Predicting Viral Host from Codon Usage Bias

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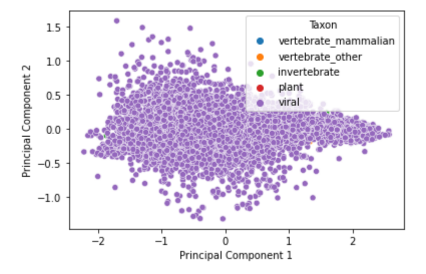
BIO 334/335 Bioinformatics Final Project

```
In [1]:
        import pandas as pd
        import numpy as np
        import matplotlib.pyplot as plt
        import seaborn as sns
        import math
        import itertools
        from sklearn.decomposition import PCA
        from sklearn.cluster import KMeans
        from sklearn.utils import shuffle
        from sklearn.svm import SVC
        from sklearn.neighbors import KNeighborsClassifier
        from sklearn.tree import DecisionTreeClassifier
        from sklearn.ensemble import RandomForestClassifier
        # all my helper functions
        from helper import *
In [2]:
        # load data
        bias pd = pd.read csv("datasets/report/cub full genome.csv")
        bias pd.shape
Out[2]: (9701, 69)
In [3]:
        bias pd.head()
             UUU
                     UUC
                             UUA
                                    UUG
                                            CUU
                                                    CUC
                                                           CUA
                                                                   CUG
                                                                           AUU
Out[3]:
         2 0.475298 0.524702 0.088171 0.135714 0.139210 0.191816 0.068511 0.376578 0.366103 C
        3 0.453638 0.546362 0.078937 0.127617 0.129814 0.198149 0.064855 0.400627
                                                                        0.351219 (
         0.449071 0.550929 0.077686 0.128257 0.129594 0.197832 0.066708 0.399924 0.351122 C
       5 rows × 69 columns
In [4]:
        # store which columns we drop to just get 59 codons
        drop columns = ["AUG", "UGA", "UAA", "UAG", "UGG", "AccessionNum", "SeqLen", "No
        # drop columns biased = drop columns + ["BiasedSeqLen", "PropBiasedRegions",
```

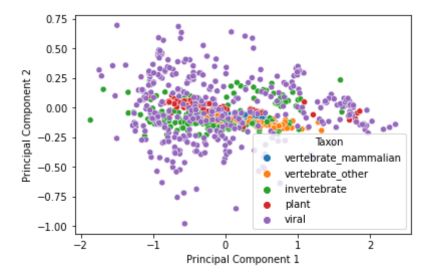
```
In [5]: # summarize entire dataset in 2d
bias_2d = reduce_dim(bias_pd, dim=2, drop_columns=drop_columns, vertebrate=False
bias_2d.shape
bias_2d.head()
```

```
Dim1
                           Dim2 AUG
                                            UGA
                                                      UAA
                                                                UAG UGG
                                                                                            AccessionNu
Out[5]:
                       -0.068775
            0.335693
                                   1.0
                                        0.498901 0.269464
                                                            0.231635
                                                                        1.0
                                                                              GCF_903992535.2_mArvAmp'
             0.342254
                       -0.079371
                                       0.500936
                                                  0.272810
                                                            0.226254
                                                                        1.0
                                                                                 GCF_000493695.1_BalAcu1
                      -0.080829
                                       0.494974
                                                                             GCF_000754665.1_Bison_UMD'
             0.307215
                                   1.0
                                                  0.280713
                                                            0.224312
                                                                        1.0
            0.436720
                       -0.076458
                                   1.0
                                        0.501819
                                                  0.267367
                                                            0.230814
                                                                        1.0
                                                                           GCF_000247795.1_Bos_indicus_1
                                                                           GCF_002288905.1_ASM228890
             0.461088
                       -0.062136
                                   1.0
                                        0.509291
                                                  0.262621 0.228088
```

```
In [6]: # plot of all CUB in the dataset - overwhelmed by the number of viruses vs. othe
    sns.scatterplot(data=bias_2d, x='Dim1', y='Dim2', hue='Taxon')
    plt.xlabel('Principal Component 1')
    plt.ylabel('Principal Component 2');
```



```
In [7]:
# plot with fewer viruses - still hard to see any meaningful differences
sns.scatterplot(data=bias_2d.head(1000), x='Dim1', y='Dim2', hue='Taxon')
plt.xlabel('Principal Component 1')
plt.ylabel('Principal Component 2');
```



Identifying virus host from CUB

Pull in extra viral host data from: https://www.ncbi.nlm.nih.gov/genomes/GenomesGroup.cgi?taxid=10239&cmd=download2 (On https://www.ncbi.nlm.nih.gov/genome/viruses/, it's the Accession List of all viral genomes)

```
In [8]: # read in viral data
hosts = pd.read_csv("datasets/viral_hosts.csv", skiprows=[0])
hosts = hosts.drop(columns=["Neighbor", "Segment name", "Representative", "Selec
# remove duplicate entries for each virus
hosts = hosts.drop_duplicates(ignore_index=True).dropna()
print(hosts.shape)
hosts.head(10)
(16095 2)
```

Out[8]:

(1	(16095, 2)									
	Host	Taxonomy name								
0	human,vertebrates	Cowpox virus								
1	human,vertebrates	Monkeypox virus								
2	human,vertebrates	Monkeypox virus Zaire-96-I-16								
3	human,vertebrates	Vaccinia virus								
4	human,vertebrates	Vaccinia virus Copenhagen								
5	human,vertebrates	Vaccinia virus Ankara								
6	human,vertebrates	Vaccinia virus Tian Tan								
7	human,vertebrates	Rabbitpox virus								
8	human,vertebrates	Horsepox virus								
9	human,vertebrates	Buffalopox virus								

```
# follow CUB dataset naming convention
hosts['Species'] = hosts['Taxonomy name'].apply(lambda x: x.replace(" ", "_"))
```

```
# column for whether human is host or not
hosts['Human_Host'] = hosts['Host'].apply(lambda x: "human" in x)
hosts.head()
```

```
Host
                                        Taxonomy name
                                                                         Species Human Host
 Out[9]:
          0 human, vertebrates
                                           Cowpox virus
                                                                     Cowpox_virus
                                                                                        True
            human,vertebrates
                                        Monkeypox virus
                                                                   Monkeypox_virus
                                                                                        True
            human, vertebrates Monkeypox virus Zaire-96-I-16 Monkeypox_virus_Zaire-96-I-16
                                                                                        True
            human, vertebrates
                                           Vaccinia virus
                                                                     Vaccinia_virus
                                                                                        True
            human,vertebrates
                                Vaccinia virus Copenhagen
                                                          Vaccinia_virus_Copenhagen
                                                                                        True
In [10]:
           # join in host info where matches any viruses in the CUB dataset
           # inner join -> only keep species w/ info in both datasets
           virus_cub = pd.merge(bias_pd, hosts, on="Species")
           virus cub.shape
Out[10]: (4660, 72)
In [11]:
           # 220 viruses w/ human host
           human_sum = np.sum(virus_cub["Human_Host"])
           human sum
Out[11]: 220
In [12]:
           # keep equal numbers human and non-human viruses
           # (otherwise non-human viruses overwhelm the dataset)
           np.random.seed(88)
           virus human = virus cub[virus cub["Human Host"]]
           virus non human = virus cub[~virus cub["Human Host"]]
           virus_non_human = shuffle(virus_non_human).head(human_sum) # keep random viruse
           virus cub short = pd.concat([virus human, virus non human])
           virus cub short.shape
Out[12]: (440, 72)
In [13]:
           viral_drop_columns = drop_columns + ["Host", "Taxonomy name", "Human_Host"]
           virus cub short.head()
                                              UUG
                  UUU
                            UUC
                                     UUA
                                                        CUU
                                                                 CUC
                                                                          CUA
                                                                                    CUG
                                                                                             AUU
Out[13]:
                        0.861386 0.009050 0.045249 0.167421 0.597285 0.027149 0.153846
           49
               0.138614
                                                                                         0.113043
           63 0.864198 0.135802 0.255319 0.212766
                                                     0.117021 0.037234 0.154255 0.223404 0.489583
```

0.176471 0.085973

121 0.607143 0.392857 0.191646 0.272727 0.154791 0.095823 0.147420 0.137592 0.419355

0.018100 0.122172 0.095023 0.482270

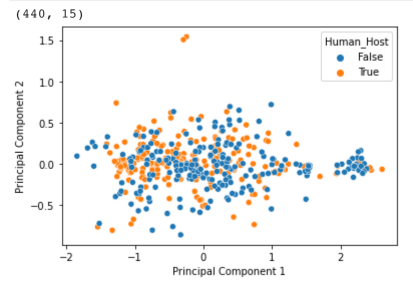
64 0.954545 0.045455 0.502262

```
        UUU
        UUC
        UUA
        UUG
        CUU
        CUC
        CUA
        CUG
        AUU

        123
        0.440678
        0.559322
        0.068627
        0.238562
        0.114379
        0.133987
        0.163399
        0.281046
        0.321839
```

5 rows × 72 columns

```
# visualize virus_cub_short
virus_2d = reduce_dim(virus_cub_short, dim=2, drop_columns=viral_drop_columns)
print(virus_2d.shape)
sns.scatterplot(data=virus_2d, x='Dim1', y='Dim2', hue='Human_Host')
plt.xlabel('Principal Component 1')
plt.ylabel('Principal Component 2');
```



In [15]: # dataset for our classifier, with just CUB and Human_Host columns
 virus_cub_short_num = virus_cub_short.drop(columns=viral_drop_columns[:-1]) # ke
 virus_cub_short_num.head()

Out[15]:		UUU	UUC	UUA	UUG	CUU	CUC	CUA	CUG	AUU
	49	0.138614	0.861386	0.009050	0.045249	0.167421	0.597285	0.027149	0.153846	0.113043
	63	0.864198	0.135802	0.255319	0.212766	0.117021	0.037234	0.154255	0.223404	0.489583
	64	0.954545	0.045455	0.502262	0.176471	0.085973	0.018100	0.122172	0.095023	0.482270
	121	0.607143	0.392857	0.191646	0.272727	0.154791	0.095823	0.147420	0.137592	0.419355
	123	0.440678	0.559322	0.068627	0.238562	0.114379	0.133987	0.163399	0.281046	0.321839

5 rows × 60 columns

and now, we build the classifier...

```
Out[22]: (396, 60)
In [17]:
          # cross-validation for model selection
          # get all cv errors for each method
          cv_errors = all_cv_errors(virus_tt, ['neighbor', 'tree', 'forest', 'SVM'])
          # store best method
          best method = cv errors[0,0]
          lowest_cv_err = cv_errors[0,1]
          cv_errors
Out[17]: array([['tree', 0.2626282051282051],
                ['neighbor', 0.29826923076923073],
                ['forest', 0.3106410256410257],
                ['SVM', 0.36903846153846154]], dtype=object)
In [18]:
          # print best model
          print('best model')
          print('method: ', best_method)
                           ', lowest_cv_err)
          print('error:
         best model
         method: tree
         error:
                   0.2626282051282051
In [19]:
          # build best type of model
          # mod_virus = KNeighborsClassifier(n_neighbors = 9)
          # mod virus = SVC(kernel="sigmoid")
          mod virus = DecisionTreeClassifier()
          # mod virus = RandomForestClassifier(n estimators=20, max features=min(3, virus
          mod virus.fit(virus tt.iloc[:,:-1], virus tt.iloc[:,-1]) # from all but the last
Out[19]: DecisionTreeClassifier(ccp alpha=0.0, class weight=None, criterion='gini',
                                max depth=None, max features=None, max leaf nodes=None,
                                min impurity decrease=0.0, min impurity split=None,
                                min samples leaf=1, min samples split=2,
                                min weight fraction leaf=0.0, presort='deprecated',
                                random state=None, splitter='best')
In [20]:
          # validate the model
          # compute validation error
          val preds = mod virus.predict(virus valid.iloc[:,:-1])
          val error = classification mse(val preds, virus valid.iloc[:,-1])
          print('validation error:
                                              ', val_error)
          print('best cross-validation error:', lowest_cv_err)
         validation error:
                                       0.25
         best cross-validation error: 0.2626282051282051
In [21]:
          # confusion matrix
          tp = np.sum((virus valid.iloc[:,-1]) & (val preds == 1)) # true positive
          tn = np.sum(-(virus_valid.iloc[:,-1]) & (val_preds == 0)) # true negative
          fp = np.sum(~(virus valid.iloc[:,-1]) & (val preds == 1)) # false positive
          fn = np.sum((virus_valid.iloc[:,-1]) & (val_preds == 0)) # false negative
                                                                                       # ty
```

```
print('\tactual class')
print(' p\t\t n')
print(f'tp: {tp/(tp+fn):.2f}\tfp: {fp/(tn+fp):.2f}')
print(f'fn: {fn/(tp+fn):.2f}\ttn: {tn/(tn+fp):.2f}')
```

```
actual class
p n
tp: 0.78 fp: 0.29
fn: 0.22 tn: 0.71
```

In our validation set, we correctly predicted whether a virus is hosted by humans or not 75% of the time. Thus, our model performs moderately well, and certainly better than random chance. The moderate success of this classifier suggests that CUB may be a useful predictor of whether a virus may infect humans, alongside other factors. Potential applications could include incorporating CUB as a factor in models that predict emerging viral threats to the human species.