

Exercise 1

1.

$$\text{let } f(x) = w^T x + b = 0$$

plugging in parameters, we get.

$$x_1 - 2x_2 + 3x_3 - 1 = 0$$

Since this equation $x_1 - 2x_2 + 3x_3 = 1$ defines a flat 2D surface and it is in 3D feature space, it is then a hyperplane.

2.

$$\nabla_x f(x) = w = n = [1 \ -2 \ 3]^T$$

normal vector $n = w$

the normal vector represents where the decision function grows most rapidly.

3.

(a)

$$f([1 \ 1 \ 1]^T) = 1 - 2 + 3 - 1 = 1$$

$$\| [1 \ -2 \ 3]^T \| = \sqrt{1 + 4 + 9} = \sqrt{14}.$$

$$d(x) = \frac{1}{\sqrt{14}}.$$

(b) since $d(x) > 0$, the point $x [1 \ 1 \ 1]^T$ is positive by the decision boundary classification, which is a prediction.

(c) The distance represent the margin, if the margin is small, it means distance is small which leads to low confidence and vice versa.

(d) from (a) we have $d(x) = \frac{1}{\sqrt{14}}$ $\|w\| = \sqrt{14}$ $w = [1, -2, 3]^T$

$$\begin{aligned}\text{Projection} &= x - d(x) \frac{w}{\|w\|} \\ &= [1 \ 1 \ 1]^T - \frac{1}{\sqrt{14}} \frac{1}{\sqrt{14}} [1 \ -2 \ 3]^T \\ &= \left[\frac{13}{14} \quad \frac{8}{7} \quad \frac{11}{14} \right]^T\end{aligned}$$

(e) Since orthogonal projection gives the distance from points to decision boundary, the maximal margin classifier use this value to find support vectors and thus define the hyperplane.

MLE25_sheet01

May 3, 2025

1 Machine Learning Essentials SS25 - Exercise Sheet 1

1.1 Instructions

- TODO's indicate where you need to complete the implementations.
- You may use external resources, but write your own solutions.
- Provide concise, but comprehensible comments to explain what your code does.
- Code that's unnecessarily extensive and/or not well commented will not be scored.

1.2 Exercise 2: The Perceptron Algorithm

```
[1]: import numpy as np
import matplotlib.pyplot as plt
from sklearn.datasets import load_breast_cancer
from sklearn.model_selection import train_test_split
from sklearn.preprocessing import StandardScaler
from sklearn.metrics import accuracy_score
```

```
[2]: # =====
# 1. Load & Visualize the Dataset
# =====

# TODO: Load dataset, print feature names
data = load_breast_cancer()

print("Feature Names:")
print(data.feature_names)

# TODO: Select features & corresponding labels
X = data.data[:, [0, 1]]
y = data.target

# Convert labels from {0,1} to {-1,1} to match Perceptron convention from sheet
y = 2 * (y - 0.5)

# TODO: Standardize the data to zero mean and unit variance, explain why it's
↪ useful
```

```

scaler = StandardScaler()
X_std = scaler.fit_transform(X)

# we standardize the features, so that features with large ranges don't
↳ dominate features with smaller ranges, even if both have the same importance

# TODO: Visualize dataset using plt.scatter()
plt.figure(figsize=(6, 6))
plt.scatter(X_std[y == -1, 0], X_std[y == -1, 1], color='red', label='Malignant',
↳ (-1)', alpha=0.6)
plt.scatter(X_std[y == 1, 0], X_std[y == 1, 1], color='blue', label='Benign',
↳ (+1)', alpha=0.6)
plt.xlabel('Standardized Mean Radius')
plt.ylabel('Standardized Mean Texture')
plt.legend()
plt.grid(True)
plt.show()

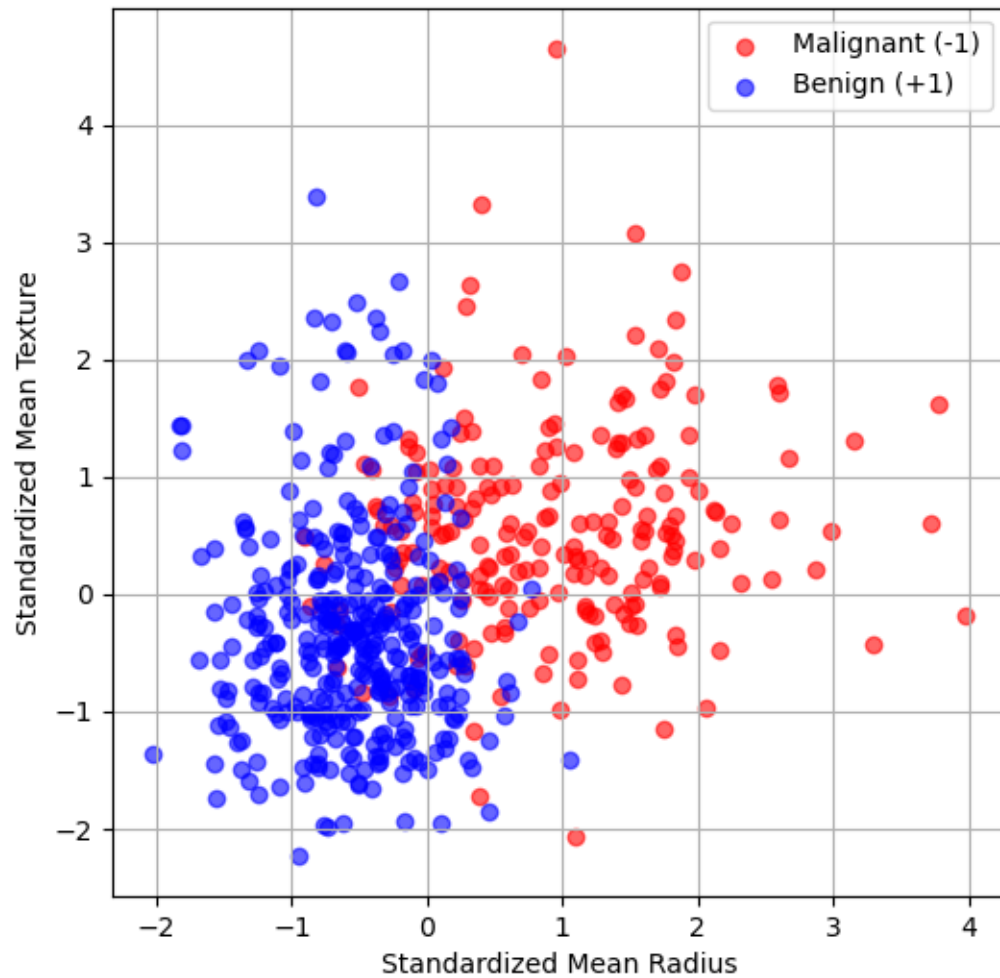
```

Feature Names:

```

['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]: # =====
# 2. Implement the Perceptron's training algorithm
# =====

class Perceptron:
    def __init__(self, learning_rate=0.1, num_epochs=10):
        self.learning_rate = learning_rate
        self.num_epochs = num_epochs
        self.w = None # Weights
        self.b = None # Bias
        self.history = [] # Store parameters for decision boundary @ each
        ↪ update for visualization

    def train(self, X, y):
        """Train the perceptron using the online Perceptron algorithm."""
        n_samples, n_features = X.shape
```

```

        # TODO: Initialize weights and bias
        self.w = np.zeros(n_features)
        self.b = 0

        # Train for num_epochs iterations
        for _ in range(self.num_epochs):
            for i in range(n_samples):
                X_i = X[i]
                # TODO: Implement the update rule
                if y[i] != self.predict(X_i):
                    self.w = self.learning_rate * y[i] * X_i
                    self.b = self.learning_rate * y[i]
                    self.history.append((self.w, self.b)) # Save state for
↪ visualization

    def predict(self, X):
        """Predict the label of a sample."""
        linear_output = np.dot(X, self.w) + self.b
        return np.sign(linear_output)

```

```

[4]: # =====
# 3. Train the Perceptron & Evaluate Performance
# =====

# TODO: Split the data into training and test set
X_train, X_test, y_train, y_test = train_test_split(
    X_std, y, test_size=0.2, random_state=42
)

# TODO: Initialize the Perceptron and train it on the training set
perceptron = Perceptron(learning_rate=0.1, num_epochs=100)
perceptron.train(X_train, y_train)

y_train_pred = perceptron.predict(X_train)
y_test_pred = perceptron.predict(X_test)
...

# TODO: Use the trained Perceptron to compute the accuracy on the training set
↪ and on the test set
...
train_acc = accuracy_score(y_train, y_train_pred)
test_acc = accuracy_score(y_test, y_test_pred)

print(f"Training Accuracy: {train_acc:.3f}")
print(f"Test Accuracy: {test_acc:.3f}")

```

Training Accuracy: 0.668
Test Accuracy: 0.649

```
[5]: # =====
# 4. Plot decision boundary evolution
# =====
print(f"Total updates in history: {len(perceptron.history)}")

# Visualize the first 5 consecutive decision boundaries for data
decision_boundaries = perceptron.history[:5] # Get the parameters of the first
↳5 decision boundaries used during training

# TODO: Plot decision boundaries for iterations 1-5
plt.figure(figsize=(10, 8))

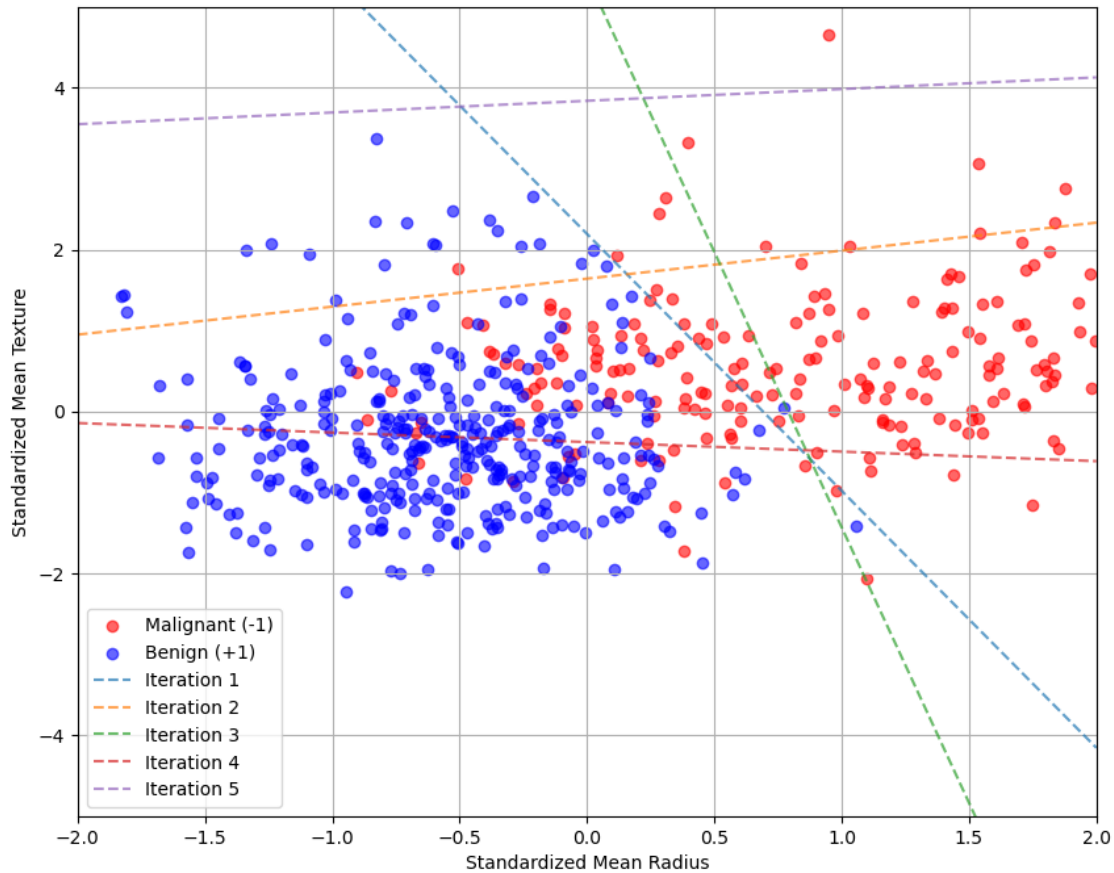
# Plot the data points
plt.scatter(X_std[y == -1, 0], X_std[y == -1, 1], color='red', label='Malignant
↳(-1)', alpha=0.6)
plt.scatter(X_std[y == 1, 0], X_std[y == 1, 1], color='blue', label='Benign
↳(+1)', alpha=0.6)

# Plot the first 5 decision boundaries
for i, (w, b) in enumerate(perceptron.history[:5]):
    x_plot = np.linspace(-2, 2, 100)
    y_plot = (-w[0] * x_plot - b) / w[1]

    plt.plot(x_plot, y_plot, label=f'Iteration {i+1}', linestyle='--', alpha=0.
↳7)

plt.xlabel('Standardized Mean Radius')
plt.ylabel('Standardized Mean Texture')
plt.legend()
plt.grid(True)
plt.xlim(-2, 2)
plt.ylim(-5, 5)
plt.show()
```

Total updates in history: 16094



1.2.1 5.

TODO: How many updates do you need until convergence (i.e. until no more model updates occur)? Explain why.

It depends on the linear separability and the learning rate. For separable data, the perceptron will converge after a finite amount of updates and stops when there are no more mistakes in the training data. We can check it using `len(perceptron.history)`.

```
[6]: # =====
# 6. Evaluate Performance Over Multiple Runs
# =====

#TODO: Evaluate performance over multiple runs. Compute and store test_
      ↳ accuracies

#TODO: Plot histogram for the test accuracies
num_runs = 100
test_accuracies = []
```



```

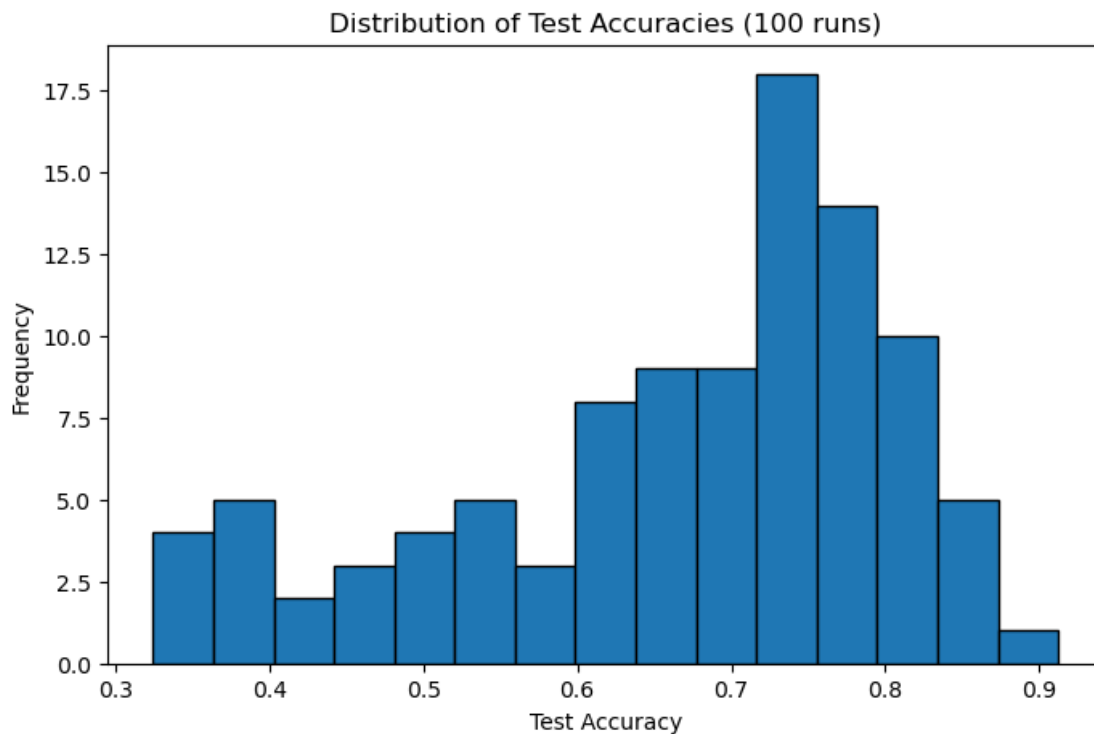
for _ in range(num_runs):
    X_train, X_test, y_train, y_test = train_test_split(
        X_std, y, test_size=0.2
    )

    perceptron = Perceptron(learning_rate=0.1, num_epochs=1000)
    perceptron.train(X_train, y_train)

    # Test accuracy
    y_test_pred = perceptron.predict(X_test)
    test_acc = accuracy_score(y_test, y_test_pred)
    test accuracies.append(test_acc)

plt.figure(figsize=(8, 5))
plt.hist(test accuracies, bins=15, edgecolor='black')
plt.xlabel('Test Accuracy')
plt.ylabel('Frequency')
plt.title('Distribution of Test Accuracies (100 runs)')
plt.show()

```



1.2.2 (a)

TODO: What does the shape of the histogram tell you?

The histogram gives us insight about the models behavior and the dataset. The higher the peak the more consistent is the accuracy. A wide histogram indicates that the model is sensitive to the training data. If it is left-skewed than our data is more problematic to separate.

```
[17]: # (b)
#TODO: Compute the sample mean and standard deviation of the test accuracy
mean_acc = np.mean(test_accuracies)
std_acc = np.std(test_accuracies)
print(f"Mean Test Accuracy: {mean_acc:.3f}")
print(f"Standard Deviation: {std_acc:.3f}")
```

Mean Test Accuracy: 0.665

Standard Deviation: 0.142

1.2.3 (c)

TODO: Given enough data points and for many training runs, what type of probability distribution would the histogram approximate and what is the reason for that?

It would approximate a right-skewed normal-distribution. This is due to the Central Limit Theorem, as the test accuracy is the average of many independent predictions with noise.

```
[8]: # (d)
p_values = [0, 10, 20, 30, 40, 50] # % of flipped training labels
#TODO: Add noise by flipping p% of labels. Visualize the effect using
↳ histograms for each p.

num_runs = 100
results = {p: [] for p in p_values}

for p in p_values:
    for _ in range(num_runs):
        # Split data
        X_train, X_test, y_train, y_test = train_test_split(X_std, y,
↳ test_size=0.2)

        # Add noise
        if p > 0:
            n_noise = int(len(y_train) * p/100)
            noise_indices = np.random.choice(len(y_train), n_noise,
↳ replace=False)
            y_train[noise_indices] *= -1 # Flip labels

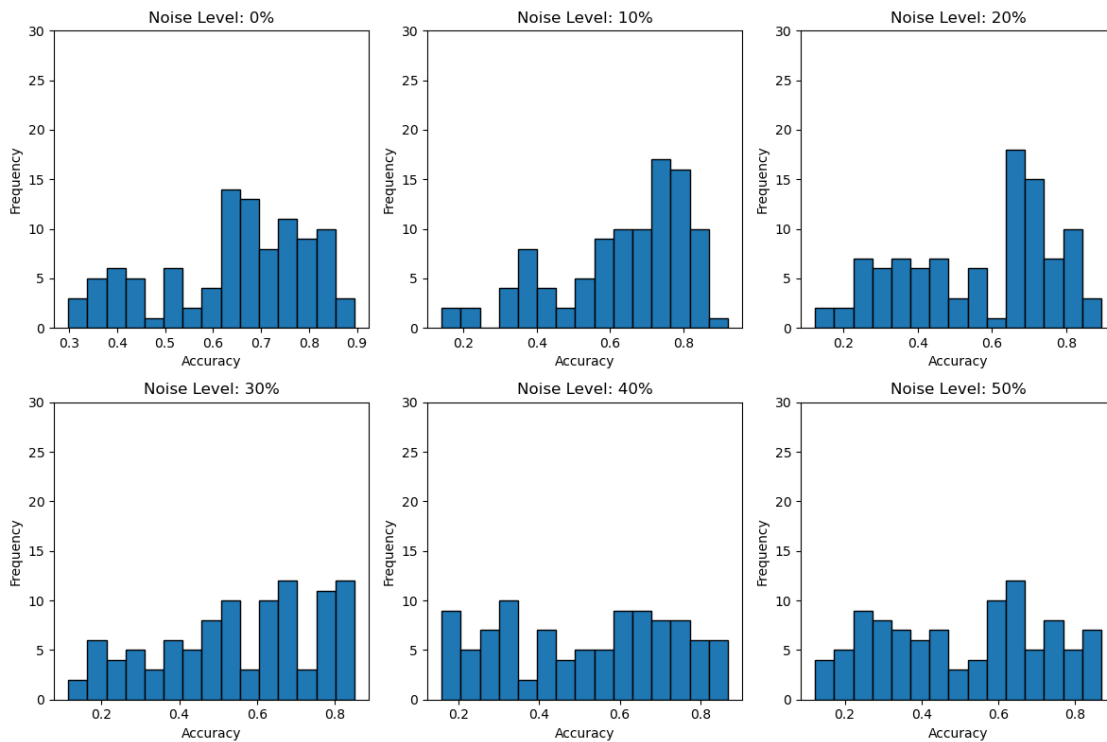
        # Train and test
        perceptron = Perceptron(learning_rate=0.1, num_epochs=100)
        perceptron.train(X_train, y_train)
```

```

y_test_pred = perceptron.predict(X_test)
test_acc = accuracy_score(y_test, y_test_pred)
results[p].append(test_acc)

# Plot histograms
plt.figure(figsize=(12, 8))
for i, p in enumerate(p_values):
    plt.subplot(2, 3, i+1)
    plt.hist(results[p], bins=15, edgecolor='black')
    plt.title(f'Noise Level: {p}%')
    plt.xlabel('Accuracy')
    plt.ylabel('Frequency')
    plt.ylim(0, 30)
plt.tight_layout()
plt.show()

```



TODO: Interpret the results

As noise increases, the test becomes less accurate (left-skewed) and the variance will become wider, because the variance increases. The perceptron assumes perfect linear separability, which deteriorates when the labels flip.

1.3 Exercise 3: SVM

```
[9]: import numpy as np
import matplotlib.pyplot as plt
from sklearn.datasets import make_circles, make_blobs
from cvxopt import matrix, solvers # Install cvxopt via "pip install cvxopt"

[10]: # =====
# 1. Complete SVM implementation
# =====

class DualSVM:
    def __init__(self, C=1.0, kernel="linear", gamma=1.0):
        self.C = C # Regularization constant
        self.kernel = kernel # Kernel type: "linear" or "rbf"
        self.gamma = gamma # Kernel parameter ("bandwidth")
        self.alpha = None # Lagrange multipliers
        self.sv_X = None # Support vectors
        self.sv_y = None # Support vector labels
        self.w = None # Weights
        self.b = None # Bias

    def linear_kernel(self, X1, X2):
        #TODO: Implement linear kernel
        return np.dot(X1, X2.T)

    def rbf_kernel(self, X1, X2):
        #TODO: Implement RBF kernel
        X1_sq = np.sum(X1**2, axis=1).reshape(-1, 1)
        X2_sq = np.sum(X2**2, axis=1).reshape(1, -1)
        dist_sq = X1_sq - 2 * np.dot(X1, X2.T) + X2_sq
        return np.exp(-self.gamma * dist_sq)

    def compute_kernel(self, X1, X2):
        if self.kernel == "linear":
            return self.linear_kernel(X1, X2)
        elif self.kernel == "rbf":
            return self.rbf_kernel(X1, X2)
        else:
            raise ValueError("Unknown kernel type.")

    def fit(self, X, y):
        n_samples, n_features = X.shape

        # Compute kernel matrix K:  $K[i, j] = K(x_i, x_j)$ 
        K = self.compute_kernel(X, X)

        """
```

```

The dual objective is:
    max sum_i alpha_i - 1/2 sum_i sum_j alpha_i alpha_j y_i y_j K(x_i,
↪x_j)
subject to:
    sum_i alpha_i y_i = 0 and 0 <= alpha_i <= C for all i.
In QP formulation:
    P = (y_i y_j K(x_i, x_j))_{i,j},    q = -1 (vector),
    A = y^T, b = 0, and G, h implement 0 <= alpha_i <= C.
"""

# TODO: Use the matrixx function of cvxopt to define the QP parameters
P = matrix(np.outer(y, y) * K, tc='d') # Use "d" flag to make sure that
↪the matrix is in double precision format
q = matrix([-1.0] * n_samples, (n_samples, 1), "d") # Use "d" flag to
↪make sure that the matrix is in double precision format
A = matrix(y, (1, n_samples), "d") # Use "d" flag to make sure that the
↪matrix is in double precision format (labels are integers)
b = matrix(0., (1, 1), "d") # Use "d" flag to make sure that the matrix
↪is in double precision format

# TODO: Implement inequality constraints by defining G and h
G = matrix(np.vstack((-np.eye(n_samples), np.eye(n_samples))), tc='d')
↪# Use "d" flag to make sure that the matrix is in double precision format
h = matrix(np.hstack((np.zeros(n_samples), np.ones(n_samples) * self.
↪C)), tc='d') # Use "d" flag to make sure that the matrix is in double
↪precision format

# Solve the QP problem using cvxopt
solvers.options["show_progress"] = False
solution = solvers.qp(P, q, G, h, A, b)
alphas = np.ravel(solution["x"]) # Get optimal alphas

# Get support vectors (i.e. data points with non-zero lagrange
↪multipliers, that are on the margin)
sv = alphas > 1e-5 # alpha > 1e-5 to account for numerical errors
self.alpha = alphas[sv]
self.sv_X = X[sv]
self.sv_y = y[sv]

# The bias corresponds to the average error over all support vectors:
# Why does the bias corresponds to the average error over all support
↪vectors?
# The answer is that the bias is the average of the differences between
↪the true labels and the predicted labels

```

```

        # for the support vectors. The predicted labels are computed by the
        ↪ decision function  $f(x) = \sum(\alpha_i y_i K(x, x_i)) + b$ .
        # The difference between the true labels and the predicted labels is
        ↪ the error for each support vector.
        # The bias is the average of these errors.
        self.b = np.mean(self.sv_y - np.sum(self.alpha * self.sv_y * K[sv][:,
        ↪ sv], axis=1))

    def predict(self, X):
        #TODO: Implement the decision function and return the corresponding
        ↪ predicted labels
        return np.sum(self.alpha * self.sv_y * self.compute_kernel(X, self.
        ↪ sv_X), axis=1) + self.b

```

```

[11]: # =====
# 2. Apply linear SVM on blobs
# =====

# TODO: Generate blobs dataset
X_linear, y_linear = make_blobs(n_samples=100, n_features=2, centers=2,
    ↪ random_state=114514)

# Convert labels from {0,1} to {-1,1}
y_linear = 2 * (y_linear - 0.5)

#TODO: Train SVM with linear kernel
svm_linear = DualSVM(C=2.5, kernel="linear")
svm_linear.fit(X_linear, y_linear)

#TODO: Plot decision boundary
def plot_decision_boundary(svm, X, y, title="Decision Boundary"):
    # Create a grid to plot the decision boundary
    x_min, x_max = X[:, 0].min() - 1, X[:, 0].max() + 1
    y_min, y_max = X[:, 1].min() - 1, X[:, 1].max() + 1
    xx, yy = np.meshgrid(np.linspace(x_min, x_max, 100), np.linspace(y_min,
    ↪ y_max, 100))

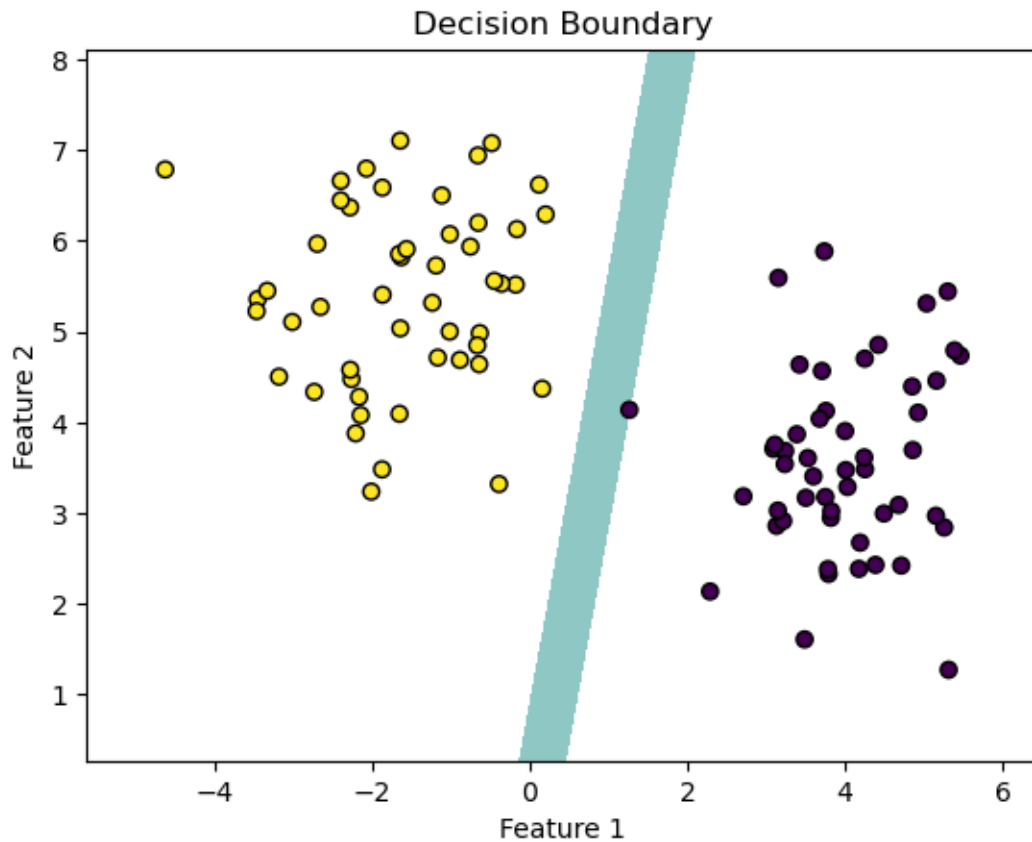
    # Compute the decision function for each point in the grid
    # .ravel(): [[], [], ..., []] |-> [...] create a flattened array
    # np.c_() concatenates the two arrays along the second axis (i.e. columns)
    # after doing this we get 100x100 pairs of points and feed it to the SVM
    Z = svm.predict(np.c_[xx.ravel(), yy.ravel()])
    Z = Z.reshape(xx.shape)

    # Plot the decision boundary and the data points
    plt.contourf(xx, yy, Z, levels=[-1, 0], alpha=0.5)
    plt.scatter(X[:, 0], X[:, 1], c=y, edgecolors='k', marker='o')

```

```
plt.title(title)
plt.xlabel("Feature 1")
plt.ylabel("Feature 2")
plt.show()

plot_decision_boundary(svm_linear, X_linear, y_linear)
```

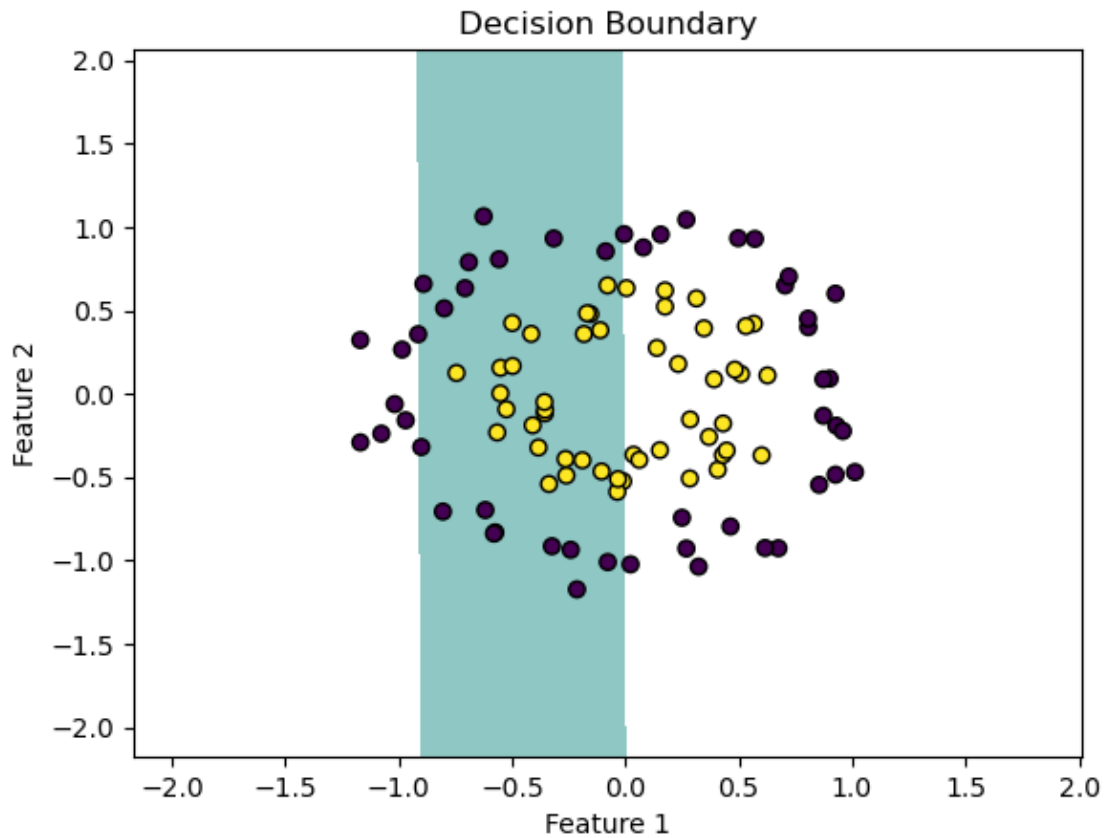


```
[12]: # =====
# 3. Apply linear SVM on circles
# =====

#TODO: Generate circles dataset
X_circles, y_circles = make_circles(n_samples=100, noise=0.1, factor=0.5,
    ↪random_state=114514)
y_circles = 2 * (y_circles - 0.5) # Convert labels from {0,1} to {-1,1}

#TODO: Train SVM with linear kernel
svm = DualSVM(C=2.5, kernel="linear")
svm.fit(X_circles, y_circles)
```

```
#TODO: Plot decision boundary
plot_decision_boundary(svm, X_circles, y_circles)
```



```
[13]: # =====
# 4. Apply feature transformation
# =====

def transform_features(X):
    # TODO: compute feature transformation:  $f(x) = [x_1, x_2, x_1^2 + x_2^2]$ 
    x1 = X[:, 0]
    x2 = X[:, 1]
    return np.column_stack((x1, x2, x1**2 + x2**2))

#TODO: Train SVM with linear kernel on transformed features
X_transformed = transform_features(X_circles)
svm_transformed = DualSVM(C=2.5, kernel="linear")
svm_transformed.fit(X_transformed, y_circles)
```

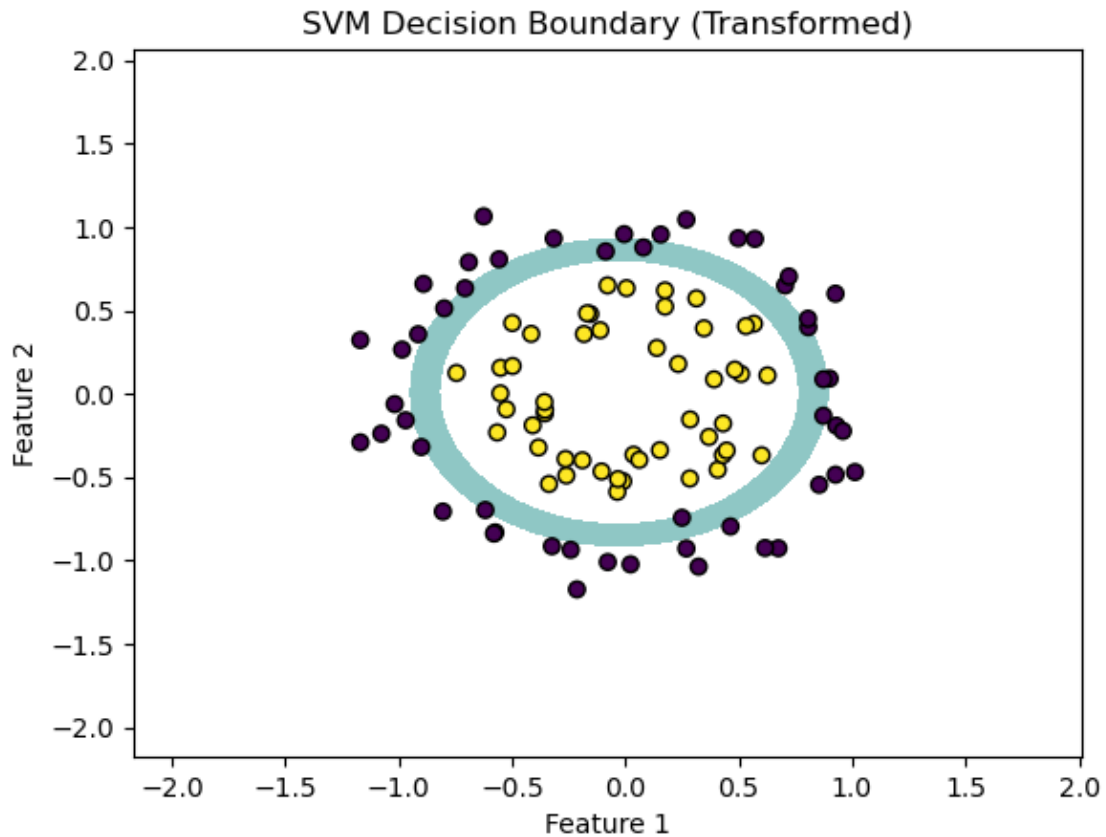


```

def plot_decision_boundary_transformed(X, y, model, title="SVM Decision_
↳Boundary (Transformed)":
    # TODO: Implement plotting function for decision boundary in the_
↳transformed feature space
    # Hint: You could do this by creating a 2D meshgrid which you transform_
↳using the feature mapping.
    # Then, after evaluating the model on it, you can plot the result as a_
↳contour plot (plt.contourf).
    x_min, x_max = X[:, 0].min() - 1, X[:, 0].max() + 1
    y_min, y_max = X[:, 1].min() - 1, X[:, 1].max() + 1
    xx, yy = np.meshgrid(np.linspace(x_min, x_max, 100), np.linspace(y_min,
↳y_max, 100))
    Z = model.predict(transform_features(np.c_[xx.ravel(), yy.ravel()]))
    Z = Z.reshape(xx.shape)
    plt.contourf(xx, yy, Z, levels=[-1, 0], alpha=0.5)
    plt.scatter(X[:, 0], X[:, 1], c=y, edgecolors='k', marker='o')
    plt.title(title)
    plt.xlabel("Feature 1")
    plt.ylabel("Feature 2")
    plt.show()

#TODO: Plot decision boundary in the transformed feature space
plot_decision_boundary_transformed(X_circles, y_circles, svm_transformed)

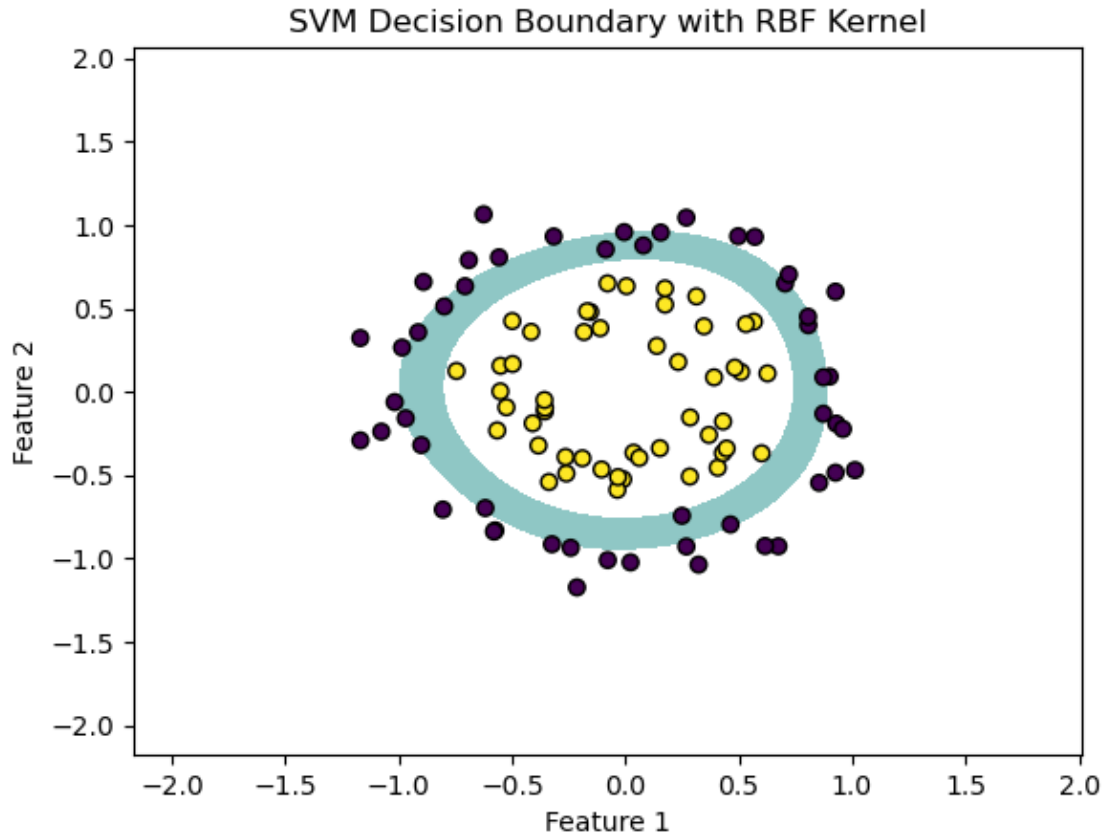
```



```
[14]: # =====
# 5. SVM with RBF Kernel on Circular Data
# =====

#TODO: Train SVM with RBF kernel on circular data
svm_rbf = DualSVM(C=2.5, kernel="rbf", gamma=1.0)
svm_rbf.fit(X_circles, y_circles)

#TODO: Plot decision boundary
plot_decision_boundary(svm_rbf, X_circles, y_circles, title="SVM Decision_
↳Boundary with RBF Kernel")
```



1.3.1 6.

TODO: Compare the decision boundaries from Tasks 3, 4, and 5. How does feature transformation differ from using an RBF kernel? When would one approach be preferable to the other?

The decision boundary of transformation looks smoother (rounder?) than radial basis function.

When we use transformation, we don't necessarily know how we could transform the data to make them separable by some hyperplanes. At the same time, we have full control and interpretability.

RBF kernel can adapt to different kind of boundaries automatically, whereas sometimes we need to tune the parameter γ .

1.3.2 7.

TODO: Besides the dual formulation, SVMs also have an equivalent primal formulation. The key factor in choosing which one to use as the optimization criterion is the dimensionality of the features. Explain why.

In high-dimensional spaces (i.e., when d is large), the primal problem becomes computationally expensive because it involves directly manipulating a high-dimensional weight vector w , and the number of operations required scales with d .

On the other hand, the dual problem only involves the kernel matrix, which is $n \times n$, where n is the number of samples. Since the kernel matrix doesn't directly depend on the feature dimension d , the dual formulation can be more efficient when d is large and n is smaller.