rbf_cv_std = svm_rbf_gs.cv_results_["std_test_score"][svm_rbf_gs.best_index_]
         rbf_val_acc = accuracy_score(y_val, svm_rbf_gs.predict(X_val))
         print("\n=== SVM (RBF) - best ===")
         print("Best params:", svm_rbf_gs.best_params_)
         print(f"CV accuracy (mean ± std): {rbf_cv_mean:.4f} ± {rbf_cv_std:.4f}")
         print(f"Validation accuracy: {rbf val acc:.4f}")
         svm_rbf_best = {"best_params": svm_rbf_gs.best_params_, "cv_mean": rbf_cv_mean,
```

```
=== SVM (RBF) - all CV results ===
                                       params mean_test_score std_test_score \
            {'svc__C': 7, 'svc__gamma': 0.01}
                                                                      0.007136
                                                  0.976556
        1
            {'svc__C': 8, 'svc__gamma': 0.01}
                                                     0.976556
                                                                      0.007136
             {'svc__C': 5, 'svc__gamma': 0.03}
        2
                                                     0.976513
                                                                      0.011797
             {'svc__C': 5, 'svc__gamma': 0.01}
        3
                                                     0.973615
                                                                      0.005842
        4
            {'svc__C': 6, 'svc__gamma': 0.01}
                                                     0.973615
                                                                      0.005842
                                                           . . .
            {'svc__C': 8, 'svc__gamma': 10}
                                                    0.627579
        105
                                                                      0.006104
              {'svc__C': 10, 'svc__gamma': 3}
        106
                                                     0.627579
                                                                      0.006104
            {'svc__C': 10, 'svc__gamma': 10}
        107
                                                     0.627579
                                                                      0.006104
            {'svc__C': 100, 'svc__gamma': 3}
        108
                                                     0.627579
                                                                      0.006104
                                               0.627579
             {'svc__C': 100, 'svc__gamma': 10}
        109
                                                                      0.006104
             rank_test_score
        0
        1
                          1
        2
                          3
        3
                          4
        4
                          4
        . .
                         . . .
       105
                         71
        106
                         71
        107
                         71
        108
                         71
        109
                         71
        [110 rows x 4 columns]
       === SVM (RBF) - best ===
        Best params: {'svc__C': 7, 'svc__gamma': 0.01}
        CV accuracy (mean \pm std): 0.9766 \pm 0.0071
       Validation accuracy:
                                 0.9649
In [62]: #Summary of all CV
         models = {
             "Decision Tree": dt_gs.best_estimator_,
             "SVM (Linear)": svm_lin_gs.best_estimator_,
             "SVM (RBF)": svm_rbf_gs.best_estimator_
         }
         # For plotting
         rows = []
         for name, gs in [
             ("Decision Tree", dt_gs),
             ("SVM (Linear)", svm_lin_gs),
             ("SVM (RBF)", svm_rbf_gs),
         1:
             mean = gs.best_score_
             std = gs.cv_results_["std_test_score"][gs.best_index_]
             val_acc = accuracy_score(y_val, gs.predict(X_val))
             rows.append({"Model": name, "CV Mean": mean, "CV Std": std, "Val Acc": val_a
         perf = pd.DataFrame(rows)
         print("\n=== Summary of Best Models ===")
         print(perf)
```

```
Model
                         CV Mean
                                     CV Std Val Acc
          Decision Tree 0.938491 0.021304 0.938596
            SVM (Linear) 0.967775 0.017064 0.964912
        2
               SVM (RBF) 0.976556 0.007136 0.964912
In [63]: # Plotting training performance
         plt.figure(figsize=(7,5))
         plt.bar(perf["Model"], perf["CV Mean"], yerr=perf["CV Std"], capsize=6)
         plt.ylabel("Cross-validation Accuracy (mean ± std)")
         plt.title("Training Performance (5-fold CV)")
         plt.ylim(0.85, 1.0)
         plt.grid(axis='y', linestyle='--', alpha=0.5)
         plt.show()
         # Plotting validation performance
         plt.figure(figsize=(7,5))
         plt.bar(perf["Model"], perf["Val Acc"], color='orange')
         plt.ylabel("Validation Accuracy")
         plt.title("Validation Performance")
         plt.ylim(0.85, 1.0)
         plt.grid(axis='y', linestyle='--', alpha=0.5)
```

=== Summary of Best Models ===



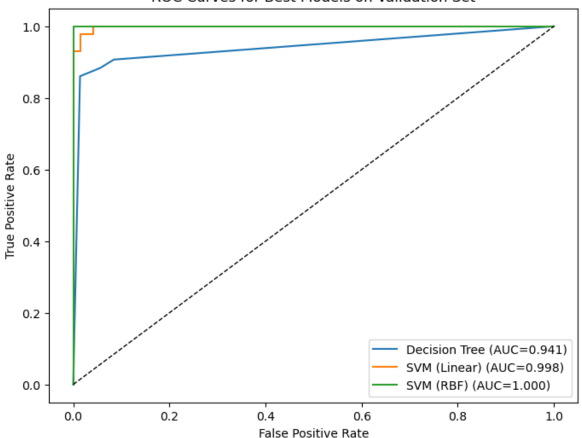


Looking at these two plots, you can see that the RBF SVM overfit a small amount to the training set, whilst the linear perform about the same which means in generalizes a little better.

```
In [64]: # ROC curves for best models on validation set

plt.figure(figsize=(8,6))
    for name, model in models.items():
        y_proba = model.predict_proba(X_val)[:, 1]
        fpr, tpr, _ = roc_curve(y_val, y_proba)
        auc_val = auc(fpr, tpr)
        plt.plot(fpr, tpr, label=f"{name} (AUC={auc_val:.3f})")

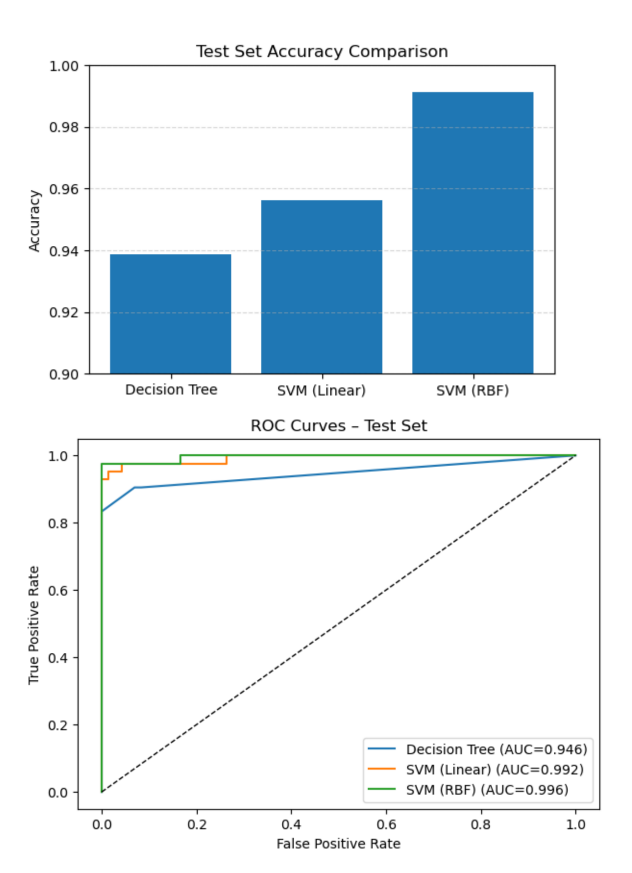
plt.plot([0,1], [0,1], 'k--', lw=1)
    plt.xlabel("False Positive Rate")
    plt.ylabel("True Positive Rate")
    plt.title("ROC Curves for Best Models on Validation Set")
    plt.legend()
    plt.show()
```

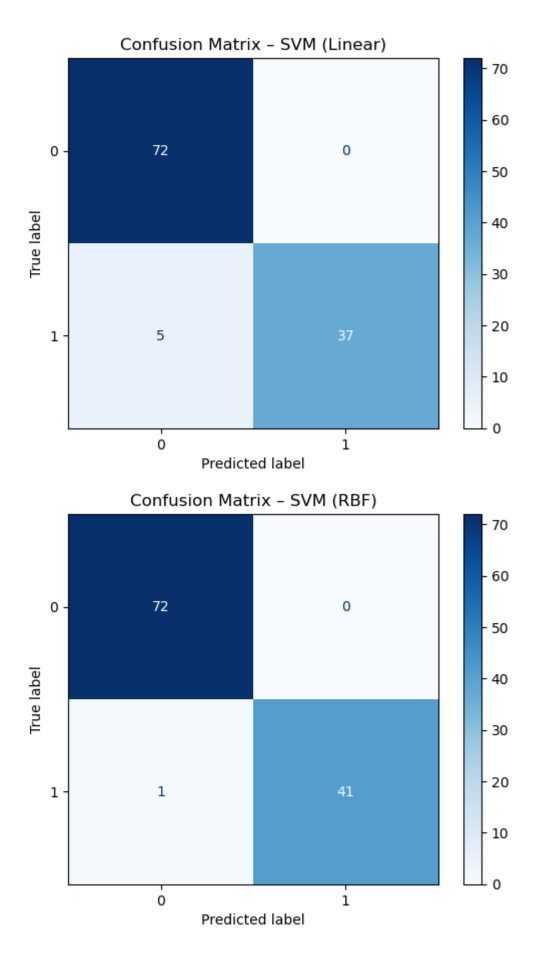


Exercise 4.4 - Compare the final test set accuracy of the best Decision Tree and best SVM. Which model performs better in practice?

```
In [65]: # Best models
         best_dt = dt_gs.best_estimator_
         best_lin = svm_lin_gs.best_estimator_
         best_rbf = svm_rbf_gs.best_estimator_
         # Final evaluation on test set
         models = {
             "Decision Tree": best_dt,
             "SVM (Linear)": best lin,
             "SVM (RBF)": best_rbf
         print("\n=== Final Test Set Performance ===")
         for name, model in models.items():
             y_pred = model.predict(X_test)
             acc = accuracy_score(y_test, y_pred)
             print(f"{name}: Test Accuracy = {acc:.4f}")
        === Final Test Set Performance ===
        Decision Tree: Test Accuracy = 0.9386
        SVM (Linear): Test Accuracy = 0.9561
        SVM (RBF): Test Accuracy = 0.9912
In [66]: from sklearn.metrics import roc_curve, auc, ConfusionMatrixDisplay
         import matplotlib.pyplot as plt
         # Best models already defined
```

```
models = {
     "Decision Tree": dt_gs.best_estimator_,
     "SVM (Linear)": svm_lin_gs.best_estimator_,
     "SVM (RBF)": svm_rbf_gs.best_estimator_,
 }
 # Test set accuracy for each model
 test_accs = {name: accuracy_score(y_test, mdl.predict(X_test))
              for name, mdl in models.items()}
 print({k: round(v, 4) for k, v in test_accs.items()})
 # bar chart
 plt.figure(figsize=(6,4))
 plt.bar(test_accs.keys(), test_accs.values())
 plt.ylabel("Accuracy")
 plt.title("Test Set Accuracy Comparison")
 plt.ylim(0.9, 1.0)
 plt.grid(axis="y", linestyle="--", alpha=0.5)
 plt.show()
 # 2. ROC curves
 plt.figure(figsize=(7,5))
 for name, model in models.items():
    y_proba = model.predict_proba(X_test)[:, 1]
    fpr, tpr, _ = roc_curve(y_test, y_proba)
     roc_auc = auc(fpr, tpr)
     plt.plot(fpr, tpr, label=f"{name} (AUC={roc_auc:.3f})")
 plt.plot([0,1],[0,1],'k--',lw=1)
 plt.xlabel("False Positive Rate")
 plt.ylabel("True Positive Rate")
 plt.title("ROC Curves - Test Set")
 plt.legend()
 plt.show()
 # Confusion matrix for best model (SVM Linear)
 linear_best_model_matrix=svm_lin_gs.best_estimator_
 ConfusionMatrixDisplay.from_estimator(linear_best_model_matrix, X_test, y_test,
 plt.title("Confusion Matrix - SVM (Linear)")
 plt.show()
 # Confusion matrix for best model (RBF)
 best_model = svm_rbf_gs.best_estimator_
 ConfusionMatrixDisplay.from_estimator(best_model, X_test, y_test, cmap="Blues")
 plt.title("Confusion Matrix - SVM (RBF)")
 plt.show()
{'Decision Tree': 0.9386, 'SVM (Linear)': 0.9561, 'SVM (RBF)': 0.9912}
```





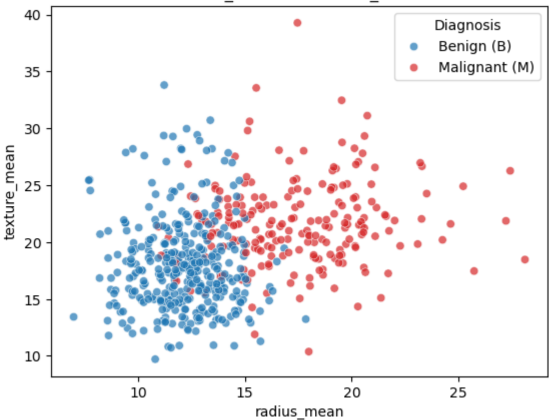
Looking at the plots you can see the RBF model performing best. The confusion matrix shows only one mis-classification, however it is a bad one. In practice, this means taht the model sees a malignant case as begign, which is very dangerous for the patient receiving this diagnosis. The accuracy of the RBF model is good, other than the mis of one case.

4.5 - The Breast Cancer Wisconsin dataset has 30 continuous features, many of which are correlated and not linearly separable. Explain why a linear SVM might fail to capture complex patterns in this dataset. How does using an RBF kernel help in this case? Discuss your answer in terms of the dataset's feature space and the geometry of the decision boundary.

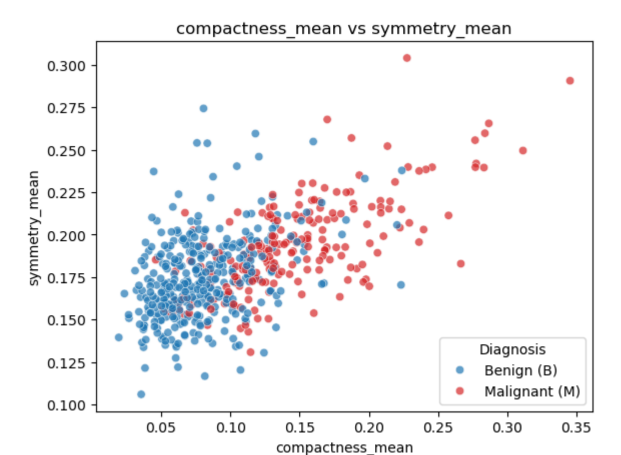
Linear SVM assumes possibility to seperate linearly

```
In [67]: # Scatter plot of two features colored by diagnosis, radius_mean and texture_mean
         feat1, feat2 = "radius_mean", "texture_mean"
         # Plot
         ax = sns.scatterplot(
            data=df,
             x=feat1, y=feat2,
             hue="diagnosis",
             hue_order=["B", "M"],
             palette=["tab:blue", "tab:red"],
             alpha=0.7
         # Customize Legend
         handles, labels = ax.get_legend_handles_labels()
         label_map = {"B": "Benign (B)", "M": "Malignant (M)"}
         labels = [label_map.get(1, 1) for 1 in labels]
         ax.legend(handles=handles, labels=labels, title="Diagnosis", loc="best")
         plt.title(f"{feat1} vs {feat2}")
         plt.show()
```

#### radius mean vs texture mean



```
In [68]: # Scatter plot of two features colored by diagnosis, compactness_mean and symmet
         feat1, feat2 = "compactness_mean", "symmetry_mean"
         # plot
         ax = sns.scatterplot(
             data=df,
             x=feat1, y=feat2,
             hue="diagnosis",
             hue_order=["B", "M"],
             palette=["tab:blue", "tab:red"],
             alpha=0.7
         # Customize Legend
         handles, labels = ax.get_legend_handles_labels()
         label_map = {"B": "Benign (B)", "M": "Malignant (M)"}
         labels = [label_map.get(1, 1) for 1 in labels]
         ax.legend(handles=handles, labels=labels, title="Diagnosis", loc="best")
         plt.title(f"{feat1} vs {feat2}")
         plt.show()
```



Since linear SVM requires the classes to be seperable with a straight line, and looking the two plots above, you can see that these features, and others, are close to impossible to separate linearly. Many of the features are not linearly independent for each other. For example, perimeter\_worst, perimeter\_mean and perimeter\_se have a dependency on each other. Every feature in the data set have 3 different categories, the same way as perimeter has.

Using a technique called similarity features, which measures how much each instance resembles a particular landmark, in combination with the RBF kernel can help make features linearly seperable. Simplest way to select landmarks positions is doing it for every instance in the dataset. This is computing heavy and creates many dimensions, but also increases chances of getting a linearly seperable training set. The hyperparameter gamma helps create a bell curve, an tuning this correctly hte RBF kernel create a decision boundary which the data set can fit in.