# Appendix 1

# Instructions for the Reproduction of the Results in the Paper

First, go to <a href="https://github.com/millanp95/DeLUCS">https://github.com/millanp95/DeLUCS</a> and clone the repository. Then run the series of commands for each test.

# **Test #1:**

- python build dp.py --data path='../data/Vertebrata/Test Files'
- 2. python get\_pairs.py --data\_path='../data/Vertebrata/Test Files/train.p'
   --k=6 --modify='mutation' --output='../data/Vertebrata/Test
   Files/testing data.p'
- 3. python EvaluateDeLUCS.py --data\_dir='../data/Vertebrata/Test Files'
   --out dir='../data/Vertebrata/Test Files'

# **Comparison Models:**

- python EvaluateComparison.py --data\_path='../data/Vertebrata/Test Files/testing data.p' --method='Supervised' --k=6
- python EvaluateComparison.py --data\_path='../data/Vertebrata/Test Files/train.p' --method='k-means++' --k=6
- python EvaluateComparison.py --data\_path='../data/Vertebrata/Test Files/train.p' --method='GMM' --k=6 --k=6

# **Test #2:**

- 1. python build dp.py --data path='../data/Fish/Test Files/Actinopterygii'
- 2. python get\_pairs.py --data\_path='../data/Fish/Test
   Files/Actinopterygii/train.p' --k=6 --modify='mutation'
   --output=../data/Fish/Test Files/Actinopterygii/testing data.p'
- 3. python EvaluateDeLUCS.py --data\_dir='../data/Fish/Test Files/Actinopterygii'
  --out dir='../data/Fish/Test Files/Actinopterygii'

- python EvaluateComparison.py --data\_path='../data/Fish/Test
   Files/Actinopterygii/testing\_data.p' --method='Supervised' --k =6
- python EvaluateComparison.py --data\_path='.../data/Fish/Test Files/Actinopterygii/train.p' --method='k-means++' --k=6
- python EvaluateComparison.py --data\_path='../data/Fish/Test Files/Actinopterygii/train.p' --method='GMM' --k=6

# **Test #3:**

- 1. python build dp.py --data path='../data/Fish/Test Files/Neopterygii'
- 2. python get\_pairs.py --data\_path='../data/Fish/Test
  Files/Neopterygii/train.p' --k=6 --modify='mutation'
  --output=../data/Fish/Test Files/Neopterygii/testing data.p'
- 3. python EvaluateDeLUCS.py --data\_dir='../data/Fish/Test Files/Neopterygii'
   --out dir='../data/Fish/Test Files/Neopterygii'

# **Comparison Models:**

- python EvaluateComparison.py --data\_path='../data/Fish/Test
   Files/Neopterygii/testing\_data.p' --method='Supervised' --k =6
- python EvaluateComparison.py --data\_path='../data/Fish/Test Files/Neopterygii/train.p' --method='k-means++' --k=6
- python EvaluateComparison.py --data\_path='../data/Fish/Test Files/Neopterygii/train.p' --method='GMM' --k=6

# **Test #4:**

- 1. python build dp.py --data path='../data/Fish/Test Files/Ostariophysi'
- 2. python get\_pairs.py --data\_path='../data/Fish/Test
   Files/Ostariophysi/train.p' --k=6 --modify='mutation'
   --output=../data/Fish/Test Files/Ostariophysi/testing\_data.p'
- 3. python EvaluateDeLUCS.py --data\_dir='../data/Fish/Test Files/Ostariophysi'
  --out dir='../data/Fish/Test Files/Ostariophysi'

# Comparison Models:

- python EvaluateComparison.py --data\_path='./data/Fish/Test Files/Ostariophysi/testing\_data.p' --method='Supervised' --k =6
- python EvaluateComparison.py --data\_path='../data/Fish/Test Files/Ostariophysi/train.p' --method='k-means++' --k=6
- python EvaluateComparison.py --data\_path='../data/Fish/Test Files/Ostariophysi/train.p' --method='GMM' --k=6

### **Test #5:**

- 1. python build\_dp.py --data\_path='../data/Fish/Test Files/Cypriniformes'
- 2. python get\_pairs.py --data\_path='../data/Fish/Test
  Files/Cypriniformes/train.p' --k=6 --modify='mutation'
  --output=../data/Fish/Test Files/Cypriniformes/testing\_data.p'
- 3. python EvaluateDeLUCS.py --data\_dir='../data/Fish/Test Files/Cypriniformes'
   --out\_dir='../data/Fish/Test Files/Cypriniformes'

- python EvaluateComparison.py --data\_path='../data/Fish/Test Files/Cypriniformes/testing\_data.p' --method='Supervised' --k =6
- python EvaluateComparison.py --data\_path='../data/Fish/Test Files/Cypriniformes/train.p' --method='k-means++' --k=6
- python EvaluateComparison.py --data\_path='../data/Fish/Test Files/Cypriniformes/train.p' --method='GMM' --k=6

# **Test #6:**

- 1. python build dp.py --data path='../data/Fish/Test Files/Cyprinidae'
- 2. python get pairs.py --data path='../data/Fish/Test Files/Cyprinidae/train.p' --k=6 --modify='mutation' --output=../data/Fish/Test Files/Cyprinidae/testing data.p'
- 3. python EvaluateDeLUCS.py --data\_dir='../data/Fish/Test Files/Cyprinidae' --out dir='../data/Fish/Test Files/Cyprinidae'

#### Comparison Models:

- python EvaluateComparison.py --data\_path='../data/Fish/Test Files/Cyprinidae/testing\_data.p' --method='Supervised' --k =6
- python EvaluateComparison.py --data\_path='../data/Fish/Test Files/Cyprinidae/train.p' --method='k-means++' --k=6
- python EvaluateComparison.py --data\_path='../data/Fish/Test Files/Cyprinidae/train.p' --method="GMM" --k=6

#### **Test #7:**

- python build\_dp.py --data\_path='../data/Bacteria/Test Files'
   python get\_pairs.py --data\_path='../data/Bacteria/Test Files/train.p' --k=6 --modify='mutation' --output='../data/Bacteria/Test Files/testing\_data.p'
- 3. python EvaluateDeLUCS.py --data dir='../data/Bacteria/Test Files' --out dir='../data/Bacteria/Test Files'

# Comparison Models:

- python EvaluateComparison.py --data path='../data/Bacteria/Test Files/testing\_data.p' --method='Supervised' --k =6
- python EvaluateComparison.py --data\_path='../data/Bacteria/Test Files/train.p' --method='k-means++' --k=6
- python EvaluateComparison.py --data path='../data/Bacteria/Test Files/train.p' --method='GMM' --k=6

## **Test #8:**

- 1. python build\_dp.py --data\_path='../data/ProteoBacteria/Test Files'
- 2. python get\_pairs.py --data\_path='../data/ProteoBacteria/Test Files/train.p' --k=6 --modify='mutation' --output='../data/ProteoBacteria/Test Files/testing data.p'
- 3. python EvaluateDeLUCS.py --data dir='.../data/ProteoBacteria/Test Files' --out dir='../data/ProteoBacteria/Test Files'

- python EvaluateComparison.py --data\_path='../data/ProteoBacteria/Test Files/testing data.p' --method='Supervised' --k =6
- python EvaluateComparison.py --data\_path='../data/ProteoBacteria/Test Files/train.p' --method='k-means++' --k=6
- python EvaluateComparison.py --data\_path='../data/ProteoBacteria/Test Files/train.p' --method='GMM' --k=6

### **Test #9:**

- 1. python build\_dp.py --data\_path='../data/Influenza-A/Test Files'
- 2. python get\_pairs.py --data\_path='../data/Influenza-A/Test Files/train.p'
   --k=6 --modify='mutation' --output='../data/Influenza-A/Test
   Files/testing data.p'
- 3. python EvaluateDeLUCS.py --data\_dir='../data/Influenza-A/Test Files'
   --out\_dir='../data/Influenza-A/Test Files'

# Comparison Models:

- python EvaluateComparison.py --data\_path='../data/Influenza-A/Test
   Files/testing data.p' --method='Supervised' --k =6
- python EvaluateComparison.py --data\_path='../data/Influenza-A/Test Files/train.p' --method='k-means++' --k=6
- python EvaluateComparison.py --data\_path=../data/Influenza-A/Test Files/train.p' --method='GMM' --k=6

# **Test #10:**

- 1. python build dp.py --data path='../data/Dengue/Test Files'
- 2. python get\_pairs.py --data\_path='../data/Dengue/Test Files/train.p' --k=6 --modify='mutation' --output='../data/Dengue/Test Files/testing\_data.p'
- 3. python EvaluateDeLUCS.py --data\_dir='../data/Dengue/Test Files'
  --out dir='../data/Dengue/Test Files'

# **Comparison Models:**

- python EvaluateComparison.py --data\_path='../data/Dengue/Test Files/testing\_data.p' --method='Supervised' --k =6
- python EvaluateComparison.py --data\_path='../data/Dengue/Test Files/train.p'
   --method='k-means++' --k=6
- python EvaluateComparison.py --data\_path='../data/Dengue/Test Files/train.p'
   --method='GMM' --k=6

# **Test #11:**

- 1. python build\_dp.py --data\_path='../data/HBV/Test Files'
- 2. python get\_pairs.py --data\_path='../data/HBV/Test Files/train.p' --k=6 --modify='mutation' --output='../data/HBV/Test Files/testing\_data.p'
- 3. python EvaluateDeLUCS.py --data\_dir='../data/HBV/Test Files' --out\_dir='../data/HBV/Test Files'

- 1. python EvaluateComparison.py --data\_path='../data/HBV/Test
  Files/testing data.p' --method='Supervised' --k =6
- 2. python EvaluateComparison.py --data\_path='../data/HBV/Test Files/train.p'
   --method='k-means++' --k=6
- 3. python EvaluateComparison.py --data\_path='../data/HBV/Test Files/train.p'
   --method='GMM' --k=6

# **DeLUCS** computational pipeline for running your own dataset:

- 1. Build the dataset:
  - o Input: Folders with the sequences in FASTA format
  - Output : file in the form (label, sequence, accession)

\*If the true label is unknown then place ALL the sequences in the same folder

```
python build dp.py --data path=<PATH sequence folder>
```

- 2. Compute the mimics.
  - Input: file in the form (label, sequence, accession)
  - Output : file in the form of (pairs, x\_test, y\_test).

```
python get_pairs.py --data_path=<PATH_dataset> --k=6 --modify='mutation'
--output=<PATH_output_file>
```

- 3. Train the model.
  - Input: file in the form of (pairs, x test).
  - o Output:
    - Image with the confusion matrix (provided the "ground truth" is available)
    - File with the misclassified sequences in the form (accession, true\_label, predicted label)

python EvaluateDeLUCS.py --data dir=<PATH of computed mimics> --out dir=<OUTPURDIR>

\*\* If the true labels are unknown for your dataset and you want to use DeLUCS for a cluster prediction without the assessment by means of the Hungarian algorithm, run:

python TrainDeLUCS.py --data dir=<PATH of computed mimics> --out dir=<OUTPURDIR>