Module2 USL

July 21, 2022

0.0.1 Grading

This week's lab doesn't have any auto-graded components. Each question in this notebook has an accompanying Peer Review question. Although the lab shows as being ungraded, you need to complete the notebook to answer the Peer Review questions. **DO NOT CHANGE VARIABLE OR METHOD SIGNATURES**

0.0.2 Validate Button

This week's lab doesn't have any auto-graded components. Each question in this notebook has an accompanying Peer Review question. Although the lab shows as being ungraded, you need to complete the notebook to answer the Peer Review questions.

You do not need to use the Validate button for this lab since there are no auto-graded components. If you hit the Validate button, it will time out given the number of visualizations in the notebook. Cells with longer execution times cause the validate button to time out and freeze. This notebook's Validate button time-out does not affect the final submission grading.

1 Clustering RNA sequences to identify cancer types

In this assignment, we will use clustering algorithms on RNA sequence data to identify cancer types. Since the whole data (from Cancer Genome Atlas Pan-Cancer project) is very big, we will use a subset data from UCI Machine Learning repository. The subset data contains only 5 labels; BRCA, KIRC, COAD, LUAD and PRAD. The meanings of those labels are as below.

Abbreviation	Cancer
LUSC	Lung squamous cell carcinoma
READ	Rectum adenocarcinoma
GBM	Glioblastoma multiforme
BLCA	Bladder Urothelial Carcinoma
UCEC	Uterine Corpus Endometrioid Carcinoma
COAD	Colon adenocarcinoma
OV	Ovarian serous cystadenocarcinoma
LAML	Acute Myeloid Leukemia
HNSC	Head and Neck squamous cell carcinoma
LUAD	Lung adenocarcinoma

Abbreviation	Cancer
BRCA	Breast invasive carcinoma
KIRC	Kidney renal clear cell carcinoma

Although we can use the data for supervised learning model training, we will not use these labels for training, but use them for evaluation.

[5]: import pandas as pd

```
import matplotlib.pyplot as plt
      import numpy as np
      from sklearn.cluster import AgglomerativeClustering, KMeans
      from sklearn.metrics import accuracy_score, confusion_matrix
      import time
 [6]: # Read data. Do not change the variable names (data, label)
      data = pd.read_csv('data/data.csv')
      label = pd.read csv('data/labels.csv')
      data=data.drop('Unnamed: 0',axis=1)
      label=label.drop('Unnamed: 0',axis=1)
[33]: data.head(5)
[33]:
         gene_0
                   gene_1
                             gene_2
                                        gene_3
                                                   gene_4
                                                           gene_5
                                                                      gene_6 \
      0
            0.0
                 2.017209
                           3.265527
                                     5.478487
                                                10.431999
                                                               0.0
                                                                    7.175175
      1
            0.0
                0.592732
                           1.588421
                                     7.586157
                                                 9.623011
                                                               0.0 6.816049
      2
            0.0
                 3.511759
                           4.327199
                                      6.881787
                                                 9.870730
                                                               0.0
                                                                    6.972130
      3
            0.0
                3.663618
                           4.507649
                                     6.659068
                                                10.196184
                                                               0.0
                                                                    7.843375
      4
            0.0
                2.655741
                           2.821547
                                     6.539454
                                                 9.738265
                                                               0.0 6.566967
                   gene_8
                           gene_9
                                       gene_20521
                                                   gene_20522
                                                               gene_20523
           gene_7
        0.591871
                      0.0
                               0.0
                                         4.926711
                                                     8.210257
                                                                  9.723516
      0
        0.000000
                      0.0
                               0.0
                                         4.593372
                                                     7.323865
      1
                                                                  9.740931
                      0.0
                               0.0
      2 0.452595
                                         5.125213
                                                     8.127123
                                                                 10.908640
      3 0.434882
                      0.0
                               0.0 ...
                                         6.076566
                                                     8.792959
                                                                 10.141520
      4 0.360982
                      0.0
                               0.0
                                         5.996032
                                                     8.891425
                                                                 10.373790
         gene_20524
                     gene_20525
                                 gene_20526
                                              gene_20527
                                                           gene_20528
                                                                       gene_20529
      0
           7.220030
                       9.119813
                                   12.003135
                                                9.650743
                                                             8.921326
                                                                         5.286759
      1
           6.256586
                       8.381612
                                   12.674552
                                               10.517059
                                                             9.397854
                                                                         2.094168
      2
           5.401607
                       9.911597
                                    9.045255
                                                9.788359
                                                            10.090470
                                                                         1.683023
      3
           8.942805
                       9.601208
                                   11.392682
                                                9.694814
                                                             9.684365
                                                                         3.292001
           7.181162
                       9.846910
                                   11.922439
                                                9.217749
                                                             9.461191
                                                                         5.110372
         gene_20530
      0
                0.0
      1
                0.0
```

```
2
            0.0
3
            0.0
            0.0
```

[5 rows x 20531 columns]

1.0.1 A. [Peer Review] Perform basic data inspection or EDA on the pandas dataframe.

• How many observations?

```
• How many features?
[7]: # perform basic data inspection such as getting the number of observations and
     → number of features
     # you can also display part of the dataframe or run data.info()
     # your code here
    print('The total amount of missing data points in the dataframe is: %d' % data.
     →isnull().sum().sum())
    print('The number of observations in the data frame is: %d' % data.shape[0])
    print('The total number of features in the dataframe is: %d' % data.shape[1])
    print('----')
    data.info()
    The total amount of missing data points in the dataframe is: 0
    The number of observations in the data frame is: 801
    The total number of features in the dataframe is: 20531
    <class 'pandas.core.frame.DataFrame'>
    RangeIndex: 801 entries, 0 to 800
    Columns: 20531 entries, gene_0 to gene_20530
    dtypes: float64(20531)
    memory usage: 125.5 MB
[8]: label.head(5)
[8]:
      Class
    O PRAD
```

- 1 LUAD
- 2 PRAD
- 3 PRAD
- 4 BRCA

[9]: data.head(5)

[9]: gene_6 \ gene_0 gene_1 gene_2 gene_3 gene_4 gene_5 0.0 2.017209 3.265527 5.478487 10.431999 0.0 7.175175

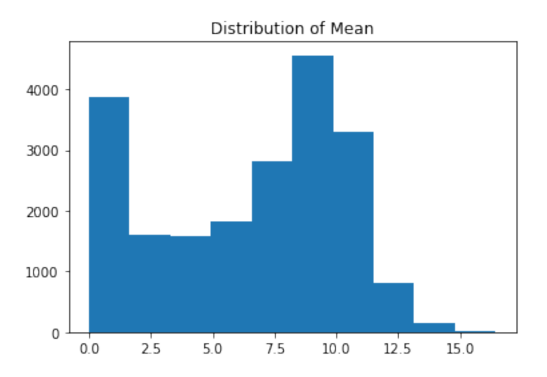
```
1
      0.0 0.592732
                      1.588421
                                7.586157
                                            9.623011
                                                         0.0 6.816049
2
      0.0 3.511759
                     4.327199
                                6.881787
                                            9.870730
                                                         0.0
                                                               6.972130
3
      0.0
          3.663618
                     4.507649
                                6.659068
                                           10.196184
                                                         0.0
                                                               7.843375
4
          2.655741
                      2.821547
                                6.539454
                                            9.738265
                                                         0.0 6.566967
             gene_8
                     gene_9
                                 gene_20521
                                              gene_20522
                                                          gene_20523 \
     gene_7
   0.591871
                0.0
                         0.0
                                   4.926711
                                                8.210257
                                                             9.723516
0
   0.000000
                0.0
                         0.0
1
                                   4.593372
                                                7.323865
                                                             9.740931
2
  0.452595
                0.0
                         0.0
                                   5.125213
                                                8.127123
                                                            10.908640
3 0.434882
                0.0
                         0.0
                                   6.076566
                                                8.792959
                                                            10.141520
4 0.360982
                0.0
                         0.0
                                   5.996032
                                                8.891425
                                                            10.373790
   gene_20524
               gene_20525
                            gene_20526
                                        gene_20527
                                                     gene_20528
                                                                  gene_20529
0
     7.220030
                 9.119813
                             12.003135
                                           9.650743
                                                       8.921326
                                                                    5.286759
     6.256586
                 8.381612
                             12.674552
                                          10.517059
                                                       9.397854
                                                                    2.094168
1
                                                                    1.683023
2
     5.401607
                 9.911597
                              9.045255
                                           9.788359
                                                      10.090470
3
     8.942805
                 9.601208
                             11.392682
                                           9.694814
                                                       9.684365
                                                                    3.292001
4
     7.181162
                 9.846910
                             11.922439
                                           9.217749
                                                       9.461191
                                                                    5.110372
   gene_20530
0
          0.0
1
          0.0
2
          0.0
3
          0.0
4
          0.0
```

[5 rows x 20531 columns]

• Draw histograms of mean, max and min values in each feature. You may see numbers around 0-20. What do those numbers mean? (We do not expect students to know or figure out the meanings, but if you do know by chance, feel free to discuss them with the class on the discussion board.) Answer the Peer Review question about this section.

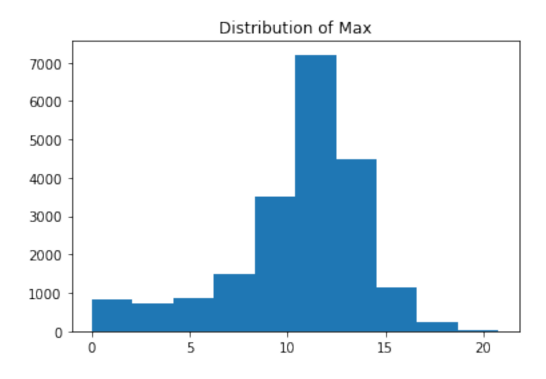
```
[10]: # draw histograms of mean, max and min values in each feature
# your code here
plt.hist(data.mean())
plt.title('Distribution of Mean')
```

[10]: Text(0.5, 1.0, 'Distribution of Mean')



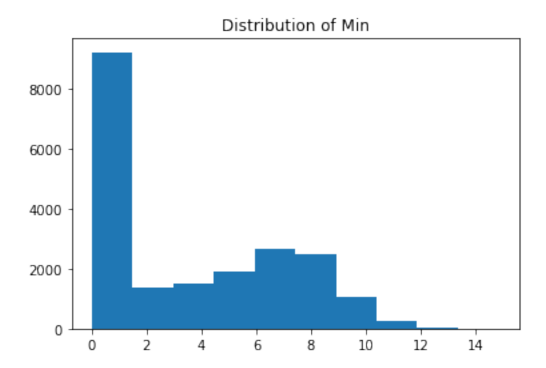
```
[11]: plt.hist(data.max())
  plt.title('Distribution of Max')
```

[11]: Text(0.5, 1.0, 'Distribution of Max')



```
[12]: plt.hist(data.min())
plt.title('Distribution of Min')
```

[12]: Text(0.5, 1.0, 'Distribution of Min')



- If we were to train a "supervised" learning model, how would you deal with such large feature dimension?
- Even after feature dimension reduction, still the number of useful features may be enormous. How it would impact performance or runtime of certain supervised learning algorithms? Which algorithms would suffer from high dimension features than others and why?
- How it would impact performance or runtime of an unsupervised learning algorithm?
- Draw histograms of mean, max and min values in each feature. You may see numbers around 0-20. What those numbers mean? (We do not expect students to know or figure out the meanings, but if you do know by chance, feel free to discuss them with the class on the discussion board.) Anwer these questions in this week's Peer Review assignment.
- If we were to train a "supervised" learning model, how would you deal with such large feature dimension? If supervised learning is used, using a model that subsamples features such as random forest might be better than other models that don't sample the features.
- Even after feature dimension reduction, still the number of useful features may be enormous. How it would impact performance or runtime of certain supervised learning algorithms? Which algorithms would suffer from high dimension features

than others and why? Additive models (gradient boosting) and linear models (e.g. logistic regression) without random feature subsampling will be especially slow (additive models) or perform poorly (logistic regression).

First, k-nearest neighbors struggle when the dimensions increase. This is because the distance measure struggles as the number of features increases. This would impact its runtime and accuracy. Logistic is not the best model for this data not only because logistic is best with a binary response but high-dimensional data can cause issues with correlation and multicollinearity. The variables within the Matrices may be linearly dependent. Therefore, we would fail to find a unique solution. Random Forests will not suffer from high dimensionality due to using a subset of features to fit different trees. Last, the support vector machine would be the best supervised model for this dataset. Support vector machine uses the C parameter, to define the boundaries. These boundaries use the 'supports' (subset of all the data points) that determines the boundaries. The high dimensions do not affect the boundary decision.

• How it would impact performance or runtime of an unsupervised learning algorithm? Dimensionality reduction such as PCA might be considered, but practically it won't work because the matrix size is too big for SVD.

1.0.2 B. [Peer Review] Build a hierarchical clustering model

Let's build a model using hierarchical clustering. Hierarchical clustering module is available from sklearn.cluster.AgglomerativeClustering. You can choose linkage type and metric. Please check its documentation for more details.

a) Number of clusters vs distance threshold Oftentimes hierarchical clustering does not need to know the number of clusters in advance. Instead, one needs to choose threshold distance/similarity to cut the dendrogram later. The AgglomerativeClustering module lets you specify either the number of clusters (n_clusters) or the threshold (distance_threshold). Based on our data, which should we choose to set to which value and why? Answer this question in the Peer Review assignment.

It's an open-ended question. The easiest guess would be to set the number of clusters to be five as we have the five labels (cancer types). However, if our goal is to explore more potential subgroups, we may want to let the number of clusters vary. In our case, for the evaluation purpose, setting n clusters=5 is the most reasonable choice.

- b) Guess which metric? Can you guess which metric to use (distance-based vs. similarity-based) and why? This question is not graded, but we encourage you to share your thoughts with the class. See the ungraded discussion prompt for this week's material.
- c) Build a model Build a model using n_clusters=5 option. Choose any metric and linkage type at first. Display the clustering result labels (you can just print out the result). Do not change the variable (model) name. Answer the question about this section in the Peer Review.

```
[39]: # build a model using n_clusters=5 option
model=None
# your code here
```

```
model = AgglomerativeClustering(n_clusters=5, linkage='complete', affinity='12')
model.fit(data)
```

```
[39]: AgglomerativeClustering(affinity='12', compute_full_tree='auto', connectivity=None, distance_threshold=None, linkage='complete', memory=None, n_clusters=5)
```

d) Label permuation In clustering, the labels get assigned randomly, so the label numbering won't match the ground truth necessarily. Write a function below to find best matching label ordering based on the accuracy. Do not change the variable names. Answer the question about this section in the Peer Review.

```
[37]: import itertools
      def label_permute_compare(ytdf,yp,n=5):
          ytdf: labels dataframe object
          yp: clustering label prediction output
          Returns permuted label order and accuracy.
          Example output: (3, 4, 1, 2, 0), 0.74
          .....
      ### BEGIN SOLUTION
          perms = list(itertools.permutations(list(range(n))))
          acc=[]
          for i in range(len(perms)):
              mapdict = dict(zip(list(label['Class'].unique()),list(perms[i])))
              yt = ytdf['Class'].apply(lambda x: mapdict[x])
              acc.append(accuracy_score(yt,yp))
          idx = np.argmax(acc)
          return perms[idx], acc[idx]
[28]: zipper = zip([1,2,3], ['Chuckie', 'penut', 'ice'])
      for i in zipper:
          print(i)
     (1, 'Chuckie')
     (2, 'penut')
     (3, 'ice')
[38]: labelorder, acc = label_permute_compare(label, model.labels_)
      print(labelorder, acc)
```

(4, 3, 1, 0, 2) 0.9313358302122348

e) Check confusion matrix Use sklearn's confusion matrix and display the results. Answer the Peer Review question about this section.

```
[44]: # display confusion matrix here
      # your code here
     mapdict = dict(zip(list(label['Class'].unique()), labelorder))
     yt = label['Class'].apply(lambda x: mapdict[x])
[45]: confusion_matrix(yt, model.labels_)
[45]: array([[299,
                    Ο,
                         Ο,
                              1,
                                   0],
            [ 0, 146,
                                   0],
                         0,
                              0,
            [ 0,
                    0, 136,
                              0,
                                   0],
            [ 2,
                    0, 0, 139,
                                  0],
                              1, 77]])
            [ 0,
                    0, 0,
```

f) Change linkage method and distance metric. Which ones lead the best performance? Print out the accuracy and confusion matrix for the best model. Answer the Peer Review questions about this section.

```
[43]: # programmatically evaluate which linkage method and distance metric lead to⊔
       \rightarrow the best performance
      for linkage in ['ward', 'complete', 'average', 'single']:
          for affinity in ['euclidean', 'l1', 'l2', 'manhattan', 'cosine']:
              acc=0
              t0=time.time()
              try:
                  model =
       →AgglomerativeClustering(n_clusters=5,linkage=linkage,affinity=affinity).
       →fit(data)
                  labelorder, acc = label_permute_compare(label, model.labels_)
                  t1=time.time()
                  print(t1-t0, linkage, affinity, labelorder, acc)
              except:
                  print(linkage, 'with', affinity, 'not allowed.')
      # Final answer
      model =
       →AgglomerativeClustering(n clusters=5,linkage='ward',affinity='euclidean').
       →fit(data)
      labelorder, acc = label_permute_compare(label,model.labels_)
      mapdict = dict(zip(list(label['Class'].unique()),list(labelorder)))
      yt = label['Class'].apply(lambda x: mapdict[x])
      print(acc)
      confusion_matrix(yt,model.labels_)
```

```
5.671780586242676 ward euclidean (2, 3, 0, 1, 4) 0.9950062421972534
     ward with 11 not allowed.
     ward with 12 not allowed.
     ward with manhattan not allowed.
     ward with cosine not allowed.
     5.622270345687866 complete euclidean (4, 3, 1, 0, 2) 0.9313358302122348
     5.719428539276123 complete 11 (4, 3, 0, 1, 2) 0.7228464419475655
     5.674745082855225 complete 12 (4, 3, 1, 0, 2) 0.9313358302122348
     5.6989336013793945 complete manhattan (4, 3, 0, 1, 2) 0.7228464419475655
     5.56896185874939 complete cosine (3, 4, 1, 2, 0) 0.7403245942571786
     5.639066696166992 average euclidean (2, 4, 0, 3, 1) 0.3645443196004994
     5.651516675949097 average l1 (1, 2, 0, 3, 4) 0.365792759051186
     5.651885509490967 average 12 (2, 4, 0, 3, 1) 0.3645443196004994
     5.70735764503479 average manhattan (1, 2, 0, 3, 4) 0.365792759051186
     5.5710837841033936 average cosine (2, 4, 1, 3, 0) 0.3645443196004994
     5.646402359008789 single euclidean (1, 2, 0, 3, 4) 0.3757802746566791
     5.696093559265137 single l1 (3, 1, 0, 2, 4) 0.37453183520599254
     5.655005931854248 single 12 (1, 2, 0, 3, 4) 0.3757802746566791
     5.696987152099609 single manhattan (3, 1, 0, 2, 4) 0.37453183520599254
     5.544532537460327 single cosine (1, 2, 0, 3, 4) 0.3757802746566791
     0.9950062421972534
[43]: array([[299,
                                   0],
                    Ο,
                         Ο,
                              1,
             [ 0, 146,
                         0,
                              0,
                                   0],
             [ 0,
                    0, 136,
                              Ο,
                                   0],
                     0, 0, 139,
             [ 2,
                                   0],
             [ 0,
                    Ο,
                         Ο,
                               1, 77]])
```

1.0.3 C. What about k-means clustering?

Can we apply kmeans clustering on this data? Which clustering methods give a better performance? Is kmeans faster or slower?

```
[46]: t0=time.time()
kmeans = KMeans(5).fit(data)
t1=time.time()
print(t1-t0)
labelorder, acc = label_permute_compare(label,kmeans.labels_)
print(labelorder, acc)
mapdict = dict(zip(list(label['Class'].unique()),list(labelorder)))
yt = label['Class'].apply(lambda x: mapdict[x])
print(acc)
confusion_matrix(yt,kmeans.labels_)
```

```
8.811512231826782
(2, 4, 1, 0, 3) 0.9925093632958801
0.9925093632958801
```

```
[46]: array([[145,
                                       0],
                            Ο,
                                  Ο,
              [
                 0, 299,
                            Ο,
                                  Ο,
                                       1],
              [ 0,
                       0, 136,
                                  Ο,
                                       0],
              [ 0,
                       Ο,
                            Ο,
                                 76,
                                       2],
              [ 0,
                       2,
                                  0, 139]])
                            Ο,
```

Hierarchical clustering with ward linkage and euclidean distance metric performs slightly better than kmeans clustering. However, the kmeans clustering also works well and is slightly faster.

[]: