PCA

May 24, 2021

If you plan to run the assignment locally: You can download the assignments and run them locally, but please be aware that as much as we would like our code to be universal, computer platform differences may lead to incorrectly reported errors even on correct solutions. Therefore, we encourage you to validate your solution in Coursera whenever this may be happening. If you decide to run the assignment locally, please: 1. Try to download the necessary data files from your home directory one at a time, 2. Don't update anything other than this Jupyter notebook back to Coursera's servers, and 3. Make sure this notebook maintains its original name after you upload it back to Coursera.

Note: You need to submit the assignment to be graded, and passing the validation button's test does not grade the assignment. The validation button's functionality is exactly the same as running all cells.

```
[3]: %matplotlib inline
%load_ext autoreload
%autoreload 2

import matplotlib.pyplot as plt
import numpy as np
import seaborn as sns
import pandas as pd
import time
import os
from sklearn.decomposition import TruncatedSVD

from aml_utils import test_case_checker, perform_computation
```

0.1 Attention:

This assignment is computationally heavy, and inefficient implementations may not pass the autograding even if they technically produce the correct results. To avoid this, make sure you read and understand all the instructions before starting to implement the tasks. Failure to follow the instructions closely will most likely cause timeouts.

It is **your responsibility** to make sure your implementation is not only **correct**, but also as **efficient** as possible. If you follow all the instructions provided, you should be able to have all the cells evaluated in under 10 minutes.

1 *Assignment Summary

CIFAR-10 is a dataset of 32x32 images in 10 categories, collected by Alex Krizhevsky, Vinod Nair, and Geoffrey Hinton. It is often used to evaluate machine learning algorithms. You can download this dataset from https://www.cs.toronto.edu/~kriz/cifar.html.

- For each category, compute the mean image and the first 20 principal components. Plot the error resulting from representing the images of each category using the first 20 principal components against the category.
- Compute the distances between mean images for each pair of classes. Use principal coordinate analysis to make a 2D map of the means of each categories. For this exercise, compute distances by thinking of the images as vectors.
- Here is another measure of the similarity of two classes. For class A and class B, define E(A | B) to be the average error obtained by representing all the images of class A using the mean of class A and the first 20 principal components of class B. Now define the similarity between classes to be (1/2)(E(A | B) + E(B | A)). If A and B are very similar, then this error should be small, because A's principal components should be good at representing B. But if they are very different, then A's principal components should represent B poorly. In turn, the similarity measure should be big. Use principal coordinate analysis to make a 2D map of the classes. Compare this map to the map in the previous exercise? are they different? why?

References:

• Textbook section 5.1 https://link.springer.com/chapter/10.1007/978-3-030-18114-7_5

2 0. Data

2.1 0.1 Description

CIFAR-10 is a dataset of 32x32 images in 10 categories, collected by Alex Krizhevsky, Vinod Nair, and Geoffrey Hinton. It is often used to evaluate machine learning algorithms. You can download this dataset from https://www.cs.toronto.edu/~kriz/cifar.html.

2.2 0.2 Information Summary

- Input/Output: This data has a set of 32 pixel rows, 32 pixel columns, and 3 color channels. Therefore, each single image, is vectorized, will consist of 32 × 32 × 3 elements (i.e., each image has 3072 dimensions). There are a total of 60000 samples labelled from 10 class. The data set is balanced with each class having exactly 6000 samples.
- Missing Data: There is no missing data.
- Final Goal: We want to understand the data using multi-dimensional scaling methods.

2.3 0.3 Loading The Data

If you are curious how the original data was obtained, we used the torchvision API to download and pre-process it. The ready-to-use data is stored in numpy format for easier access.

```
[4]: if os.path.exists('../PCA-lib/cifar10.npz'):
    np_file = np.load('../PCA-lib/cifar10.npz')
```

```
train_images_raw = np_file['train_images_raw']
    train_labels = np_file['train_labels']
    eval_images_raw = np_file['eval_images_raw']
    eval_labels = np_file['eval_labels']
else:
    import torchvision
    download_ = not os.path.exists('../PCA-lib/cifar10/')
    data_train = torchvision.datasets.CIFAR10('.../PCA-lib/cifar10', train=True,_
 →transform=None, target_transform=None, download=download_)
    data_eval = torchvision.datasets.CIFAR10('.../PCA-lib/cifar10', train=False,__
 →transform=None, target_transform=None, download=download_)
    train_images_raw = data_train.data
    train_labels = np.array(data_train.targets)
    eval_images_raw = data_eval.data
    eval_labels = np.array(data_eval.targets)
    np.savez('.../PCA-lib/cifar10.npz', train_images_raw=train_images_raw,_
 →train_labels=train_labels,
             eval_images_raw=eval_images_raw, eval_labels=eval_labels)
```

```
[6]: images_raw = np.concatenate([train_images_raw, eval_images_raw], axis=0)
    labels = np.concatenate([train_labels, eval_labels], axis=0)
    images_raw.shape, labels.shape
```

[6]: ((60000, 32, 32, 3), (60000,))

3 1. Principal Component Analysis

0. Let's say we have Data Matrix X with N rows (i.e., data points) and d columns (i.e., features).

$$X = [\cdots]_{N \times d}$$

1. Let's perform SVD on the X.

$$X = U_x S_x V_x^T$$

Let's assume N>d (We have 6000 data points per class, which is more than the 3072 dimenstions).

By the way SVD works, we should have

$$U_x = [\cdots]_{N \times d}$$

$$S_x = [\cdots]_{d \times d}$$

$$V_x = [\cdots]_{d \times d}$$

and

$$U_x^T U_x = I_{d \times d}$$

$$V_r^T V_x = I_{d \times d}$$

2. The textbook says we need the following decomposition for the covariance matrix Σ :

$$\Sigma \mathcal{U} = \mathcal{U} \Lambda$$

3. We assume that X has mean zero (i.e., we already subtracted the feature averages). If X has N rows (i.e., data items), we have

$$\Sigma = \frac{1}{N} X^T X$$

4. Let's find Σ in terms of U_x , S_x , and V_x

$$\Sigma = \frac{1}{N} X^{T} X = \frac{1}{N} V_{x} S_{x} U_{x}^{T} U_{x} S_{x} V_{x}^{T} = V_{x} \frac{S_{x}^{2}}{N} V_{x}^{T}$$

$$\Rightarrow \Sigma V_x = V_x \frac{S_x^2}{N}$$

5. By comparison, we have

$$\mathcal{U} = V_x$$

$$\Lambda = \frac{S_x^2}{N}$$

3.0.1 Considering the above:

- 1. There is no need to compute the covariance matrix Σ and then find its diagonalization; You can easily perform SVD on the data matrix X, and get what you need!
- 2. In fact, you do not even need the matrices V_x and U_x for computing the mean squared error; You can infer the mean squared error using only the S_x matrix.
 - Numpy's SVD function np.linalg.svd has an argument compute_uv that turns off returning the *U* and *V* matrices for better efficiency. Therefore, you may be able to save some runtime in large data sets if you only care about the mean squared error!

4 Task 1

Write a function pca_mse that takes two arguments as input

1. data_raw: a numpy array with the shape (N, \dots) , where N is the number of samples, and there may be many excess dimensions denoted by \dots . You will have to reshape this input data_raw matrix to obtain a shape of (N, d), where d is the vectorized data's dimension. For example, data_raw could have an input shape of (6000, 50, 50, 3). In this case you will have to reshape the input data to have a shape of (6000, 7500).

2. $num_components$: This is the number of PCA components that we want to retain. This variable is denoted by r in the PCA definition in the textbook.

and returns the variable mse which is the mean squared error of the PCA projection into the designated number of principal components.

Important Note: Make sure you use np.linalg.svd for the SVD operation. Do not use any other matrix factorization function for this question (such as np.linalg.eig).

Important Note: Make sure you read and understand the notes from the previous cells before you start implementing. Failing to properly set the arguments for np.linalg.svd or trying to find the mean squared error by calculating the reconstruction error may cause extreme delays and timeouts for your implementation.

Hint: If you don't know how to extract the mean squared error of the PCA projection, or don't have a fresh probability and statistics memory, take a look at the Principal Component Analysis chapter of the most recent version of the textbook; the subsection titled "The error in a low-dimensional representation" explains how to find the mean squared error of the PCA projection as a function of the eigenvalues that were dropped.

```
[7]: def pca_mse(data_raw, num_components=20):
    # your code here
    shapetuple = data_raw.shape
    nn = len(shapetuple)
    r=1
    for i in range(1,nn):
        r = r * shapetuple[i]
        data_raw.reshape(shapetuple[0],r)
        new_data_raw = data_raw - np.mean(data_raw,axis=0)
        s = np.linalg.svd(new_data_raw, full_matrices=True,compute_uv=False)

lam = (s**2)/shapetuple[0]
    mse = sum(lam[num_components:r])
```

```
[8]: # Performing sanity checks on your implementation
    some_data = (np.arange(35).reshape(5,7) ** 13) % 20
    some_mse = pca_mse(some_data, num_components=2)
    assert some_mse.round(3) == 37.903

# Checking against the pre-computed test database
    test_results = test_case_checker(pca_mse, task_id=1)
    assert test_results['passed'], test_results['message']
```

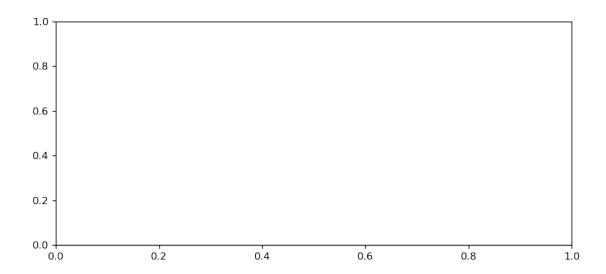
```
[31]: #Task 1 Test Cell
```

```
[9]: if perform_computation:
          class_names = []
          class_mses = []
          for cls_name, cls_label in class_to_idx.items():
              data_raw = images_raw[labels == cls_label,:,:,:]
              start_time = time.time()
              print(f'Processing Class {cls_name}', end='')
              cls_mse = pca_mse(data_raw, num_components=20)
              print(f' (The SVD operation took %.3f seconds)' % (time.
       →time()-start_time))
              class_names.append(cls_name)
              class_mses.append(cls_mse)
     Processing Class airplane (The SVD operation took 0.609 seconds)
     Processing Class automobile (The SVD operation took 0.605 seconds)
     Processing Class bird (The SVD operation took 0.619 seconds)
     Processing Class cat (The SVD operation took 0.597 seconds)
     Processing Class deer (The SVD operation took 0.608 seconds)
     Processing Class dog (The SVD operation took 0.601 seconds)
     Processing Class frog (The SVD operation took 0.609 seconds)
     Processing Class horse (The SVD operation took 0.607 seconds)
     Processing Class ship (The SVD operation took 0.597 seconds)
     Processing Class truck (The SVD operation took 0.600 seconds)
[10]: if perform computation:
          fig, ax = plt.subplots(figsize=(9,4.), dpi=120)
          sns.barplot(class_names, class_mses, ax=ax)
          ax.set_title('The Mean Squared Error of Representing Each Class by the 
       → Principal Components')
          ax.set_xlabel('Class')
          _ = ax.set_ylabel('Mean Squared Error')
             TypeError
                                                        Traceback (most recent call
      →last)
             <ipython-input-10-9fa5465e8c7e> in <module>
               1 if perform computation:
                     fig, ax = plt.subplots(figsize=(9,4.), dpi=120)
                     sns.barplot(class_names, class_mses, ax=ax)
                     ax.set_title('The Mean Squared Error of Representing Each Class_
      →by the Principal Components')
               5
                     ax.set_xlabel('Class')
```

```
→inner_f(*args, **kwargs)
                           44
                            45
                                                                  kwargs.update({k: arg for k, arg in zip(sig.parameters,__
→args)})
          ---> 46
                                                                  return f(**kwargs)
                            47
                                                    return inner_f
                            48
                         /opt/conda/lib/python3.8/site-packages/seaborn/categorical.py in_
→barplot(x, y, hue, data, order, hue_order, estimator, ci, n_boot, units, seed, __
→orient, color, palette, saturation, errcolor, errwidth, capsize, dodge, ax, u
→**kwargs)
                     3177 ):
                     3178
          -> 3179
                                                   plotter = _BarPlotter(x, y, hue, data, order, hue_order,
                                                                                                                                  estimator, ci, n boot, units, seed,
                     3180
                     3181
                                                                                                                                  orient, color, palette, saturation,
                         /opt/conda/lib/python3.8/site-packages/seaborn/categorical.py in in incomparison of the condary 
→__init__(self, x, y, hue, data, order, hue_order, estimator, ci, n_boot, u
→units, seed, orient, color, palette, saturation, errcolor, errwidth, capsize,
→dodge)
                     1582
                                                                                                   errwidth, capsize, dodge):
                     1583
                                                                   """Initialize the plotter."""
          -> 1584
                                                                  self.establish_variables(x, y, hue, data, orient,
                                                                                                                                                           order, hue_order, units)
                     1585
                     1586
                                                                  self.establish_colors(color, palette, saturation)
                         /opt/conda/lib/python3.8/site-packages/seaborn/categorical.py in in incomparison of the condary 
→establish_variables(self, x, y, hue, data, orient, order, hue_order, units)
                         154
                         155
                                                                                 # Figure out the plotting orientation
                                                                                 orient = infer_orient(
          --> 156
                         157
                                                                                              x, y, orient, require_numeric=self.require_numeric
                         158
                        /opt/conda/lib/python3.8/site-packages/seaborn/_core.py in_
→infer_orient(x, y, orient, require_numeric)
                                                    elif require_numeric and "numeric" not in (x_type, y_type):
                     1343
                     1344
                                                                  err = "Neither the `x` nor `y` variable appears to be_{\sqcup}
→numeric."
```

```
-> 1345 raise TypeError(err)
1346
1347 else:
```

TypeError: Neither the `x` nor `y` variable appears to be numeric.



5 2. Principal Coordinate Analysis

```
[11]: class_mean_list = []
for cls_label in sorted(class_to_idx.values()):
    data_raw = images_raw[labels == cls_label,:,:,:]
    class_mean = np.mean(data_raw, axis=0).reshape(1,-1)
    class_mean_list.append(class_mean)
class_means = np.concatenate(class_mean_list, axis=0)
```

6 Task 2

Write a function mean_image_squared_distances that takes the matrix class_means as an input and return the SquaredDistances matrix as output.

class_means is a number array like a traditional data matrix; it has a shape of (N, d) where there are N individual data-points where each is stored in a single d dimensional row. (N, d) could be anything, so do not make assumptions about it.

Your job is to produce the numby array SquaredDistances whose i^{th} row and j^{th} column is the squared Euclidean distance between the i^{th} row of class_means and j^{th} row of class_means. Obviously * The diagonal elements should be zero. * The SquaredDistances should be symmetric.

```
[12]: def mean_image_squared_distances(class_means):
    # your code here
    M = class_means.shape[0]

A_dots = (class_means*class_means).sum(axis=1).reshape((M,1))*np.
    ones(shape=(1,M))
    B_dots = (class_means*class_means).sum(axis=1)*np.ones(shape=(M,1))
    D_squared = A_dots + B_dots -2*class_means.dot(class_means.T)

return D_squared
```

```
[13]: #Performing sanity checks
     some_{data} = ((np.arange(35).reshape(5,7) ** 13) % 20) / 7.
     some_dist = mean_image_squared_distances(some_data)
     assert np.array_equal(some_dist.round(3), np.array([[ 0. , 4.551, 18.204, 8.
      306, 14.041,
                                                         [4.551, 0., 12.714,
      \rightarrow918, 12.551],
                                                         [18.204, 12.714, 0.
      633, 8.735],
                                                         [8.306, 3.918, 8.633, 0.
      → 7.49],
                                                         [14.041, 12.551, 8.735, 7.
      49 , 0. ]]))
      # Checking against the pre-computed test database
     test_results = test_case_checker(mean_image_squared_distances, task_id=2)
     assert test_results['passed'], test_results['message']
```

[14]: # Task 2 Test Cell

7 Task 3

Read and implement the Principal Coordinate Analysis procedure from your textbook by writing the function PCoA which takes the following arguments: 1. SquaredDistances: A numpy array which is square in shape, symmetric, and is the square of a distance matrix of some unknown set of points. The output of the mean_image_squared_distances function you wrote previously will be fed as this argument.

2. \mathbf{r} : This is the dimension of the visualization space, and corresponds to the same r variable in the textbook procedure.

Things to keep in mind: 1. There is an erratum in the textbook's description of the PCoA procedure. There is a missing negative sign when computing the matrix W; the correct definition of W is $W := -\frac{1}{2}\mathcal{A}\mathcal{D}^{(2)}\mathcal{A}^T$. 2. It is **vital** to make sure that eigenvalues are sorted as the textbook mentioned, and the eigenvectors are also ordered accordingly. Some decomposition functions such as numpy's np.linalg.eig do not guarantee to return the eigenvalues and eigenvectors in any sorted way, and np.linalg.eigh guarantees to return them in ascending order; you will have to make sure they are sorted as the textbook says.

Note: You should only use np.linalg.eigh for matrix factorization in this question since we're dealing with a symmetric matrix; do not use np.linalg.eig, np.linalg.svd, or any other matrix decomposition function in this question.

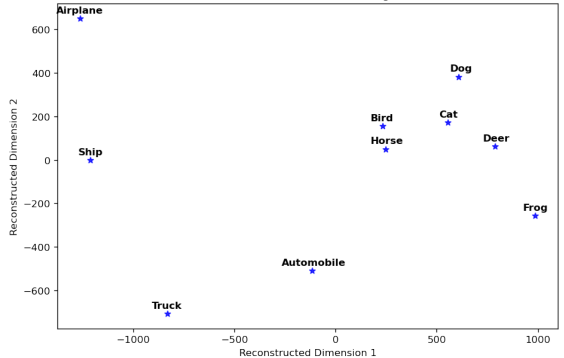
```
[29]: def PCoA(SquaredDistances, r=2):
    assert SquaredDistances.shape[0] == SquaredDistances.shape[1]
    num_points = SquaredDistances.shape[0]
    A = np.identity(num_points) - (np.ones([num_points, num_points])/num_points)
    W = -(0.5 * np.dot(np.dot(A, SquaredDistances), A.T))
    eigval, eigvec = np.linalg.eigh(W)
    eigval_mat = np.diag(np.flip(eigval))
    eigvec_f = np.flip(eigvec, axis=1)
    eigvec_r = eigvec_f[:,:r]
    eigval_matr = np.sqrt(eigval_mat[:r,:r])
    VT = np.dot(eigvec_r,eigval_matr)

assert VT.shape[0] == num_points
    assert VT.shape[1] == r
    return VT
[30]: some_data = ((np.arange(35).reshape(5,7) ** 13) % 20) / 7.
```

VT = PCoA(SquaredDistances, r=2)

```
[33]: if perform_computation:
         class_names_list = sorted(list(class_to_idx.keys()))
         fig, ax = plt.subplots(figsize=(9,6.), dpi=120)
         x_components = VT[:,0]
         y_components = VT[:,1]
         sns.regplot(x=x_components, y=y_components, fit_reg=False, marker="*",_
      for class_idx in range(VT.shape[0]):
             num_letters = len(class_names_list[class_idx])
             ax.text(x_components[class_idx]-num_letters*15,__
      →y_components[class_idx]+25,
                     class_names_list[class_idx].capitalize(),
                     horizontalalignment='left', size='medium', color='black', u
      →weight='semibold')
         ax.set_xlabel('Reconstructed Dimension 1')
         ax.set_ylabel('Reconstructed Dimension 2')
          _ = ax.set_title('PCoA on Mean CIFAR-10 Images')
```





8 3. Generalized PCoA with Non-Metric Similarities

9 Task 4

Write a function principal_components_precise_svd that returns the principal components of a data matrix and takes the following arguments as input

- 1. data_raw: a numpy array with the shape (N, \dots) , where N is the number of samples, and there may be many excess dimensions denoted by \dots . You will have to reshape this input data_raw matrix to obtain a shape of (N, d), where d is the vectorized data's dimension. For example, data_raw could have an input shape of (6000, 50, 50, 3). In this case you will have to reshape the input data to have a shape of (6000, 7500).
- 2. $num_components$: This is the number of PCA components that we want to retain. This variable is denoted by r in the PCA definition in the textbook.

and returns the variable V_x which is a numpy array with the shape (d, num_components). The columns are the unitary principal components sorted descendingly with respect to the eigenvalues.

Important Note: Do not try to recover the covariance matrix Σ and then find its eigenvalues. This can prove to be both inefficient and unnecessary. As the theoretical review before the first task concluded, **There is no need to compute the covariance matrix** Σ . Instead, all you need to do is to find the SVD of the data matrix, and extract the principal components from it.

Important Note: Do not use any matrix factorization function other than np.linalg.svd for this task; incorporating any other matrix factorization function (such as np.linalg.eig) may not be compatible with the results we expect and may even be inefficient.

```
[38]: def principal components precise svd(data raw, num components=20):
          # your code here
          shapetuple = data_raw.shape
          nn = len(shapetuple)
          r=1
          for i in range(1,nn):
              r = r * shapetuple[i]
          data_raw.reshape(shapetuple[0],r)
          new_data_raw = data_raw - np.mean(data_raw,axis=0)
          U,sigma,vh = np.linalg.svd(new_data_raw, full_matrices=True)
          vh = vh.T
          V_x = vh[:,:num_components]
          # Don't mind/change the following lines.
          # This is for mitigating the ambiguity up to -/+1 factor in PCs.
          # (i.e., if x is a unitary PC, then -x is also a unitary PC).
          # We multiply each column by the sign of the largest element (in absolute _{f L}
       →value) of that column
          sign_unambiguity = np.sign(V_x[np.abs(V_x).argmax(axis=0), np.arange(V_x.
       \rightarrowshape[1])]).reshape(1,-1)
          V_x *= sign_unambiguity
```

```
return V_x
[39]: some_data = (np.arange(35).reshape(5,7) ** 13) % 20
      some_pcs = principal_components_precise_svd(some_data, num_components=2)
      assert np.array_equal(some_pcs.round(3), np.array([[-0.123, -0.114],
                                                          [-0.43, 0.119],
                                                          [-0.021, 0.41],
                                                          [-0.603, -0.164],
                                                          [ 0.084, 0.491],
                                                          [-0.223, 0.724],
                                                          [ 0.616, 0.109]]))
      # Checking against the pre-computed test database
      test_results = test_case_checker(principal_components_precise_svd, task_id=4)
      assert test_results['passed'], test_results['message']
[40]: #Task 4 Test Cell
     The following cell will run your principal_components_precise_svd function on a single class of
     data, and provide you with some running time estimate.
[41]: if perform_computation:
          first_class_features = images_raw[labels == 0, :, :, :]
          starting time = time.time()
          first_class_pcs = principal_components_precise_svd(first_class_features,_
       →num_components=20)
          end_time = time.time()
          print('Finding the principal components on a single class took %.3f seconds.
       →'%(end time-starting time))
             IndexError
                                                        Traceback (most recent call⊔
      →last)
             <ipython-input-41-11e8b4af82da> in <module>
               4
                     starting_time = time.time()
```

→principal_components_precise_svd(first_class_features, num_components=20)

first class pcs = ___

7

end_time = time.time()

IndexError: shape mismatch: indexing arrays could not be broadcast \rightarrow together with shapes (3,32,6000) (3,)

Although, this performance is extremely hardware-dependent, it's certainly not negligible. Keep in mind that we will have to run this function about 100 times on data of the same size to construct a similarity matrix in later tasks; any speedup may very well be appreciated.

Most of the computation time in the previous task was spent on the SVD factorization. Essentially, we found all the singular values and directions, ignored most of them, and only kept the top 20. This can be a very good place to start saving on computation; if only there was an SVD variant which you could tell in advance that you're only interested in the top 20 components, so that it wouldn't waste your time computing non-important singular values and directions...

Spoiler Alert: Such an efficient SVD variant exists, and sometimes is referred to as the "Truncated SVD" in application. Next task will be a redo of the previous task using this fast factroization.

10 Task 5

Similar to principal_components_precise_svd, write a function principal_components that uses scikit-learn's TruncatedSVD decomposition instead of the precise np.linalg.svd decomposition that was used in the previous task. As in the previous task, principal_components should return the principal components of a data matrix and take the following arguments as input

- 1. data_raw: a numpy array with the shape (N, \dots) , where N is the number of samples, and there may be many excess dimensions denoted by \dots . You will have to reshape this input data_raw matrix to obtain a shape of (N,d), where d is the vectorized data's dimension. For example, data_raw could have an input shape of (6000, 50, 50, 3). In this case you will have to reshape the input data to have a shape of (6000, 7500).
- 2. $num_components$: This is the number of PCA components that we want to retain. This variable is denoted by r in the PCA definition in the textbook.

principal_components should return the variable V_x which is a numpy array with the shape (d, num_components). The columns are the unitary principal components sorted descendingly with respect to the eigenvalues.

Important Note: You should only use scikit-learn's TruncatedSVD decomposition for this task. You can read about this function at https://scikit-

learn.org/stable/modules/generated/sklearn.decomposition.TruncatedSVD.html.

- You must use the randomized algorithm implementation as it is more efficient.
- Since this heuristic is stochastic, you must provide random_state=12345 as an input argument to this object's constructor for reproducibility.
- Use exactly 5 iterations for this heuristic (i.e., specify n_iter to be exactly 5).

Important Note: Do not try to recover the covariance matrix Σ and then find its eigenvalues. This can prove to be both inefficient and unnecessary. As the theoretical review before the first task concluded, **There is no need to compute the covariance matrix** Σ . Instead, all you need to do is to find the SVD of the data matrix, and extract the principal components from it.

```
[48]: def principal_components(data_raw, num_components=20):
          shapetuple = data_raw.shape
          nn = len(shapetuple)
          for i in range(1,nn):
              r = r * shapetuple[i]
          data_raw.reshape(shapetuple[0],r)
          new_data_raw = data_raw - np.mean(data_raw,axis=0)
          svd = TruncatedSVD(n_components = num_components, algorithm='randomized',__
       →n_iter=5, random_state=12345 )
          transformed = svd.fit transform(new data raw)
          V_x = svd.components_.T
          assert V_x.ndim==2
          # Don't mind/change the following lines.
          # This is for mitigating the ambiguity up to -/+1 factor in PCs.
          # (i.e., if x is a unitary PC, then -x is also a unitary PC).
          # We multiply each column by the sign of the largest element (in absolute_
       →value) of that column
          sign_unambiguity = np.sign(V_x[np.abs(V_x).argmax(axis=0), np.arange(V_x.
       \hookrightarrowshape[1])]).reshape(1,-1)
          V_x *= sign_unambiguity
          return V_x
```

```
# Checking against the pre-computed test database
      test_results = test_case_checker(principal_components, task_id=5)
      assert test_results['passed'], test_results['message']
[50]: #Task 5 Test Cell
[51]: if perform_computation:
          first_class_features = images_raw[labels == 0, :, :, :]
          starting_time = time.time()
          first_class_pcs = principal_components(first_class_features,_
       →num_components=20)
          end_time = time.time()
          print('Finding the principal components on a single class took %.3f seconds.

        '%(end_time-starting_time))

             ValueError
                                                        Traceback (most recent call_
      →last)
             <ipython-input-51-b137cd82a5bb> in <module>
               4
                     starting_time = time.time()
         ---> 5
                     first_class_pcs = principal_components(first_class_features,_
      →num_components=20)
                     end_time = time.time()
               7
             <ipython-input-48-74a798387a83> in principal_components(data_raw,_
      →num_components)
               9
                     new_data_raw = data_raw - np.mean(data_raw,axis=0)
                     svd = TruncatedSVD(n_components = num_components,__
      →algorithm='randomized', n_iter=5, random_state=12345 )
         ---> 11
                     transformed = svd.fit_transform(new_data_raw)
                     V_x = svd.components_.T
              12
              13
             /opt/conda/lib/python3.8/site-packages/sklearn/decomposition/
      →_truncated_svd.py in fit_transform(self, X, y)
             162
                             Reduced version of X. This will always be a dense array.
```

```
11 11 11
       163
   --> 164
                   X = self._validate_data(X, accept_sparse=['csr', 'csc'],
                                            ensure_min_features=2)
       165
       166
                   random_state = check_random_state(self.random_state)
       /opt/conda/lib/python3.8/site-packages/sklearn/base.py in__
→_validate_data(self, X, y, reset, validate_separately, **check_params)
       419
                       out = X
       420
                   elif isinstance(y, str) and y == 'no_validation':
                       X = check_array(X, **check_params)
   --> 421
       422
                       out = X
       423
                   else:
       /opt/conda/lib/python3.8/site-packages/sklearn/utils/validation.py in u
→inner_f(*args, **kwargs)
        61
                       extra_args = len(args) - len(all_args)
                       if extra_args <= 0:</pre>
        62
   ---> 63
                            return f(*args, **kwargs)
        64
        65
                       # extra_args > 0
       opt/conda/lib/python3.8/site-packages/sklearn/utils/validation.py in ∪
→check_array(array, accept_sparse, accept_large_sparse, dtype, order, copy, u

→force_all_finite, ensure_2d, allow_nd, ensure_min_samples,

□
→ensure_min_features, estimator)
       657
                                "into decimal numbers with dtype='numeric'")
⊶from e
                   if not allow_nd and array.ndim >= 3:
       658
                       raise ValueError("Found array with dim %d. %s expected_
   --> 659
<= 2."</p>
       660
                                         % (array.ndim, estimator_name))
       661
```

ValueError: Found array with dim 4. Estimator expected <= 2.

Using this principal_components function, and the images_raw array, you could reconstruct an arbitrary image using a small number of components, see the effect of the number of components on the reconstructed image's quality, and share your results on Piazza!

11 Task 6

Write the function $E_A_given_B$ that computes the E[A|B] and takes the two matrices $class_A_data$ and $class_B_data$.

- 1. class_A_data is a number arrays with the shape (N, \dots) , where N is the number of samples, and there may be many excess dimensions denoted by \dots . You will have to reshape this input matrix to obtain a shape of (N, d), where d is the vectorized data's dimension.
- 2. class_B_data has the same data structure as class_A_data.

To compute E[A|B]: 1. First, do whatever reshaping you have to do. 2. Subtract Class A's mean from its data 3. Use the principal_components function you wrote before to extract the 20 principal components of class_B_data. 4. Project Class A's data onto the mentioned principal components and get back to the original space. 5. Compute Class A's residuals (i.e., the difference between the original and the projection). 5. Find the squared residual sizes for each sample, and then return their mean as the E_A_cond_B scalar. In other words, square class A's residuals, sum them over each sample (which should reduce the squared residual matrix to only N elements), and then report the mean of them as E_A_cond_B.

```
[55]: def E_A_given_B(class_A_data, class_B_data):
          shapetuple = class_A_data.shape
          nn = len(shapetuple)
          r=1
          for i in range(1,nn):
              r = r * shapetuple[i]
          class_A_data.reshape(shapetuple[0],r)
          shapetupleII = class_B_data.shape
          mm = len(shapetupleII)
          f=1
          for i in range(1,mm):
              f = f * shapetupleII[i]
          class_B_data.reshape(shapetupleII[0],f)
          new_cAd = class_A_data - np.mean(class_A_data,axis=0)
          VxB = principal_components(class_B_data,num_components=20)
          class_A_projection = np.dot( VxB, np.dot(VxB.T, new_cAd.T)).T
          E_A_cond_B = np.mean(np.sum(np.square(class_A_projection - new_cAd),_
       →axis=1))
          return E_A_cond_B
```

```
[56]: some_data = ((np.arange(35).reshape(5,7) ** 13) % 20) / 7.
some_data = np.repeat(some_data, 8, axis=1)
some_E = E_A_given_B(some_data, (some_data**1.02))
assert some_E.round(3)==0.001

# Checking against the pre-computed test database
test_results = test_case_checker(E_A_given_B, task_id=6)
assert test_results['passed'], test_results['message']
```

```
[57]: #Task 6 Test Cell
[58]: if perform computation:
          num_classes = class_means.shape[0]
          SimilarityMatrix = np.zeros((num classes, num classes))
          for row in range(num_classes):
              print(f'Row {row}', end='')
              row_st_time = time.time()
              for col in range(row+1):
                  class_A_data = images_raw[labels == row, :, :, :]
                  class_B_data = images_raw[labels == col, :, :, :]
                  E_A_cond_B = E_A_given_B(class_A_data, class_B_data)
                  E_B_cond_A = E_A_given_B(class_B_data, class_A_data)
                  SimilarityMatrix[col, row] = (E_A_cond_B + E_B_cond_A)/2.
                  SimilarityMatrix[row, col] = (E_A_cond_B + E_B_cond_A)/2.
              print(f' (This row took %.3f seconds to finish)'%(time.time() -
       →row_st_time))
     Row 0
             ValueError
                                                        Traceback (most recent call_
      →last)
             <ipython-input-58-c9e323725cb3> in <module>
                             class_A_data = images_raw[labels == row, :, :, :]
               9
                             class_B_data = images_raw[labels == col, :, :, :]
         ---> 10
                             E_A_cond_B = E_A_given_B(class_A_data, class_B_data)
                             E_B_cond_A = E_A_given_B(class_B_data, class_A_data)
              11
              12
                             SimilarityMatrix[col, row] = (E_A_cond_B + E_B_cond_A)/2.
             <ipython-input-55-ed65683ec264> in E_A_given_B(class_A_data,__
      →class_B_data)
                     class_B_data.reshape(shapetupleII[0],f)
              13
              14
                     new_cAd = class_A_data - np.mean(class_A_data,axis=0)
                     VxB = principal components(class B data,num components=20)
         ---> 15
                     class_A_projection = np.dot( VxB, np.dot(VxB.T, new_cAd.T)).T
              16
              17
                     E A cond B = np.mean(np.sum(np.square(class A projection -
      →new_cAd), axis=1))
             <ipython-input-48-74a798387a83> in principal_components(data_raw,_
      →num_components)
```

```
new_data_raw = data_raw - np.mean(data_raw,axis=0)
        10
               svd = TruncatedSVD(n_components = num_components,__
→algorithm='randomized', n_iter=5, random_state=12345 )
   ---> 11
               transformed = svd.fit_transform(new_data_raw)
        12
               V x = svd.components .T
        13
       /opt/conda/lib/python3.8/site-packages/sklearn/decomposition/
→_truncated_svd.py in fit_transform(self, X, y)
       162
                       Reduced version of X. This will always be a dense array.
       163
   --> 164
                   X = self._validate_data(X, accept_sparse=['csr', 'csc'],
       165
                                            ensure_min_features=2)
       166
                   random state = check random state(self.random state)
       /opt/conda/lib/python3.8/site-packages/sklearn/base.py in⊔
→ validate_data(self, X, y, reset, validate_separately, **check_params)
       419
                       out = X
       420
                   elif isinstance(y, str) and y == 'no \ validation':
  --> 421
                       X = check_array(X, **check_params)
       422
                       out = X
       423
                   else:
       opt/conda/lib/python3.8/site-packages/sklearn/utils/validation.py in in incomplete.
→inner_f(*args, **kwargs)
        61
                       extra_args = len(args) - len(all_args)
                       if extra_args <= 0:</pre>
        62
   ---> 63
                           return f(*args, **kwargs)
        64
        65
                       # extra_args > 0
       /opt/conda/lib/python3.8/site-packages/sklearn/utils/validation.py in u
→check_array(array, accept_sparse, accept_large_sparse, dtype, order, copy,

→force_all_finite, ensure_2d, allow_nd, ensure_min_samples,

→ensure_min_features, estimator)
       657
                                "into decimal numbers with dtype='numeric'")
→from e
                   if not allow_nd and array.ndim >= 3:
       658
   --> 659
                       raise ValueError("Found array with dim %d. %s expected ⊔
→<= 2."
                                         % (array.ndim, estimator name))
       660
       661
```

ValueError: Found array with dim 4. Estimator expected <= 2.

If you apply any general SimilarityMatrix variable to the previously defined PCoA function, you may get NaN entries due to the fact that they may not generally be a metric distance matrix (i.e., having non-zero diagonal elements and the triangle inequality not alway holding).

This issue can be best seen when having a similarity measure that is extremely uneven (i.e., when the small entries are extremely small and the large entries are extremely large). This will make it difficult for the triangle inequality to hold. It is a good idea to ammend the PCoA in a way that can deal with such non-metric similarity measures.

```
[59]: VT = None
    if perform_computation:
        VT = PCoA(SimilarityMatrix**40, r=10)
VT

[59]: array([[0., 0., 0., 0., 0., 0., 0., 0., 0., 0.],
        [0., 0., 0., 0., 0., 0., 0., 0., 0.],
        [0., 0., 0., 0., 0., 0., 0., 0., 0.],
        [0., 0., 0., 0., 0., 0., 0., 0., 0.],
        [0., 0., 0., 0., 0., 0., 0., 0., 0.],
        [0., 0., 0., 0., 0., 0., 0., 0., 0.],
        [0., 0., 0., 0., 0., 0., 0., 0., 0.],
        [0., 0., 0., 0., 0., 0., 0., 0.],
        [0., 0., 0., 0., 0., 0., 0., 0.],
        [0., 0., 0., 0., 0., 0., 0., 0.],
        [0., 0., 0., 0., 0., 0., 0., 0.]]
```

12 Task 7

Write the function Lingoes_PreProcessing that does some pre-processing to the SimilarityMatrix to make it have the Euclidean property and the triangles to close.

Here is a very brief and to the point description from the r documentation page (https://www.rdocumentation.org/packages/ape/versions/5.2/topics/pcoa).

"In the Lingoes (1971) procedure, a constant c1, equal to twice absolute value of the largest negative eigenvalue of the original principal coordinate analysis, is added to each original squared distance in the distance matrix, except the diagonal values. A new principal coordinate analysis, performed on the modified distances, has at most (n-2) positive eigenvalues, at least 2 null eigenvalues, and no negative eigenvalue."

If you're interested, you can read more about correction for negative eigenvalues in http://biol09.biol.umontreal.ca/PLcourses/Ordination_sections_1.3+1.4_PCoA_Eng.pdf.

The function Lingoes_PreProcessing takes the numpy array SimilarityMatrix as input, and returns ProcessedSimilarityMatrix based on the following condition: 1. If all eigenvalues computed during PCoA are non-negative, then ProcessedSimilarityMatrix should be the same as

the SimilarityMatrix. 2. Otherwise, follow the instructions to perform the Lingoes correction on the SimilarityMatrix and return ProcessedSimilarityMatrix.

In other words, this is what you're supposed to do: 1. Perform the PCoA analysis on SimilarityMatrix right up to the point where you find the eigenvalues. Do not go any further. More precisely, you should only find the eigenvalues of the matrix \mathcal{W} corresponding to SimilarityMatrix in the PCoA analysis. 2. Find the minimum eigenvalue and call it λ_{\min} . 3. If $\lambda_{\min} \geq 0$, then stop and return SimilarityMatrix as it was without any change. 4. If $\lambda_{\min} < 0$, then add $2|\lambda_{\min}|$ to all the non-diagonal elements of SimilarityMatrix and return the resulting matrix.

Important Note: Do not call the PCoA function on SimilarityMatrix. You should not call the whole PCoA function on SimilarityMatrix, as you do not care about the output reconstructions of PCoA. Instead, you need the eigenvalues for further processing, which are not returned by the PCoA function.

Note: You do not need a for loop for adding a scalar to the non-diagonal elements of a matrix; you can add the scalar to all the elements of the matrix, and then subtract it from the same scalar multiple of the identity matrix (i.e., using a function like np.eye for instance).

```
[62]: def Lingoes_PreProcessing(SimilarityMatrix):
          assert SimilarityMatrix.shape[0] == SimilarityMatrix.shape[1]
          num_points = SimilarityMatrix.shape[0]
          A = np.identity(num_points) - (np.ones([num_points, num_points])/num_points)
          W = -(0.5 * np.dot(np.dot(A, SimilarityMatrix), A.T))
          eigval, eigvec = np.linalg.eigh(W)
          eigval_min = np.amin(eigval)
          if (eigval_min >=0):
              ProcessedSimilarityMatrix = SimilarityMatrix
          elif (eigval_min<0):</pre>
              ProcessedSimilarityMatrix = SimilarityMatrix.copy()
              icr = 2* (abs(eigval_min))
              for i in range(0,num_points):
                  for j in range(0,num_points):
                      if (i!=j):
                          ProcessedSimilarityMatrix[i][j]+=icr
          return ProcessedSimilarityMatrix
```

```
[2896177.9, 1229280.9,
         0., 944977.4, 947878.7],
                                                            [ 936570.7, 897958.5, _
      944977.4, 0., 920604.3],
                                                            [1442744.7, 1208489.7, ]
      947878.7, 920604.3, 0. 11))
      # Checking against the pre-computed test database
     test_results = test_case_checker(Lingoes_PreProcessing, task_id=7)
     assert test results['passed'], test results['message']
[64]: # Task 7 Test Cell
[65]: def PCoA lingoes(SimilarityMatrix, r=2):
         ProcessedSimilarityMatrix = Lingoes_PreProcessing(SimilarityMatrix)
         return PCoA(ProcessedSimilarityMatrix, r=r)
[66]: VT = None
     if perform_computation:
         VT = PCoA lingoes(SimilarityMatrix, r=2)
     VT
[66]: array([[0., 0.],
            [0., 0.],
            [0., 0.],
            [0., 0.],
            [0., 0.],
            [0., 0.],
            [0., 0.],
            [0., 0.],
            [0., 0.],
            [0., 0.]])
[67]: if perform_computation:
         class_names_list = sorted(list(class_to_idx.keys()))
         fig, ax = plt.subplots(figsize=(9,6.), dpi=120)
         x_components = VT[:,0]
         y_components = VT[:,1]
         sns.regplot(x=x_components, y=y_components, fit_reg=False, marker="*",_
      for class_idx in range(VT.shape[0]):
             num letters = len(class names list[class idx])
             ax.text(x_components[class_idx]-num_letters*8,_
      →y_components[class_idx]+10,
                     class_names_list[class_idx].capitalize(),
                     horizontalalignment='left', size='medium', color='black', L
       →weight='semibold')
```

```
_ = ax.set_title('Generalized PCoA on CIFAR-10 Images')
                  ValueError
                                                                                                                                   Traceback (most recent call_
→last)
                   /opt/conda/lib/python3.8/site-packages/IPython/core/formatters.py in_
→__call__(self, obj)
                   339
                                                                        pass
                   340
                                                             else:
        --> 341
                                                                        return printer(obj)
                   342
                                                             # Finally look for special method names
                   343
                                                             method = get_real_method(obj, self.print_method)
                   /opt/conda/lib/python3.8/site-packages/IPython/core/pylabtools.py in_
→<lambda>(fig)
                   246
                   247
                                        if 'png' in formats:
                                                   png_formatter.for_type(Figure, lambda fig: print_figure(fig,_
        --> 248
→'png', **kwargs))
                   249
                                        if 'retina' in formats or 'png2x' in formats:
                                                   png_formatter.for_type(Figure, lambda fig:_u
                   250
→retina_figure(fig, **kwargs))
                   /opt/conda/lib/python3.8/site-packages/IPython/core/pylabtools.py in_
→print_figure(fig, fmt, bbox_inches, **kwargs)
                   130
                                                  FigureCanvasBase(fig)
                   131
                                        fig.canvas.print_figure(bytes_io, **kw)
        --> 132
                                       data = bytes_io.getvalue()
                   133
                   134
                                        if fmt == 'svg':
                   opt/conda/lib/python3.8/site-packages/matplotlib/backend_bases.py in in in incomplete in incomplete
→print figure(self, filename, dpi, facecolor, edgecolor, orientation, format,
→bbox_inches, pad_inches, bbox_extra_artists, backend, **kwargs)
                2208
                2209
                                                             try:
        -> 2210
                                                                        result = print_method(
                2211
                                                                                   filename,
```

ax.set_xlabel('Reconstructed Dimension 1')
ax.set_ylabel('Reconstructed Dimension 2')

2212 dpi=dpi,

```
/opt/conda/lib/python3.8/site-packages/matplotlib/backend_bases.py in u
→wrapper(*args, **kwargs)
      1637
                       kwargs.pop(arg)
      1638
  -> 1639
                   return func(*args, **kwargs)
      1640
      1641
               return wrapper
       /opt/conda/lib/python3.8/site-packages/matplotlib/backends/backend_agg.
→py in print_png(self, filename_or_obj, metadata, pil_kwargs, *args)
       507
                       *metadata*, including the default 'Software' key.
       508
   --> 509
                   FigureCanvasAgg.draw(self)
       510
                   mpl.image.imsave(
       511
                       filename_or_obj, self.buffer_rgba(), format="png",_
→origin="upper",
       /opt/conda/lib/python3.8/site-packages/matplotlib/backends/backend agg.
→py in draw(self)
               def draw(self):
       400
       401
                   # docstring inherited
                   self.renderer = self.get_renderer(cleared=True)
   --> 402
       403
                   # Acquire a lock on the shared font cache.
       404
                   with RendererAgg.lock, \
       /opt/conda/lib/python3.8/site-packages/matplotlib/backends/backend_agg.
→py in get_renderer(self, cleared)
                                     and getattr(self, "_lastKey", None) == key)
       416
                   if not reuse_renderer:
       417
   --> 418
                       self.renderer = RendererAgg(w, h, self.figure.dpi)
                       self._lastKey = key
       419
       420
                   elif cleared:
       /opt/conda/lib/python3.8/site-packages/matplotlib/backends/backend_agg.
→py in __init__(self, width, height, dpi)
                   self.width = width
        94
        95
                   self.height = height
   ---> 96
                   self._renderer = _RendererAgg(int(width), int(height), dpi)
                   self._filter_renderers = []
        97
        98
```

<Figure size 1080x720 with 1 Axes>

[]: