Anomaly Detection

In this exercise, you will implement the anomaly detection algorithm and apply it to detect failing servers on a network.

Outline

- 1 Packages
- 2 Anomaly detection
 - 2.1 Problem Statement
 - 2.2 Dataset
 - 2.3 Gaussian distribution
 - Exercise 1
 - o Exercise 2
 - 2.4 High dimensional dataset

1 - Packages

First, let's run the cell below to import all the packages that you will need during this assignment.

- numpy is the fundamental package for working with matrices in Python.
- matplotlib is a famous library to plot graphs in Python.
- utils.py contains helper functions for this assignment. You do not need to modify code in this file.

```
In [1]: import numpy as np
import matplotlib.pyplot as plt
from utils import *

%matplotlib inline
```

2 - Anomaly detection

2.1 Problem Statement

In this exercise, you will implement an anomaly detection algorithm to detect anomalous behavior in server computers.

The dataset contains two features -

- throughput (mb/s) and
- latency (ms) of response of each server.

While your servers were operating, you collected m=307 examples of how they were behaving, and thus have an unlabeled dataset $\{x^{(1)},\dots,x^{(m)}\}$.

• You suspect that the vast majority of these examples are "normal" (non-anomalous) examples of the servers operating normally, but there might also be some examples of servers acting anomalously within this dataset.

You will use a Gaussian model to detect anomalous examples in your dataset.

- You will first start on a 2D dataset that will allow you to visualize what the algorithm is doing.
- On that dataset you will fit a Gaussian distribution and then find values that have very low probability and hence can be considered anomalies.
- After that, you will apply the anomaly detection algorithm to a larger dataset with many dimensions.

2.2 Dataset

You will start by loading the dataset for this task.

- The load_data() function shown below loads the data into the variables
 X_train, X_val and y_val
 - You will use X_train to fit a Gaussian distribution
 - You will use X_val and y_val as a cross validation set to select a threshold and determine anomalous vs normal examples

```
In [2]: # Load the dataset
X_train, X_val, y_val = load_data()
```

View the variables

Let's get more familiar with your dataset.

A good place to start is to just print out each variable and see what it contains.

The code below prints the first five elements of each of the variables

```
In [3]: # Display the first five elements of X_train
print("The first 5 elements of X_train are:\n", X_train[:5])
```

```
The first 5 elements of X train are:
        [[13.04681517 14.74115241]
        [13.40852019 13.7632696 ]
        [14.19591481 15.85318113]
        [14.91470077 16.17425987]
        [13.57669961 14.04284944]]
In [4]: # Display the first five elements of X_val
        print("The first 5 elements of X_val are\n", X_val[:5])
       The first 5 elements of X_val are
        [[15.79025979 14.9210243 ]
        [13.63961877 15.32995521]
        [14.86589943 16.47386514]
        [13.58467605 13.98930611]
        [13.46404167 15.63533011]]
In [5]: # Display the first five elements of y val
        print("The first 5 elements of y_val are\n", y_val[:5])
       The first 5 elements of y val are
        [0 0 0 0 0]
```

Check the dimensions of your variables

Another useful way to get familiar with your data is to view its dimensions.

The code below prints the shape of X_train, X_val and y_val.

```
In [6]: print ('The shape of X_train is:', X_train.shape)
    print ('The shape of X_val is:', X_val.shape)
    print ('The shape of y_val is: ', y_val.shape)

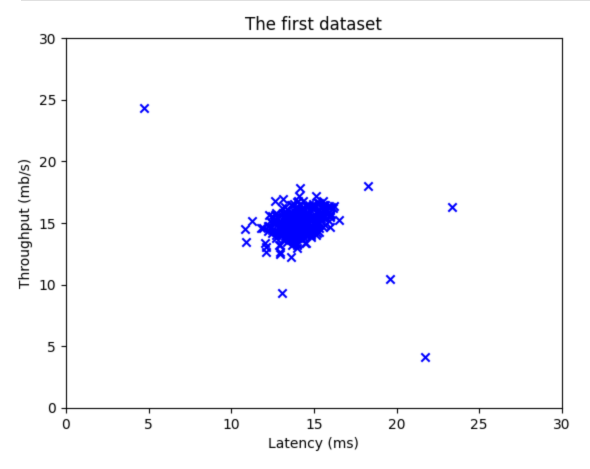
The shape of X_train is: (307, 2)
    The shape of X_val is: (307, 2)
    The shape of y_val is: (307,)
```

Visualize your data

Before starting on any task, it is often useful to understand the data by visualizing it.

- For this dataset, you can use a scatter plot to visualize the data (X_train), since it has only two properties to plot (throughput and latency)
- Your plot should look similar to the one below
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```
plt.ylabel('Throughput (mb/s)')
# Set the x-axis label
plt.xlabel('Latency (ms)')
# Set axis range
plt.axis([0, 30, 0, 30])
plt.show()
```



2.3 Gaussian distribution

To perform anomaly detection, you will first need to fit a model to the data's distribution.

- Given a training set $\{x^{(1)}, \dots, x^{(m)}\}$ you want to estimate the Gaussian distribution for each of the features x_i .
- Recall that the Gaussian distribution is given by

$$p(x;\mu,\sigma^2) = rac{1}{\sqrt{2\pi\sigma^2}} \mathrm{exp}^{-rac{(x-\mu)^2}{2\sigma^2}}$$

where μ is the mean and σ^2 is the variance.

• For each feature $i=1\dots n$, you need to find parameters μ_i and σ_i^2 that fit the data in the i-th dimension $\{x_i^{(1)},\dots,x_i^{(m)}\}$ (the i-th dimension of each example).

2.3.1 Estimating parameters for a Gaussian distribution

Implementation:

Your task is to complete the code in estimate_gaussian below.

Exercise 1

Complete the estimate_gaussian function below to calculate mu (mean for each feature in X) and var (variance for each feature in X).

You can estimate the parameters, (μ_i, σ_i^2) , of the i-th feature by using the following equations. To estimate the mean, you will use:

$$\mu_i = rac{1}{m} \sum_{j=1}^m x_i^{(j)}$$

and for the variance you will use:

$$\sigma_i^2 = rac{1}{m} \sum_{j=1}^m (x_i^{(j)} - \mu_i)^2$$

You can check if your implementation is correct by running the following test code:

```
In [9]: # Estimate mean and variance of each feature
mu, var = estimate_gaussian(X_train)

print("Mean of each feature:", mu)
print("Variance of each feature:", var)

# UNIT TEST
from public_tests import *
estimate_gaussian_test(estimate_gaussian)
```

Mean of each feature: [14.11222578 14.99771051] Variance of each feature: [1.83263141 1.70974533] All tests passed!

Expected Output:

Mean of each feature: [14.11222578 14.99771051]

Variance of each feature: [1.83263141 1.70974533]

Now that you have completed the code in estimate_gaussian, we will visualize the contours of the fitted Gaussian distribution.

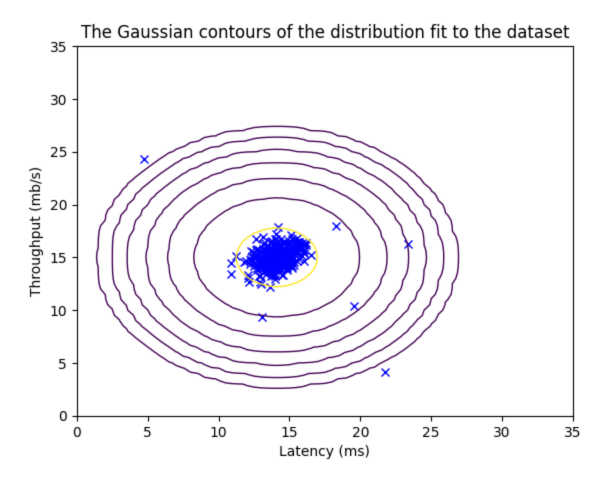
You should get a plot similar to the figure below.

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From your plot you can see that most of the examples are in the region with the highest probability, while the anomalous examples are in the regions with lower probabilities.

```
In [10]: # Returns the density of the multivariate normal
    # at each data point (row) of X_train
    p = multivariate_gaussian(X_train, mu, var)

#Plotting code
visualize_fit(X_train, mu, var)
```



2.3.2 Selecting the threshold ϵ

Now that you have estimated the Gaussian parameters, you can investigate which examples have a very high probability given this distribution and which examples have a very low probability.

- The low probability examples are more likely to be the anomalies in our dataset.
- One way to determine which examples are anomalies is to select a threshold based on a cross validation set.

In this section, you will complete the code in select_threshold to select the threshold ε using the F_1 score on a cross validation set.

- For this, we will use a cross validation set $\{(x_{\rm cv}^{(1)},y_{\rm cv}^{(1)}),\ldots,(x_{\rm cv}^{(m_{\rm cv})},y_{\rm cv}^{(m_{\rm cv})})\}$, where the label y=1 corresponds to an anomalous example, and y=0 corresponds to a normal example.
- For each cross validation example, we will compute $p(x_{ ext{cv}}^{(i)})$. The vector of all of these probabilities $p(x_{ ext{cv}}^{(1)}),\dots,p(x_{ ext{cv}}^{(m_{ ext{cv}})})$ is passed to <code>select_threshold</code> in the vector <code>p_val</code>.
- The corresponding labels $y_{\rm cv}^{(1)},\dots,y_{\rm cv}^{(m_{\rm cv})}$ are passed to the same function in the vector <code>y_val</code> .

Exercise 2

Complete the select_threshold function below to find the best threshold to use for selecting outliers based on the results from the validation set (p_val) and the ground truth (y_val).

- In the provided code select_threshold, there is already a loop that will try many different values of ε and select the best ε based on the F_1 score.
- You need to implement code to calculate the F1 score from choosing epsilon as the threshold and place the value in F1.
 - lacktriangledown Recall that if an example x has a low probability p(x)<arepsilon, then it is classified as an anomaly.
 - Then, you can compute precision and recall by:

$$egin{aligned} prec = & & rac{tp}{tp+fp} \ rec = & & rac{tp}{tp+fn}, \end{aligned}$$

where

- \circ tp is the number of true positives: the ground truth label says it's an anomaly and our algorithm correctly classified it as an anomaly.
- \circ fp is the number of false positives: the ground truth label says it's not an anomaly, but our algorithm incorrectly classified it as an anomaly.
- \circ fn is the number of false negatives: the ground truth label says it's an anomaly, but our algorithm incorrectly classified it as not being anomalous.
- The F_1 score is computed using precision (prec) and recall (rec) as follows:

$$F_1 = rac{2 \cdot prec \cdot rec}{prec + rec}$$

Implementation Note: In order to compute tp, fp and fn, you may be able to use a vectorized implementation rather than loop over all the examples.

```
p_val (ndarray): Results on validation set
Returns:
    epsilon (float): Threshold chosen
    F1 (float):
                  F1 score by choosing epsilon as threshold
best_epsilon = 0
best F1 = 0
F1 = 0
step\_size = (max(p\_val) - min(p\_val)) / 1000
for epsilon in np.arange(min(p_val), max(p_val), step_size):
    ### START CODE HERE ###
    predictions = (p_val < epsilon).astype(int) # Classify as anomaly i</pre>
    tp = np.sum((predictions == 1) & (y_val == 1)) # True positives
    fp = np.sum((predictions == 1) & (y_val == 0)) # False positives
    fn = np.sum((predictions == 0) & (y_val == 1)) # False negatives
    if tp + fp == 0 or tp + fn == 0:
        continue # Avoid division by zero
    precision = tp / (tp + fp)
    recall = tp / (tp + fn)
    F1 = (2 * precision * recall) / (precision + recall) # Compute F1 s
    ### END CODE HERE ###
    if F1 > best_F1:
        best F1 = F1
        best epsilon = epsilon
return best_epsilon, best_F1
```

You can check your implementation using the code below

All tests passed!

Expected Output:

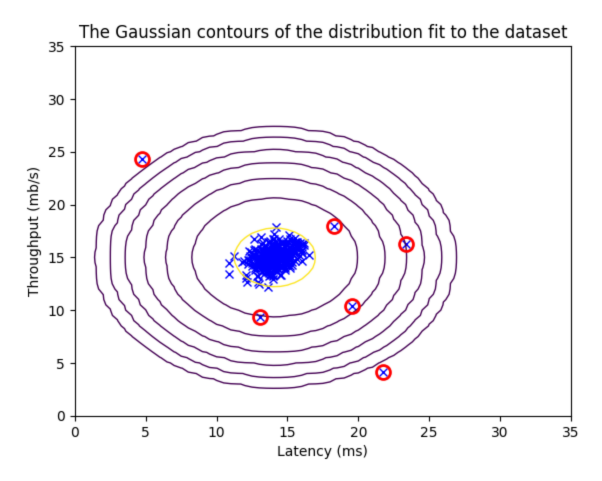
Best epsilon found using cross-validation: 8.99e-05

Best F1 on Cross Validation Set: 0.875

Now we will run your anomaly detection code and circle the anomalies in the plot (Figure 3 below).

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Out[13]: [<matplotlib.lines.Line2D at 0x772d7d25dca0>]



2.4 High dimensional dataset

Now, we will run the anomaly detection algorithm that you implemented on a more realistic and much harder dataset.

In this dataset, each example is described by 11 features, capturing many more properties of your compute servers.

Let's start by loading the dataset.

- The load_data() function shown below loads the data into variables
 X_train_high, X_val_high and y_val_high
 - _high is meant to distinguish these variables from the ones used in the previous part
 - We will use X_train_high to fit Gaussian distribution
 - We will use X_val_high and y_val_high as a cross validation set to select a threshold and determine anomalous vs normal examples

```
In [14]: # load the dataset
    X_train_high, X_val_high, y_val_high = load_data_multi()
```

Check the dimensions of your variables

Let's check the dimensions of these new variables to become familiar with the data

```
In [15]: print ('The shape of X_train_high is:', X_train_high.shape)
    print ('The shape of X_val_high is:', X_val_high.shape)
    print ('The shape of y_val_high is: ', y_val_high.shape)

The shape of X_train_high is: (1000, 11)
    The shape of X_val_high is: (100, 11)
    The shape of y_val_high is: (100,)
```

Anomaly detection

Now, let's run the anomaly detection algorithm on this new dataset.

The code below will use your code to

- Estimate the Gaussian parameters (μ_i and σ_i^2)
- Evaluate the probabilities for both the training data X_train_high from which you estimated the Gaussian parameters, as well as for the the cross-validation set X_val_high.
- Finally, it will use select_threshold to find the best threshold ε .

```
In [16]: # Apply the same steps to the larger dataset

# Estimate the Gaussian parameters
mu_high, var_high = estimate_gaussian(X_train_high)

# Evaluate the probabilites for the training set
p_high = multivariate_gaussian(X_train_high, mu_high, var_high)

# Evaluate the probabilites for the cross validation set
p_val_high = multivariate_gaussian(X_val_high, mu_high, var_high)

# Find the best threshold
epsilon_high, F1_high = select_threshold(y_val_high, p_val_high)

print('Best epsilon found using cross-validation: %e'% epsilon_high)
print('Best F1 on Cross Validation Set: %f'% F1_high)
print('# Anomalies found: %d'% sum(p_high < epsilon_high))</pre>
```

Best epsilon found using cross-validation: 1.377229e-18
Best F1 on Cross Validation Set: 0.615385
Anomalies found: 117

Expected Output:

Best epsilon found using cross-validation: 1.38e-18
Best F1 on Cross Validation Set: 0.615385
anomalies found: 117

```
In []:
```