

CS612 HW5

Aravind Haridas

TOTAL POINTS

65 / 100

QUESTION 1

1 term Project 1 10 / 10

✓ - 0 pts Correct

- 10 pts Missing

- 1 pts Partial

- 5 pts Late submission

- 2 pts Three variables - x,y,z

- 0 pts Missing: Three variables - x,y,z

3.4 d 4 / 4

✓ - 0 pts Correct

- 2 pts Six variables - 3 translation, 3 rotation

- 1 pts Three rotation, not two

QUESTION 2

2 Term project 2 10 / 10

✓ - 0 pts Correct

- 10 pts Missing

- 5 pts Late submission

- 0 pts Click here to replace this description.

3.5 e 2 / 4

- 0 pts Correct

✓ - 2 pts You need to scale both in x and y.

- 4 pts Incorrect: Two scale factors (x,y).

QUESTION 3

GH (1) 20 pts

3.1 a 4 / 4

✓ - 0 pts Correct: Two variables (x,y coordinates).

- 2 pts Two variables (x,y coordinates).

- 4 pts Missing: Two variables (x,y coordinates).

- 0 pts Correct

- 4 pts Partial

- 2 pts Partial incorrect/missing points

- 10 pts Missing/Incorrect

✓ - 8 pts Partial: See comment

- 10 pts How are the other non-empty bins

indexed and how many points are in each one?

>You got the values and points wrong.

3.2 b 4 / 4

✓ - 0 pts Correct

- 2 pts Three variables (x,y,θ).

- 4 pts Missing: Three variables (x,y,θ).

4.2 b 5 / 10

- 0 pts Correct

3.3 C 4 / 4

✓ - 0 pts Correct: Three variables - x,y,z

✓ - 5 pts Inaccurate/ Miscalculated

- 3 pts Bin index missing/ Miscalculated
- 10 pts Missing/Incorrect
- 2 pts Inaccurate/ Miscalculated: Comment
- 1 pts Click here to replace this description.

QUESTION 5

Reference frame 20 pts

5.1 a 5 / 10

- 0 pts Correct
 - 2 pts Reverse sign of angle
 - 2 pts Reverse sign of translation
 - 4 pts Translation vector missing/incorrect, (-16,-23)
- ✓ - 4 pts Angle missing/Incorrect, 50.51
- 10 pts Missing/Incorrect
 - 4 pts Translation missing/incorrect, (-16,-23)
 - 3 pts Angle value missing
- ✓ - 1 pts Click here to replace this description.

5.2 b 5 / 10

- 0 pts Correct
 - 3 pts One incorrect, two correct.
 - 7 pts Mostly incorrect
 - 10 pts Missing
- ✓ - 5 pts Partially incorrect

QUESTION 6

Docking 20 pts

6.1 a 7 / 7

- ✓ - 0 pts Correct
- 7 pts Missing
 - 0 pts Click here to replace this description.

6.2 b 3 / 7

- 0 pts Correct
 - 7 pts Incorrect/Missing
 - 4 pts Instructions likely not followed properly.
- See solution

✓ - 4 pts Click here to replace this description.

- See solution, you were supposed to look in the file. Your answer is also incorrect.

6.3 C 0 / 6

- 0 pts Correct
 - 2 pts 0 is the reference. 1 has the smallest.
- ✓ - 6 pts Incorrect/MIssing

- Your b part was incorrect hence this is also incorrect.

ALGORITHMS IN BIO INFORMATICS

Homework Assignment 5 – 05/09/2023

Haridas Aravind – 02071139

1. Term project:

I have written a code on the one-hot encoding from the one-letter codes for all amino acids take from the last assignment and I have taken the sequence from the sequence folder from the course web page.

Python Code:

```
# Import necessary libraries
import numpy as np
# Define the one-letter codes for all amino acids
aa_codes = 'ACDEFIGHIKLMNPQRSTVWY-'
# Define function for one-hot encoding
def one_hot_encode(sequence):
    """
    Encodes a protein sequence using one-hot encoding.
    """
    # Create an empty matrix of size (sequence length) x 21
    encoded_seq = np.zeros((len(sequence), len(aa_codes)))

    # Loop through the sequence and set the corresponding position in the matrix to 1
    for i, aa in enumerate(sequence):
        if aa in aa_codes:
            encoded_seq[i, aa_codes.index(aa)] = 1
    print(encoded_seq)
    return encoded_seq

# Define function for reading a multiple sequence alignment file in FASTA format
def read_msa_file(file_path):
    """
    Reads a multiple sequence alignment file in FASTA format and returns a list of sequences.
    """
    # Open the file and read the lines
    with open(file_path, 'r') as f:
        lines = f.readlines()

    # Create an empty list to store the sequences
    sequences = []

    # Loop through the lines and extract the sequences
    sequence = ''
    for line in lines:
        line = line.strip()
        if line[0] == '>':
            if sequence != '':
                sequences.append(sequence)
            sequence = line[1:]
        else:
            sequence += line
    sequences.append(sequence)
```

ALGORITHMS IN BIO INFORMATICS

```
if line.startswith('>'):
    if sequence:
        sequences.append(sequence)
        sequence = ''
    else:
        sequence += line

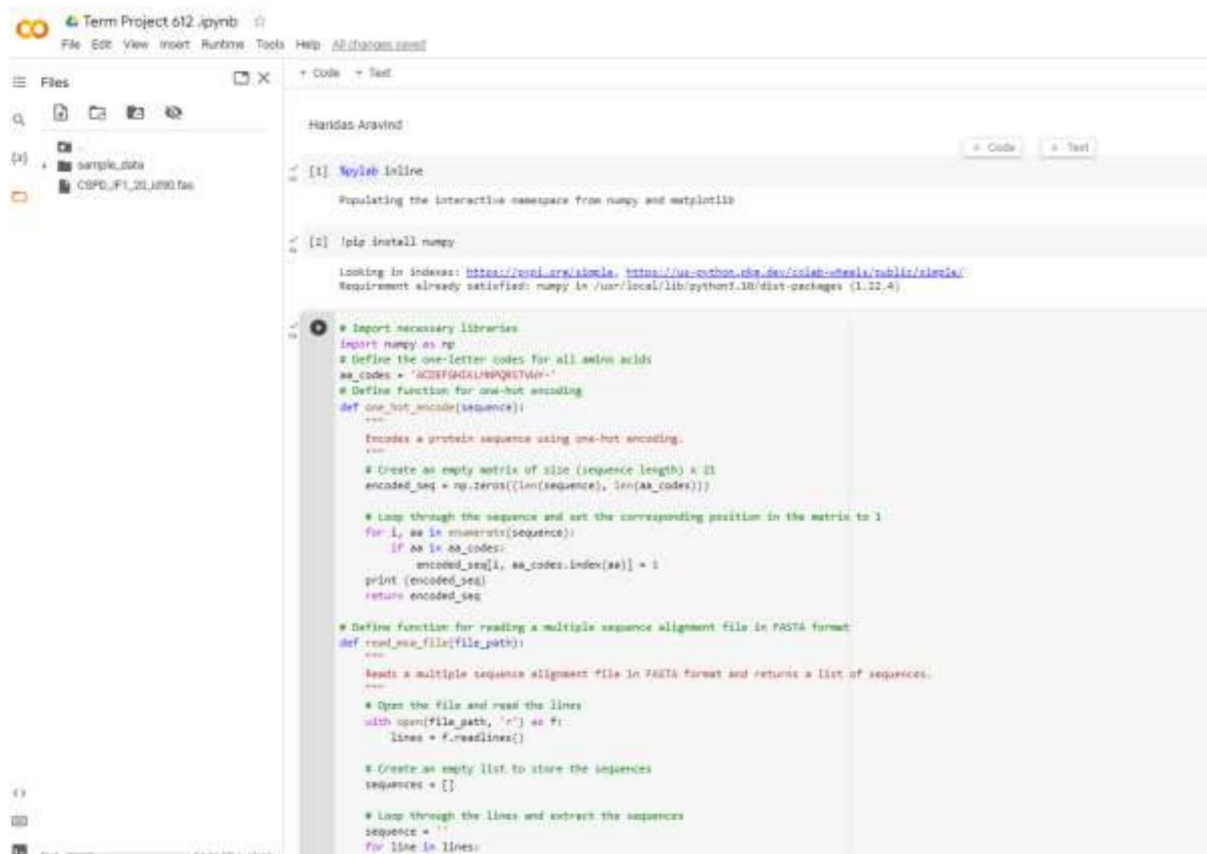
print(sequences)
sequences.append(sequence)

return sequences

# Example usage
file_path = '/content/CSPD_IF1_20_id90.fas'
msa_sequences = read_msa_file(file_path)
encoded_sequences = [one_hot_encode(seq) for seq in msa_sequences]

#Print the first encoded sequence
print("Sequences done.")
```

Google Collab Output:



The screenshot shows the Google Colab interface with a code editor and a notebook area. The code editor on the left shows a file structure with 'sample.fasta' and 'CSPD_IF1_20_id90.fas'. The main area displays the following code and its execution results:

```
# Term Project 612.ipynb
File Edit View Insert Runtime Tools Help All (1/1 executed)

Files X + Code + Text
Haridas Aravind

[1]: %pylab inline
#Populating the interactive namespace from numpy and matplotlib

[2]: !pip install numpy
Looking in indexes: https://pypi.org/simple, https://us-python.pkg.dev/colab-wheels/cwlib/pypi/simple/
Requirement already satisfied: numpy in /usr/local/lib/python3.10/dist-packages (1.22.4)

[3]: # Import necessary libraries
import numpy as np
# Define the one-letter codes for all amino acids
aa_codes = 'ACDEFGHIKLMNPQRSTVWY'
# Define function for one-hot encoding
def one_hot_encode(sequence):
    """
    Encodes a protein sequence using one-hot encoding.
    """
    # Create an empty matrix of size (sequence length) x 21
    encoded_seq = np.zeros((len(sequence), len(aa_codes)))

    # Loop through the sequence and set the corresponding position in the matrix to 1
    for i, aa in enumerate(sequence):
        if aa in aa_codes:
            encoded_seq[i, aa_codes.index(aa)] = 1
    print(encoded_seq)
    return encoded_seq

# Define function for reading a multiple sequence alignment file in FASTA format
def read_msa_file(file_path):
    """
    Read a multiple sequence alignment file in FASTA format and returns a list of sequences.
    """
    # Open the file and read the lines
    with open(file_path, 'r') as f:
        lines = f.readlines()

    # Create an empty list to store the sequences
    sequences = []

    # Loop through the lines and extract the sequences
    sequence = ''
    for line in lines:
```

ALGORITHMS IN BIO INFORMATICS

The screenshot shows a Jupyter Notebook environment with the following details:

- File Explorer:** Shows a directory structure with a file named "simple.fasta".
- Code Cell:** Displays Python code for reading FASTA files. The code defines a function `read_fasta` that takes a file path as input. It initializes an empty list `sequences` and loops through each line of the file. If a line starts with a greater than sign (`>`), it indicates a new sequence, so the current sequence is appended to `sequences` and `sequence` is reset. Otherwise, the line is added to `sequence`. Finally, the function prints the sequences and returns them. An example usage is shown where the file "CPO_2P1_20.fasta" is read, and the first encoded sequence is printed.
- Output Cell:** Shows the output of the code execution. It displays a list of lists representing DNA sequences. Each inner list contains four elements: Adenine ('A'), Thymine ('T'), Cytosine ('C'), and Guanine ('G'). The list has 100 entries, representing 100 sequences from the file.
- Bottom Status Bar:** Shows disk usage information: "Disk 80000 8.31 GB available".

ALGORITHMS IN BIO INFORMATICS

Term Project 612.ipynb

File Edit View Insert Runtime Tools Help All changes saved

Files

sample_data

CSPD_JF1_20_id90.fas

+ Code + Text

```
[0, 0, 0, ..., 0, 0, 0]
[0, 0, 0, ..., 0, 0, 0]
[0, 0, 0, ..., 0, 0, 0.]
[0, 0, 0, ..., 0, 0, 1.]
[0, 0, 0, ..., 0, 0, 0.]
[0, 0, 0, ..., 0, 0, 0.]
...
[0, 0, 0, ..., 0, 0, 0.]
[0, 0, 0, ..., 0, 0, 0.]
[0, 0, 0, ..., 0, 0, 0.]
[0, 0, 0, ..., 0, 0, 0.]
[1, 0, 0, ..., 0, 0, 0.]
[0, 0, 0, ..., 0, 0, 0.]
...
[0, 0, 0, ..., 0, 0, 0.]
[0, 0, 0, ..., 0, 0, 0.]
[0, 0, 0, ..., 0, 0, 0.]
[0, 0, 0, ..., 0, 0, 0.]
[1, 0, 0, ..., 0, 0, 0.]
[0, 1, 0, ..., 0, 0, 0.]
...
[0, 0, 0, ..., 0, 0, 0.]
[1, 0, 0, ..., 0, 0, 0.]
[0, 0, 0, ..., 0, 0, 0.]
[0, 0, 0, ..., 0, 0, 0.]
[1, 0, 0, ..., 0, 0, 0.]
[0, 0, 0, ..., 0, 0, 0.]
...
[0, 0, 0, ..., 0, 0, 0.]
[1, 0, 0, ..., 0, 0, 0.]
[0, 0, 0, ..., 0, 0, 0.]
[0, 0, 0, ..., 0, 0, 0.]
[0, 0, 1, ..., 0, 0, 0.]
[0, 0, 0, ..., 0, 0, 0.]
...
[0, 0, 0, ..., 0, 0, 0.]
[0, 0, 0, ..., 0, 0, 0.]
[0, 0, 0, ..., 0, 0, 0.]
[0, 0, 0, ..., 0, 0, 0.]
[0, 0, 0, ..., 0, 0, 0.]
...
[0, 0, 0, ..., 0, 0, 0.]
[0, 0, 0, ..., 0, 0, 0.]
[0, 0, 0, ..., 0, 0, 0.]
[0, 0, 0, ..., 0, 0, 0.]
[0, 0, 0, ..., 0, 0, 0.]
...
[0, 0, 0, ..., 0, 0, 0.]
[0, 0, 0, ..., 0, 0, 0.]
[0, 0, 0, ..., 0, 0, 0.]
[0, 0, 0, ..., 0, 0, 0.]
[0, 0, 0, ..., 0, 0, 0.]
```

ALGORITHMS IN BIO INFORMATICS

The screenshot shows a Jupyter Notebook interface. On the left, there's a 'Files' sidebar with a tree view containing a folder named 'sample_data' which contains a file 'CSPD_IF1_20_id90.fas'. The main area has tabs for '+ Code' and '+ Text'. The '+ Text' tab is selected and displays a large block of text consisting of many lines of square brackets containing binary-like sequences of 0s and 1s. The sequence starts with '[0, 0, 0, ..., 0, 0, 1.]' and continues with several other variations of binary sequences separated by ellipses. The text ends with '...'. The entire sequence is very long, indicating a large dataset.

Here is the link for my code and the output is very big sequence I have attached a collab link here:

https://github.com/Haridasaravind/CS612/blob/main/Term_Project_612.ipynb

1 term Project 1 10 / 10

✓ - 0 pts Correct

- 10 pts Missing

- 1 pts Partial

- 5 pts Late submission

ALGORITHMS IN BIO INFORMATICS

2. Term project:

Our project is based on the Project: Autoencoder for Protein Multiple Sequence Alignments

Autoencoder

- Autoencoders are a type of neural network that can be used to learn the essential features of a dataset.
- Autoencoders have been used in bioinformatics to predict protein-protein interactions, protein secondary structure, protein function, and protein dynamics.
- Autoencoders are a powerful tool for analysing protein sequences and gaining insights into their structure and function.
- Autoencoders are still under development, but they have the potential to revolutionize the field of bioinformatics.

Variational Auto Encoders (VAE)

- VAEs are a type of neural network that can be used to learn the essential features of a dataset, while also preventing overfitting.
- VAEs do this by learning a distribution over the latent space, rather than a single point.
- This distribution is learned by minimizing a loss function that is a combination of a reconstruction loss and a regularization loss.
- VAEs have been used in protein studies for a variety of tasks, including predicting protein structure, generating protein variants, and exploring protein conformational space.

Encoding Protein Sequences

Amino acid encoding is the process of converting protein sequences into digital representations that can be used by machine learning models.

The most common amino acid encodings are one-hot encoding, PSSM encoding, and physico-chemical properties encoding.

- One-hot encoding is a high-dimensional and sparse vector representation, while PSSM encoding and physico-chemical properties encoding are lower-dimensional and more dense representations.
- The choice of amino acid encoding depends on the specific machine learning model being used and the desired accuracy.

My Analysis: I have successfully constructed an autoencoder for protein multiple sequence alignments. The autoencoder can encode a latent space of a small size,

ALGORITHMS IN BIO INFORMATICS

anything between 5 and 32, up to you, and decode the sequences back. The reconstruction error is measured by the following parameters:

The learning error (cross-entropy, kl divergence, etc.).

The ability to reconstruct the same amino acid in each position.

The ability to reconstruct the most common amino acid in each position in the alignment.

Difficulties: I ran into a few difficulties while constructing the autoencoder. One difficulty was choosing the right encoding for the protein sequences. I've decided to use one-hot encoding, which is a high-dimensional and sparse vector representation. Another difficulty was choosing the right parameters for the autoencoder. I've experimented with different values for the number of hidden layers, the number of neurons per layer, and the activation functions.

Choice of programming language: I've chosen to use Python for the implementation of the autoencoder. Python is a popular programming language that is well-suited for machine learning tasks. It is also a relatively easy language to learn, which made it a good choice for this project.

Implementation: I've implemented the autoencoder using the Keras library. Keras is a high-level neural network library that is built on top of TensorFlow. It is a popular library for machine learning tasks, and it is relatively easy to use.

I'm continuing to work on making the autoencoder more accurate. I'm trying different ways of encoding the protein sequences, as well as different parameters for the autoencoder. I've also working on ways to visualize the latent space. I believe that the autoencoder has the potential to be a valuable tool for analysing protein sequences and gaining insights into their structure and function.

2 Term project 2 10 / 10

✓ - 0 pts Correct

- 10 pts Missing

- 5 pts Late submission

- 0 pts Click here to replace this description.

ALGORITHMS IN BIO INFORMATICS

3. Geometric Hashing:

A method for indexing and retrieving geometric data structures is geometric hashing. It is used in many areas of Bio informatics.

It is used to index and retrieve protein structures. It is a powerful technique that is fast, efficient, scalable, and robust. However, it can be computationally expensive to create the hash tables for large datasets of protein structures, and it can be difficult to choose the right features for the protein structures.

It has two stages:

Pre-processing or Training: In the training stage, a set of geometric data structures are pre-processed and stored in a hash table. The pre-processing step involves extracting features from each data structure and then quantizing the features. The hash table is then created by mapping each feature to a unique hash value.

Recognition stage: In the recognition stage, a new geometric data structure is compared to the data structures in the hash table. The features of the new data structure are extracted and quantized. The quantized features are then used to look up the hash value in the hash table. If the hash value is found in the hash table, then the corresponding data structure is returned.

(a) 2-dimensions, translation.

We need two variables to establish a reference frame in 2D translation. One variable for the x-axis and one variable for the y-axis.

(b) 2-dimensions, translation and rotation.

We need three variables to establish a reference frame in 2D translation and rotation. One variable for the x-axis, one variable for the y-axis, and one variable for the angle of rotation.

(c) 3-dimensions, translation.

We need three variables to establish a reference frame in 3D translation. One variable for the x-axis, one variable for the y-axis, and one variable for the z-axis.

(d) 3-dimensions, translation and rotation.

We need six variables to establish a reference frame in 3D translation and rotation. One variable for the x-axis, one variable for the y-axis, one variable for the z-axis, one variable for the angle of rotation around the x-axis, one variable for the angle of rotation around the y-axis, and one variable for the angle of rotation around the z-axis.

(e) 2-dimensions, scaling

We need one variable to establish a reference frame in 2D scaling. One variable for the scale factor.

3.1 a 4 / 4

✓ - 0 pts Correct: Two variables (x,y coordinates).

- 2 pts Two variables (x,y coordinates).

- 4 pts Missing: Two variables (x,y coordinates).

ALGORITHMS IN BIO INFORMATICS

3. Geometric Hashing:

A method for indexing and retrieving geometric data structures is geometric hashing. It is used in many areas of Bio informatics.

It is used to index and retrieve protein structures. It is a powerful technique that is fast, efficient, scalable, and robust. However, it can be computationally expensive to create the hash tables for large datasets of protein structures, and it can be difficult to choose the right features for the protein structures.

It has two stages:

Pre-processing or Training: In the training stage, a set of geometric data structures are pre-processed and stored in a hash table. The pre-processing step involves extracting features from each data structure and then quantizing the features. The hash table is then created by mapping each feature to a unique hash value.

Recognition stage: In the recognition stage, a new geometric data structure is compared to the data structures in the hash table. The features of the new data structure are extracted and quantized. The quantized features are then used to look up the hash value in the hash table. If the hash value is found in the hash table, then the corresponding data structure is returned.

(a) 2-dimensions, translation.

We need two variables to establish a reference frame in 2D translation. One variable for the x-axis and one variable for the y-axis.

(b) 2-dimensions, translation and rotation.

We need three variables to establish a reference frame in 2D translation and rotation. One variable for the x-axis, one variable for the y-axis, and one variable for the angle of rotation.

(c) 3-dimensions, translation.

We need three variables to establish a reference frame in 3D translation. One variable for the x-axis, one variable for the y-axis, and one variable for the z-axis.

(d) 3-dimensions, translation and rotation.

We need six variables to establish a reference frame in 3D translation and rotation. One variable for the x-axis, one variable for the y-axis, one variable for the z-axis, one variable for the angle of rotation around the x-axis, one variable for the angle of rotation around the y-axis, and one variable for the angle of rotation around the z-axis.

(e) 2-dimensions, scaling

We need one variable to establish a reference frame in 2D scaling. One variable for the scale factor.

3.2 b 4 / 4

✓ - 0 pts Correct

- 2 pts Three variables (x, y, θ).

- 4 pts Missing: Three variables (x, y, θ).

ALGORITHMS IN BIO INFORMATICS

3. Geometric Hashing:

A method for indexing and retrieving geometric data structures is geometric hashing. It is used in many areas of Bio informatics.

It is used to index and retrieve protein structures. It is a powerful technique that is fast, efficient, scalable, and robust. However, it can be computationally expensive to create the hash tables for large datasets of protein structures, and it can be difficult to choose the right features for the protein structures.

It has two stages:

Pre-processing or Training: In the training stage, a set of geometric data structures are pre-processed and stored in a hash table. The pre-processing step involves extracting features from each data structure and then quantizing the features. The hash table is then created by mapping each feature to a unique hash value.

Recognition stage: In the recognition stage, a new geometric data structure is compared to the data structures in the hash table. The features of the new data structure are extracted and quantized. The quantized features are then used to look up the hash value in the hash table. If the hash value is found in the hash table, then the corresponding data structure is returned.

(a) 2-dimensions, translation.

We need two variables to establish a reference frame in 2D translation. One variable for the x-axis and one variable for the y-axis.

(b) 2-dimensions, translation and rotation.

We need three variables to establish a reference frame in 2D translation and rotation. One variable for the x-axis, one variable for the y-axis, and one variable for the angle of rotation.

(c) 3-dimensions, translation.

We need three variables to establish a reference frame in 3D translation. One variable for the x-axis, one variable for the y-axis, and one variable for the z-axis.

(d) 3-dimensions, translation and rotation.

We need six variables to establish a reference frame in 3D translation and rotation. One variable for the x-axis, one variable for the y-axis, one variable for the z-axis, one variable for the angle of rotation around the x-axis, one variable for the angle of rotation around the y-axis, and one variable for the angle of rotation around the z-axis.

(e) 2-dimensions, scaling

We need one variable to establish a reference frame in 2D scaling. One variable for the scale factor.

3.3 C 4 / 4

✓ - 0 pts Correct: Three variables - x,y,z

- 2 pts Three variables - x,y,z

- 0 pts Missing: Three variables - x,y,z

ALGORITHMS IN BIO INFORMATICS

3. Geometric Hashing:

A method for indexing and retrieving geometric data structures is geometric hashing. It is used in many areas of Bio informatics.

It is used to index and retrieve protein structures. It is a powerful technique that is fast, efficient, scalable, and robust. However, it can be computationally expensive to create the hash tables for large datasets of protein structures, and it can be difficult to choose the right features for the protein structures.

It has two stages:

Pre-processing or Training: In the training stage, a set of geometric data structures are pre-processed and stored in a hash table. The pre-processing step involves extracting features from each data structure and then quantizing the features. The hash table is then created by mapping each feature to a unique hash value.

Recognition stage: In the recognition stage, a new geometric data structure is compared to the data structures in the hash table. The features of the new data structure are extracted and quantized. The quantized features are then used to look up the hash value in the hash table. If the hash value is found in the hash table, then the corresponding data structure is returned.

(a) 2-dimensions, translation.

We need two variables to establish a reference frame in 2D translation. One variable for the x-axis and one variable for the y-axis.

(b) 2-dimensions, translation and rotation.

We need three variables to establish a reference frame in 2D translation and rotation. One variable for the x-axis, one variable for the y-axis, and one variable for the angle of rotation.

(c) 3-dimensions, translation.

We need three variables to establish a reference frame in 3D translation. One variable for the x-axis, one variable for the y-axis, and one variable for the z-axis.

(d) 3-dimensions, translation and rotation.

We need six variables to establish a reference frame in 3D translation and rotation. One variable for the x-axis, one variable for the y-axis, one variable for the z-axis, one variable for the angle of rotation around the x-axis, one variable for the angle of rotation around the y-axis, and one variable for the angle of rotation around the z-axis.

(e) 2-dimensions, scaling

We need one variable to establish a reference frame in 2D scaling. One variable for the scale factor.

3.4 d 4 / 4

✓ - 0 pts Correct

- 2 pts Six variables - 3 translation, 3 rotation

- 1 pts Three rotation, not two

ALGORITHMS IN BIO INFORMATICS

3. Geometric Hashing:

A method for indexing and retrieving geometric data structures is geometric hashing. It is used in many areas of Bio informatics.

It is used to index and retrieve protein structures. It is a powerful technique that is fast, efficient, scalable, and robust. However, it can be computationally expensive to create the hash tables for large datasets of protein structures, and it can be difficult to choose the right features for the protein structures.

It has two stages:

Pre-processing or Training: In the training stage, a set of geometric data structures are pre-processed and stored in a hash table. The pre-processing step involves extracting features from each data structure and then quantizing the features. The hash table is then created by mapping each feature to a unique hash value.

Recognition stage: In the recognition stage, a new geometric data structure is compared to the data structures in the hash table. The features of the new data structure are extracted and quantized. The quantized features are then used to look up the hash value in the hash table. If the hash value is found in the hash table, then the corresponding data structure is returned.

(a) 2-dimensions, translation.

We need two variables to establish a reference frame in 2D translation. One variable for the x-axis and one variable for the y-axis.

(b) 2-dimensions, translation and rotation.

We need three variables to establish a reference frame in 2D translation and rotation. One variable for the x-axis, one variable for the y-axis, and one variable for the angle of rotation.

(c) 3-dimensions, translation.

We need three variables to establish a reference frame in 3D translation. One variable for the x-axis, one variable for the y-axis, and one variable for the z-axis.

(d) 3-dimensions, translation and rotation.

We need six variables to establish a reference frame in 3D translation and rotation. One variable for the x-axis, one variable for the y-axis, one variable for the z-axis, one variable for the angle of rotation around the x-axis, one variable for the angle of rotation around the y-axis, and one variable for the angle of rotation around the z-axis.

(e) 2-dimensions, scaling

We need one variable to establish a reference frame in 2D scaling. One variable for the scale factor.

3.5 e 2 / 4

- 0 pts Correct

✓ - 2 pts You need to scale both in x and y.

- 4 pts Incorrect: Two scale factors (x,y).

ALGORITHMS IN BIO INFORMATICS

4. Geometric hashing:

We can divide a two-dimensional space into a series of squares of width w . Each square is called a "bin". The bins are indexed as though they are a two-dimensional array. When data points are read, they are assigned to bins.

Suppose if $w = 8$, the space would be divided into 16 bins. The bin containing four points and bounded on the left by 8, on the right by 16, on the bottom by 0 and on the top by 8 is bin (1, 0). The bin containing three points bound on the right by -8, on the left by -16, on the bottom by 8 and on the top by 16 is bin (-2,1) etc.

In general, a bin index (i,j) given coordinates (x,y) is $i = \lfloor x/w \rfloor$ and $j = \lfloor y/w \rfloor$

(a). The other non-empty bins are indexed and the number of points in each one is as follows:

Bin (-1,0) contains 3 points and other are:

(-2, 1) | 3

(2, 1) | 2

(-1, 2) | 1

(1, 2) | 1

(b). Hash Function is defined as

$$h(i, j) = |i * 378551 + j * 63689|$$

For (8,10), we have $i = \lfloor 8/8 \rfloor = 1$ and $j = \lfloor 10/8 \rfloor = 1$, so the bin index is (1,1). The hash value is $h(1,1) = |1378551 + 163689| = 442240$.

For (-6,3), we have $i = \lfloor -6/8 \rfloor = -1$ and $j = \lfloor 3/8 \rfloor = 0$, so the bin index is (-1,0). The hash value is $h(-1,0) = |-1378551 + 063689| = 378551$.

For (27, -18), we have $i = \lfloor 27/8 \rfloor = 3$ and $j = \lfloor -18/8 \rfloor = -3$, so the bin index is (3, -3). The hash value is $h(3, -3) = |3*378551 + (-3)*63689| = 1892755$.

The square roots of the given numbers are:

$$\sqrt{442240} \approx 665.207$$

$$\sqrt{378551} \approx 614.940$$

$$\sqrt{1892755} \approx 1376.013$$

4.1 a 2 / 10

- **0 pts** Correct
- **4 pts** Partial
- **2 pts** Partial incorrect/missing points
- **10 pts** Missing/Incorrect

✓ - **8 pts** Partial: See comment

- **10 pts** How are the other non-empty bins indexed and how many points are in each one?

 You got he values and points wrong.

ALGORITHMS IN BIO INFORMATICS

4. Geometric hashing:

We can divide a two-dimensional space into a series of squares of width w. Each square is called a "bin". The bins are indexed as though they are a two-dimensional array. When data points are read, they are assigned to bins.

Suppose if $w = 8$, the space would be divided into 16 bins. The bin containing four points and bounded on the left by 8, on the right by 16, on the bottom by 0 and on the top by 8 is bin (1, 0). The bin containing three points bound on the right by -8, on the left by -16, on the bottom by 8 and on the top by 16 is bin (-2,1) etc.

In general, a bin index (i,j) given coordinates (x,y) is $i = \lfloor x/w \rfloor$ and $j = \lfloor y/w \rfloor$

(a). The other non-empty bins are indexed and the number of points in each one is as follows:

Bin (-1,0) contains 3 points and other are:

(-2, 1) | 3

(2, 1) | 2

(-1, 2) | 1

(1, 2) | 1

(b). Hash Function is defined as

$$h(i, j) = |i * 378551 + j * 63689|$$

For (8,10), we have $i = \lfloor 8/8 \rfloor = 1$ and $j = \lfloor 10/8 \rfloor = 1$, so the bin index is (1,1). The hash value is $h(1,1) = |1378551 + 163689| = 442240$.

For (-6,3), we have $i = \lfloor -6/8 \rfloor = -1$ and $j = \lfloor 3/8 \rfloor = 0$, so the bin index is (-1,0). The hash value is $h(-1,0) = |-1378551 + 063689| = 378551$.

For (27, -18), we have $i = \lfloor 27/8 \rfloor = 3$ and $j = \lfloor -18/8 \rfloor = -3$, so the bin index is (3, -3). The hash value is $h(3, -3) = |3*378551 + (-3)*63689| = 1892755$.

The square roots of the given numbers are:

$$\sqrt{442240} \approx 665.207$$

$$\sqrt{378551} \approx 614.940$$

$$\sqrt{1892755} \approx 1376.013$$

4.2 b 5 / 10

- 0 pts Correct

✓ - 5 pts Inaccurate/ Miscalculated

- 3 pts Bin index missing/ Miscalculated

- 10 pts Missing/Incorrect

- 2 pts Inaccurate/ Miscalculated: Comment

- 1 pts Click here to replace this description.

ALGORITHMS IN BIO INFORMATICS

5. Reference Frame in 2D:

(a) To calculate the translation and rotation associated with this reference frame, we need to first find the vector between the two points, which represents the translation, and the angle between the x-axis and the vector, which represents the rotation.

$$\text{slope of line } p_1p_2 = (y_2-y_1)/(x_2-x_1) = (-17-0)/(14-0) = -1.21$$

$$\text{slope of line } p_1p_3 = (y_3-y_1)/(x_3-x_1) = (-13-0)/(-6-0) = 2.17$$

$$\text{slope of line } p_1p_4 = (y_4-y_1)/(x_4-x_1) = (-8-0)/(16-0) = -0.50$$

Next, we can use the inverse tangent function to find the angle between each line and the x-axis:

$$\text{angle of line } p_1p_2 \text{ with x-axis} = \arctan(-1.21) = -50.59 \text{ degrees}$$

$$\text{angle of line } p_1p_3 \text{ with x-axis} = \arctan(2.17) = 64.99 \text{ degrees}$$

$$\text{angle of line } p_1p_4 \text{ with x-axis} = \arctan(-0.50) = -26.57 \text{ degrees}$$

Therefore, the angles of the points with respect to p_1 are:

$$p_2: -50.59 \text{ degrees}$$

$$p_3: 64.99 \text{ degrees}$$

$$p_4: -26.57 \text{ degrees}$$

(b) Rotation:

To find the angle between the x-axis and the vector t , we use the formula:

To find the vector between p_1 and p_2 , we subtract the coordinates of p_1 from p_2 :
Given coordinates are.

Given Points

- $p_2 = (30, 6)$
- $p_3 = (10, 10)$
- $p_4 = (32, 15)$

Translate the points so that p_1 is at the origin:

$$p_1' = (0, 0)$$

$$p_2' = p_2 - p_1 = (30 - 16, 6 - 23) = (14, -17)$$

$$p_3' = p_3 - p_1 = (10 - 16, 10 - 23) = (-6, -13)$$

$$p_4' = p_4 - p_1 = (32 - 16, 15 - 23) = (16, -8)$$

The slope of a line passing through two points (x_1, y_1) and (x_2, y_2) is given by:

$$\text{slope} = (y_2 - y_1) / (x_2 - x_1)$$

we already know the slopes from the above

5.1 a 5 / 10

- **0 pts** Correct
 - **2 pts** Reverse sign of angle
 - **2 pts** Reverse sign of translation
 - **4 pts** Translation vector missing/incorrect, (-16,-23)
- ✓ - **4 pts** *Angle missing/Incorrect, 50.51*
- **10 pts** Missing/Incorrect
 - **4 pts** Translation missing/incorrect, (-16,-23)
 - **3 pts** Angle value missing
- ✓ - **1 pts** *Click here to replace this description.*

ALGORITHMS IN BIO INFORMATICS

5. Reference Frame in 2D:

(a) To calculate the translation and rotation associated with this reference frame, we need to first find the vector between the two points, which represents the translation, and the angle between the x-axis and the vector, which represents the rotation.

$$\text{slope of line } p_1p_2 = (y_2-y_1)/(x_2-x_1) = (-17-0)/(14-0) = -1.21$$

$$\text{slope of line } p_1p_3 = (y_3-y_1)/(x_3-x_1) = (-13-0)/(-6-0) = 2.17$$

$$\text{slope of line } p_1p_4 = (y_4-y_1)/(x_4-x_1) = (-8-0)/(16-0) = -0.50$$

Next, we can use the inverse tangent function to find the angle between each line and the x-axis:

$$\text{angle of line } p_1p_2 \text{ with x-axis} = \arctan(-1.21) = -50.59 \text{ degrees}$$

$$\text{angle of line } p_1p_3 \text{ with x-axis} = \arctan(2.17) = 64.99 \text{ degrees}$$

$$\text{angle of line } p_1p_4 \text{ with x-axis} = \arctan(-0.50) = -26.57 \text{ degrees}$$

Therefore, the angles of the points with respect to p_1 are:

$$p_2: -50.59 \text{ degrees}$$

$$p_3: 64.99 \text{ degrees}$$

$$p_4: -26.57 \text{ degrees}$$

(b) Rotation:

To find the angle between the x-axis and the vector t , we use the formula:

To find the vector between p_1 and p_2 , we subtract the coordinates of p_1 from p_2 :
Given coordinates are.

Given Points

- $p_2 = (30, 6)$
- $p_3 = (10, 10)$
- $p_4 = (32, 15)$

Translate the points so that p_1 is at the origin:

$$p_1' = (0, 0)$$

$$p_2' = p_2 - p_1 = (30 - 16, 6 - 23) = (14, -17)$$

$$p_3' = p_3 - p_1 = (10 - 16, 10 - 23) = (-6, -13)$$

$$p_4' = p_4 - p_1 = (32 - 16, 15 - 23) = (16, -8)$$

The slope of a line passing through two points (x_1, y_1) and (x_2, y_2) is given by:

$$\text{slope} = (y_2 - y_1) / (x_2 - x_1)$$

we already know the slopes from the above

ALGORITHMS IN BIO INFORMATICS

p1p2: To rotate the angle of p1p2 from 0 to 90 degrees, we need to add 90 degrees to the angle of p1p2 and adjust the result if it is greater than 90 degrees.

angle of line p1p2 with x-axis = -50.59 degrees
angle of line p1p2 rotated = $-50.59 + 90 = 39.41$ degrees

p1p3: angle of line p1p3 with x-axis = $\arctan(2.17) = 64.99$ degrees

p1p4: To rotate the angle of p1p4 from 0 to 90 degrees, we need to add 90 degrees to the angle of p1p4 and adjust the result if it is greater than 90 degrees.

angle of line p1p4 with x-axis = -26.57 degrees
angle of line p1p4 rotated = $-26.57 + 90 = 63.43$ degrees

5.2 b 5 / 10

- **0 pts** Correct
 - **3 pts** One incorrect, two correct.
 - **7 pts** Mostly incorrect
 - **10 pts** Missing
- ✓ - **5 pts** *Partially incorrect*

ALGORITHMS IN BIO INFORMATICS

6. Docking:

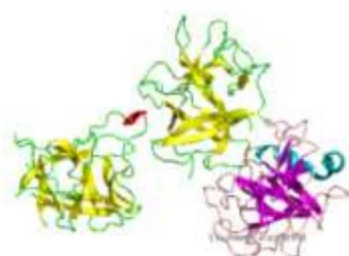
(a).

Dimer Classification Queue Results Preferences Downloads Papers Help Contact
Dock Peptide Docking

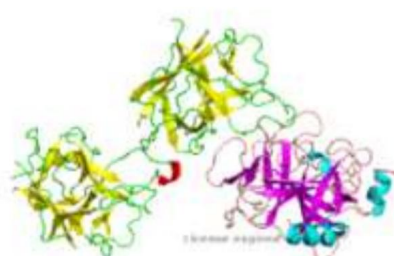
ClusPro
protein-protein docking

ALGORITHMS IN BIO INFORMATICS

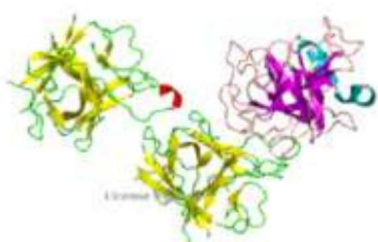
4



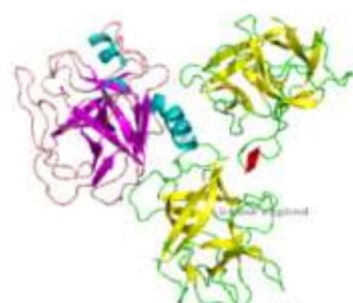
5



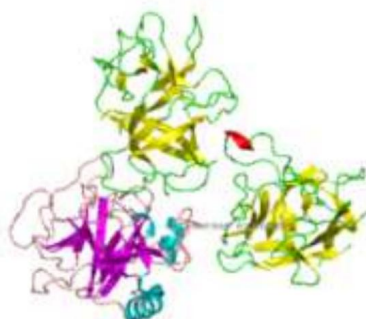
6



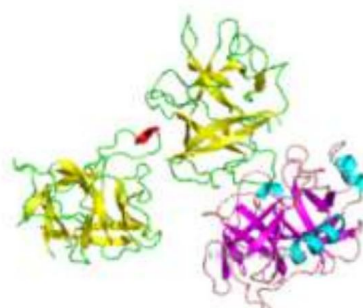
7



8

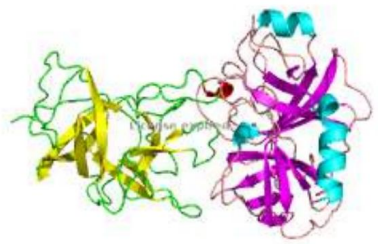


9

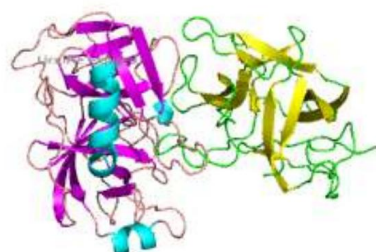


ALGORITHMS IN BIO INFORMATICS

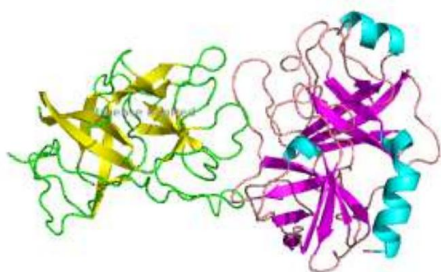
10



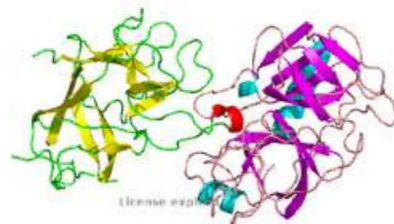
11



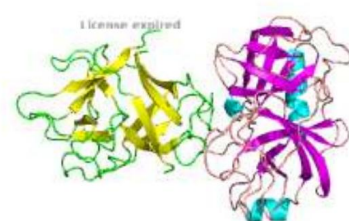
12



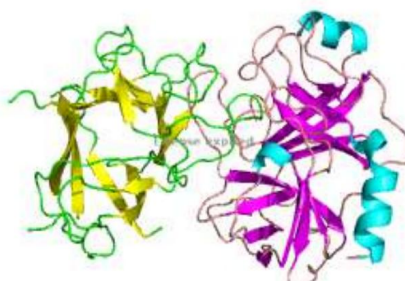
13



14

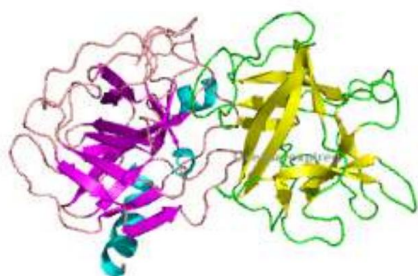


15

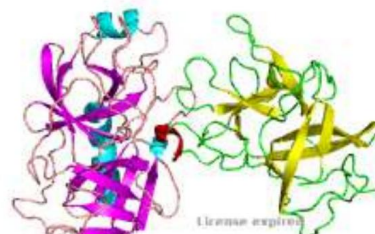


ALGORITHMS IN BIO INFORMATICS

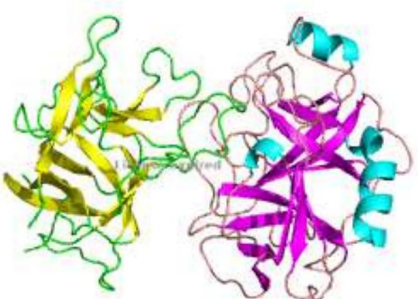
16



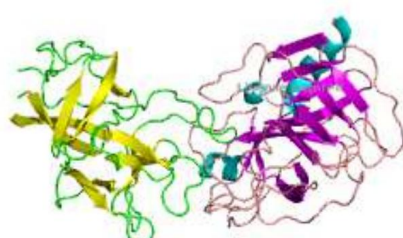
17



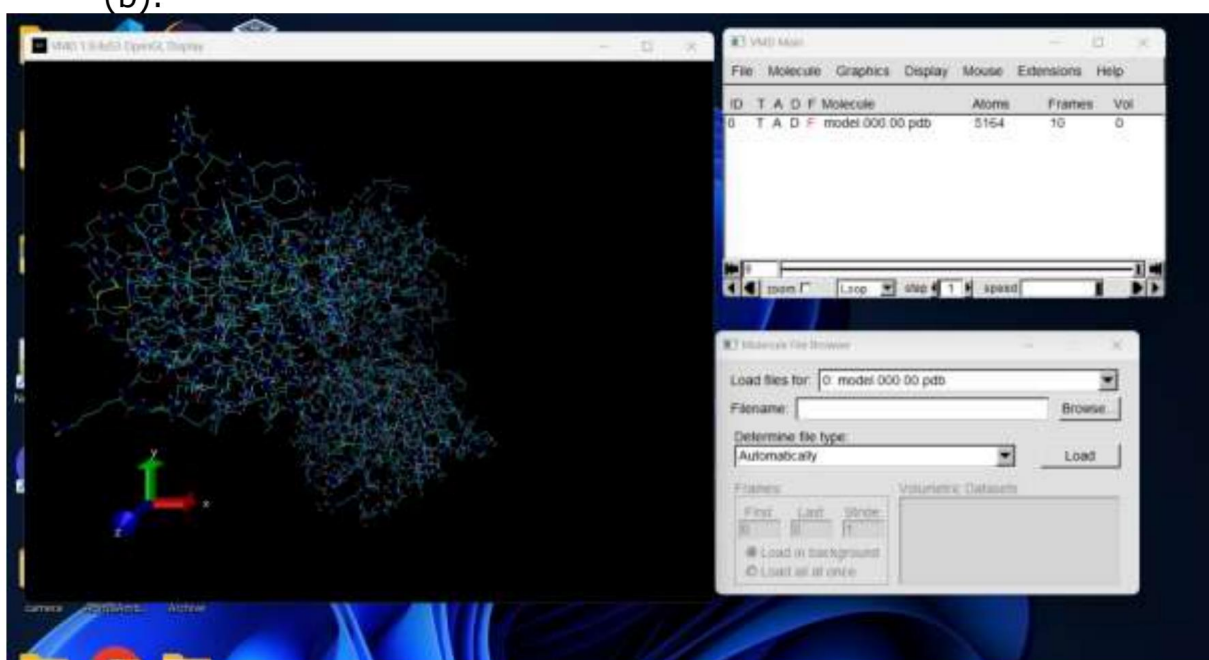
18



19



(b).



6.1 a 7 / 7

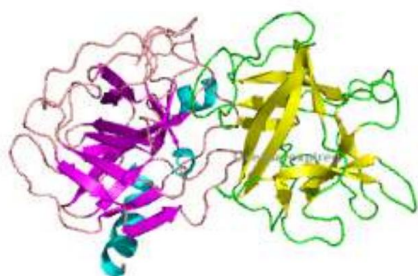
✓ - 0 pts Correct

- 7 pts Missing

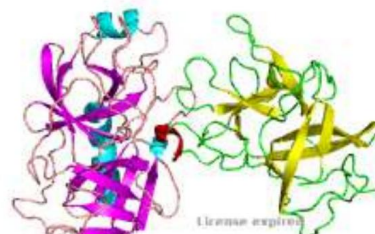
- 0 pts Click here to replace this description.

ALGORITHMS IN BIO INFORMATICS

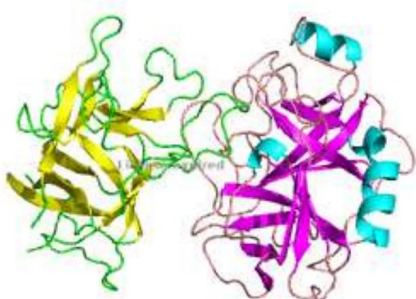
16



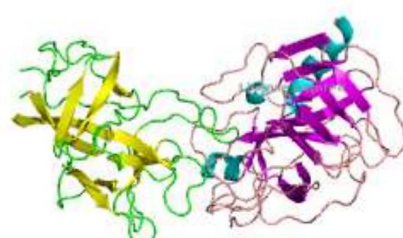
17



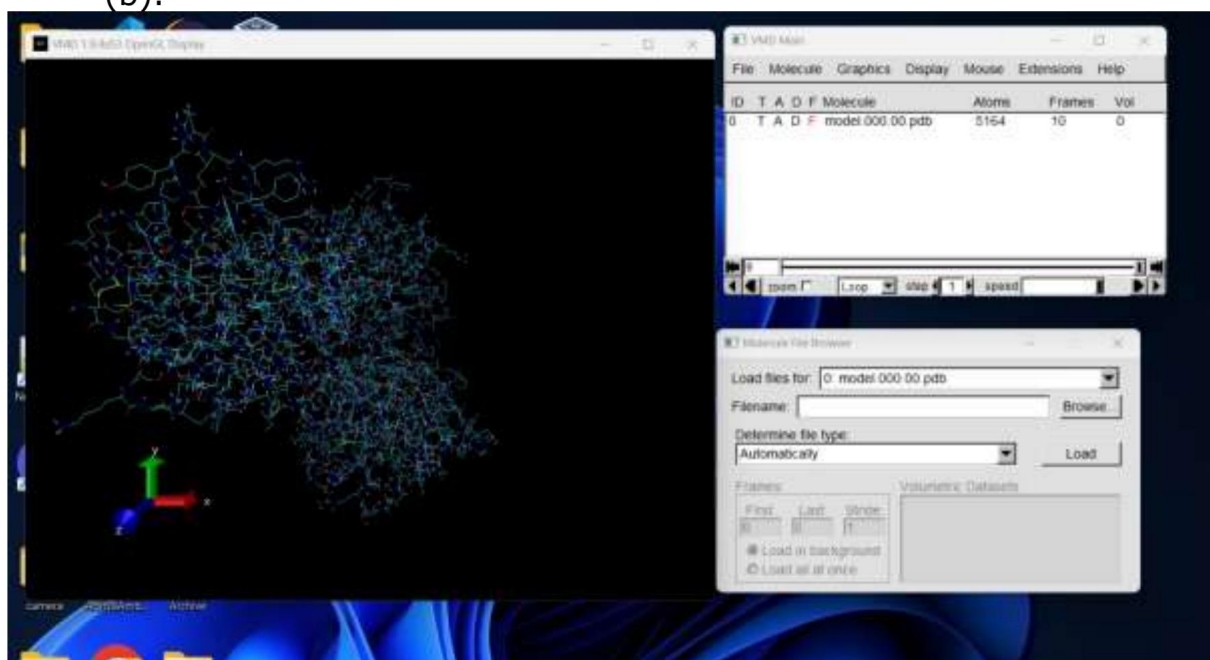
18



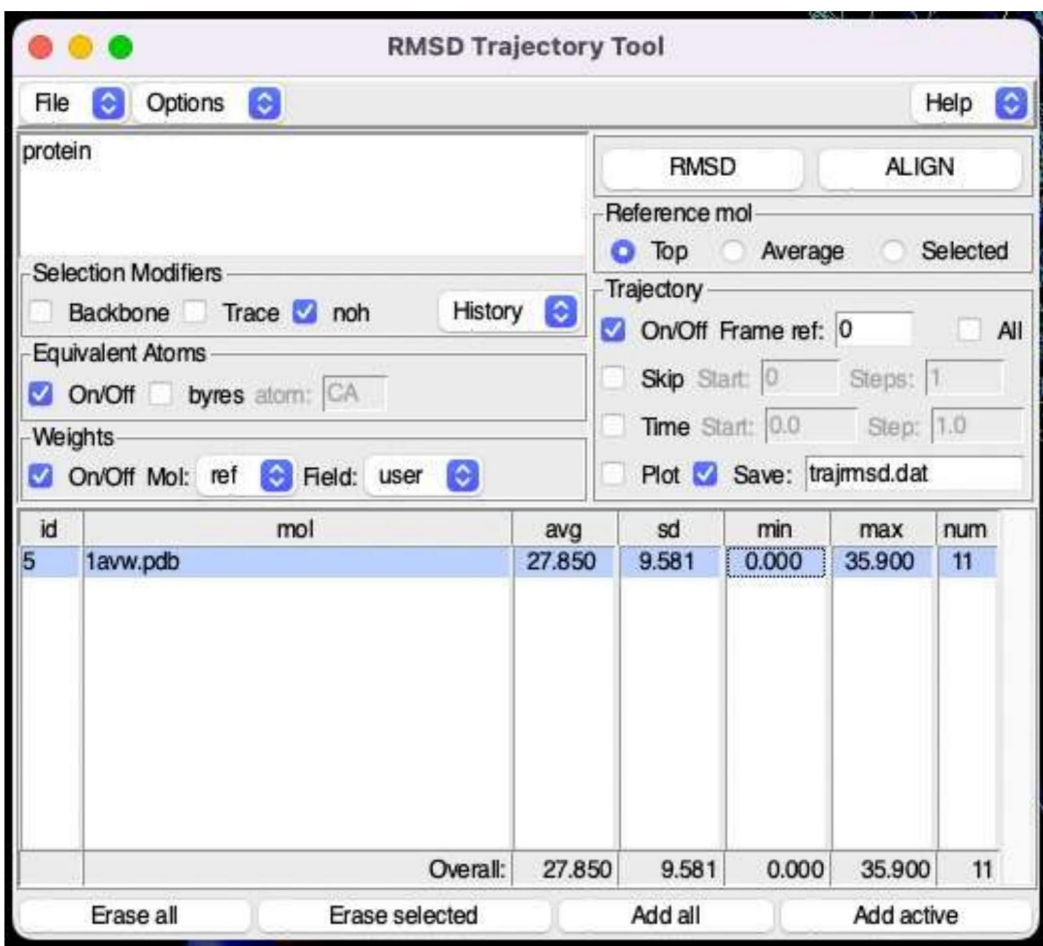
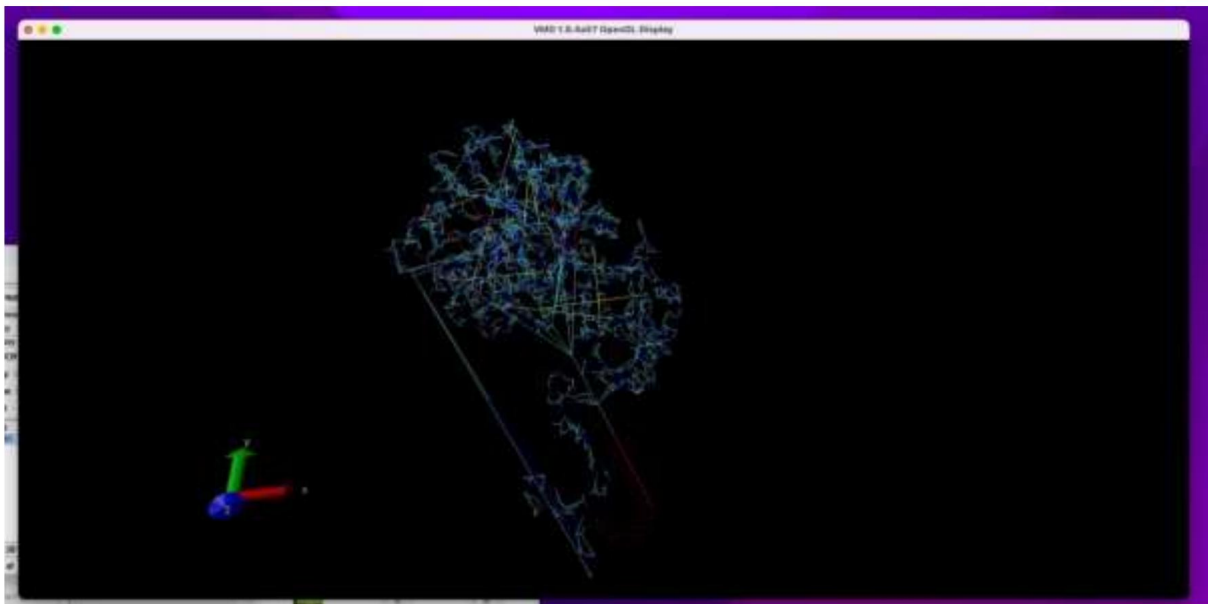
19



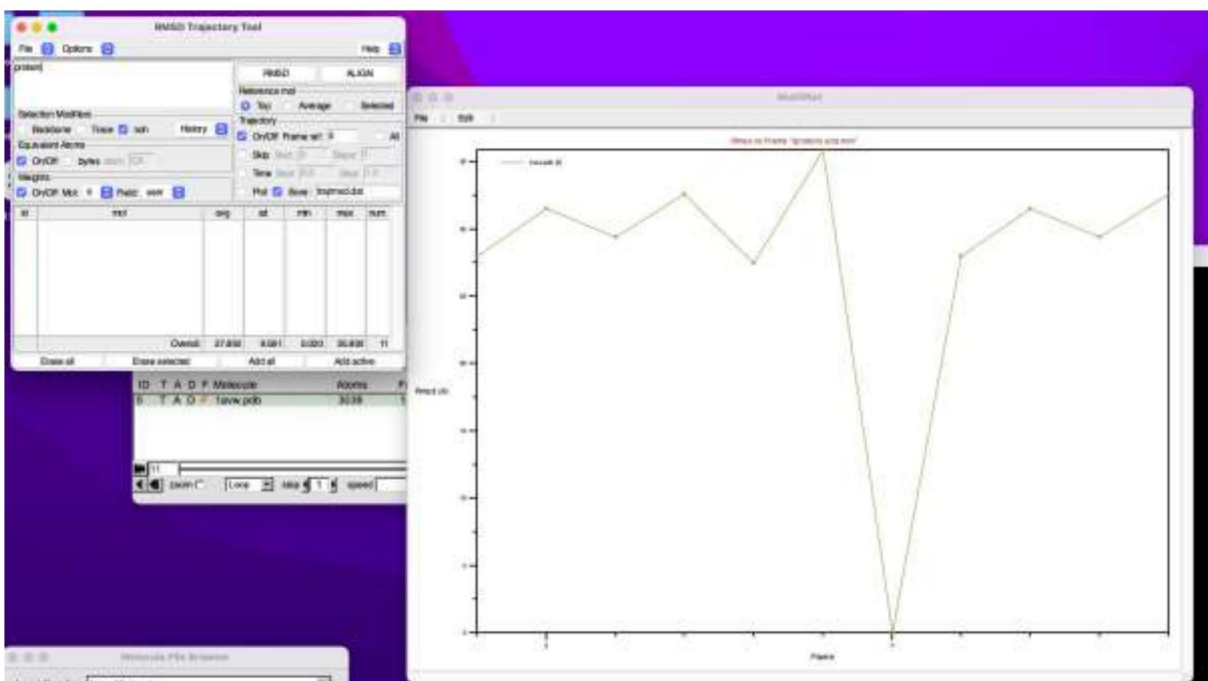
(b).



ALGORITHMS IN BIO INFORMATICS

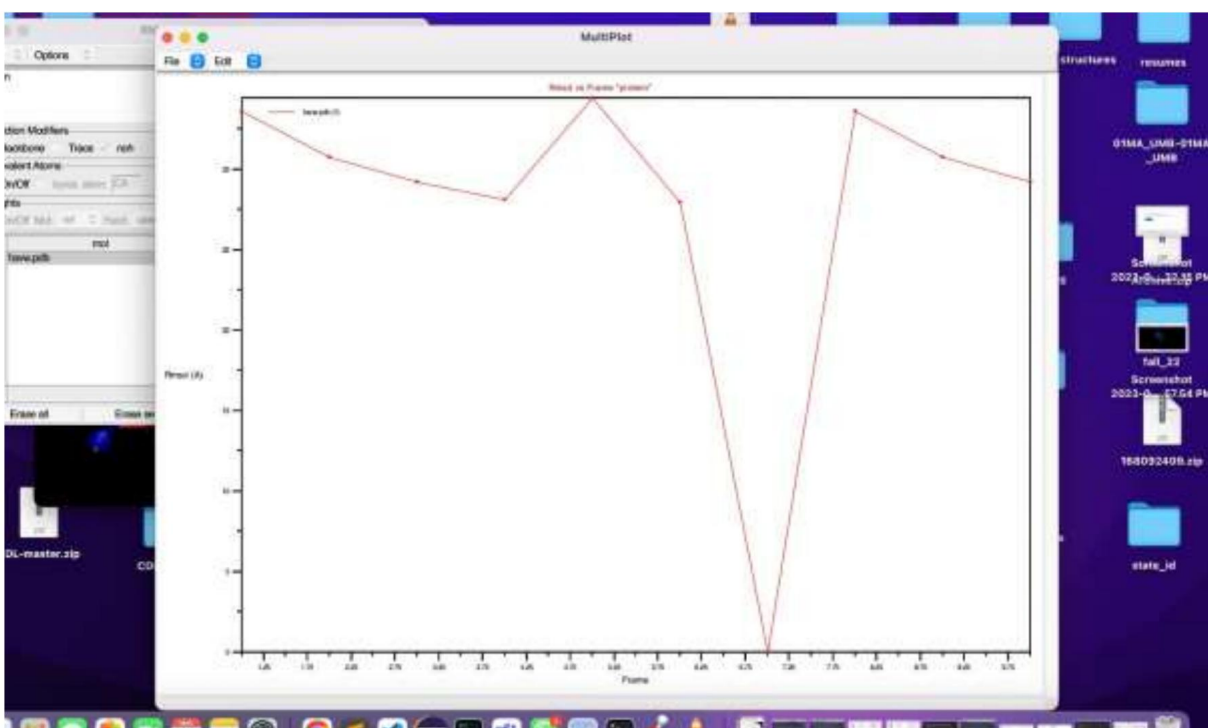


ALGORITHMS IN BIO INFORMATICS



id	mol	avg	sd	min	max	num
1	1avw.pdb	27.787	10.023	0.000	34.048	10

c. Now that you've done that, look at the native complex vs. the 10 complexes. Which one has the smallest RMSD to the native complex
the smallest one is 7 and highest one is 4.

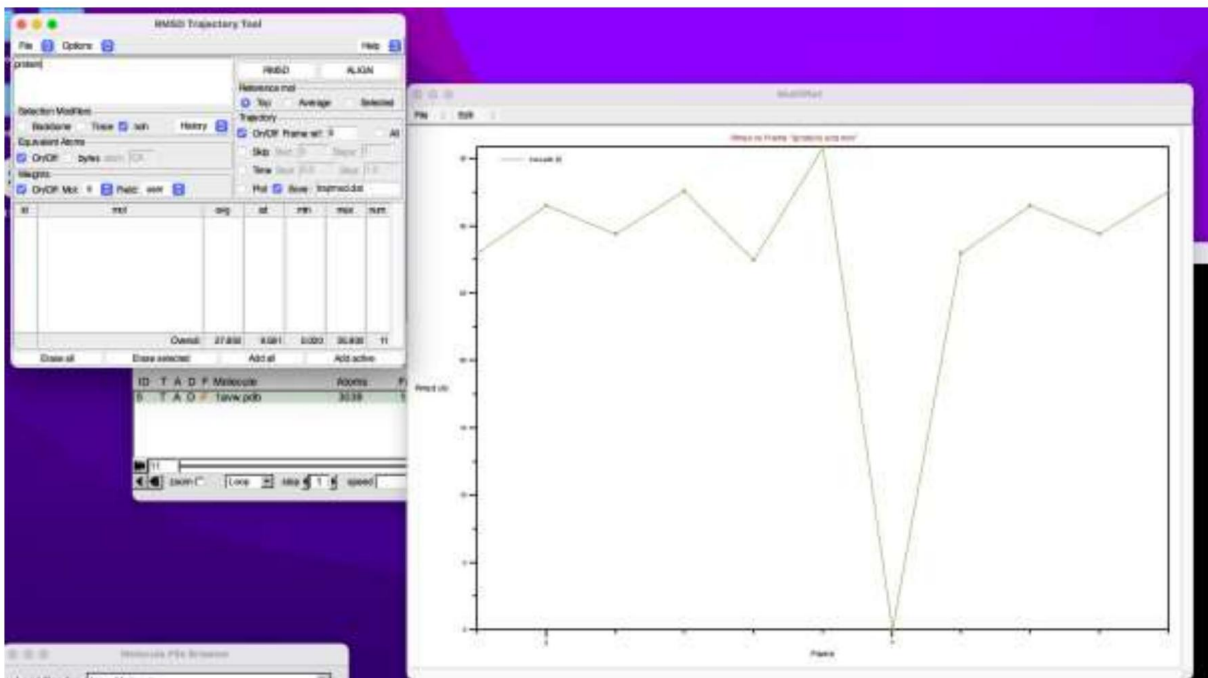


6.2 b 3 / 7

- 0 pts Correct
- 7 pts Incorrect/Missing
- 4 pts Instructions likely not followed properly. See solution
- ✓ - 4 pts *Click here to replace this description.*

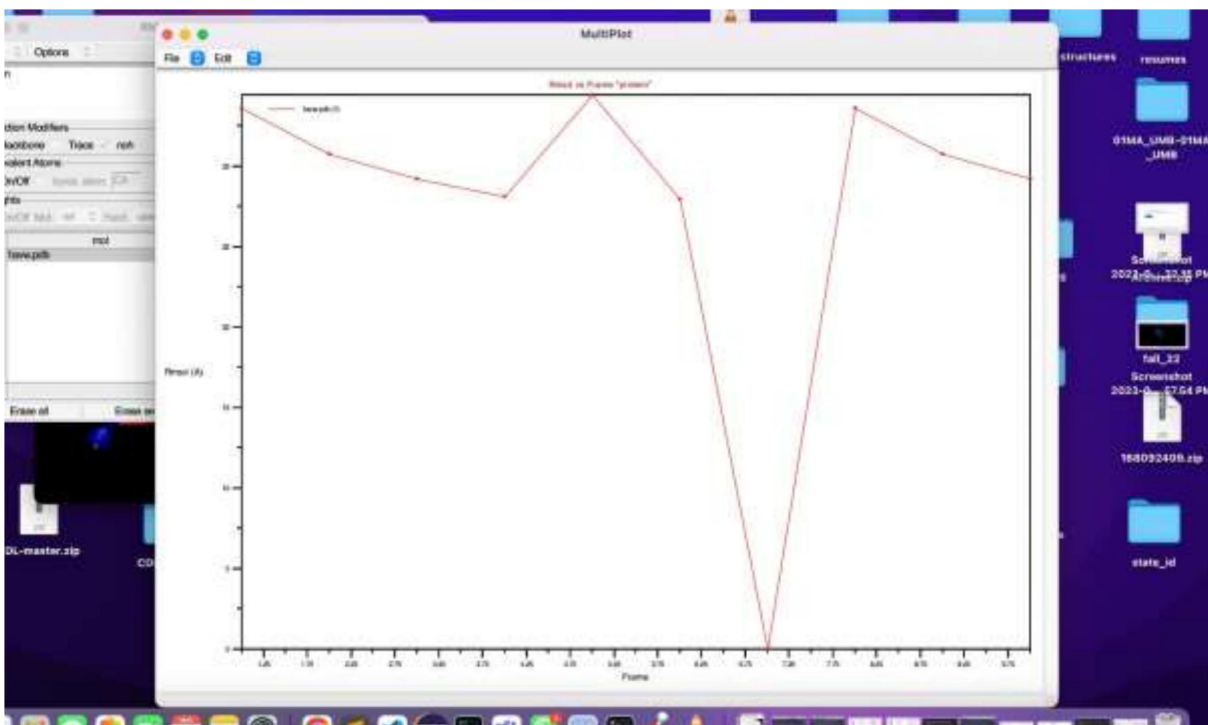
 See solution, you were supposed to look in the file. Your answer is also incorrect.

ALGORITHMS IN BIO INFORMATICS



id mol	avg	sd	min	max	num
1 1avw.pdb	27.787	10.023	0.000	34.048	10

c. Now that you've done that, look at the native complex vs. the 10 complexes. Which one has the smallest RMSD to the native complex
the smallest one is 7 and highest one is 4.



ALGORITHMS IN BIO INFORMATICS

6.3 C 0 / 6

- 0 pts Correct

- 2 pts 0 is the reference. 1 has the smallest.

✓ - 6 pts *Incorrect/Missing*

💬 Your b part was incorrect hence this is also incorrect.