# SeqPlotter: Python package for sequence data analysis and visualization

SeqPlotter is a Python package developed to streamline the visualization and analysis of biological sequence data. It provides seamlessly integration into your Python workflow, with robust data handling capabilities, intuitive visualization, and useful analysis features. From parsing raw sequence files to creating stunning plots and charts.

# 1. Working with DNA sequences

#### 1.1. Importing DNA module

```
In [1]: # Importing module
    from seqplotter.nucl import DNA
```

#### 1.2. Creating DNA objects

```
>>> dna seq1 = DNA("seqID", "DNA sequence")
```

Output: retruns a DNA object with seqid and seq pair

```
In [2]: # creating DNA object with seqID and sequence pair
seq1 = DNA("seq1", "GTGTTTTGACTAATAATTGGTCAAGCCTAC")
```

#### 1.3. Sequence length

```
>>> dna seq1.length()
```

Output: returns sequence length (int) in bp

```
In [3]: # print sequence length
seq1.length()
```

Out[3]: 30

#### 1.4. Sequence composition

```
>>> dna_seq1.comp()
```

Output: returns a dict object with sequence composition i.e. count of each nucleotides

```
In [4]: # print sequence composition
    seq1.comp()
```

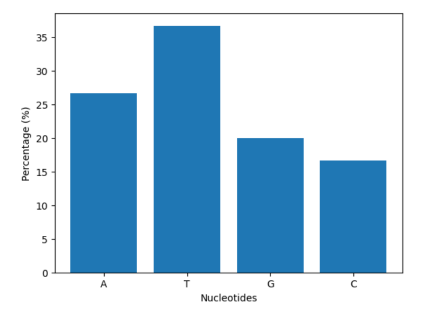
```
Out[4]: {'A': 26.666666666666666,
 'T': 36.6666666666666664,
 'G': 20.0,
 'C': 16.6666666666666664}
```

#### 1.5 Sequence composition barplot

```
>>> seq1_comp = dna_seq1.comp()
>>> DNA.barplot(seq1_comp)
```

Output: generate barplot showing sequence composition

```
In [5]: # plot sequence composition
    comp = seq1.comp()
    DNA.barplot(comp)
```



# 1.6. Sequence slicing

```
>>> # returns nucleotide for the given position
>>> dna_seq1.slice(pos)
>>> # returns nucleotides between start to end position
>>> dna_seq1.slice(start_pos, end_pos)
```

Output: returns a string that contain the sliced sequence

#### 1.7. GC content

```
>>> dna_seq1.gc_percent()
```

Output: returns GC percent (string)

```
In [8]: # GC percent
seq1.gc_percent()
```

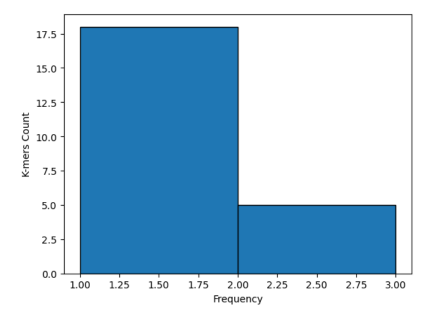
Out[8]: '36.67'

# 1.8. K-mer abundance plot

```
# generated k-mer plot with default kmer size (k=3)
>>> dna_seq1..kmer_abundance_plot()
# generate k-mer plot with k=6
>>> dna_seq1..kmer_abundance_plot(6)
```

Output: generate k-mer plot (histogram)

```
In [9]: # kmer abundance plot
     seq1.kmer_abundance_plot()
```



#### 1.9. Translate gene sequence

```
>>> dna_seq1.translate()
```

Output: translate DNA sequence into amino acid sequence

Out[10... 'VF\*LIIGQAY'

## 1.10. Readling FASTA file (DNA Sequence)

```
>>> fasta_records = DNA.read_fasta("/path/to/fasta/file")
```

Output: returns a list containing DNA object(s)

#### 1.11. Using head() and tail() functions

```
>>> # return summary of first 5 sequence (default)
>>> DNA.head()
>>> # return summary of first 10 sequence
>>> DNA.head(10)
>>> # return summary of last 5 sequence (default)
>>> DNA.tail()
>>> # return summary of last 10 sequence
>>> DNA.tail(10)
```

Output: print head and tail summary of parsed FASTA file

```
In [12... # print first five records (by default)
         DNA.head()
        seqID
                 [\texttt{CTTTTCTTTAGTGTTTTGACTAATAATTGGTCAAGCCTACCATTACAAACTATGTTCCATTACCAGTACA}]
        seq1:
                 [CTTTTCTTTAGTGTTTTGACTAATAATTGGTCAAGCCTACCATTACAAACTATGTTCCATTACCAGTACA]
        seq2:
                 [CTTTTCTTTAGTGTTTTGACTAATAATTGGTCAAGCCTACCATTACAAACTATGTTCCATTACCAGTACA]
        sea3:
                 [{\tt CTTTTCTTTAGTGTTTTGACTAATAATTGGAAACTATGTTCCATTACCAGTACA}]
        seq4:
        seq5:
                 [TCACAGAAGAGCAAGAGGCTCTTGTAGTGAAGTCTTGGAGTGTCATGAAGAAAAACTCAGCTGAATTAGG] \\
In [13...
         # print first two records
         DNA.head(2)
        seqID
```

seq1: [CTTTTCTTTAGTGTTTTGACTAATAATTGGTCAAGCCTACCATTACAAACTATGTTCCATTACCAGTACA]
seq2: [CTTTTCTTTAGTGTTTTGACTAATAATTGGTCAAGCCTACCATTACAAACTATGTTCCATTACCAGTACA]

```
In [14... # print last five records (by default)
       DNA.tail()
      seqID
             [TCACAGAAGAGCAAGAGGCTCTTGTAGTGAAGTCTTGGAGTGTCATGAAGAAAAACTCAGCTGAATTAGG] \\
      seq5:
             sea6:
             [TGAGATATGAACTATTTTGAACTGTAGGATCTTTGAGATTGCACCAACAACGA]
      seq7:
             [TTCTTGAGAGACTCACCAATTCCTGCTGAGCAAAATCCAAAGCTCAAGCCTCACGCAATGTCTGT]
      seq8:
      seq9:
             [TCATGGTAATAATCAATATCAAATAACATGATTTT]
In [15... # print last six records
       DNA.tail(3)
      seqID
             sea
      seq7:
             [TTCTTGAGAGACTCACCAATTCCTGCTGAGCAAAATCCAAAGCTCAAGCCTCACGCAATGTCTGT]
      seq8:
      seq9:
             [TCATGGTAATAATCAATATCAAATAACATGATTTT]
```

#### 1.12. Count records

>>> DNA.count(fasta\_records)

Output: returns sequence count (int) i.e. number of records

```
In [16...
        # print record counts
         DNA.count(data)
Out[16...
```

# 1.13. Find record

>>> DNA.where(seqid="seq1")

Output: returns sequence for a given sequenceID

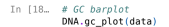
```
In [17...
         # find sequence
         DNA.where(seqid="seq1")
```

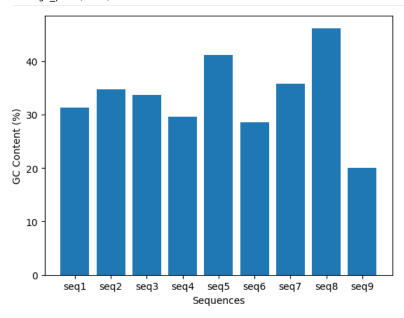
Out[17... CTTTTCTTTAGTGTTTTTGACTAATAATTGGTCAAGCCTACCATTACAAACTATGTTCCATTACCAGTACACTTTTCTTTAGTGTTTTTGACTAATAATTGGATTACCA

#### 1.14. GC percent per sequence plot

>>> DNA.gc\_plot(fasta\_records)

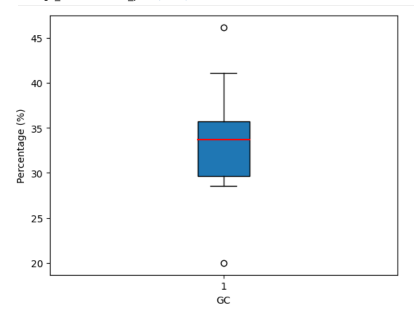
Output: generate barplot showing per sequence GC percent





#### 1.15. Boxplot showing GC distribution

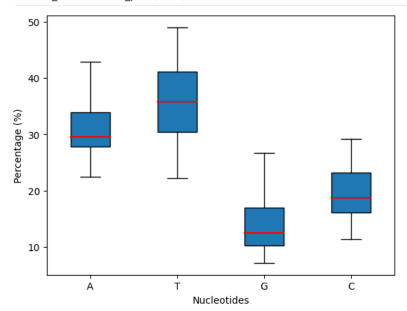
>>> DNA.gc\_distribution\_plot(fasta\_records)



# 1.16. Boxplot showing nucleotides distribution

>>> DNA.nt\_distribution\_plot(fasta\_records)

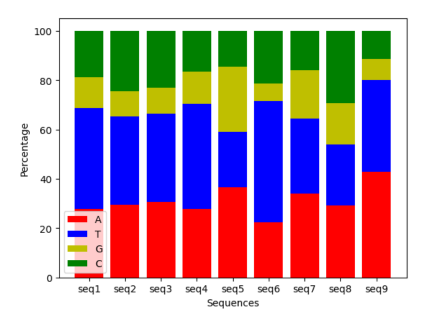
Output: generate boxplot showing overall nucleotides distribution across all the sequences



# 1.17. Stacked barplot showing per sequence base composition

>>> DNA.per\_sequence\_comp(fasta\_records)

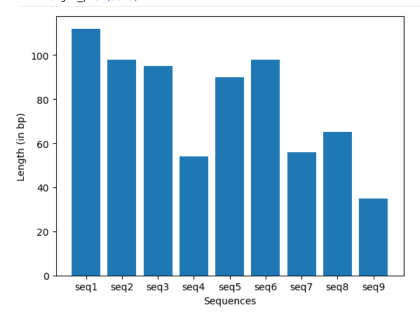
Output: generate stacked barplot showing per sequence nucleotide composition



# 1.18. Barplot showing per sequence length

>>> DNA.length\_plot(fasta\_records)

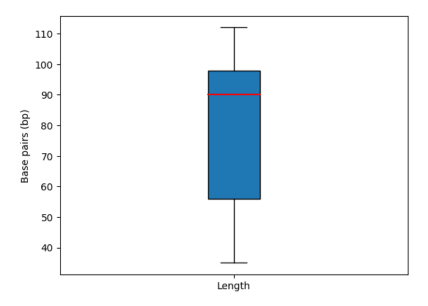
Output: generate barplot showing per sequence length distribution



# 1.19. Boxplot showing overall length distribution

>>> DNA.length\_distribution\_plot(fasta\_records)

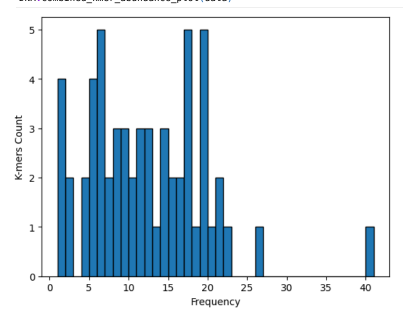
Output: generate boxplot showing overall length distribution across all the sequences



## 1.20. Overall k-mer abundance pattern

```
# default k-mer size (k=3)
>>> DNA.combined_kmer_abundance_plot(fasta_records)
# kmer size (k=7)
>>> DNA.combined_kmer_abundance_plot(fasta_records, 7)
```

Output: generate combined k-mer abundance plot (histogram) for all the sequence



# 1.21. Converting sequence object to feature matrix

```
>>> from seqplotter_analysis import seq2feature
>>> feature_matrix = seq2feature(fasta_records) # with default options (k=3, min_sample=1)
>>> feature_matrix = seq2feature(fasta_records, 6, 2) # with (k=6, min_sample=2)
```

#### Options:

- k: kmer size
- min\_sample: threshold of considering a k-mer as feature

Output: returns a dict object containing following keys ("matrix", "sample", "kmers"), where "kmers" are columns and "sample" are rows < dict{"matrix": numpy\_array, "sample": seqID\_list, "kmers": kmers\_list} >

```
In [25... # Importing module
          from seqplotter_analysis import seq2feature
          # Converting sequence to feature matrix
          feature_matrix = seq2feature(data)
          # Visualizing feature matrix
          import pandas as pd
          pd.DataFrame(feature_matrix["matrix"], index=feature_matrix["sample"], columns=feature_matrix["kmers"])
Out[25...
                CTT TTT TTC TCT TTA TAG AGT GTG TGT GTT ... GAA AGA GAG GCA AGG GCT CTC CTG TGC ACG
          seq1
                  4
                       10
                             3
                                  2
                                       5
                                            2
                                                  4
                                                       2
                                                             3
                                                                  3 ...
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                                                                                                  0
                                                                                                       0
                                                                                                             0
                                                                                                                  0
                                                                                                                       0
                                                                                                                             0
          seq2
                  2
                        5
                             3
                                       4
                                                  2
                                                            3
                                                                  3 ...
                                                                           0
                                                                                0
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                                                                                            0
                                                                                                  0
                                                                                                       0
                                                                                                             0
                                                                                                                  0
                                                                                                                        0
                                                                                                                             0
          seq3
                  2
                        5
                             3
                                       4
                                                 3
                                                            3
                                                                  3 ...
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                                                                                                             0
                                                                                                                  0
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          seq4
                  2
                        5
                             2
                                       2
                                                  2
                                                             2
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                  2
                        0
                            0
                                  2
                                       2
                                            3
                                                 3
                                                       2
                                                             2
                                                                  0 ...
                                                                           7
                                                                                4
                                                                                      3
                                                                                                  3
                                                                                                       3
                                                                                                             3
                                                                                                                  2
                                                                                                                        0
                                                                                                                             0
          seq5
          seq6
                  3
                       10
                             4
                                 12
                                       2
                                                  2
                                                       0
                                                                  0 ...
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                                                                                      0
                                                                                                  0
                                                                                                       0
                                                                                                             8
                                                                                                                  0
                                                                                                                        0
                                                                                                                             0
          seq7
                       3
                            0
                                       0
                                                 0
                                                       0
                                                                  0 ...
                                                                           2
                                                                                2
                                                                                      2
                                                                                                       0
                                                                                                             0
                                  1
                                            1
                                                                                            1
                                                                                                                       1
                                                                                                                             1
          seq8
                       0
                                  2
                                       0
                                            0
                                                  0
                                                       0
                                                                  0 ...
                                                                           0
                                                                                      3
                                                                                            2
                                                                                                  0
                                                                                                             3
                                                                                                                  3
                                                                                                                        1
          seq9
                  0
                       2
                            0
                                  0
                                       0
                                            0
                                                 0
                                                       0
                                                            0
                                                                  0 ...
                                                                           0
                                                                                0
                                                                                      0
                                                                                            0
                                                                                                  0
                                                                                                       0
                                                                                                             0
                                                                                                                  0
                                                                                                                       0
                                                                                                                             0
         9 rows × 54 columns
          1.22. Principal component analysis
              >>> from seqplotter_analysis import PCA
              >>> pca matrix = PC\overline{A}(feature matrix) # return pca matrix with first 2 PCs
              >>> pca_matrix = PCA(feature_matrix, 3) # return pca matrix with first 3 PCs
          Output: returns a dict object containing following keys ("matrix", "sample", "components"), where "components" are columns and
          "sample" are rows
          < dict{"matrix": numpy_array, "sample": seqID_list, "components": PCA_components} >
In [26... # Importing module
          from seqplotter_analysis import PCA
          # Performing PCA
          pca_matrix = PCA(feature_matrix)
          # Visualizing PCA matrix
          import pandas as pd
          pd.DataFrame(pca_matrix["matrix"], index=pca_matrix["sample"], columns=pca_matrix["components"])
```

PC-1 PC-2 seq1 9.629055 2.243708 seq2 5.947428 3.774627 6.225862 4.230590 sea3 0.776820 sea4 1.919865 **seq5** -9.639273 0.927017 3.503045 -15 066248 sea6 seq7 -4.161328 0.626651 seq8 -7.607692 0.437185 seq9 -4.673917 0.906603

Out[26...

### 1.23. Plot scatter plot using PCs

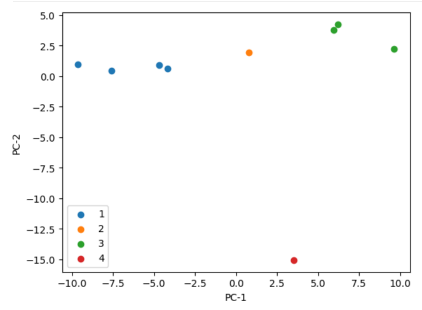
```
>>> from seqplotter_analysis import plot_pca
# Sequence IDs will be treated as class label (default)
>>> plot_pca(pca_matrix)
```

```
# Providing class labels as list
>>> labels = [3, 3, 3, 2, 1, 4, 1, 1, 1]
>>> plot_pca(pca_matrix, labels)
```

Output: generate PCA plot (2D scatter plot using first two components i.e. PC-1 and PC-2)

```
In [27... # Importing module
    from seqplotter_analysis import plot_pca

# Plotting 2D PCA plot
    labels = [3, 3, 3, 2, 1, 4, 1, 1, 1]
    plot_pca(pca_matrix, labels)
```



#### 1.24. Create sequence logo

```
>>> from seqplotter_analysis import seq_logo
# Reading motif sequences file
>>> logo_data = DNA.read_fasta("motif_seq.fasta")
# Generate sequence logo
>>> seq_logo(logo_data, seq_type)
```

Output: generate DNA sequence logo using motif sequences

```
In [28... # Importing module
         from seqplotter_analysis import seq_logo
         # Reading motif sequences file
         data = DNA.read_fasta("dna_motif_seq.fasta")
         # Generate sequence logo
         seq_logo(data, "DNA")
           2.00
           1.75
           1.50
           1.25
        £ 1.00
           0.75
           0.50
           0.25
           0.00
                                                           11
                                                               12 13 14
                                                                            15
```

Base positions

#### 1.25 Generate random nucleotide sequence

```
# Generating random sequences with given length and GC content
>>> DNA.generate_random_seq(lenght, GC_frequency, count)
```

Output: append random sequences object of given length and GC content to the DNA object

```
In [29...
        # Clearing records
        DNA.clear()
        \# Generating 10 random sequences with 70 bp length and 30% GC content
        new_data = DNA.generate_random_seq(70, 0.3, 10)
        # Printing first 5 random sequences
        DNA.head()
       seqID
              sea
                     [\mathsf{TCTATTACACTGAATAAATTTAAACCCGTGGTGTATTATATTTTACCTCTTTAAATGAATTTCACTATAA]
       random_seq1:
       random_seq2:
                     random_seq3:
                      [ACATGTTTATCAGTTTGTCCTATTGATGTTATCATTATTATTGATGATATTACCAAGGGCAATTGCAAAT]
       random_seq4:
                     [GTAAATCATTTACTCTCAACAGAATAAATTGTAATAATCTGTATGATAGAATACATGTTAATCGTTGATA]
       random_seq5:
                     [ATTAGTGATACAAACAGATTCATACTAATTAAAAGTTAACCTGTTCATTATTGAACTAAGTAATTCAATG]
```

# 2. Working with Protein sequences

#### 2.1. Importing PROT module

```
In [30... # Importing module from seqplotter.prot import PROT
```

# 2.2 Creating PROT object

```
>>> prot_seq1 = PROT("seqID", "protein_sequence")
```

Output: retruns a DNA object with seqid and seq pair

```
In [31... # Creating Protein object with seqID and sequence pair
seq1 = PROT("seq1", "VDEVGGEALGRLLVVYPWTQRFFESFGDLSTPDAV")
```

#### 2.3. Sequence length

```
>>> prot_seq1.length()
```

Output: returns sequence length (int) in bp

Out[32... 35

#### 2.4. Sequence composition

```
>>> prot_seq1.comp()
```

Output: returns a dict object with sequence composition i.e. count of each amino acids

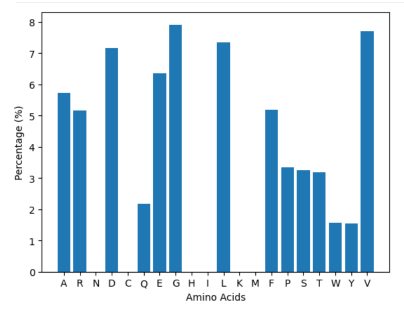
```
In [33... # print sequence composition
         seq1.comp()
Out[33... {'A': 5.714285714285714,
           'R': 5.166051660516605,
           'N': 0.0,
           'D': 7.163266076755573,
           'C': 0.0,
           'Q': 2.1718543403148107,
           'E': 6.353851345091472,
           'G': 7.909936022986237,
           'H': 0.0,
           'I': 0.0,
           'L': 7.342245635482005,
           'K': 0.0,
           'M': 0.0,
           'F': 5.188382310331203,
           'P': 3.3327849177003355,
           'S': 3.260373880823663,
           'T': 3.1947334484658114,
           'W': 1.567453020447904,
           'Y': 1.5536341292308162,
           'V': 7.701922950367426}
```

# 2.5. Sequence composition plot

```
>>> prot_comp = prot_seq1.comp()
>>> PROT.barplot(prot comp)
```

Output: generate barplot showing sequence composition

```
In [34... # plot sequence composition
    comp = seq1.comp()
    PROT.barplot(comp)
```



#### 2.6. Molecular weight in KDa

```
>>> prot_seq1.mol_weight()
```

Output: return molecular weight (int) of a protein sequence in KDa

#### 2.7. Sequence slicing

```
>>> # returns amino acid for the given position
>>> prot_seq1.slice(pos)
>>> # returns amino acids between start to end position
>>> prot_seq1.slice(start_pos, end_pos)
```

Output: returns a string that contain the sliced sequence

```
In [36... # slice particular position seq1.slice(6)

Out[36... 'G'

In [37... # slice with start and end index seq1.slice(4,8)

Out[37... 'VGGEA'
```

#### 2.8. Reading FASTA file (Protein sequence)

```
>>> fasta_records = PROT.read_fasta("/path/to/fasta/file")
```

Output: returns a list containing PROT object(s)

#### 2.9. Using head() and tail() functions

```
>>> # return summary of first 10 sequence
             >>> PROT.head(10)
             >>> # return summary of last 5 sequence (default)
             >>> PROT.tail()
             >>> # return summary of last 10 sequence
             >>> PROT.tail(10)
         Output: print head and tail summary of parsed FASTA file
In [39... # Head
         PROT.head(5)
        seqID
        seq1:
                 [MVHLTPEEKSAVTALWGKVNVDEVGGEALGRLLVVYPWTQRFFESFGDLSTPDAVMGNPKVKAHGKKVLG] \\
                 [MLPFWKRLLYAAVIAGALVGADAQFWKTAGTAGSIQDSVKHYNRNEPKFPIDDSYDIVDSAGVARGDLPP]
        seq2:
                 [TG0KGDRGEPGLNGLPGNPG0KGEPGRAGATGKPGLLGPPGPPGGGRGTPGPPGPKGPRGYVGAPGP0GL]
        seq3:
                [LPGATGEPGKPALCDLSLIEPLKGDKGYPGAPGAKGVQGFKGAEGLPGPGFKGEFGFKGEKGLSGAPGN]
        seq4:
In [40... # Tail
         PROT.tail()
        seqID
                . [MVHLTPEEKSAVTALWGKVNVDEVGGEALGRLLVVYPWTQRFFESFGDLSTPDAVMGNPKVKAHGKKVLG]
        seq1:
                 [MLPFWKRLLYAAVIAGALVGADAQFWKTAGTAGSIQDSVKHYNRNEPKFPIDDSYDIVDSAGVARGDLPP]
        seq2:
                 [TGQKGDRGEPGLNGLPGNPGQKGEPGRAGATGKPGLLGPPGPPGGGRGTPGPPGPKGPRGYVGAPGPQGL] \\
        seq3:
        seq4:
                [LPGATGEPGKPALCDLSLIEPLKGDKGYPGAPGAKGVQGFKGAEGLPGIPGPKGEFGFKGEKGLSGAPGN]
         2.10. Count records
             >>> PROT.count(fasta_records)
         Output: returns sequence count (int) i.e. number of records
         # Sequence count
         PROT.count(data)
         2.11. Find record
             >>> PROT.where(seqid="seq2")
         Output: returns sequence for the given sequenceID
         # Find sequence
         PROT.where(seqid="seq2")
          'MLPFWKRLLYAAVIAGALVGADAOFWKTAGTAGSIODSVKHYNRNEPKFPIDDSYDIVDSAGVARGDLPPKNCTAGYAGCVPKCIAEKGNRGLPGPLGPTGLKGEMG
         FPGMEGPSGDKGQKGDPGPYGQRGDKGERGSPGLHGQAGVPGVQGPAGNPGAPGINGKDGCDGQDGIPGLEGLSGMPGPRGYAGQLGSKGEKGEPAKENGDYA'
         2.12. Amino acids distribution plot
             >>> PROT.aa_distribution_plot(fasta_records)
```

>>> # return summary of first 5 sequence (default)

Output: generate boxplot showing overall amino acids distribution across all the sequences

# Boxplot aa distribution plot

PROT.aa\_distribution\_plot(data)

>>> PROT.head()

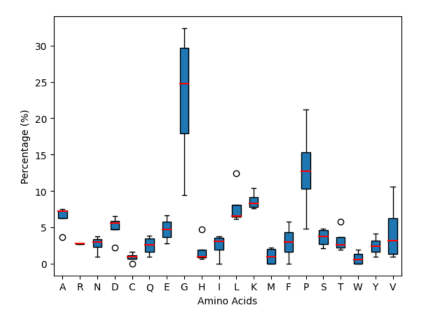
In [41...

Out[41...

In [42...

Out [42...

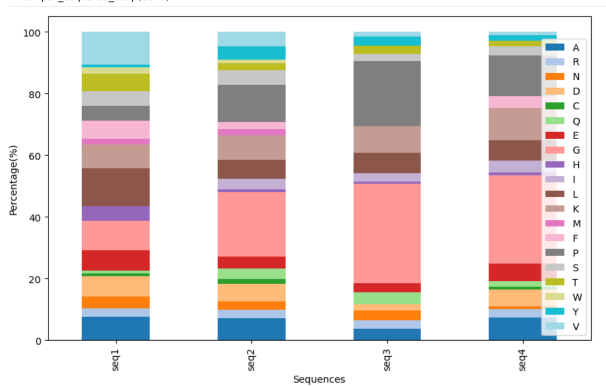
In [43...



# 2.13. Stacked barplot showing per sequence amino acid composition

>>> PROT.per\_sequence\_comp(fasta\_records)

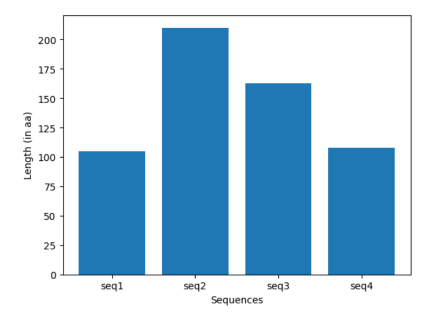
Output: generate stacked barplot showing per sequence amino acids composition



# 2.14. Barplot showing per sequence length

PROT.length\_plot(fasta\_records)

Output: generate barplot showing per sequence length distribution

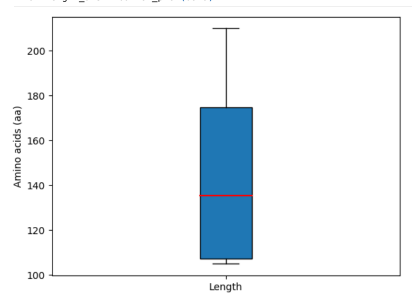


#### 2.15. Boxplot showing overall length distribution

>>> PROT.length\_distribution\_plot(fasta\_records)

Output: generate boxplot showing overall length distribution across all the sequences

In [46... PROT.length\_distribution\_plot(data)



# 2.16. Converting sequence object to feature matrix

```
>>> from seqplotter_analysis import seq2feature
>>> feature_matrix = seq2feature(fasta_records) # with default options (k=3, min_sample=1)
>>> feature_matrix = seq2feature(fasta_records, 6, 2) # with (k=6, min_sample=2)
```

#### Options:

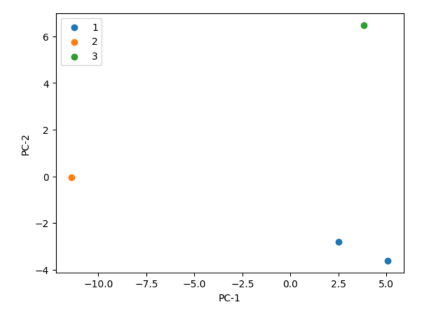
- k: kmer size
- min\_sample: threshold of considering a k-mer as feature

< dict{"matrix": numpy\_array, "sample": seqID\_list, "kmers": kmers\_list} >

Output: returns a dict object containing following keys ("matrix", "sample", "kmers"), where "kmers" are columns and "sample" are rows

```
# Visualizing feature matrix
         import pandas as pd
         pd.DataFrame(feature_matrix["matrix"], index=feature_matrix["sample"], columns=feature_matrix["kmers"])
Out[47...
               RLL GDL DLS GNP GKK DGL LKG KGT FWK AGA ... DAG PGY GYP YPG GAK AKG GFK FKG GRD RDG
                                                                  0 ...
                                                                                               0
                                                                                                               0
          seq1
                  1
                                                       1
                                                            0
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                                                                                     Ω
                                                                                          0
                                                                                                     0
                                                                                                          0
                                                                                                                     0
                                                                                                                          0
          seq2
                  1
                            0
                                       0
                                            0
                                                       0
                                                            2
                                                                          0
                                                                               0
                                                                                     0
                                                                                          0
                                                                                                0
                                                                                                     0
                                                                                                          0
                                                                                                               0
                                                                                                                     0
                                                                                                                          0
          seq3
                  0
                       0
                            0
                                  1
                                       1
                                            1
                                                  1
                                                       1
                                                            0
                                                                  1 ...
                                                                          2
                                                                               2
                                                                                          1
                                                                                               0
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                                                                                                                          0
          seq4
                  0
                       0
                                  0
                                       O
                                            0
                                                       0
                                                            0
                                                                  0 ...
                                                                          0
                                                                               0
                                                                                     2
                                                                                          2
                                                                                                2
                                                                                                     2
                                                                                                          2
                                                                                                               2
                                                                                                                     2
                                                                                                                          2
         4 rows × 84 columns
         2.17 Principal component analysis
             >>> from segplotter analysis import PCA
             >>> pca_matrix = PCA(feature_matrix) # return pca matrix with first 2 PCs
             >>> pca_matrix = PCA(feature_matrix, 3) # return pca matrix with first 3 PCs
         Output: returns a dict object containing following keys ("matrix", "sample", "components"), where "components" are columns and
          "sample" are rows
          < dict{"matrix": numpy_array, "sample": seqID_list, "components": PCA_components} >
In [48...
         # Importing module
         from seqplotter_analysis import PCA
         # Performing PCA
         pca_matrix = PCA(feature_matrix)
         # Visualizing PCA matrix
         import pandas as pd
         pd.DataFrame(pca_matrix["matrix"], index=pca_matrix["sample"], columns=pca_matrix["components"])
Out[48...
                    PC-1
                             PC-2
          seq1
                5.069106 -3.616282
          seq2
                 2.496495 -2.811089
          seq3 -11.398262 -0.046961
                 3.832661 6.474332
          seq4
         2.18. Plot scatter plot using PCs
             >>> from seqplotter_analysis import plot_pca
             # Sequence IDs will be treated as class label (default)
             >>> plot_pca(pca_matrix)
             # Providing class labels as list
             >>> labels = [3, 3, 3, 2, 1, 4, 1, 1, 1]
             >>> plot_pca(pca_matrix, labels)
         Output: generate PCA plot (2D scatter plot using first two components i.e. PC-1 and PC-2)
In [49...
         # Importing module
         from seqplotter_analysis import plot_pca
         # Plotting 2D PCA plot
```

labels = [1, 1, 2, 3]
plot\_pca(pca\_matrix, labels)



#### 2.19. Create sequence logo

```
>>> from seqplotter_analysis import seq_logo

# Reading motif sequences file
>>> logo_data = PROT.read_fasta("motif_seq.fasta")

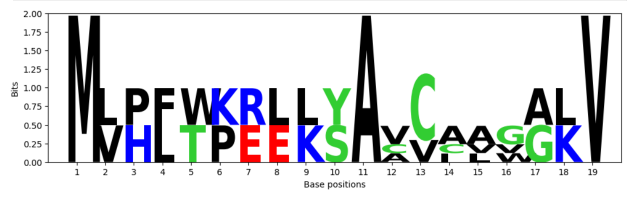
# Generate sequence logo
>>> seq_logo(logo_data, seq_type)
```

Output: generate protein sequence logo using motif sequences

```
In [50... # Importing module
    from seqplotter_analysis import seq_logo

# Reading motif sequences file
    data = PROT.read_fasta("prot_motif_seq.fasta")

# Generate sequence logo
    seq_logo(data, "PROT")
```



# 2.20. Generate hydrophobicity plot

```
>>> from seqplotter_analysis import hydrophobicity_plot
# Amino acid sequence for seq2 seqID
>>> seq = PROT.where(seqid="seq2")
# Create hydrophobicity plot
>>> hydrophobicity_plot(seq, window_size)
```

Output: generate hydrophobicity plot for a given protein sequence

```
In [51... from seqplotter_analysis import hydrophobicity_plot
    # Amino acid sequence for seq2 seqID
    seq = PROT.where(seqid="seq2")
```

