

(2) Softmax: =
$$\frac{e^{\lambda_{i}^{T}} x + bi}{\sum_{k=1}^{E} e^{\lambda_{k}^{T}} x + bk}$$
 let
$$\begin{cases} \lambda_{i}^{T} x + bi = \lambda_{i}^{T} x \\ \lambda_{k}^{T} x + bk = \lambda_{k}^{T} x \end{cases}$$
(1) $\log - k k e l \cdot hood : (t steel)$

(1)
$$\log - kke! hood : (tstub)$$

$$= - arg \max_{\theta} \frac{1}{m} \sum_{i=1}^{m} \log \left(\frac{e^{i\lambda_i} \hat{\chi}^{(i)}}{\sum_{k=1}^{m} e^{i\lambda_k} \hat{\chi}^{(i)}} \right)$$

$$= arg \min_{i=1}^{m} \sum_{k=1}^{m} \left(- \log e^{i\lambda_k} \frac{1}{\chi^{(i)}} + \log \sum_{k=1}^{m} e^{i\lambda_k} \frac{1}{\chi^{(i)}} \right)$$

$$= arg \min_{i=1}^{m} \sum_{k=1}^{m} \left(- i\lambda_i \sum_{k=1}^{m} \frac{1}{\chi^{(i)}} + \log \sum_{k=1}^{m} e^{i\lambda_k} \frac{1}{\chi^{(i)}} \right)$$

$$= arg \min_{i=1}^{m} \sum_{k=1}^{m} \left(- i\lambda_i \sum_{k=1}^{m} \frac{1}{\chi^{(i)}} + \log \sum_{k=1}^{m} e^{i\lambda_k} \frac{1}{\chi^{(i)}} \right)$$

$$\frac{1}{2} \sum_{i=1}^{M} \frac{1}{2} \frac{1}{2}$$

This is the k-nearest neighbors workbook for ECE C147/C247 Assignment #2

Please follow the notebook linearly to implement k-nearest neighbors.

Please print out the workbook entirely when completed.

The goal of this workbook is to give you experience with the data, training and evaluating a simple classifier, k-fold cross validation, and as a Python refresher.

Import the appropriate libraries

num classes = len(classes) samples per class = 7

for y, cls in enumerate(classes):

idxs = np.flatnonzero(y train == y)

plt idx = i * num classes + y + 1

for i, idx in enumerate(idxs):

```
In [1]:
         from google.colab import drive
         drive.mount("/content/drive", force remount=True)
        Mounted at /content/drive
In [2]:
         %cd /content/drive/MyDrive/ColabNotebooks/c247/hw2/
         import numpy as np # for doing most of our calculations
         import matplotlib.pyplot as plt# for plotting
         from utils.data utils import load CIFAR10 # function to load the CIFAR-10 dataset.
         # Load matplotlib images inline
         %matplotlib inline
         # These are important for reloading any code you write in external .py files.
         # see http://stackoverflow.com/questions/1907993/autoreload-of-modules-in-ipython
         %load ext autoreload
         %autoreload 2
        /content/drive/MyDrive/ColabNotebooks/c247/hw2
In [3]:
         # Set the path to the CIFAR-10 data
         cifar10 dir = 'cifar-10-batches-py' # You need to update this line
         X train, y train, X test, y test = load CIFAR10(cifar10 dir)
         # As a sanity check, we print out the size of the training and test data.
         print('Training data shape: ', X train.shape)
```

```
print('Training labels shape: ', y train.shape)
         print('Test data shape: ', X test.shape)
         print('Test labels shape: ', y test.shape)
        Training data shape: (50000, 32, 32, 3)
        Training labels shape: (50000,)
        Test data shape: (10000, 32, 32, 3)
        Test labels shape: (10000,)
In [4]:
         # Visualize some examples from the dataset.
         # We show a few examples of training images from each class.
         classes = ['plane', 'car', 'bird', 'cat', 'deer', 'dog', 'frog', 'horse', 'ship', 'truc'
```

idxs = np.random.choice(idxs, samples per class, replace=False)

```
plt.subplot(samples_per_class, num_classes, plt_idx)
    plt.imshow(X_train[idx].astype('uint8'))
    plt.axis('off')
    if i == 0:
        plt.title(cls)
plt.show()
```

```
plane car bird cat deer dog frog horse ship truck

| Same and a second a se
```

```
In [5]: # Subsample the data for more efficient code execution in this exercise
   num_training = 5000
   mask = list(range(num_training))
   X_train = X_train[mask]
   y_train = y_train[mask]

   num_test = 500
   mask = list(range(num_test))
   X_test = X_test[mask]
   y_test = y_test[mask]

# Reshape the image data into rows
   X_train = np.reshape(X_train, (X_train.shape[0], -1))
   X_test = np.reshape(X_test, (X_test.shape[0], -1))
   print(X_train.shape, X_test.shape)
```

(5000, 3072) (500, 3072)

K-nearest neighbors

In the following cells, you will build a KNN classifier and choose hyperparameters via k-fold cross-validation.

```
In [13]: # Import the KNN class
    from nndl import KNN

In [14]: # Declare an instance of the knn class.
    knn = KNN()

# Train the classifier.
    # We have implemented the training of the KNN classifier.
    # Look at the train function in the KNN class to see what this does.
    knn.train(X=X_train, y=y_train)
```

Questions

- (1) Describe what is going on in the function knn.train().
- (2) What are the pros and cons of this training step?

Answers

- (1) Knn.train would store all the training data inside the class.
- (2) Pros: It is very simple and fast. Cons: it require lots of memory to store all the training data.

KNN prediction

In the following sections, you will implement the functions to calculate the distances of test points to training points, and from this information, predict the class of the KNN.

```
In [15]: # Implement the function compute_distances() in the KNN class.
# Do not worry about the input 'norm' for now; use the default definition of the norm
# in the code, which is the 2-norm.
# You should only have to fill out the clearly marked sections.

import time
time_start =time.time()

dists_L2 = knn.compute_distances(X=X_test)

print('Time to run code: {}'.format(time.time()-time_start))
print('Frobenius norm of L2 distances: {}'.format(np.linalg.norm(dists_L2, 'fro')))
```

```
Time to run code: 42.59095358848572
Frobenius norm of L2 distances: 7906696.077040902
```

Really slow code

Note: This probably took a while. This is because we use two for loops. We could increase the speed via vectorization, removing the for loops.

If you implemented this correctly, evaluating np.linalg.norm(dists_L2, 'fro') should return: ~7906696

KNN vectorization

The above code took far too long to run. If we wanted to optimize hyperparameters, it would be timeexpensive. Thus, we will speed up the code by vectorizing it, removing the for loops.

```
In [16]: # Implement the function compute_L2_distances_vectorized() in the KNN class.
# In this function, you ought to achieve the same L2 distance but WITHOUT any for loops.
# Note, this is SPECIFIC for the L2 norm.

time_start =time.time()
dists_L2_vectorized = knn.compute_L2_distances_vectorized(X=X_test)
print('Time to run code: {}'.format(time.time()-time_start))
print('Difference in L2 distances between your KNN implementations (should be 0): {}'.fo

Time to run code: 0.5911593437194824
```

Difference in L2 distances between your KNN implementations (should be 0): 0.0

Speedup

Depending on your computer speed, you should see a 10-100x speed up from vectorization. On our computer, the vectorized form took 0.36 seconds while the naive implementation took 38.3 seconds.

Implementing the prediction

Now that we have functions to calculate the distances from a test point to given training points, we now implement the function that will predict the test point labels.

```
In [18]:
      # Implement the function predict labels in the KNN class.
      # Calculate the training error (num incorrect / total samples)
      # from running knn.predict labels with k=1
      error = 1
      # ------ #
      # YOUR CODE HERE:
        Calculate the error rate by calling predict labels on the test
        data with k = 1. Store the error rate in the variable error.
      y pred = knn.predict labels(dists L2 vectorized, k= 1)
      error = 1 - (np.sum(y test == y pred) / len(y test))
      # END YOUR CODE HERE
      print(error)
      0.726
```

If you implemented this correctly, the error should be: 0.726.

This means that the k-nearest neighbors classifier is right 27.4% of the time, which is not great, considering that chance levels are 10%.

Optimizing KNN hyperparameters

In this section, we'll take the KNN classifier that you have constructed and perform cross-validation to choose a best value of k, as well as a best choice of norm.

Create training and validation folds

First, we will create the training and validation folds for use in k-fold cross validation.

Optimizing the number of nearest neighbors hyperparameter.

In this section, we select different numbers of nearest neighbors and assess which one has the lowest k-fold cross validation error.

```
In [21]:
        time start =time.time()
        ks = [1, 2, 3, 5, 7, 10, 15, 20, 25, 30]
        # YOUR CODE HERE:
          Calculate the cross-validation error for each k in ks, testing
        # the trained model on each of the 5 folds. Average these errors
        # together and make a plot of k vs. cross-validation error. Since
          we are assuming L2 distance here, please use the vectorized code!
          Otherwise, you might be waiting a long time.
        # ----- #
        fold size = len(X train) // num folds
        training fold size = int((num folds-1) * fold size)
        x size = np.array(X train folds).shape[2]
        errors = []
        for k val in ks:
           error total = 0
           for fold idx in range(num folds):
               # train on training fold
               X training fold = np.array(X train folds[:fold idx] + X train folds[fold idx+1:
               y training fold = np.array(y train folds[:fold idx] + y train folds[fold idx+1:
               knn.train(X=X training fold, y=y training fold)
               # compute L2 distance and predict y on test fold
               dists = knn.compute L2 distances vectorized(X=X train folds[fold idx])
               y pred = knn.predict labels(dists, k=k val)
               # add error to error total
               error total += np.mean(y train folds[fold idx] != y pred)
            error_avg = error_total / num_folds
           print('k={}: error={}'.format(k val, round(error avg, 4)))
            errors.append(error avg)
        # ------ #
        # END YOUR CODE HERE
        print('Computation time: %.2f'%(time.time()-time start))
```

```
k=1: error=0.7344
k=2: error=0.7626
k=3: error=0.7504
k=5: error=0.7268
k=7: error=0.7256
k=10: error=0.7198
k=15: error=0.725
k=20: error=0.721
k=25: error=0.7242
k=30: error=0.7266
Computation time: 63.21
```

Questions:

- (1) What value of k is best amongst the tested k's?
- (2) What is the cross-validation error for this value of k?

Answers:

- (1) K equal to 10 has the vest testing performance.
- (2) The cross-validation error for this value is 0.7198

Optimizing the norm

Next, we test three different norms (the 1, 2, and infinity norms) and see which distance metric results in the best cross-validation performance.

```
In [23]:
         time start =time.time()
         L1 norm = lambda x: np.linalg.norm(x, ord=1)
         L2 norm = lambda x: np.linalg.norm(x, ord=2)
         Linf norm = lambda x: np.linalg.norm(x, ord= np.inf)
         norms = [L1 norm, L2 norm, Linf norm]
         # YOUR CODE HERE:
           Calculate the cross-validation error for each norm in norms, testing
           the trained model on each of the 5 folds. Average these errors
           together and make a plot of the norm used vs the cross-validation error
           Use the best cross-validation k from the previous part.
           Feel free to use the compute distances function. We're testing just
           three norms, but be advised that this could still take some time.
           You're welcome to write a vectorized form of the L1- and Linf- norms
           to speed this up, but it is not necessary.
         k best = 10
         errors = []
         for norm idx, norm in enumerate(norms):
            error total = 0
            for fold idx in range(num folds):
                # train on training fold
                X training fold = np.array(X train folds[:fold idx] + X train folds[fold idx+1:
                y training fold = np.array(y train folds[:fold idx] + y train folds[fold idx+1:]
                knn.train(X=X training fold, y=y training fold)
                # compute distance and predict y on test fold
```

```
norm=<function <lambda> at 0x7fd7ab8074d0>: error=0.6886 norm=<function <lambda> at 0x7fd7ab7f9170>: error=0.7198 norm=<function <lambda> at 0x7fd7ab7ed170>: error=0.837 Computation time: 805.66
```

Questions:

- (1) What norm has the best cross-validation error?
- (2) What is the cross-validation error for your given norm and k?

Answers:

- (1) L1 norm has the best cross-validation error.
- (2) Using L1 norm and K= 10, the error is 0.6886.

Evaluating the model on the testing dataset.

Now, given the optimal k and norm you found in earlier parts, evaluate the testing error of the k-nearest neighbors model.

Error rate achieved: 0.716

Question:

How much did your error improve by cross-validation over naively choosing k=1 and using the L2-norm?

Answer:

```
It improve by 0.01 (0.726-0.716 = 0.01)
```

knn.py code

```
In [ ]:
       import numpy as np
       import pdb
       class KNN(object):
         def init (self):
          pass
         def train(self, X, y):
          Inputs:
          - X is a numpy array of size (num examples, D)
           - y is a numpy array of size (num examples, )
          self.X train = X
          self.y train = y
         def compute distances(self, X, norm=None):
           Compute the distance between each test point in X and each training point
          in self.X train.
          Inputs:
           - X: A numpy array of shape (num test, D) containing test data.
           - norm: the function with which the norm is taken.
          Returns:
           - dists: A numpy array of shape (num test, num train) where dists[i, j]
            is the Euclidean distance between the ith test point and the jth training
            point.
           if norm is None:
            norm = lambda x: np.sqrt(np.sum(x**2))
            \#norm = 2
          num test = X.shape[0]
          num train = self.X train.shape[0]
           dists = np.zeros((num test, num train))
           for i in np.arange(num test):
            for j in np.arange(num train):
              # ------ #
              # YOUR CODE HERE:
              # Compute the distance between the ith test point and the jth
              # training point using norm(), and store the result in dists[i, j].
              # ================== #
              #Calculating the L1-norm
              dists[i,j] = norm(self.X train[j] - X[i])
              # END YOUR CODE HERE
              # ----- #
```

```
return dists
def compute L2 distances vectorized(self, X):
 Compute the distance between each test point in X and each training point
 in self.X train WITHOUT using any for loops.
 Inputs:
 - X: A numpy array of shape (num test, D) containing test data.
 - dists: A numpy array of shape (num test, num train) where dists[i, j]
   is the Euclidean distance between the ith test point and the jth training
 num test = X.shape[0]
 num train = self.X train.shape[0]
 dists = np.zeros((num test, num train))
 # ----- #
 # YOUR CODE HERE:
   Compute the L2 distance between the ith test point and the jth
    training point and store the result in dists[i, j]. You may
    NOT use a for loop (or list comprehension). You may only use
    numpy operations.
    HINT: use broadcasting. If you have a shape (N,1) array and
    a shape (M,) array, adding them together produces a shape (N, M)
    array.
  # ------ #
 train sqr = np.sum(np.square(self.X train), axis = 1)
 test sqr = np.sum(np.square(X), axis= 1)
 test sqr = test sqr.reshape((test sqr.shape[0], 1))
 trainSqr testSqr = X @ self.X train.T
 dists = np.sqrt(train sqr + test sqr - 2 * trainSqr testSqr)
 # ----- #
 # END YOUR CODE HERE
 return dists
def predict labels(self, dists, k=1):
 Given a matrix of distances between test points and training points,
 predict a label for each test point.
 Inputs:
 - dists: A numpy array of shape (num test, num train) where dists[i, j]
   gives the distance betwen the ith test point and the jth training point.
 Returns:
 - y: A numpy array of shape (num test,) containing predicted labels for the
   test data, where y[i] is the predicted label for the test point X[i].
 num test = dists.shape[0]
 y pred = np.zeros(num test)
 for i in np.arange(num test):
   \# A list of length k storing the labels of the k nearest neighbors to
   # the ith test point.
   closest y = []
```

```
# YOUR CODE HERE:
 # Use the distances to calculate and then store the labels of
 # the k-nearest neighbors to the ith test point. The function
   numpy.argsort may be useful.
 # After doing this, find the most common label of the k-nearest
 # neighbors. Store the predicted label of the ith training example
   as y pred[i]. Break ties by choosing the smaller label.
 closest idxs = np.argsort(dists[i])[:k]
 closest y = [self.y train[idx] for idx in closest idxs]
 unique, counts = np.unique(closest y, return counts=True)
 closest y = sorted(dict(zip(unique, counts)).items(), key=lambda x: <math>(-x[1], x[0])
 y pred[i] = closest y[0][0]
 # ----- #
    # END YOUR CODE HERE
    # ----- #
return y pred
```

This is the softmax workbook for ECE C147/C247 Assignment #2

Please follow the notebook linearly to implement a softmax classifier.

Please print out the workbook entirely when completed.

X val -= mean image

The goal of this workbook is to give you experience with training a softmax classifier.

```
In [1]:
         from google.colab import drive
         drive.mount("/content/drive", force remount=True)
         %cd /content/drive/MyDrive/ColabNotebooks/c247/hw2/
        Mounted at /content/drive
        /content/drive/MyDrive/ColabNotebooks/c247/hw2
In [2]:
         import random
         import numpy as np
         from utils.data utils import load CIFAR10
         import matplotlib.pyplot as plt
         %matplotlib inline
         %load ext autoreload
         %autoreload 2
In [4]:
         def get CIFAR10 data(num training=49000, num validation=1000, num test=1000, num dev=500
             Load the CIFAR-10 dataset from disk and perform preprocessing to prepare
             it for the linear classifier. These are the same steps as we used for the
             SVM, but condensed to a single function.
             # Load the raw CIFAR-10 data
             cifar10 dir = 'cifar-10-batches-py' # You need to update this line
             X train, y train, X test, y test = load CIFAR10(cifar10 dir)
             # subsample the data
             mask = list(range(num training, num training + num validation))
             X val = X train[mask]
             y val = y train[mask]
             mask = list(range(num training))
             X train = X train[mask]
             y_train = y_train[mask]
             mask = list(range(num test))
             X test = X test[mask]
             y test = y test[mask]
             mask = np.random.choice(num training, num dev, replace=False)
             X dev = X train[mask]
             y dev = y train[mask]
             # Preprocessing: reshape the image data into rows
             X train = np.reshape(X train, (X train.shape[0], -1))
             X \text{ val} = \text{np.reshape}(X \text{ val}, (X \text{ val.shape}[0], -1))
             X test = np.reshape(X test, (X test.shape[0], -1))
             X_{dev} = np.reshape(X_{dev}, (X_{dev.shape}[0], -1))
             # Normalize the data: subtract the mean image
             mean image = np.mean(X train, axis = 0)
             X train -= mean image
```

```
X test -= mean image
    X dev -= mean image
     # add bias dimension and transform into columns
    X train = np.hstack([X train, np.ones((X train.shape[0], 1))])
    X \text{ val} = \text{np.hstack}([X \text{ val, np.ones}((X \text{ val.shape}[0], 1))])
    X test = np.hstack([X test, np.ones((X test.shape[0], 1))])
    X dev = np.hstack([X dev, np.ones((X dev.shape[0], 1))])
    return X train, y train, X val, y val, X test, y test, X dev, y dev
 # Invoke the above function to get our data.
X train, y train, X val, y val, X test, y test, X dev, y dev = get CIFAR10 data()
print('Train data shape: ', X_train.shape)
print('Train labels shape: ', y train.shape)
print('Validation data shape: ', X val.shape)
print('Validation labels shape: ', y val.shape)
print('Test data shape: ', X test.shape)
print('Test labels shape: ', y test.shape)
print('dev data shape: ', X dev.shape)
print('dev labels shape: ', y dev.shape)
Train data shape: (49000, 3073)
Train labels shape: (49000,)
```

```
Train data shape: (49000, 3073)
Train labels shape: (49000,)
Validation data shape: (1000, 3073)
Validation labels shape: (1000,)
Test data shape: (1000, 3073)
Test labels shape: (1000,)
dev data shape: (500, 3073)
dev labels shape: (500,)
```

Training a softmax classifier.

The following cells will take you through building a softmax classifier. You will implement its loss function, then subsequently train it with gradient descent. Finally, you will choose the learning rate of gradient descent to optimize its classification performance.

Softmax loss

```
In [7]: ## Implement the loss function of the softmax using a for loop over
# the number of examples
loss = softmax.loss(X_train, y_train)
```

```
In [8]:
```

```
print(loss)
```

2.3277607028048966

Question:

You'll notice the loss returned by the softmax is about 2.3 (if implemented correctly). Why does this make sense?

Answer:

We initialize the weight randomly, so the probability that the classifier can predict correctly is only 10%. Therefore, loss of the softmadx would be -log(1/10) which is equal to 2.3....

Softmax gradient

```
In [9]: ## Calculate the gradient of the softmax loss in the Softmax class.
# For convenience, we'll write one function that computes the loss
# and gradient together, softmax.loss_and_grad(X, y)
# You may copy and paste your loss code from softmax.loss() here, and then
# use the appropriate intermediate values to calculate the gradient.

loss, grad = softmax.loss_and_grad(X_dev,y_dev)

# Compare your gradient to a gradient check we wrote.
# You should see relative gradient errors on the order of 1e-07 or less if you implement softmax.grad_check_sparse(X_dev, y_dev, grad)

numerical: -0.115047 analytic: -0.115047, relative error: 1.156804e-07
```

```
numerical: -0.113047 analytic: -0.113047, relative error: 1.136004e-07 numerical: 2.963360 analytic: 2.963359, relative error: 1.842207e-08 numerical: 0.826104 analytic: 0.826104, relative error: 5.298482e-08 numerical: 1.493036 analytic: 1.493036, relative error: 9.809412e-09 numerical: 1.652761 analytic: 1.652761, relative error: 3.783540e-08 numerical: 2.158975 analytic: 2.158975, relative error: 1.272820e-08 numerical: -0.258691 analytic: -0.258691, relative error: 2.130137e-07 numerical: -0.313621 analytic: -0.313621, relative error: 1.161359e-07 numerical: -0.310841 analytic: -0.310841, relative error: 1.787894e-08 numerical: -3.010592 analytic: -3.010592, relative error: 9.172221e-09
```

A vectorized version of Softmax

To speed things up, we will vectorize the loss and gradient calculations. This will be helpful for stochastic gradient descent.

```
In [10]: import time

In [11]: ## Implement softmax.fast_loss_and_grad which calculates the loss and gradient
# WITHOUT using any for loops.

# Standard loss and gradient
tic = time.time()
loss, grad = softmax.loss_and_grad(X_dev, y_dev)
toc = time.time()
print('Normal loss / grad_norm: {} / {} computed in {}s'.format(loss, np.linalg.norm(gratic = time.time())
loss_vectorized, grad_vectorized = softmax.fast_loss_and_grad(X_dev, y_dev)
```

```
toc = time.time()
print('Vectorized loss / grad: {} / {} computed in {}s'.format(loss_vectorized, np.lina)
# The losses should match but your vectorized implementation should be much faster.
print('difference in loss / grad: {} /{} '.format(loss - loss_vectorized, np.linalg.norr
# You should notice a speedup with the same output.
```

```
Normal loss / grad_norm: 2.340247274470396 / 349.5738796202033 computed in 0.13048052787 780762s
Vectorized loss / grad: 2.340247274470397 / 349.5738796202033 computed in 0.043454170227 05078s
difference in loss / grad: -8.881784197001252e-16 /3.250136140987487e-13
```

Stochastic gradient descent

We now implement stochastic gradient descent. This uses the same principles of gradient descent we discussed in class, however, it calculates the gradient by only using examples from a subset of the training set (so each gradient calculation is faster).

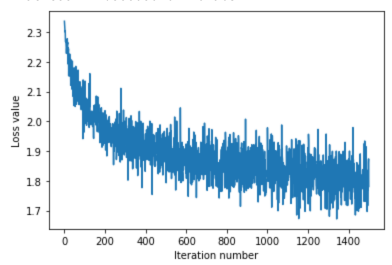
Question:

How should the softmax gradient descent training step differ from the svm training step, if at all?

Answer:

They use the same way to calculate the gradient and update their parameter, which would give a very similar training steps. However, since they are using different loss function, each of their outure would be different.

```
iteration 0 / 1500: loss 2.336592660663754
iteration 100 / 1500: loss 2.0557222613850827
iteration 200 / 1500: loss 2.0357745120662813
iteration 300 / 1500: loss 1.9813348165609888
iteration 400 / 1500: loss 1.9583142443981614
iteration 500 / 1500: loss 1.862265307354135
iteration 600 / 1500: loss 1.8532611454359382
iteration 700 / 1500: loss 1.8353062223725827
iteration 800 / 1500: loss 1.8293892468827635
iteration 900 / 1500: loss 1.899215853035748
iteration 1000 / 1500: loss 1.97835035402523
iteration 1100 / 1500: loss 1.8470797913532633
iteration 1200 / 1500: loss 1.8411450268664082
iteration 1300 / 1500: loss 1.7910402495792102
iteration 1400 / 1500: loss 1.8705803029382257
That took 14.388030767440796s
```



Evaluate the performance of the trained softmax classifier on the validation data.

```
In [13]: ## Implement softmax.predict() and use it to compute the training and testing error.

y_train_pred = softmax.predict(X_train)
print('training accuracy: {}'.format(np.mean(np.equal(y_train,y_train_pred), )))
y_val_pred = softmax.predict(X_val)
print('validation accuracy: {}'.format(np.mean(np.equal(y_val, y_val_pred)), ))

training accuracy: 0.3811428571428571
```

Optimize the softmax classifier

validation accuracy: 0.398

You may copy and paste your optimization code from the SVM here.

```
- The best validation accuracy corresponding to the best validation error.
   Select the SVM that achieved the best validation error and report
     its error rate on the test set.
 # ----- #
 lr = [1e-1, 1e-2, 1e-3, 1e-4, 1e-5, 1e-6, 1e-7, 1e-8]
 val acc = []
 clf = Softmax()
 best rate = -1
 best accuracy = 0
 for i in np.arange(len(lr)):
   clf.train(X train, y train, learning rate= lr[i],
           num iters= 1500, verbose= False)
   pred = clf.predict(X val)
  acc = np.mean((pred == y val))
  val acc.append(acc)
  print('Learning Rate= {}, Accuracy= {}'.format(lr[i], acc))
   if (best accuracy < acc):</pre>
    best rate = lr[i]
    best accuracy = acc
 clf.train(X train, y train, learning rate= best rate,
          num iters= 1500, verbose= False)
 test error = 1 - np.mean((clf.predict(X test) == y test))
 print()
 print("Best Learning Rate:", best rate)
 print("Best Validation accuracy:", best accuracy)
 print("Test error with best learning rate:", test error)
 # END YOUR CODE HERE
 /content/drive/MyDrive/ColabNotebooks/c247/hw2/nndl/softmax.py:136: RuntimeWarning: over
flow encountered in exp
  e A = np.exp(A)
/content/drive/MyDrive/ColabNotebooks/c247/hw2/nndl/softmax.py:147: RuntimeWarning: inva
lid value encountered in true divide
  S = e A / summed e A.reshape(num train, 1)
Learning Rate= 0.1, Accuracy= 0.087
Learning Rate= 0.01, Accuracy= 0.087
Learning Rate= 0.001, Accuracy= 0.087
Learning Rate= 0.0001, Accuracy= 0.293
Learning Rate= 1e-05, Accuracy= 0.315
Learning Rate= 1e-06, Accuracy= 0.403
Learning Rate= 1e-07, Accuracy= 0.391
Learning Rate= 1e-08, Accuracy= 0.298
Best Learning Rate: 1e-06
Best Validation accuracy: 0.403
Test error with best learning rate: 0.617
softmax.py code
 import numpy as np
```

In []:

class Softmax(object):

def init (self, dims=[10, 3073]):

```
self.init weights(dims=dims)
def init weights(self, dims):
 Initializes the weight matrix of the Softmax classifier.
 Note that it has shape (C, D) where C is the number of
 classes and D is the feature size.
 self.W = np.random.normal(size=dims) * 0.0001
def loss(self, X, y):
 11 11 11
 Calculates the softmax loss.
 Inputs have dimension D, there are C classes, and we operate on minibatches
 of N examples.
 Inputs:
 - X: A numpy array of shape (N, D) containing a minibatch of data.
 - y: A numpy array of shape (N,) containing training labels; y[i] = c means
   that X[i] has label c, where 0 \le c \le C.
 Returns a tuple of:
 - loss as single float
 # Initialize the loss to zero.
 loss = 0.0
 # ----- #
 # YOUR CODE HERE:
 # Calculate the normalized softmax loss. Store it as the variable loss.
   (That is, calculate the sum of the losses of all the training
    set margins, and then normalize the loss by the number of
 # training examples.)
 # ------ #
 num train = X.shape[0]
 for i in range(num train):
     A i = np.matmul(self.W, X[i])
     log summed e A = np.log(np.sum(np.exp(A i)))
     A y = np.matmul(self.W[y[i]], X[i])
     loss += log summed e A - A y
 loss /= num train
 # ------ #
 # END YOUR CODE HERE
 return loss
def loss and grad(self, X, y):
 Same as self.loss(X, y), except that it also returns the gradient.
 Output: grad -- a matrix of the same dimensions as W containing
  the gradient of the loss with respect to W.
 # Initialize the loss and gradient to zero.
 loss = 0.0
 grad = np.zeros like(self.W)
 # ------ #
  # YOUR CODE HERE:
  # Calculate the softmax loss and the gradient. Store the gradient
   as the variable grad.
```

```
# ------ #
 num train = X.shape[0]
 num classes = self.W.shape[0]
 for i in range(num train):
     A i = np.matmul(self.W, X[i])
     e A = np.exp(A i)
     summed e A = np.sum(e A)
     log summed e A = np.log(summed e A)
     A y = np.matmul(self.W[y[i]], X[i])
     loss += log summed e A - A y
     for j in range(num classes):
        grad[j] += X[i] * e A[j] / summed e A
     grad[y[i]] -= X[i]
 loss /= num train
 grad /= num train
 # END YOUR CODE HERE
 return loss, grad
def grad check sparse(self, X, y, your grad, num checks=10, h=1e-5):
 sample a few random elements and only return numerical
 in these dimensions.
 for i in np.arange(num checks):
   ix = tuple([np.random.randint(m) for m in self.W.shape])
   oldval = self.W[ix]
   self.W[ix] = oldval + h # increment by h
   fxph = self.loss(X, y)
   self.W[ix] = oldval - h # decrement by h
   fxmh = self.loss(X, y) # evaluate f(x - h)
   self.W[ix] = oldval # reset
   grad numerical = (fxph - fxmh) / (2 * h)
   grad analytic = your grad[ix]
   rel error = abs(grad numerical - grad analytic) / (abs(grad numerical) + abs(grad
   print('numerical: %f analytic: %f, relative error: %e' % (grad numerical, grad analytic)
def fast loss and grad(self, X, y):
 A vectorized implementation of loss and grad. It shares the same
 inputs and ouptuts as loss and grad.
 loss = 0.0
 grad = np.zeros(self.W.shape) # initialize the gradient as zero
 # ----- #
 # YOUR CODE HERE:
 # Calculate the softmax loss and gradient WITHOUT any for loops.
 num train = X.shape[0]
 # Loss
 A = np.matmul(X, self.W.T)
 e A = np.exp(A)
 summed e A = np.sum(e A, axis=1)
```

```
log_summed_e_A = np.log(summed e A)
 pred = np.sum(np.multiply(self.W[y], X), axis=1)
 loss matrix = log summed e A - pred
 loss = np.sum(loss matrix) / num train
 # Gradient
 S = e A / summed e A.reshape(num train, 1)
 S[np.arange(num train), y] -= 1
 S = np.matmul(S.T, X)
 grad = S / num train
 # END YOUR CODE HERE
 # ------ #
 return loss, grad
def train(self, X, y, learning rate=1e-3, num iters=100,
        batch size=200, verbose=False):
 Train this linear classifier using stochastic gradient descent.
 Inputs:
 - X: A numpy array of shape (N, D) containing training data; there are N
  training samples each of dimension D.
 - y: A numpy array of shape (N,) containing training labels; y[i] = c
   means that X[i] has label 0 \le c \le C for C classes.
 - learning rate: (float) learning rate for optimization.
 - num iters: (integer) number of steps to take when optimizing
 - batch size: (integer) number of training examples to use at each step.
 - verbose: (boolean) If true, print progress during optimization.
 Outputs:
 A list containing the value of the loss function at each training iteration.
 num train, dim = X.shape
 num classes = np.max(y) + 1 # assume y takes values 0...K-1 where K is number of classes
 self.init weights (dims=[np.max(y) + 1, X.shape[1]]) # initializes the weights of sel
 # Run stochastic gradient descent to optimize W
 loss history = []
 for it in np.arange(num iters):
   X batch = None
   y batch = None
   # YOUR CODE HERE:
     Sample batch size elements from the training data for use in
       gradient descent. After sampling,
        - X batch should have shape: (dim, batch size)
       - y batch should have shape: (batch size,)
     The indices should be randomly generated to reduce correlations
     in the dataset. Use np.random.choice. It's okay to sample with
     replacement.
   # ----- #
   idxs = np.random.choice(num train, batch size, replace=True)
   X \text{ batch} = X[idxs]
   y \text{ batch} = y[idxs]
   # END YOUR CODE HERE
```

```
# evaluate loss and gradient
  loss, grad = self.fast loss and grad(X batch, y batch)
  loss history.append(loss)
  # ----- #
  # YOUR CODE HERE:
    Update the parameters, self. W, with a gradient step
  # ----- #
  self.W -= learning rate * grad
  # END YOUR CODE HERE
  # ----- #
  if verbose and it % 100 == 0:
   print('iteration {} / {}: loss {}'.format(it, num iters, loss))
 return loss history
def predict(self, X):
 11 11 11
 Inputs:
 - X: N x D array of training data. Each row is a D-dimensional point.
 Returns:
 - y_pred: Predicted labels for the data in X. y pred is a 1-dimensional
 array of length N, and each element is an integer giving the predicted
  class.
 y pred = np.zeros(X.shape[1])
 # ----- #
 # YOUR CODE HERE:
  Predict the labels given the training data.
 # ============ #
 y pred = np.argmax(np.matmul(X, self.W.T), axis=1)
 # ----- #
 # END YOUR CODE HERE
 return y pred
```