Name: Barathkrishna Satheeshkumar AndrewID: bsathees

Problem 1

Option 2: 6 x 6

Problem 2

i = 0, j = 0, k = 1

Problem 3

- PC1
- PC2

Problem 4

Option 3: 10 x 10

<u>Problem 5</u>

4. t-SNE is a non-linear dimensionality technique that can learn embeddings of manifolds

M12-L1 Problem 1

This problem is intended to demonstrate PCA on a small 2D dataset. This will emphasize how PCs are computed and what they mean.

Computing the Principal Components

First, compute the principal components of the dataset by following these steps:

- 1. Compute M (1×2) , the mean of each dimension in X
- 2. Compute S (2×2) , the covariance matrix of X (see np.cov)
- 3. Report w, the 2 eigenvalues of S (see np.linalg.eig)
- 4. Get e1 and e2, the eigenvectors corresponding to the elements of w

The principal components in this problem are then e1 and e2.

```
In [3]: print('X:\n', X)
        # YOUR CODE GOES HERE: Compute M
        M = np.mean(X, axis=0)
        print('\nMean of each dimension:\n', M)
        # YOUR CODE GOES HERE: Compute S
        S = np.cov(X, rowvar=False)
        print('\nCovariance Matrix:\n', S)
        # YOUR CODE GOES HERE: Compute w
        w, V = np.linalg.eig(S)
        print('\nEigenvalues of covariance matrix:\n',w)
        # YOUR CODE GOES HERE: Compute e1, e2
        e1 = V[:, 0]
        e2 = V[:, 1]
        print('\nPrincipal Components:')
        print('e1:',e1)
        print('e2:',e2)
       Χ:
        [[2.5 2.4]
        [0.5 \ 0.7]
        [2.2 2.9]
        [1.9 2.2]
        [3.1 3. ]
        [2.3 2.7]
        [2. 1.6]
        [1. 1.1]
        [1.5 1.6]
        [1.1 0.9]]
       Mean of each dimension:
        [1.81 1.91]
       Covariance Matrix:
        [[0.61655556 0.61544444]
        [0.61544444 0.71655556]]
       Eigenvalues of covariance matrix:
        [0.0490834 1.28402771]
       Principal Components:
       e1: [-0.73517866 0.6778734 ]
       e2: [-0.6778734 -0.73517866]
```

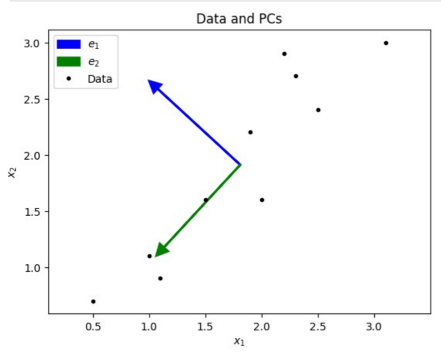
Plotting data with principal components

Complete the code below to plot the original data with principal components represented as unit vector arrows.

```
In [4]: plt.figure()
  plt.title("Data and PCs")
```

```
e1, e2 = e1.flatten(), e2.flatten()
plt.arrow(M[0],M[1],e1[0],e1[1], color="blue", linewidth=2, head_width=0.1, head_length=0.1, label="$e_1$")
plt.arrow(M[0],M[1],e2[0],e2[1], color="green", linewidth=2, head_width=0.1, head_length=0.1, label="$e_2$")
plt.plot(X[:,0],X[:,1],'.',color="black", label="Data")

plt.xlabel("$x_1$")
plt.ylabel("$x_2$")
plt.legend()
plt.axis("equal")
plt.show()
```



Plotting transformed data

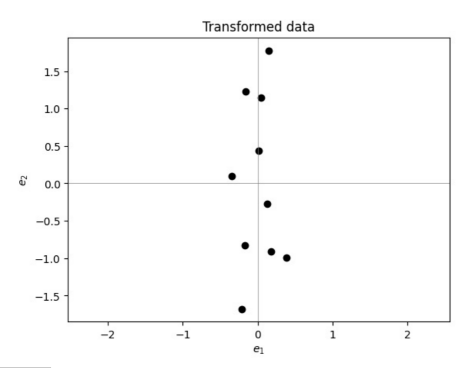
Now, transform the data with the formula $a_i = (x - \mu) \cdot e_i$.

Print the transformed data matrix columns a1 and a2.

0.09910944 1.14457216 0.43804614 1.22382056]

Then plot the transformed data on $e_1 - e_2$ axes.

```
In [5]: # YOUR CODE GOES HERE: Compute a1, a2
        X centered = X - M
        a1 = np.dot(X centered, e1)
        a2 = np.dot(X centered, e2)
        print("a_1 = ",a1)
print("a_2 = ",a2)
        plt.figure()
        plt.title("Transformed data")
        e1, e2 = e1.flatten(), e2.flatten()
        # YOUR CODE GOES HERE: Plot transformed data
        plt.scatter(a1, a2, color="black", label="Transformed Data")
        plt.axhline(0, color='gray', linewidth=0.5)
        plt.axvline(0, color='gray', linewidth=0.5)
        plt.xlabel("$e_1$")
        plt.ylabel("$e_2$")
        plt.axis("equal")
        plt.show()
       a\_1 = \begin{bmatrix} -0.17511531 & 0.14285723 & 0.38437499 & 0.13041721 & -0.20949846 & 0.17528244 \end{bmatrix}
        a = [-0.82797019 \ 1.77758033 \ -0.99219749 \ -0.27421042 \ -1.67580142 \ -0.9129491
```



Processing math: 100%

M12-L1 Problem 2

Sometimes the dimensionality is greater than the number of samples. For example, in this problem X has 19 features, but there are only 4 data points. You will need to use the alternate PCA formulation in this case. Follow the steps in the cells below to implement this method.

Computing Principal Components

The A matrix

First, you should compute the A matrix, where A is $(X - \mu)'$. (Note the transpose)

Print this matrix below. It should have size 19×4 .

```
In [13]: # YOUR CODE GOES HERE
        mu = np.mean(X, axis=0)
        X_{centered} = X - mu
        A = X_centered.T
        print("A = \n", A)
        [[-2.75 0.25 0.25 2.25]
        [ 1.25 2.25 -2.75 -0.75]
        [ 2. -4. 2.
                         0. ]
        [-3.5 1.5 0.5 1.5]
        [ 3.5 -4.5 -0.5
                         1.5
        [ 2.25 3.25 -1.75 -3.75]
              6. -4. -3. ]
        [ 2.25 1.25 -1.75 -1.75]
        [-1.75 0.25 1.25 0.25]
        [ 0.75 -0.25 1.75 -2.25]
        [ 1.25 -2.75 -1.75 3.25]
        [ 0.5 -0.5 -3.5 3.5 ]
        [ 1. -1. -3.
                         3. ]
        [ 1.75 -0.25 -2.25 0.75]
        [-1.5 -1.5 1.5 1.5]
        [-3.
              1. 0.
                        2. ]
        [ 0.5 -0.5 3.5 -3.5 ]
        [ 0.25 -3.75 3.25 0.25]
        [ 0.25 -1.75 2.25 -0.75]]
```

"Small" covariance matrix

By transposing $X - \mu$ to get A, now we can compute a smaller covariance matrix with A'A. Compute this matrix, C, below and print the result.

```
In [14]: # YOUR CODE GOES HERE
    n = A.shape[0]
    C = (1 / (n - 1)) * np.dot(A.T, A)
    print("C = \n", C)

C =
    [[ 3.88194444 -1.04861111 -1.46527778 -1.36805556]
    [-1.04861111 6.74305556 -2.95138889 -2.74305556]
    [-1.46527778 -2.95138889 5.46527778 -1.04861111]
    [-1.36805556 -2.74305556 -1.04861111 5.15972222]]
```

Finding nonzero eigenvectors

Next, find the useful (nonzero) eigenvectors of C.

For validation purposes, there should be 3 useful eigenvectors, and the first one is [-0.06628148 -0.79038331 0.47285044 0.38381435].

Keep these eigenvectors in a 4×3 array e.

```
In [21]: # YOUR CODE GOES HERE
    eigenvalues, eigenvectors = np.linalg.eig(C)

sorted_indices = np.argsort(eigenvalues)[::-1]
    eigenvalues = eigenvalues[sorted_indices]
    eigenvectors = eigenvectors[:, sorted_indices]

e = eigenvectors[:, :-1]
    print("Eigenvectors, e:\n", e)

Eigenvectors, e:
    [[-0.06628148    0.04124587   -0.86249959]
    [-0.79038331   -0.06822502    0.34733208]
    [ 0.47285044   -0.69123739    0.22046165]
```

Calculating "eigenfaces"

[0.38381435 0.71821654 0.29470586]]

Now, we have all we need to compute U, the matrix of eigenfaces.

```
\mathbf{U_i} = \mathbf{Ae_i}
(19 \times 3) = (19 \times 4)(4 \times 3)
```

Compute and print U. Be sure to normalize your eigenvectors e before using the above equation.

```
In [16]: # YOUR CODE GOES HERE
    e_normalized = e / np.linalg.norm(e, axis=0)
    U = np.dot(A, e_normalized)
    print("Eigenfaces, U:\n",U)
```

```
Eigenfaces, U:
[-3.44941376 1.26029144 -1.12392626]
[ 3.97467115 -1.02708294 -2.67340418]
[-0.14144304 0.48500804 4.0920363 ]
[-4.98467116 -1.61257473 -2.30274969]
[-7.85142613 0.24219565 -0.54447132]
[-2.63627584 -0.0396916 -2.40800213]
[ 0.6054134 -0.77372912 1.94546084]
[ 0.11179069 -2.77766199 -1.010988 ]
[ 2.51061063  3.78304534 -1.46130156]
[ 0.05042462  4.98782419 -0.34506112]
[ 0.45699358  4.33783268 -0.98709906]
[-0.69444948 2.18318305 -1.87121663]
[ 2.56999436  0.08093747  1.54550254]
[ 0.67367721 -4.8783533 -0.86477055]
[ 4.58008454 -1.80081206 -0.72794335]
[ 2.14265314 -1.96424127 -0.5484467 ]]
```

Projecting data into 3D

Now project your data into 3 dimensions with the formula:

```
\Omega=U^{
m T}A (3	imes4)=(3	imes19)(19	imes4) Call the projected data \Omega " W ". Print W.T
```

Reconstructing data in 19-D

[67.37906326 82.10439798

We can project the transformed data W back into the original 19-D space using:

27.29912392]]

$$\Gamma_f = U\Omega + \Psi$$
 where: $\Gamma_f =$ reconstructed data $U =$ eigenfaces $\Omega =$ Reduced data $\Psi =$ Means

Do this, and compute the MSE between each reconstructed sample and corresponding original points. Report all 4 MSE values.

```
In [35]: # YOUR CODE GOES HERE
    Psi = np.mean(X, axis=0).reshape(-1, 1)
    Gamma_f = np.dot(U, W) + Psi
    mse = np.mean((X - Gamma_f.T) ** 2, axis=1)

    for i in range(4):
        print("MSE for sample %d: %e" %(i+1,mse[i]))

MSE for sample 1: 1.371806e+03
    MSE for sample 2: 1.762219e+05
    MSE for sample 3: 9.985760e+04
    MSE for sample 4: 8.132361e+04
```

2-D Reconstruction

What if we had only used the first 2 eigenvectors to compute the eigenfaces? Below, redo the earlier calculations, but use only two eigenfaces. Compute the 4 MSE values that you would get in this case.

(You should get an MSE of 3.626 for the first sample.)

```
In [37]: # YOUR CODE GOES HERE
    e = eigenvectors[:, :2]
    e_normalized = e / np.linalg.norm(e, axis=0)
    U = np.dot(A, e_normalized)
    U = U/np.linalg.norm(U, axis=0)

W = np.dot(U.T, A)

Psi = np.mean(X, axis=0).reshape(-1, 1)
Gamma_f = np.dot(U, W) + Psi
mse = np.mean((X - Gamma_f.T) ** 2, axis=1)

print("Using only 2 eigenvectors:")
for i in range(4):
    print("MSE for sample %d: %e" %(i+1,mse[i]))
```

Using only 2 eigenvectors:
MSE for sample 1: 3.626804e+00
MSE for sample 2: 5.881609e-01
MSE for sample 3: 2.369586e-01
MSE for sample 4: 4.234322e-01

Problem 1

Problem Description

In this problem you will use PCA and TSNE to apply dimensionality reduction to 64x64 images of signed distance fields (SDFs) on parts belonging to 8 different classes. Each class is topologically similar, with some variation in void size and shape. These signed distance fields are helpful in the prediction of internal stress fields in the parts. You will also apply KNN to predict the class of the part with the reduced space.

Fill out the notebook as instructed, making the requested plots and printing necessary values.

You are welcome to use any of the code provided in the lecture activities.

Summary of deliverables:

- 3x8 subplot visualization of the first 3 samples from each of the 8 classes
- Bar plot of the variance explained for the first 25 PCs and the number of PCs required to explain > 90% of the variance in the training
- 4x8 subplot visualization of reconstructed samples using 3, 10, 50 and all PCs on the first sample from each of the 8 classes in the
 test set
- Test accuracy of KNN classifier trained on the 3D, 10D, and 50D PCA reduced feature spaces
- Plot of the 2D TSNE reduced feature space
- · Test accuracy of the KNN classifier trained on the 2D TSNE reduced feature space
- Discussion questions 1 and 2

Imports and Utility Functions:

```
In [1]: import numpy as np
        import matplotlib.pyplot as plt
        from scipy import io
        from sklearn.decomposition import PCA
        from sklearn.manifold import TSNE
        from sklearn.neighbors import KNeighborsClassifier
        from sklearn.model selection import train test split
        def dataLoader(filepath):
            # Load and flatten the SDF dataset
            mat = io.loadmat(filepath)
            data = []
            for i in range(800):
                sdf = mat["sdf"][i][0].T
                data.append(sdf.flatten())
            data = np.vstack(data)
            # Assign labels
            labels = np.repeat(np.arange(8), 100)
            return data, labels
        def plot_sdf(data, ax = None, title = None):
            # If no axes, make them
            if ax is None:
                ax = plt.gca()
            # Reshape image data into square
            sdf = data.reshape(64,64)
            # Plot image, with bounds of the SDF values for the entire dataset
            ax.imshow(sdf, vmin=-0.31857, vmax=0.206349, cmap="jet")
            ax.axis('off')
            # If there is a title, add it
            if title:
                ax.set title(title)
```

Visualization

Using the provided dataLoader() function, load the data and labels from sdf_images.mat. The returned data will contain 800 samples, with 4096 features. Then, using the provided plot_sdf() function, generate a 3x8 subplot figure containing visualizations of the first 3 SDFs in each class.

```
In [3]: # YOUR CODE GOES HERE
  data, labels = dataLoader('data/sdf_images.mat')
  fig, axes = plt.subplots(3, 8, figsize=(15, 6))

for i in range(8):
```

```
idx = i * 100 + j
           plot sdf(data[idx], ax=axes[j, i], title=f"Class {i+1}, Sample {j+1}")
plt.tight layout()
plt.show()
Class 1, Sample 1
                   Class 2, Sample 1
                                       Class 3, Sample 1
                                                          Class 4, Sample 1
                                                                             Class 5, Sample 1
                                                                                                 Class 6, Sample 1
                                                                                                                     Class 7, Sample 1
                                                                                                                                        Class 8, Sample 1
                                      Class 3, Sample 2
                                                          Class 4, Sample 2
                                                                             Class 5, Sample 2
                                                                                                                     Class 7, Sample 2
                                                                                                                                        Class 8, Sample 2
Class 1, Sample 2
                   Class 2, Sample 2
                                                                                                 Class 6, Sample 2
Class 1, Sample 3
                   Class 2, Sample 3
                                       Class 3, Sample 3
                                                          Class 4, Sample 3
                                                                              Class 5, Sample 3
                                                                                                 Class 6, Sample 3
                                                                                                                     Class 7, Sample 3
                                                                                                                                        Class 8, Sample 3
```

Explained Variance

for j in range(3):

Use $train_test_split()$ to partition the data and labels into a training and test set with $test_size = 0.2$ and $test_size = 0.2$. Then train a PCA model on the training data and generate a bar plot of the variance explained for the first 25 principal components. Determine the number of principal components required to explain > 90% of the variance in the training data.

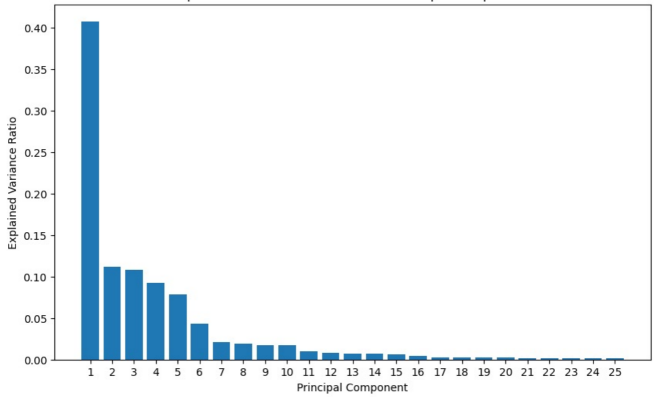
```
In []: # YOUR CODE GOES HERE
   X_train, X_test, y_train, y_test = train_test_split(data, labels, test_size=0.2, random_state=0)

pca = PCA(n_components=25)
pca.fit(X_train)

plt.figure(figsize=(10, 6))
plt.bar(range(1, 26), pca.explained_variance_ratio_)
plt.xlabel('Principal Component')
plt.ylabel('Explained Variance Ratio')
plt.title('Explained Variance for the First 25 Principal Components')
plt.xticks(range(1, 26))
plt.show()

cumulative_variance = np.cumsum(pca.explained_variance_ratio_)
n_components_90 = np.argmax(cumulative_variance > 0.90) + 1 # Add 1 to include the first component
print(f"Number of components required to explain > 90% of the variance: {n_components_90}")
```

Explained Variance for the First 25 Principal Components

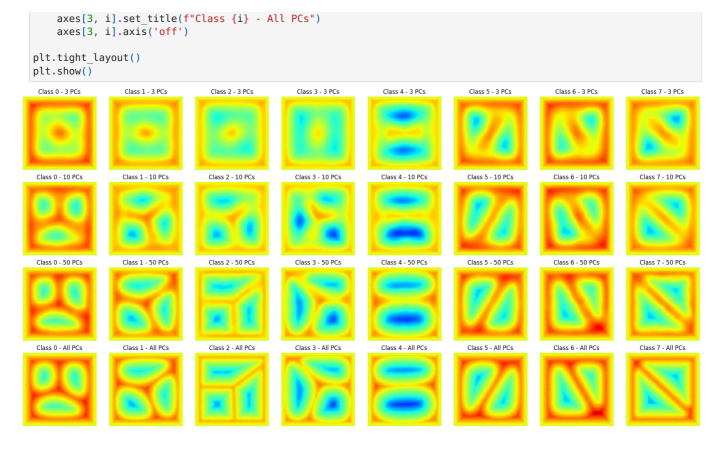


Number of components required to explain > 90% of the variance: 9

PCA Reconstruction

Using the training data, generate 4 PCA models using 3, 10, 50, and all of the principal components. Use these models to transform the test data into the reduced space, and then reconstruct the data from the reduced space. Plot the reconstruction for each model, on the first occurence of each class in the test set. Your generated plot should be a 4x8 subplot figure, with each subplot title containing the class and the number of PCs used.

```
In [6]: # YOUR CODE GOES HERE
        pca 3 = PCA(n components=3)
        pca_3.fit(X_train)
        pca_10 = PCA(n_components=10)
        pca_10.fit(X_train)
        pca 50 = PCA(n components=50)
        pca_50.fit(X_train)
        pca_all = PCA(n_components=None)
        pca all.fit(X train)
        first occurrence = []
        for i in range(8):
            first_occurrence.append(np.where(y_test == i)[0][0])
        fig, axes = plt.subplots(4, 8, figsize=(20, 10))
        for i, idx in enumerate(first occurrence):
            sample = X_test[idx]
            recon_3 = pca_3.inverse_transform(pca_3.transform(sample.reshape(1, -1)))
            axes[0, i].imshow(recon_3.reshape(64, 64), cmap="jet", vmin=-0.31857, vmax=0.206349)
            axes[0, i].set_title(f"Class {i} - 3 PCs")
            axes[0, i].axis('off')
            recon 10 = pca 10.inverse transform(pca 10.transform(sample.reshape(1, -1)))
            axes[1, i].imshow(recon_10.reshape(64, 64), cmap="jet", vmin=-0.31857, vmax=0.206349)
            axes[1, i].set title(f"Class {i} - 10 PCs")
            axes[1, i].axis('off')
            recon_50 = pca_50.inverse_transform(pca_50.transform(sample.reshape(1, -1)))
            axes[2, i].imshow(recon_50.reshape(64, 64), cmap="jet", vmin=-0.31857, vmax=0.206349)
            axes[2, i].set_title(f"Class {i} - 50 PCs")
            axes[2, i].axis('off')
            recon_all = pca_all.inverse_transform(pca_all.transform(sample.reshape(1, -1)))
            axes[3, i].imshow(recon all.reshape(64, 64), cmap="jet", vmin=-0.31857, vmax=0.206349)
```



KNN on PCA Reduced Data

Now train a KNN classifier to predict the class of the 3D, 10D, and 50D PCA reduced data. You should train the KNN on the reduced training data, and report the prediction accuracy on the test set. You will also need to determine the <code>n_neighbors</code> parameter for your KNN classifier that gives good results.

```
In [7]: # YOUR CODE GOES HERE
         from sklearn.model selection import GridSearchCV
         X train 3D = pca 3.transform(X train)
         X_{test_3D} = pca_3.transform(X_{test_3D})
         X_train_10D = pca_10.transform(X_train)
         X test 10D = pca 10.transform(X test)
         X train 50D = pca 50.transform(X train)
         X_{\text{test}}_{50D} = pca_{50}.transform(X_{\text{test}})
         knn = KNeighborsClassifier()
         param_grid = {'n_neighbors': range(1, 21)}
         def train_and_evaluate_knn(X_train, X_test, y_train, y_test, param_grid):
             grid search = GridSearchCV(knn, param grid, cv=5)
             grid_search.fit(X_train, y_train)
             best_n_neighbors = grid_search.best_params_['n_neighbors']
             best_knn = grid_search.best_estimator_
             accuracy = best_knn.score(X_test, y_test)
             return best n neighbors, accuracy
         best\_n\_3D, \ accuracy\_3D = train\_and\_evaluate\_knn(X\_train\_3D, \ X\_test\_3D, \ y\_train, \ y\_test, \ param\_grid)
         best n 10D, accuracy 10D = train and evaluate knn(X train 10D, X test 10D, y train, y test, param grid)
         best_n_50D, accuracy_50D = train_and_evaluate_knn(X_train_50D, X_test_50D, y_train, y_test, param_grid)
         print(f"Best n_neighbors for 3D PCA: {best_n_3D}, Test accuracy: {accuracy_3D * 100:.2f}%")
         print(f"Best n_neighbors for 10D PCA: {best_n_10D}, Test accuracy: {accuracy_10D * 100:.2f}%")
print(f"Best n_neighbors for 50D PCA: {best_n_50D}, Test accuracy: {accuracy_50D * 100:.2f}%")
        Best n neighbors for 3D PCA: 15, Test accuracy: 70.62%
        Best n_neighbors for 10D PCA: 14, Test accuracy: 91.25\%
        Best n neighbors for 50D PCA: 14, Test accuracy: 92.50%
```

TSNE Visualization

with a scatter plot, coloring each point according to its class.

```
In [10]: # YOUR CODE GOES HERE
pca_50 = PCA(n_components=50)
X_pca_50 = pca_50.fit_transform(data)

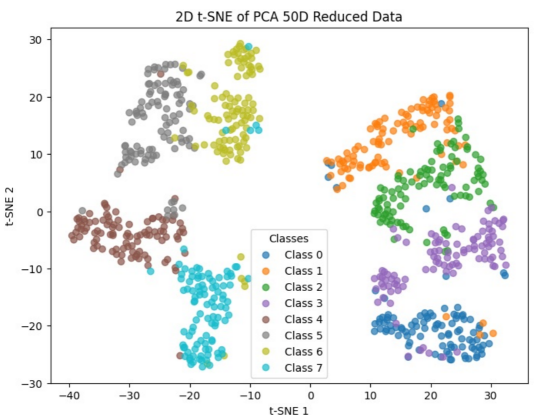
tsne = TSNE(n_components=2, random_state=0)
X_tsne_2D = tsne.fit_transform(X_pca_50)

plt.figure(figsize=(8, 6))
scatter = plt.scatter(X_tsne_2D[:, 0], X_tsne_2D[:, 1], c=labels, cmap='tab10', alpha=0.7)

plt.title('2D t-SNE of PCA 50D Reduced Data')
plt.xlabel('t-SNE 1')
plt.ylabel('t-SNE 2')

plt.legend(handles=scatter.legend_elements()[0], labels=[f'Class {i}' for i in range(8)], title='Classes')

plt.show()
```



KNN on PCA/TSNE Reduced Data

Using the same 2D PCA/TSNE data, split the data into train and test data and labels using train_test_split with a random_state = 0 parameter so you have the same train/test partition as before. Then, train a KNN on this 2D feature space with the training set, and report the KNN classifier accuracy on the test set. Again, you will need to determine the n_neighbors parameter in the KNN classifier that gives good results.

```
In [12]: # YOUR CODE GOES HERE
from sklearn.metrics import accuracy_score

X_train, X_test, y_train, y_test = train_test_split(X_tsne_2D, labels, test_size=0.2, random_state=0)

neighbors_range = range(1, 21)
best_accuracy = 0
best_k = 1

for k in neighbors_range:
    knn = KNeighborsClassifier(n_neighbors=k)
    knn.fit(X_train, y_train)
    y_pred = knn.predict(X_test)
    accuracy = accuracy_score(y_test, y_pred)

if accuracy > best_accuracy:
    best_accuracy = accuracy
    best_k = k

knn_final = KNeighborsClassifier(n_neighbors=best_k)
```

```
knn_final.fit(X_train, y_train)
y_pred_final = knn_final.predict(X_test)

final_accuracy = accuracy_score(y_test, y_pred_final)
print(f"Best n_neighbors: {best_k}")
print(f"Test accuracy of KNN classifier: {final_accuracy:.4f}")
```

Best n neighbors: 15

Test accuracy of KNN classifier: 0.9187

Discussion

- 1. Discuss how the number of principal components relates to the quality of reconstruction of the data. Using all of the principal components, should there be any error in the reconstruction of a sample from the training data? What about in the reconstruction of an unseen sample from the testing data?
- 2. Discuss how you determined k, the number of neighbors in your KNN models. Why do we perform dimensionality reduction to our data before feeding it to our KNN classifier?

Your response goes here

- 1. Using all principal components results in no reconstruction error, as it is a lossless transformation. With fewer components, the reconstruction is approximate, and the error increases, especially for unseen test data.
- 2. k is chosen by testing different values and selecting the one that gives the best accuracy. Dimensionality reduction reduces noise, mitigates the curse of dimensionality, and improves KNN's efficiency and accuracy.

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