HW4

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\mathbf{a}

True. This is because the rank of the matrix (k = rank (X)) represents the maximum number of linearly independent vectors in the matrix. PCA finds the best linear subspace of a given dimensionality (in this case, k) that captures the most variance in the data. If the dimensionality of the subspace is equal to the rank of the data matrix, it means that the subspace can capture all the variations in the data, leading to no information loss in the projection.

h

False. $X = USV^T$ then $X^TX = VS^TU^TUSV^T = VS^TSV^T$, due to X^TX being symmetric, it guarantees it has eigenvalue and eigenvectors such that $X^TXv = \lambda v$, therefore $VS^TSV^Tv = \lambda v$, if we let v = V, then $VS^TS = \lambda V$. Since S^TS is diagonal, it shows that λV is equivalent as scaling each column of V with S^TS . Therefore it is the column of V that are equal to the eigenvectors of X^TX not rows

\mathbf{c}

False. Minimizing the k-means objective function by increasing k can lead to overfitting, where clusters may become arbitrarily small and not necessarily meaningful. It is better to use methods like the elbow method or silhouette score to determine an appropriate k that balances the granularity of clustering with the overall cluster quality.

\mathbf{d}

False, if there are repeated value s in S, then the row and column in U and V associated with s can have multiple valid order, this shows that USV decomposition are not unique.

\mathbf{e}

False, the rank of a square matrix is determined by the number of linearly independent rows or columns. However, a linearly independent matrix may have repeated eigenvalues and results in number of unique eigenvalues less than its rank,

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Data Pre-processing Steps:

- Data Cleaning: First, Handle missing values, duplicates, and outliers in the dataset.
- Encoding Categorical Data: Then, Convert categorical variables into numerical values using one-hot encoding or label encoding.
- Normalization/Standardization: Scale numerical data to ensure fair contribution to the model.
- Missing Data Interpolation: For missing entries, use methods like k-nearest neighbors, mean/mode imputation, or model-based imputation where appropriate.
- Balancing Classes: Once dataset is complete, check if the dataset is imbalanced, apply oversampling, undersampling, or bootstraping.

Machine Learning Pipeline Steps:

- Train Test Validation Split: split train, test, validation with 70%, 20%, 10%.
- Feature Selection: Employ L1 regularization (Lasso) to help in feature selection and to create a sparse model emphasizing important risk factors.
- Model Selection: Start with logistic regression for binary natured classification (disease/no disease). If non-linear relationships are suspected, consider kernel for quicker compute, and deep learning techniques.
- Validation Technique: Use k-fold cross-validation to assess the generalizability of the model.
- Hyperparameter Tuning: Use grid search or random search with cross-validation to find optimal hyperparameters.

Acknowledging Constraints and Measuring Results:

- Accuracy over Efficiency: Focus on a model that provides more accurate result as they are preciseness of infomation creates huge matter to the user.
- Privacy Considerations: Ensure that the model does not require or store more personal data than necessary, respecting user privacy.

b

Shortcomings of Training Process:

Limited representation in the training dataset can lead to skewed accuracy across different demographic groups. Features may inadvertently emphasize specific characteristics of the majority population in the dataset. Moreover, Over-representation of certain classes (e.g., more non-disease cases) might result in biased predictions.

Addressing the Shortcomings:

1. Include data from varied demographics to ensure representativeness. 2. plot the data base one the feature and remove data that are overly populated in the same cluster, make sure the training data is equally balanced before training 3. Apply regularization and stratified k-fold cross-validation for robust performance evaluation.

 \mathbf{c}

Ignoring shortcomings in crime datasets can lead to biased predictive policing, disproportionately affecting minority communities. This can misguide resource allocation, favoring increased law enforcement over necessary social services in areas needing them. Such practices erode public trust in law enforcement, potentially reducing crime reporting and exacerbating data inaccuracies. Relying on skewed data raises legal and ethical issues due to the unfair targeting of specific demographics and perpetuates stereotypes, negatively impacting the socio-economic fabric of communities.

a b

Center 1

Center 2

Center 3

Center 4

Center 5

3

```
Center 6
                       Center 7
                                        Center 8
                                                         Center 9
from typing import List, Tuple
import numpy as np
import itertools
from utils import problem
Oproblem.tag("hw4-A")
def calculate_centers(
    data: np.ndarray, classifications: np.ndarray, num_centers: int
) -> np.ndarray:
    11 11 11
    Sub-routine of Lloyd's algorithm that calculates the centers given datapoints and their respective
    num_centers is additionally provided for speed-up purposes.
    Args:
        data (np.ndarray): Array of shape (n, d). Training data set.
        classifications (np.ndarray): Array of shape (n,) full of integers in range {0, 1, ..., num_
            Data point at index i is assigned to classifications[i].
        num_centers (int): Number of centers for reference.
            Might be usefull for pre-allocating numpy array (Faster that appending to list).
    Returns:
        np.ndarray: Array of shape (num_centers, d) containing new centers.
    d = data.shape[1]
    cluster_sums = np.zeros((num_centers, d))
    cluster_counts = np.zeros(num_centers)
```

for i in range(len(data)):

cluster_sums[classifications[i]] += data[i]
cluster_counts[classifications[i]] += 1

Avoid division by zero for empty clusters

```
cluster_counts[cluster_counts == 0] = 1
   new_centers = cluster_sums / cluster_counts[:, None]
   return new_centers
@problem.tag("hw4-A")
def cluster_data(data: np.ndarray, centers: np.ndarray) -> np.ndarray:
    Sub-routine of Lloyd's algorithm that clusters datapoints to centers given datapoints and centers
    Args:
        data (np.ndarray): Array of shape (n, d). Training data set.
        centers (np.ndarray): Array of shape (k, d). Each row is a center to which a datapoint can be
        np.ndarray: Array of integers of shape (n,), with each entry being in range \{0, 1, 2, \ldots, k\}
            Entry j at index i should mean that j^th center is the closest to data[i] datapoint.
    distances = np.zeros((data.shape[0], centers.shape[0]))
    for idx, center in enumerate(centers):
        distances[:, idx] = np.sqrt(np.sum((data - center) ** 2, axis=1))
    return np.argmin(distances, axis=1)
def calculate_error(data: np.ndarray, centers: np.ndarray) -> float:
    """ This method has been implemented for you.
    Calculates error/objective function on a provided dataset, with trained centers.
    Args:
        data (np.ndarray): Array of shape (n, d). Dataset to evaluate centers on.
        centers (np.ndarray): Array of shape (k, d). Each row is a center to which a datapoint can be
            These should be trained on training dataset.
    Returns:
        float: Single value representing mean objective function of centers on a provided dataset.
    distances = np.zeros((data.shape[0], centers.shape[0]))
    for idx, center in enumerate(centers):
        distances[:, idx] = np.sqrt(np.sum((data - center) ** 2, axis=1))
    return np.mean(np.min(distances, axis=1))
Oproblem.tag("hw4-A")
def lloyd_algorithm(
    data: np.ndarray, num_centers: int, epsilon: float = 10e-3
) -> Tuple[np.ndarray, List[float]]:
    """Main part of Lloyd's Algorithm.
    Args:
        data (np.ndarray): Array of shape (n, d). Training data set.
        num_centers (int): Number of centers to train/cluster around.
        epsilon (float, optional): Epsilon for stopping condition.
```

```
Defaults to 10e-3.
    Returns:
        np.ndarray: Tuple of 2 numpy arrays:
            Element at index 0: Array of shape (num_centers, d) containing trained centers.
            Element at index 1: List of floats of length # of iterations
                containing errors at the end of each iteration of lloyd's algorithm.
                You should use the calculate_error() function that has been implemented for you.
    Note:
        - For initializing centers please use the first `num_centers` data points.
   last_trained_center = np.ones((num_centers, data.shape[1]))
    trained_center = data[np.random.choice(np.arange(data.shape[0]), num_centers, False)]
    errors = []
    while (np.max(abs(trained_center - last_trained_center)) > epsilon):
        clustered_data = cluster_data(data, trained_center)
        last_trained_center = trained_center
        trained_center = calculate_centers(data=data, classifications=clustered_data, num_centers=num
        errors.append(calculate_error(data=data, centers=trained_center))
   return (trained_center, errors)
if __name__ == "__main__":
   from k_means import lloyd_algorithm # type: ignore
else:
   from .k_means import lloyd_algorithm
import matplotlib.pyplot as plt
import numpy as np
from utils import load_dataset, problem
Oproblem.tag("hw4-A", start_line=1)
def main():
    """Main function of k-means problem
   Run Lloyd's Algorithm for k=10, and report 10 centers returned.
    NOTE: This code might take a while to run. For debugging purposes you might want to change:
        x_train to x_train[:10000]. CHANGE IT BACK before submission.
    (x_train, _), _ = load_dataset("mnist")
    centers = lloyd_algorithm(x_train, num_centers=10, epsilon=10e-3)
    # Reshape centers to 28x28 (assuming MNIST images are 28x28)
    centers_images = centers[0].reshape(-1, 28, 28)
    # Plotting the centers
    fig, axes = plt.subplots(2, 5, figsize=(10, 4))
    for i, ax in enumerate(axes.flatten()):
        ax.imshow(centers_images[i], cmap='gray')
        ax.axis('off')
        ax.set_title(f'Center {i+1}')
```

Training should stop when max(abs(centers - previous_centers)) is smaller or equal to eps

```
plt.tight_layout()
  plt.show()

if __name__ == "__main__":
  main()
```

4

 \mathbf{a}

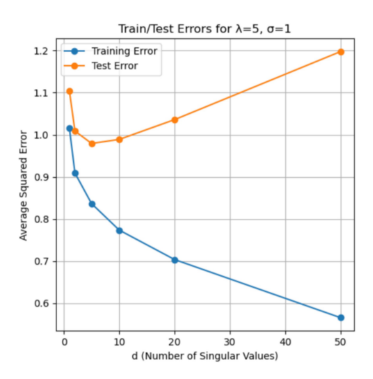
$$\hat{R} = rac{\sum_{alli}(R_{ij}|R_{ij_1} = R_{ij_2})}{num(R_{ij}|R_{ij_1} = R_{ij_2})} forj = 0, 1, 2, \dots, movies - 1$$

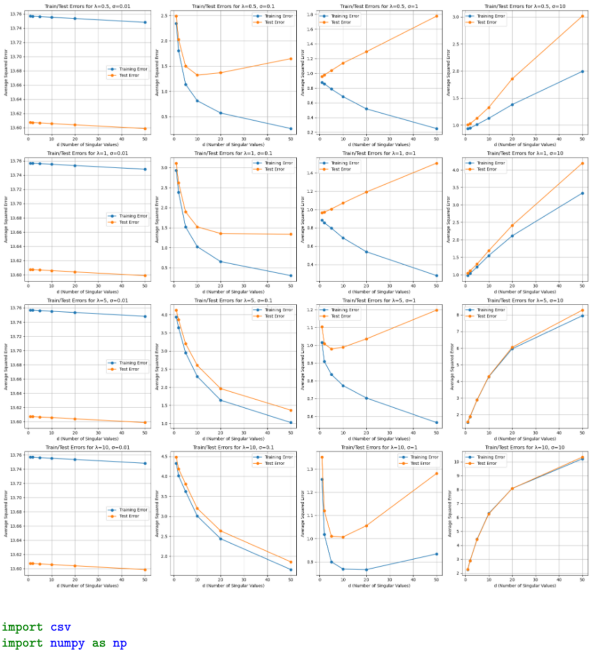
Test Error (MSE) for the estimator $\hat{R}{:}\ 1.063564200567445$

b



 \mathbf{c}





```
import csv
import numpy as np
from scipy.sparse.linalg import svds
import matplotlib.pyplot as plt
import torch
data = []
with open('u.data') as csvfile:
    spamreader = csv.reader(csvfile, delimiter='\t')
    for row in spamreader:
        data.append([int(row[0])-1, int(row[1])-1, int(row[2])])
data = np.array(data)

num_observations = len(data)  # num_observations = 100,000
num_users = max(data[:,0])+1  # num_users = 943, indexed 0,...,942
num_items = max(data[:,1])+1  # num_items = 1682 indexed 0,...,1681

np.random.seed(1)
num_train = int(0.8*num_observations)
```

```
perm = np.random.permutation(data.shape[0])
train = data[perm[0:num_train],:]
test = data[perm[num_train::],:]
print(f"Successfully loaded 100K MovieLens dataset with",
      f"{len(train)} training samples and {len(test)} test samples")
# Your code goes here
# Compute estimate
print(np.max(train[:, 1]))
# Calculate average rating for each movie
movie_ratings_sum = np.zeros(num_items)
movie_ratings_count = np.zeros(num_items)
for user, movie, rating in train:
    movie_ratings_sum[movie] += rating
    movie_ratings_count[movie] += 1
# Avoid division by zero
movie_ratings_count[movie_ratings_count == 0] = 1
average_movie_ratings = movie_ratings_sum / movie_ratings_count
# Construct rank-one matrix R_hat
R_hat = np.tile(average_movie_ratings, (num_users, 1))
# Evaluate test error
test_error = 0
for user, movie, actual_rating in test:
    predicted_rating = R_hat[user, movie]
    test_error += (predicted_rating - actual_rating) ** 2
test_error /= len(test)
print("Test Error (MSE) for the estimator R_hat:", test_error)
# Your code goes here
# Create the matrix R twiddle (\widetilde{R}).
R_tilde = np.zeros((num_users, num_items))
for user, movie, rating in train:
    R_tilde[user, movie] = rating
  # Your code goes here
def construct_estimator(d, r_twiddle):
    U, s, Vt = svds(r_twiddle, k = d)
    R_hat_d = U @ np.diag(s) @ Vt
    return R_hat_d
def get_error(d, r_twiddle, dataset):
    R_hat_d = construct_estimator(d, r_twiddle)
    error = 0
    for user, movie, actual_rating in dataset:
        predicted_rating = R_hat_d[user, movie]
        error += (predicted_rating - actual_rating) ** 2
```

```
return error / len(dataset)
# Your code goes here
# Evaluate train and test error for: d = 1, 2, 5, 10, 20, 50.
d_{values} = [1, 2, 5, 10, 20, 50]
usv_train_errors = []
usv_test_errors = []
for d in d_values:
    usv_train_error=get_error(d, R_tilde, train)
    usv_test_error=get_error(d, R_tilde, test)
    usv_train_errors.append(usv_train_error)
    usv_test_errors.append(usv_test_error)
# Plot both train and test error as a function of d on the same plot.
plt.figure(figsize=(10, 6))
plt.plot(d_values, usv_train_errors, label='Training Error', marker='o')
plt.plot(d_values, usv_test_errors, label='Test Error', marker='o')
plt.xlabel('d (Number of Singular Values)')
plt.ylabel('Average Squared Error')
plt.title('Training and Test Errors vs. Rank of Approximation')
plt.legend()
plt.grid(True)
plt.show()
# Your code goes here
def closed_form_u(V, U, 1, R_tilde):
  print(V.shape, U.shape)
  for i in range(U.shape[0]):
    R_i_tile = R_tilde[i]
    indices = R_i_tile > 0
    V_j = V[indices, :]
    R_i = R_i_tile[indices]
    A = V_j.T @ V_j + 1 * np.eye(V_j.shape[1])
    b = (V_j.T @ R_i).reshape(-1, 1)
    U[i,:] = np.linalg.solve(A, b).flatten()
  return U
def closed_form_v(V, U, 1, R_tilde):
  for j in range(V.shape[0]):
    R_j_tile = R_tilde[:,j]
    indices = R_j_tile > 0
    U_i = U[indices, :]
    R_j = R_j_tile[indices]
    A = U_i.T @ U_i + 1 * np.eye(U_i.shape[1])
    b = (U_i.T @ R_j).reshape(-1, 1)
    V[j,:] = np.linalg.solve(A, b).flatten()
  return V
def construct_alternating_estimator(
    d, r_twiddle, l=0.0, delta=1e-2, sigma=0.1, U=None, V=None
):
  old_U, old_V = np.zeros((num_users, d)), np.zeros((num_items, d))
```

```
if U is None:
   U = np.random.rand(num_users, d) * sigma
  if V is None:
    V = np.random.rand(num_items, d) * sigma
  while (np.max(np.abs(V - old_V)) > delta and np.max(np.abs(U - old_U)) > delta):
    old_U, old_V = U, V
   U = closed_form_u(V, U, 1, r_twiddle)
   V = closed_form_v(V, U, 1, r_twiddle)
  return U, V
def calc_uv_error(dataset, U, V):
   user = dataset[:,0]
   item = dataset[:,1]
   score = dataset[:,2]
    # print(U.shape, V.shape)
   pred = np.einsum('ij,ij->i', U[user], V[item])
   mse_error = np.mean((score-pred) ** 2)
   return mse_error
from itertools import product
d_{vals} = [1, 2, 5, 10, 20, 50]
lambdas = [0.5, 1, 5, 10]
sigmas = [0.01, 0.1, 1, 10]
# Prepare the plots
fig, axes = plt.subplots(4, 4, figsize=(20, 20)) # 4x4 grid for 16 combinations
axes = axes.flatten()
# Iterate over all combinations of lambdas and sigmas
for index, (lambda_val, sigma_val) in enumerate(product(lambdas, sigmas)):
   uv_train_errors = []
   uv_test_errors = []
    for d in d_vals:
        U, V = construct_alternating_estimator(d=d, r_twiddle=R_tilde, l=lambda_val, sigma=sigma_val)
        uv_train_errors.append(calc_uv_error(train, U, V))
        uv_test_errors.append(calc_uv_error(test, U, V))
    # Plot the train and test error for each combination
    ax = axes[index]
    ax.plot(d_vals, uv_train_errors, label='Training Error', marker='o')
    ax.plot(d_vals, uv_test_errors, label='Test Error', marker='o')
    ax.set_xlabel('d (Number of Singular Values)')
    ax.set_ylabel('Average Squared Error')
    ax.set_title(f'Train/Test Errors for lambda={lambda_val}, sigma={sigma_val}')
    ax.legend()
    ax.grid(True)
# Adjust layout
plt.tight_layout()
plt.show()
```

5

 \mathbf{a}

 $lrs = [0.001,\ 0.01,\ 0.03,\ 0.04,\ 0.05]\ momentums = [0.5,\ 0.7,\ 0.9]\ Ms = [128,\ 256,\ 512,\ 1024,\ 2048]\ momentum = [20,\ 60,\ 100]$

I used the combination of random and grid search, I first hand searched few number and found out the higher momentum and learning rate around 0.03 with momentum over 60 has the best validation accuracy. After I have a sense of how accuracy is affected, then I grid searched the hyperparameters and found below are the three best models performance.

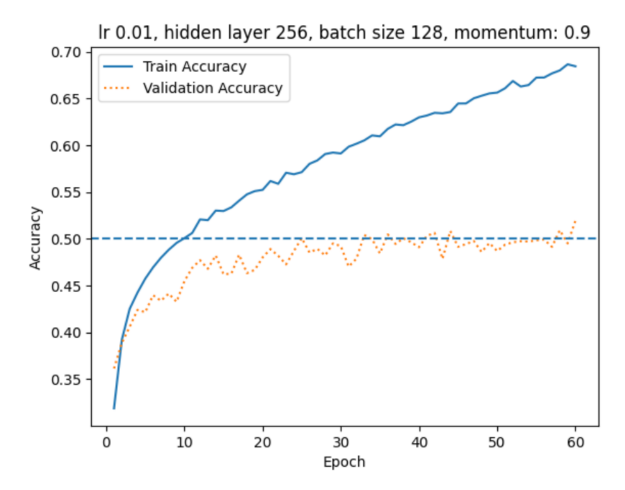


Figure 1: accuracy: 0.512

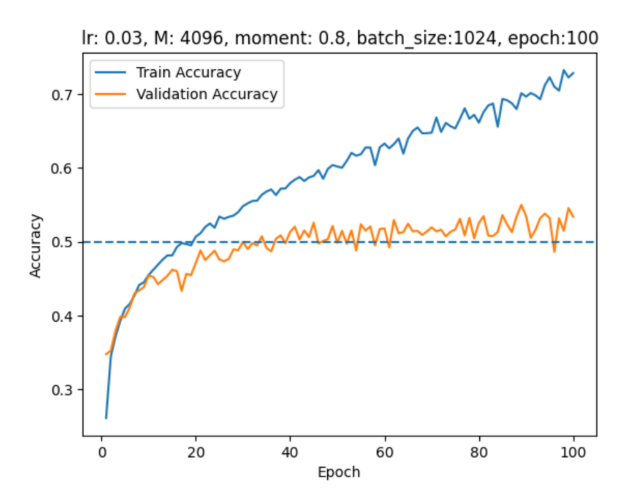


Figure 2: accuracy: 0.532

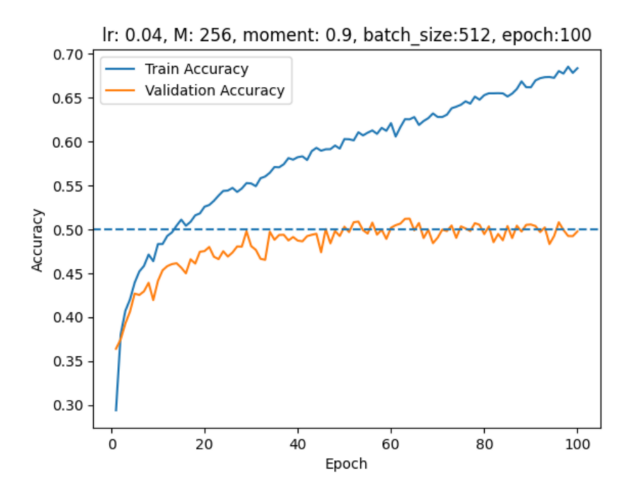


Figure 3: accuracy: 0.503

b

searched over parameters lrs = [0.001, 0.01] momentums = [0.5, 0.7, 0.9] Ms = [64, 128, 256] ks = [3, 5, 7] Ns = [2, 4, 7]

I used grid search, I used for loop to find the combinatorial of all posible hyperparameter pair and searched the best performing model.

Ir 0.01, hidden layer 64, kernel 5, pool size:4, batch size 128, momentum: 0.9

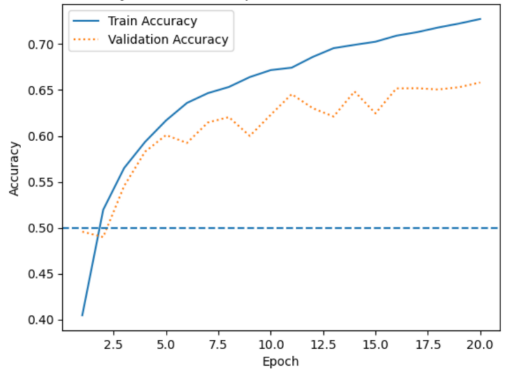


Figure 4: accuracy: 0.634

Ir 0.01, hidden layer 128, kernel 3, pool size:4, batch size 128, momentum: 0.9

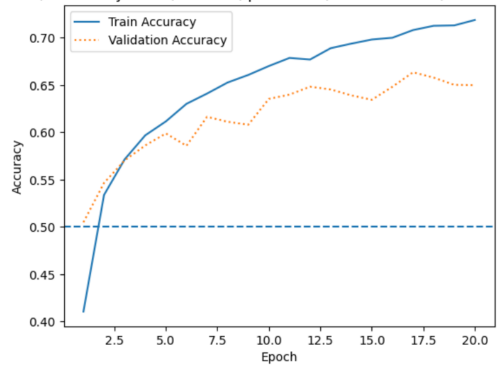


Figure 5: accuracy: 0.63

Ir 0.01, hidden layer 128, kernel 5, pool size:2, batch size 128, momentum: 0.9

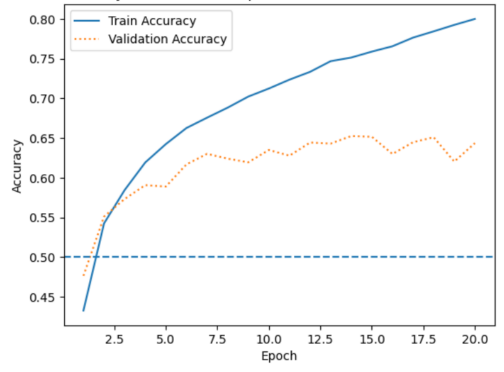


Figure 6: accuracy: 0.623

```
import torch
from torch import nn
import numpy as np
from typing import Tuple, Union, List, Callable
from torch.optim import SGD
import torchvision
from torch.utils.data import DataLoader, TensorDataset, random_split
import matplotlib.pyplot as plt
from tqdm.notebook import tqdm
assert torch.cuda.is_available(), "GPU is not available, check the directions above (or disable this
DEVICE = "cuda" if torch.cuda.is_available() else "cpu"
print(DEVICE) # this should print out CUDA
train_dataset = torchvision.datasets.CIFAR10("./data", train=True, download=True, transform=torchvisi
test_dataset = torchvision.datasets.CIFAR10("./data", train=False, download=True, transform=torchvisi
batch_size = 128
# batch_size= 4
train_dataset, val_dataset = random_split(train_dataset, [int(0.9 * len(train_dataset)), int( 0.1 * 1
# Create separate dataloaders for the train, test, and validation set
train_loader = DataLoader(
    train_dataset,
    batch_size=batch_size,
    shuffle=True,
    num_workers=8,
    pin_memory=True
)
val_loader = DataLoader(
    val_dataset,
    batch_size=batch_size,
    shuffle=True,
    num_workers=8,
    pin_memory=True
test_loader = DataLoader(
    test_dataset,
    batch_size=batch_size,
    shuffle=True,
    num_workers=8,
    pin_memory=True
)
imgs, labels = next(iter(train_loader))
print(f"A single batch of images has shape: {imgs.size()}")
example_image, example_label = imgs[0], labels[0]
c, w, h = example_image.size()
print(f"A single RGB image has {c} channels, width {w}, and height {h}.")
# This is one way to flatten our images
```

```
batch_flat_view = imgs.view(-1, c * w * h)
print(f"Size of a batch of images flattened with view: {batch_flat_view.size()}")
# This is another equivalent way
batch_flat_flatten = imgs.flatten(1)
print(f"Size of a batch of images flattened with flatten: {batch_flat_flatten.size()}")
# The new dimension is just the product of the ones we flattened
d = example_image.flatten().size()[0]
print(c * w * h == d)
# View the image
t = torchvision.transforms.ToPILImage()
plt.imshow(t(example_image))
# These are what the class labels in CIFAR-10 represent. For more information,
# visit https://www.cs.toronto.edu/~kriz/cifar.html
classes = ["airplane", "automobile", "bird", "cat", "deer", "dog", "frog",
           "horse", "ship", "truck"]
print(f"This image is labeled as class {classes[example_label]}")
def linear_model() -> nn.Module:
    """Instantiate a linear model and send it to device."""
    model = nn.Sequential(
            nn.Flatten(),
            nn.Linear(d, 10)
         )
    return model.to(DEVICE)
def fully_connected_model(M, input_dim=3072, output_dim=10):
    """Instantiate a fully connected model with one hidden layer and send it to device."""
    model = nn.Sequential(
       nn.Flatten(),
        nn.Linear(input_dim, M), # First fully-connected layer with size M
                                  # ReLU activation function
        nn.ReLU(),
        nn.Linear(M, output_dim) # Second fully-connected layer with size 10 (output layer)
    return model.to(DEVICE)
def conv_model(M, k, N, output_dim=10, input_size=(3, 32, 32)):
    """Instantiate a convolutional neural network model with one conv layer followed by max-pooling of
    # Define the model
    model = nn.Sequential(
        nn.Conv2d(input_size[0], M, k, padding=k//2), # Convolutional layer with 'same' padding
        nn.ReLU(), # Activation function
        nn.MaxPool2d(N), # Max pooling layer
        nn.Flatten(), # Flatten the output
        # No need to calculate the size manually, nn.Linear will infer it
        nn.Linear(M * ((input_size[1] // N) ** 2), output_dim) # Fully connected layer
    return model.to(DEVICE)
def train(
    model: nn.Module, optimizer: SGD,
    train_loader: DataLoader, val_loader: DataLoader,
    epochs: int = 20
```

```
)-> Tuple[List[float], List[float], List[float]]:
    Trains a model for the specified number of epochs using the loaders.
    Returns:
    Lists of training loss, training accuracy, validation loss, validation accuracy for each epoch.
   loss = nn.CrossEntropyLoss()
   train_losses = []
   train_accuracies = []
   val_losses = []
   val_accuracies = []
    for e in tqdm(range(epochs)):
       model.train()
       train_loss = 0.0
       train_acc = 0.0
       # Main training loop; iterate over train_loader. The loop
        # terminates when the train loader finishes iterating, which is one epoch.
       for (x_batch, labels) in train_loader:
            x_batch, labels = x_batch.to(DEVICE), labels.to(DEVICE)
            optimizer.zero_grad()
            labels_pred = model(x_batch)
            batch_loss = loss(labels_pred, labels)
            train_loss = train_loss + batch_loss.item()
            labels_pred_max = torch.argmax(labels_pred, 1)
            batch_acc = torch.sum(labels_pred_max == labels)
            train_acc = train_acc + batch_acc.item()
            batch_loss.backward()
            optimizer.step()
       train_losses.append(train_loss / len(train_loader))
       train_accuracies.append(train_acc / (batch_size * len(train_loader)))
       # Validation loop; use .no_grad() context manager to save memory.
       model.eval()
       val_loss = 0.0
       val_acc = 0.0
       with torch.no_grad():
            for (v_batch, labels) in val_loader:
                v_batch, labels = v_batch.to(DEVICE), labels.to(DEVICE)
               labels_pred = model(v_batch)
                v_batch_loss = loss(labels_pred, labels)
               val_loss = val_loss + v_batch_loss.item()
                v_pred_max = torch.argmax(labels_pred, 1)
                batch_acc = torch.sum(v_pred_max == labels)
                val_acc = val_acc + batch_acc.item()
            val_losses.append(val_loss / len(val_loader))
            val_accuracies.append(val_acc / (batch_size * len(val_loader)))
```

return train_losses, train_accuracies, val_losses, val_accuracies

```
def parameter_search(
   train_loader: DataLoader,
   val_loader: DataLoader,
   model_fn:Callable[[], nn.Module]
) -> float:
    Parameter search for our linear model using SGD.
   Args:
    train_loader: the train dataloader.
   val_loader: the validation dataloader.
    model_fn: a function that, when called, returns a torch.nn.Module.
   Returns:
    The learning rate with the least validation loss.
   NOTE: you may need to modify this function to search over and return
    other parameters beyond learning rate.
   num_iter = 4
   best_loss = torch.tensor(np.inf)
   best_lr = 0.0
   best_M = 0.0
   best_k=0.0
   best_N=0.0
   num_epoch = 100
   lrs = torch.linspace(0.02, 0.1, num_iter)
    # Possible learning rates to try
   lrs = [0.001, 0.01]
    # Possible momentum values to try
   momentums = [0.5, 0.7, 0.9]
    # Possible numbers of filters to try in the convolutional layer
   Ms = [64, 128, 256]
    # Possible kernel sizes to try for the convolutional layer
   ks = [3, 5, 7]
    # Possible pooling sizes to try after the convolutional layer
   Ns = [2, 4]
    # Number of epochs for each training run
   num_epoch = 20
   for M in Ms:
        for k in ks:
            for N in Ns:
                for moment in momentums:
                    for lr in lrs:
                        model = model_fn(M, k, N)
                        optim = SGD(model.parameters(), lr, momentum=moment)
                        train_loss, train_acc, val_loss, val_acc = train(
                            model,
                            optim,
                            train_loader,
```

```
val_loader,
                            epochs=num_epoch
                        print(f"result for learning rate {lr}, hidden layer {M}, kernel {k}, pool siz
                        print(f'val_loss: {min(val_loss)}')
                        if min(val_loss) < best_loss:</pre>
                            best_loss = min(val_loss)
                            best_lr = lr
                            best_M = M
                            best_moment = moment
                            best_k = k
                            best_N = N
    return train_loss, train_acc, val_loss, val_acc, best_lr, best_M, best_k, best_N, best_moment, nu
train_loss, train_accuracy, val_loss, val_accuracy, best_lr, best_M, best_k, best_N, best_moment, num
model = fully_connected_model(best_M, best_k, best_N)
optimizer = SGD(model.parameters(), best_lr, momentum=best_moment)
# We are only using 20 epochs for this example. You may have to use more.
train_loss, train_accuracy, val_loss, val_accuracy = train(
    model, optimizer, train_loader, val_loader, num_epoch
)
epochs = range(1, num_epoch+1)
plt.plot(epochs, train_accuracy, label="Train Accuracy")
plt.plot(epochs, val_accuracy, label="Validation Accuracy", linestyle='dotted')
plt.axhline(y=0.5, linestyle="--")
plt.xlabel("Epoch")
plt.ylabel("Accuracy")
plt.legend()
plt.title(f"lr {best_lr}, hidden layer {best_M}, kernel {best_k}, pool size:{best_N}, batch size {bat
plt.show()
def evaluate(
    model: nn.Module, loader: DataLoader
) -> Tuple[float, float]:
    """Computes test loss and accuracy of model on loader."""
    loss = nn.CrossEntropyLoss()
    model.eval()
    test_loss = 0.0
    test_acc = 0.0
    with torch.no_grad():
        for (batch, labels) in loader:
            batch, labels = batch.to(DEVICE), labels.to(DEVICE)
            y_batch_pred = model(batch)
            batch_loss = loss(y_batch_pred, labels)
            test_loss = test_loss + batch_loss.item()
            pred_max = torch.argmax(y_batch_pred, 1)
            batch_acc = torch.sum(pred_max == labels)
            test_acc = test_acc + batch_acc.item()
        test_loss = test_loss / len(loader)
        test_acc = test_acc / (batch_size * len(loader))
        return test_loss, test_acc
```

```
test_loss, test_acc = evaluate(model, test_loader)
print(f"Test Accuracy: {test_acc}")
```

entire days