k-Nearest Neighbor (kNN) exercise

Complete and hand in this completed worksheet (including its outputs and any supporting code outside of the worksheet) with your assignment submission. For more details see the <u>assignments page</u> (http://vision.stanford.edu/teaching/cs231n/assignments.html) on the course website.

The kNN classifier consists of two stages:

- During training, the classifier takes the training data and simply remembers it
- During testing, kNN classifies every test image by comparing to all training images and transfering the labels of the k most similar training examples
- · The value of k is cross-validated

In this exercise you will implement these steps and understand the basic Image Classification pipeline, cross-validation, and gain proficiency in writing efficient, vectorized code.

```
In [1]:
        # Run some setup code for this notebook.
        import random
        import numpy as np
        from cs231n.data utils import load CIFAR10
        import matplotlib.pyplot as plt
        from __future__ import print_function
        # This is a bit of magic to make matplotlib figures appear inline in the noteb
        # rather than in a new window.
        %matplotlib inline
        plt.rcParams['figure.figsize'] = (10.0, 8.0) # set default size of plots
        plt.rcParams['image.interpolation'] = 'nearest'
        plt.rcParams['image.cmap'] = 'gray'
        # Some more magic so that the notebook will reload external python modules;
        # see http://stackoverflow.com/questions/1907993/autoreload-of-modules-in-ipyt
        hon
        %load ext autoreload
        %autoreload 2
```

```
In [2]: # Load the raw CIFAR-10 data.
        cifar10 dir = 'cs231n/datasets/cifar-10-batches-py'
        # Cleaning up variables to prevent loading data multiple times (which may caus
        e memory issue)
        try:
           del X_train, y_train
           del X test, y test
           print('Clear previously loaded data.')
        except:
           pass
        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 [3]: # 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', 'shi
        p', 'truck']
        num classes = len(classes)
        samples_per_class = 7
        for y, cls in enumerate(classes):
            idxs = np.flatnonzero(y train == y)
            idxs = np.random.choice(idxs, samples_per_class, replace=False)
            for i, idx in enumerate(idxs):
                plt_idx = i * num_classes + y + 1
                 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()
```



```
In [4]: # 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]
```

```
In [5]: # 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)

```
In [6]: from cs231n.classifiers import KNearestNeighbor

# Create a kNN classifier instance.
# Remember that training a kNN classifier is a noop:
# the Classifier simply remembers the data and does no further processing classifier = KNearestNeighbor()
    classifier.train(X_train, y_train)
```

We would now like to classify the test data with the kNN classifier. Recall that we can break down this process into two steps:

- 1. First we must compute the distances between all test examples and all train examples.
- 2. Given these distances, for each test example we find the k nearest examples and have them vote for the label

Lets begin with computing the distance matrix between all training and test examples. For example, if there are **Ntr** training examples and **Nte** test examples, this stage should result in a **Nte** x **Ntr** matrix where each element (i,j) is the distance between the i-th test and j-th train example.

First, open cs231n/classifiers/k_nearest_neighbor.py and implement the function compute_distances_two_loops that uses a (very inefficient) double loop over all pairs of (test, train) examples and computes the distance matrix one element at a time.

```
In [7]: # Open cs231n/classifiers/k nearest neighbor.py and implement
         # compute distances two loops.
         # Test your implementation:
         dists = classifier.compute distances two loops(X test)
         print(dists.shape)
         (500, 5000)
        # We can visualize the distance matrix: each row is a single test example and
In [8]:
         # its distances to training examples
         plt.imshow(dists, interpolation='none')
         plt.show()
           0
          250
          500
                    1000
                              2000
                                       3000
                                                4000
                                                         5000
```

Inline Question #1: Notice the structured patterns in the distance matrix, where some rows or columns are visible brighter. (Note that with the default color scheme black indicates low distances while white indicates high distances.)

- What in the data is the cause behind the distinctly bright rows?
- · What causes the columns?

Your Answer: fill this in.

```
In [9]: # Now implement the function predict_labels and run the code below:
    # We use k = 1 (which is Nearest Neighbor).
    y_test_pred = classifier.predict_labels(dists, k=1)

# Compute and print the fraction of correctly predicted examples
    num_correct = np.sum(y_test_pred == y_test)
    accuracy = float(num_correct) / num_test
    print('Got %d / %d correct => accuracy: %f' % (num_correct, num_test, accuracy))

Got 137 / 500 correct => accuracy: 0.274000
```

You should expect to see approximately 27% accuracy. Now lets try out a larger k, say k = 5:

```
In [10]: y_test_pred = classifier.predict_labels(dists, k=5)
    num_correct = np.sum(y_test_pred == y_test)
    accuracy = float(num_correct) / num_test
    print('Got %d / %d correct => accuracy: %f' % (num_correct, num_test, accuracy))

Got 139 / 500 correct => accuracy: 0.278000
```

You should expect to see a slightly better performance than with k = 1.

Inline Question 2 We can also other distance metrics such as L1 distance. The performance of a Nearest Neighbor classifier that uses L1 distance will not change if (Select all that apply.):

- 1. The data is preprocessed by subtracting the mean.
- 2. The data is preprocessed by subtracting the mean and dividing by the standard deviation.
- 3. The coordinate axes for the data are rotated.
- 4. None of the above.

Your Answer:

Your explanation:

```
In [11]: # Now lets speed up distance matrix computation by using partial vectorization
         # with one loop. Implement the function compute_distances_one_loop and run the
         # code below:
         dists one = classifier.compute distances one loop(X test)
         # To ensure that our vectorized implementation is correct, we make sure that i
         # agrees with the naive implementation. There are many ways to decide whether
         # two matrices are similar; one of the simplest is the Frobenius norm. In case
         # you haven't seen it before, the Frobenius norm of two matrices is the square
         # root of the squared sum of differences of all elements; in other words, resh
         # the matrices into vectors and compute the Euclidean distance between them.
         difference = np.linalg.norm(dists - dists_one, ord='fro')
         print('Difference was: %f' % (difference, ))
         if difference < 0.001:</pre>
             print('Good! The distance matrices are the same')
         else:
             print('Uh-oh! The distance matrices are different')
```

Difference was: 0.000000 Good! The distance matrices are the same

```
In [12]: # Now implement the fully vectorized version inside compute_distances_no_loops
# and run the code
dists_two = classifier.compute_distances_no_loops(X_test)

# check that the distance matrix agrees with the one we computed before:
difference = np.linalg.norm(dists - dists_two, ord='fro')
print('Difference was: %f' % (difference, ))
if difference < 0.001:
    print('Good! The distance matrices are the same')
else:
    print('Uh-oh! The distance matrices are different')</pre>
```

X2: (500,) Xtrain_shape: (5000,) XtrainXtes_shape (500, 5000) Difference was: 0.000000 Good! The distance matrices are the same

```
In [13]: # Let's compare how fast the implementations are
         def time function(f, *args):
             Call a function f with args and return the time (in seconds) that it took
          to execute.
             import time
             tic = time.time()
             f(*args)
             toc = time.time()
             return toc - tic
         two_loop_time = time_function(classifier.compute_distances_two_loops, X_test)
         print('Two loop version took %f seconds' % two_loop_time)
         one_loop_time = time_function(classifier.compute_distances_one_loop, X_test)
         print('One loop version took %f seconds' % one loop time)
         no_loop_time = time_function(classifier.compute_distances_no_loops, X_test)
         print('No loop version took %f seconds' % no_loop_time)
         # you should see significantly faster performance with the fully vectorized im
         plementation
```

```
Two loop version took 55.754738 seconds
One loop version took 167.268579 seconds
X2: (500,) Xtrain_shape: (5000,) XtrainXtes_shape (500, 5000)
No loop version took 0.946822 seconds
```

Cross-validation

We have implemented the k-Nearest Neighbor classifier but we set the value k = 5 arbitrarily. We will now determine the best value of this hyperparameter with cross-validation.

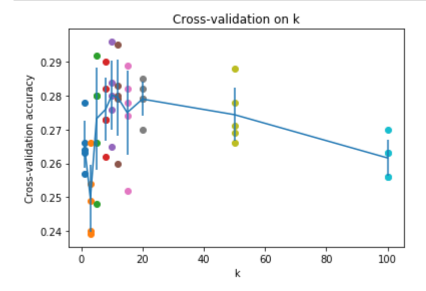
```
In [55]:
      num folds = 5
       k choices = [1, 3, 5, 8, 10, 12, 15, 20, 50, 100]
       X train folds = []
       y train folds = []
       # TODO:
       # Split up the training data into folds. After splitting, X_train_folds and
       # y_train_folds should each be lists of length num_folds, where
       # y train folds[i] is the label vector for the points in X train folds[i].
       # Hint: Look up the numpy array_split function.
       ##
       # Your code
       X train folds=np.split(X train, num folds)# split array to list
       y_train_folds=np.split(y_train, num_folds)
       ##
       #
                                END OF YOUR CODE
       #
       ##
       # A dictionary holding the accuracies for different values of k that we find
       # when running cross-validation. After running cross-validation,
       # k to accuracies[k] should be a list of length num folds giving the different
       # accuracy values that we found when using that value of k.
       k to accuracies = {}
       for i in k choices:
          k to accuracies[i]=[]
       ##
       # TODO:
       #
       # Perform k-fold cross validation to find the best value of k. For each
       # possible value of k, run the k-nearest-neighbor algorithm num folds times,
       # where in each case you use all but one of the folds as training data and the
       # last fold as a validation set. Store the accuracies for all fold and all
       # values of k in the k to accuracies dictionary.
       ##
          for val fold in range(num folds):
```

```
#
        Method 1: hornor code
       X_train_crossval=[]
       Y train crossval=[]
       X validation=[]
       Y validation=[]
       X_train_crossval+= [item for t in range(num_folds) if t!= val_fold fo
r item in X train folds[t]]#list
       Y train crossval+= [item for t in range(num folds) if t!= val fold fo
r item in y_train_folds[t]]
       X validation+= [item for item in X train folds[val fold]]
       Y validation+= [item for item in y train folds[val fold]]
         # Method 2: reference- better
#
         X train crossval= np.concatenate([X train folds[i] for i in range (n
um_folds) if i!=val_fold]) # already change to array
         Y train crossval= np.concatenate([y train folds[i] for i in range (n
um folds) if i!=val fold])
         X validation= X train folds[val fold]
         Y_validation= y_train_folds[val_fold]
       classifier = KNearestNeighbor()
       classifier.train(np.array(X train crossval), np.array(Y train crossval
))# remember the data
       #Calculate distance between test and all train data
       dists = classifier.compute_distances_no_loops(np.array(X_validation))
#Method 2, no need to use np.array to transfrom list to array
       # We use k = number of nearest neighbor (which is Nearest Neighbor).
       y test pred cross val = classifier.predict labels(dists, i)
         print ('i',i)
         print ('y test pred cross val.shape',len(y test pred cross val))
         print ('Y_validation.shape',len(Y_validation))
#
         print ('y_test_pred_cross_val',y_test_pred_cross_val)
       # Compute and print the fraction of correctly predicted examples
       num correct = np.sum(y test pred cross val == np.array(Y validation))
         print ('num correct', num correct)
         print ('number of test:',Y_validation.shape[0])
       accuracy = float(num_correct) / np.array(Y_validation).shape[0]
         print('Got %d / %d correct => accuracy: %f' % (num correct, num tes
t, accuracy))
       #Store the value into dictionary
         k to accuracies.update({i:accuracy}) #NOTE: THIS ONLY UPDATE THE OLD
VALUE OF THE KEY TO THE NEWEST, THE KEY ONLY HAVE 1 VALUE
       k to accuracies[i].append(accuracy) #NOTE: EXTEND THE VALUE OF A KEY
         print ('results', k to accuracies)
         print ('X_Train', X_Train[1].shape)
         print ('X Train', len(X Train))
#
##
#
                               END OF YOUR CODE
```

```
# Print out the computed accuracies
for k in sorted(k_to_accuracies):
    for accuracy in k to accuracies[k]:
        print('k = %d, accuracy = %f' % (k, accuracy))
k = 1, accuracy = 0.263000
k = 1, accuracy = 0.257000
k = 1, accuracy = 0.264000
k = 1, accuracy = 0.278000
k = 1, accuracy = 0.266000
k = 3, accuracy = 0.239000
k = 3, accuracy = 0.249000
k = 3, accuracy = 0.240000
k = 3, accuracy = 0.266000
k = 3, accuracy = 0.254000
k = 5, accuracy = 0.248000
k = 5, accuracy = 0.266000
k = 5, accuracy = 0.280000
k = 5, accuracy = 0.292000
k = 5, accuracy = 0.280000
k = 8, accuracy = 0.262000
k = 8, accuracy = 0.282000
k = 8, accuracy = 0.273000
k = 8, accuracy = 0.290000
k = 8, accuracy = 0.273000
k = 10, accuracy = 0.265000
k = 10, accuracy = 0.296000
k = 10, accuracy = 0.276000
k = 10, accuracy = 0.284000
k = 10, accuracy = 0.280000
k = 12, accuracy = 0.260000
k = 12, accuracy = 0.295000
k = 12, accuracy = 0.279000
k = 12, accuracy = 0.283000
k = 12, accuracy = 0.280000
k = 15, accuracy = 0.252000
k = 15, accuracy = 0.289000
k = 15, accuracy = 0.278000
k = 15, accuracy = 0.282000
k = 15, accuracy = 0.274000
k = 20, accuracy = 0.270000
k = 20, accuracy = 0.279000
k = 20, accuracy = 0.279000
k = 20, accuracy = 0.282000
k = 20, accuracy = 0.285000
k = 50, accuracy = 0.271000
k = 50, accuracy = 0.288000
k = 50, accuracy = 0.278000
k = 50, accuracy = 0.269000
k = 50, accuracy = 0.266000
k = 100, accuracy = 0.256000
k = 100, accuracy = 0.270000
k = 100, accuracy = 0.263000
k = 100, accuracy = 0.256000
k = 100, accuracy = 0.263000
```

```
In [56]: # plot the raw observations
    for k in k_choices:
        accuracies = k_to_accuracies[k]
        plt.scatter([k] * len(accuracies), accuracies)

# plot the trend line with error bars that correspond to standard deviation
    accuracies_mean = np.array([np.mean(v) for k,v in sorted(k_to_accuracies.items
        ())])
    accuracies_std = np.array([np.std(v) for k,v in sorted(k_to_accuracies.items
        ())])
    plt.errorbar(k_choices, accuracies_mean, yerr=accuracies_std)
    plt.title('Cross-validation on k')
    plt.xlabel('k')
    plt.ylabel('Cross-validation accuracy')
    plt.show()
```



```
In [57]: # Based on the cross-validation results above, choose the best value for k,
    # retrain the classifier using all the training data, and test it on the test
    # data. You should be able to get above 28% accuracy on the test data.
    best_k = 10

classifier = KNearestNeighbor()
    classifier.train(X_train, y_train)
    y_test_pred = classifier.predict(X_test, k=best_k)

# Compute and display the accuracy
    num_correct = np.sum(y_test_pred == y_test)
    accuracy = float(num_correct) / num_test
    print('Got %d / %d correct => accuracy: %f' % (num_correct, num_test, accuracy))
```

Got 141 / 500 correct => accuracy: 0.282000

Inline Question 3 Which of the following statements about k-Nearest Neighbor (k-NN) are true in a classification setting, and for all k? Select all that apply.

- 1. The training error of a 1-NN will always be better than that of 5-NN.
- 2. The test error of a 1-NN will always be better than that of a 5-NN.
- 3. The decision boundary of the k-NN classifier is linear.
- 4. The time needed to classify a test example with the k-NN classifier grows with the size of the training set.
- 5. None of the above.

Your Ai	nswer
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Your explanation: