Project1

November 11, 2024

1 Project 1

The following packages are used in this project

```
[1]: import numpy as np
from scipy.sparse import csr_matrix
import scipy.sparse.linalg as la
import matplotlib.pyplot as plt
import seaborn as sns
import scipy
import time
```

1. Write a function that loads this file and returns the matrix X and vector y. How many malicious data points are there? What can you say about the sparsity of the data? Do you think it makes sense to use one-hot-coding for some of the columns?

we use the following code to loads the data file, returns the matrix X and vector y, and count the number of malicious data points:

```
class DataLoader:
    def __init__(self, file_path= r".\data\data.csv"):
        self.X = None
        self.y = None
        self.num = 0

        self._read(file_path)

def __read(self, file_path):
        ori_data = np.loadtxt(file_path, skiprows=1, delimiter=',', dtype=int)
        self.X = ori_data[:,:-1]
        self.y = ori_data[:,-1]
        self.y[self.y == 0] = -1
        self.num = self.X.shape[0]

def split(self, ratio = 0.7):
    # the number of training data
        train_num = int(self.num * ratio)
```

```
# shuffling the original data
rand_idx = np.arange(self.num)
np.random.shuffle(rand_idx)
X = self.X[rand_idx]
y = self.y[rand_idx]

# split the data set
X_train, X_test = X[:train_num], X[train_num:]
y_train, y_test = y[:train_num], y[train_num:]

return X_train, X_test, y_train, y_test

mydata = DataLoader(r".\data\data.csv")
print(f'there are {(mydata.y == 1).sum()} \
malicious data points in the total {mydata.num} data points')
```

there are 14700 malicious data points in the total 29332 data points

From summing the ones in the data, we get that there are 14700 malicious data points in the total 29332 data points.

The dataset is highly sparse, as most feature values are zero.

For onw-hot encoding, it is reasonable to apply it when the features represent categorical data without intrinsic ordering. However, for binary features in our project, one-hot encoding may be redundant unless further categorization is needed.

2. Write a function that splits the data into a training and test set according to some fraction 0 < r < 1. Make sure to use randomization; that is, it should not be the case that the training set consists of the first data points and the test set of the remaining data points. Your function should return matrices X train and X test and vectors y train and y test.

We used the split method from the DataLoader class shown in Question 1 to divide the dataset according to the given ratio, where shuffling is implemented before spliting to avoid the case that the training set consists of the first data points and the test set of the remaining data points.

```
[3]: np.random.seed(1234)
X_train, X_test, y_train, y_test = mydata.split(ratio = 0.5)
```

3. Write a function that, given the matrix X, the vector y, and a weight vector w defining a hyperplane, returns the number of correctly classified points. Verify that the output makes sense for random weight vectors.

We used a randomly generated weight vector $\mathbf{w}_{\mathtt{init}}$ to define a hyperplane. The following function is used to compute predictions and the number of correctly classified points for the given input matrix X, output yand weight vector \mathbf{w} .

```
[4]: def evaluator(X: np.ndarray, y: np.ndarray, w: np.ndarray):
    y_pred = X.dot(w)
    y_pred[y_pred < 0] = -1</pre>
```

```
y_pred[y_pred >= 0] = 1
return (y_pred == y).sum()

np.random.seed(1234)
w_init = np.random.uniform(low = -1, high = 1, size=(mydata.X.shape[1]))
pred_corr_num = evaluator(mydata.X, mydata.y, w_init)
print(f'{pred_corr_num} data points in total {mydata.num} samples are correct_u
susing random generated weights')
```

11754 data points in total 29332 samples are correct using random generated weights

4. Consider the cost function for logistic regression as defined in the lectures. Write down a symbolic formula for the gradient of this function.

The cost function J(w) for logistic regression with regularization is defined as:

$$J(w) = \sum_{i=1}^n L(y_i x_i^T w) + \frac{\lambda}{2} \|w\|^2$$

where:

$$L(s) = \log\left(1 + e^{-s}\right)$$

The gradient of the loss term for a single data point x_i with respect to w is:

$$\nabla_w L(y_i x_i^T w) = -\frac{y_i x_i}{1 + e^{y_i x_i^T w}}$$

Therefore, the gradient of the total loss term is:

$$\sum_{i=1}^n \nabla_w L(y_i x_i^T w) = -\sum_{i=1}^n \frac{y_i x_i}{1 + e^{y_i x_i^T w}}$$

The gradient of the regularization term $\frac{\lambda}{2} ||w||^2$ with respect to w is:

$$\nabla_w \left(\frac{\lambda}{2} \|w\|^2 \right) = \lambda w$$

Thus, the gradient of J(w) with respect to w is:

$$\nabla_w J(w) = -\sum_{i=1}^n \frac{y_i x_i}{1 + e^{y_i x_i^T w}} + \lambda w$$

5. • Write a straightforward implementation for logistic regression using gradient descent with a fixed step size α . Your function should take as arguments the data matrix X and data vector y, the step size α , the regularization constant

 λ , and an integer K indicating the number of gradient descent steps. The function should return a weight vector w.

- Experiment with the hyperparameters, using dense and sparse linear algebra, on random splits of training and test data sets. (If you know about writing allocation-free code, you can also experiment with this.)
- Given for instance a 50/50 split between test and training data, what is the best classification performance you can obtain on the test set?

In this logistic regression implementation, the constants $\alpha = 0.01$ (step size), $\lambda = 0.01$ (regularization constant), and K = 100 (gradient descent steps) are chosen initially to test our code.

We implement our code in 3 different ways: 1. The logistic_regression_entrywise method calculates the gradient individually for each data point in every iteration.

- 2. The logistic_regression_vectorize method utilizes vectorizing and broadcasting to avoid explicitly calculating the gradient for each data point individually, show as below:
 - s = y * (X @ w)
 z = y / (1 + np.exp(s))
 w_grad = (X.T @ z) + lambda_ * w
- 3. The logistic_regression_sparse method is further optimized by converting training data to a CSR type sparse matrix and using sparse linear algebra, due to the sparsity in our training data.

The code implement above is shown as below:

```
[5]: class LogisticClassifier:
         def __init__(self, X_train, y_train, w_init):
             self.X_train = X_train
             self.y_train = y_train
             self.w_init = w_init
             self.w = None
         def logistic_regression(self, K, alpha, lambda_reg, flag = 0):
             if flag == 0:
                 self.logistic_regression_entrywise(K, alpha, lambda_reg)
             if flag == 1:
                 self.logistic_regression_vectorize(K, alpha, lambda_reg)
             if flag == 2:
                 self.logistic_regression_sparse(K, alpha, lambda_reg)
         def logistic_regression_entrywise(self, K, alpha, lambda_reg):
             self.w = self.w_init
             for k in range(K):
                 w_grad = lambda_reg * self.w
                 for i in range(self.X train.shape[0]):
                     # if k == 5:
                          print(i)
                     w_grad -=(self.y_train[i] * self.X_train[i])\
                     /(1 + np.exp(self.y_train[i] * self.X_train[i].dot(self.w)))
```

```
self.w -= w_grad * alpha
    return self.w
def logistic regression_vectorize(self, K, alpha, lambda_reg):
    self.w = self.w_init
   for _ in range(K):
        s = self.y_train[:,None] * self.X_train @ self.w
        z = self.y_train / (1 + np.exp(s))
        w_grad = - self.X_train.T @ z + lambda_reg * self.w
        self.w -= w_grad * alpha
    return self.w
def logistic_regression_sparse(self, K, alpha, lambda_reg):
   self.w = self.w_init
   self.X_train = csr_matrix(self.X_train)
    for _ in range(K):
        s_product = self.X_train @ self.w
        s = self.y_train * s_product
        z = self.y_train / (1 + np.exp(s))
        w_grad = - self.X_train.T @ z + lambda_reg * self.w
        self.w -= w_grad * alpha
   return self.w
def predict(self, X_test, y_test):
   w = self.w
   y_true = evaluator(X_test, y_test, w)
    acc = y_true / y_test.shape[0]
   return acc
```

Entry-Wise Implement:

```
Accuracy on the test set: 94.81%, the time cost is 14.146091222763062s
```

Dense Implement:

```
Accuracy on the test set: 94.74%, the time cost is 2.2567808628082275s
```

Sparse Implement:

```
Accuracy on the test set: 95.30%, the time cost is 0.31095409393310547s
```

From the above experiment, we have 1. The three implements achieve similar accuracy with different runing times.

- 2. The logistic_regression_entrywise method is quite slow for large datasets because of the iterative sample-by-sample gradient computation, taking approximately 14.15 seconds for 100 iteration.
- 3. The logistic_regression_vectorize method reduces computation time to roughly 2.26 second for 100 iteration.
- 4. The logistic_regression_sparse method is the fastest with a rapid runtime of 0.31 seconds for 100 iteration, making it ideal for large, sparse datasets by reducing unnecessary calculations.

By experimenting on hyperparameters, the best classification performance we can obtain the best accuracy of 95.58% in 1000 iterations on the test set with a 50/50 split with $\alpha = 0.03$ and $\lambda = 0.1$ among hyperparameters we tested.

C:\Users\11787\AppData\Local\Temp\ipykernel_6352\3012149010.py:43:
RuntimeWarning: overflow encountered in exp

```
z = self.y_train / (1 + np.exp(s))
best_acc = 0.9558161734624301, best_alpha = 0.03, best_lambda_reg = 0.1
```

6. Download the file data2.csv. This is the same as the previous data file, except that 2000 fake data points have been appended to the data set. Use the singular value transform as explained in the lecture to detect and remove most of these outliers without removing too many other data points. (For the singular value transform you can use a library function, you do not have to implement this yourself.)

We use the following function to whitening our data:

```
[8]: def data_whitening(X, k):
    # Centering
    M = X - X.mean(axis=0)

# SVD Decomposition
    U, S, _ = la.svds(M, k=k)

# Whitening
    Xw = U[:,::-1] @ scipy.sparse.diags(S[::-1])
    Xw = Xw / np.std(Xw, axis=0)[None,:]

return Xw
```

Then we compute the norm of whitenning data to detect the outliers in our data, Within the noise_removal function, we employed the scipy.sparse.linalg.svds method to compute the top k singular values and their corresponding singular vectors:

```
[9]: def noise_removal(X, k, percentile = 0.9, vis_flag = False):
    # Whitening
    Xw = data_whitening(X, k)

    norm_list = np.array([np.linalg.norm(Xw[i, :]) for i in range(Xw.shape[0])])
    outlier_mask = norm_list >= np.quantile(norm_list, percentile)

    return outlier_mask
```

Let - TP (True positive) = number of fake samples be detected - FN (False Negative) = number of fake samples be identified as true data points - TN (True Negative) = number of true data points be correctly identified - FP (False Positive) = number of true data point be wrongly identified as fake samples.

We use Precision, Recall and F1-score to assess the performance of our method

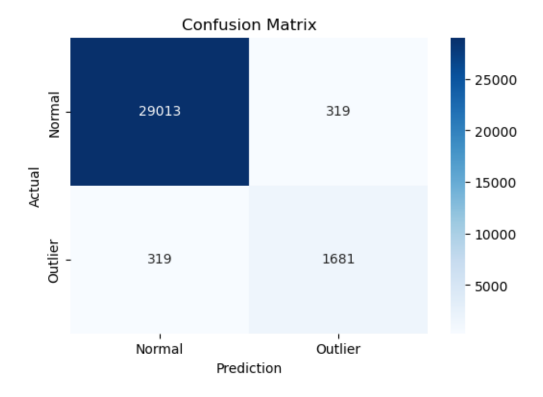
```
if vis_flag:
       visualize_confusion(*confusion_matrix)
       print(f'Accuracy: {acc:.4f}')
       print(f"Precision: {P:.4f}")
       print(f"Recall: {R:.4f}")
       print(f"F1-score: {F1:.4f}")
   return F1, (P, R), acc
def F1_score(outlier_mask, outlier_num = 2000):
   The outlier is in the last #outlier num rows.
   11 11 11
   total_num = len(outlier_mask)
   TP = (outlier_mask[-outlier_num:].sum())
   FP = (outlier_mask[:-outlier_num].sum())
   TN = total_num - outlier_num - FP
   FN = outlier num - TP
   Precision = TP / (TP + FP)
   Recall = TP / (TP + FN)
   →Recall) > 0 else 0
   return (TP, FP, TN, FN), (Precision, Recall), F1_score
def visualize_confusion(TP, FP, TN, FN):
   confusion_matrix = np.array([[TN, FP],
                               [FN, TP]])
   categories = ['Normal', 'Outlier']
   # plot confusion matrix
   plt.figure(figsize=(6, 4))
   sns.heatmap(confusion_matrix, annot=True, fmt='d', cmap='Blues',
               xticklabels=categories, yticklabels=categories)
   plt.xlabel('Prediction')
   plt.ylabel('Actual')
   plt.title('Confusion Matrix')
   plt.show()
```

The we have the following result:

```
[11]: # read data
my_data = DataLoader(r".\data\data2.csv")
X = my_data.X.astype('float64')

# Outlier rate
q = 1 - 2000 / X.shape[0]

# data whitening & noise removal
F1, _, _ = assess_performance(noise_removal(X, 81, q), vis_flag=True)
```



Accuracy: 0.9796 Precision: 0.8405 Recall: 0.8405 F1-score: 0.8405

In the plot of confusion matrix, we found that: - The diagonal elements (29,013 normal points and 1,681 outliers) indicate correctly classified instances.

- Off-diagonal elements (319 misclassified as normal and 319 asoutliers) indicate errors
- Majority of outliers was removed and only a few truer data point was misclassifieds.

By adjusting the threshold for classifying a data point as an outlier or not, we obtain different classification results, focusing either on detecting more true positives (TP) or on detecting true positives with greater accuracy. The precision-recall curve is shown below

```
[14]: def plot_PRC(P_list, R_list):
          # data clean
          valid_indices = (~np.isnan(P_list)) & (~np.isnan(R_list)) & (~np.
       →isinf(P_list)) & (~np.isinf(R_list))
          P_array = P_list[valid_indices]
          R_array = R_list[valid_indices]
          # sort data
          sorted_indices = np.argsort(R_array)
          R_sorted = R_array[sorted_indices]
          P_sorted = P_array[sorted_indices]
          # plot data
          plt.figure(figsize=(8, 6))
          plt.plot(R_sorted, P_sorted, linestyle='-', color='b')
          plt.xlabel('Recall', fontsize=12)
          plt.ylabel('Precision', fontsize=12)
          plt.title('Precision-Recall Curve', fontsize=14)
          plt.grid(True)
          plt.show()
      P list = []
      R_list = []
      for q in np.arange(0, 1, 0.01):
          print(f"Progress: {q * 100:.0f}, (q = {q:.2f})", end='\r')
          _, (P, R), _ = assess_performance(noise_removal(X, 81, q), vis_flag=False)
          P_list.append(P)
          R_list.append(R)
      # plot PR-Curve
      plot_PRC(np.array(P_list), np.array(R_list))
```

Progress: 99% (q = 0.99)

