

Practical No: 1

Aim: Write a Python/Java code to perform pairwise alignment. Take 2 sequences from user and calculate the score.

Code:

```
se1=input("Enter the first sequence::")
se2=input("Enter the second sequence::")
seq1=list(se1)
seq2=list(se2)
score=[]
```

```
def Pairwise_alignment(a,b):
    gap(a,b)
    print(a)
    print(b)
    value=0
    length=len(a)
    for i in range(0,length):
        if(a[i]==b[i]):
            score.append('1')
            value=value+1
        else:
            score.append('0')
    print(score)
    print(value)
```

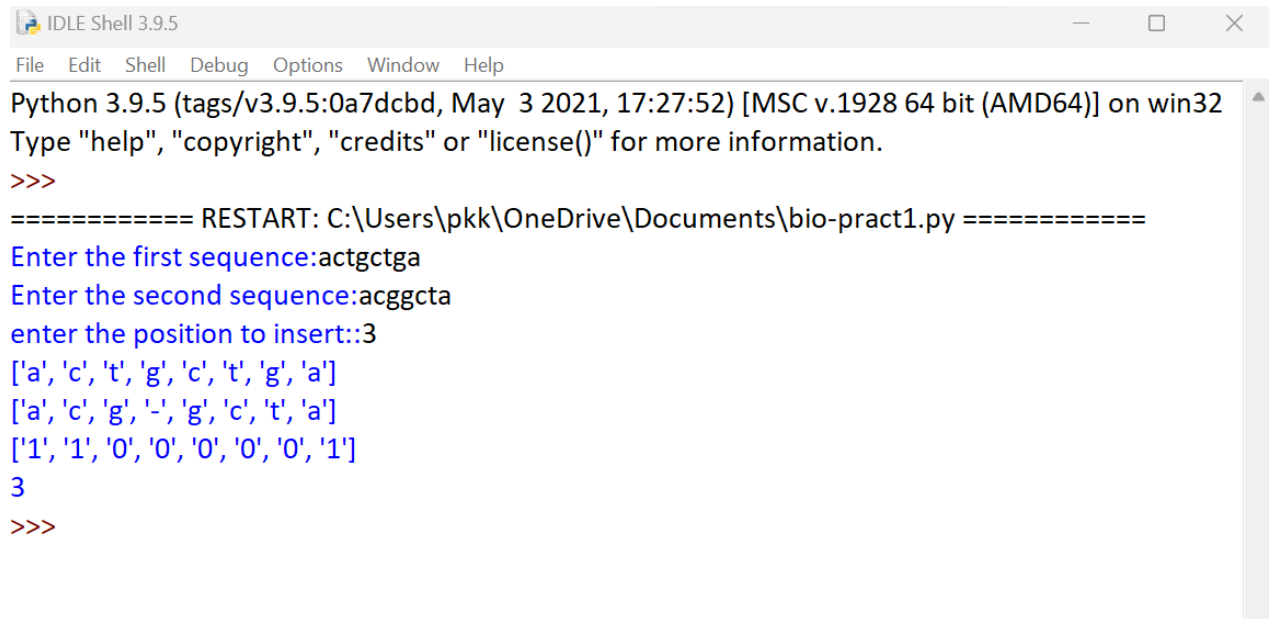
```
def gap(a,b):
    if(len(a)==len(b)):
        print()
    else:
        k=int(input("enter the position to insert::"))
        if (len(a)<len(b)):
            a.insert(k,'-')
        else:
            b.insert(k,'-')
    return(a,b)
```

```
Pairwise_alignment(seq1,seq2)
```

Output:

Name: Anjalee Pagare.

RollNo:525



```
IDLE Shell 3.9.5
File Edit Shell Debug Options Window Help
Python 3.9.5 (tags/v3.9.5:0a7dcdbd, May 3 2021, 17:27:52) [MSC v.1928 64 bit (AMD64)] on win32
Type "help", "copyright", "credits" or "license()" for more information.
>>>
===== RESTART: C:\Users\pkk\OneDrive\Documents\bio-pract1.py =====
Enter the first sequence:actgctga
Enter the second sequence:acggcta
enter the position to insert::3
['a', 'c', 't', 'g', 'c', 't', 'g', 'a']
['a', 'c', 'g', '-', 'g', 'c', 't', 'a']
['1', '1', '0', '0', '0', '0', '0', '1']
3
>>>
```

Practical No: 2

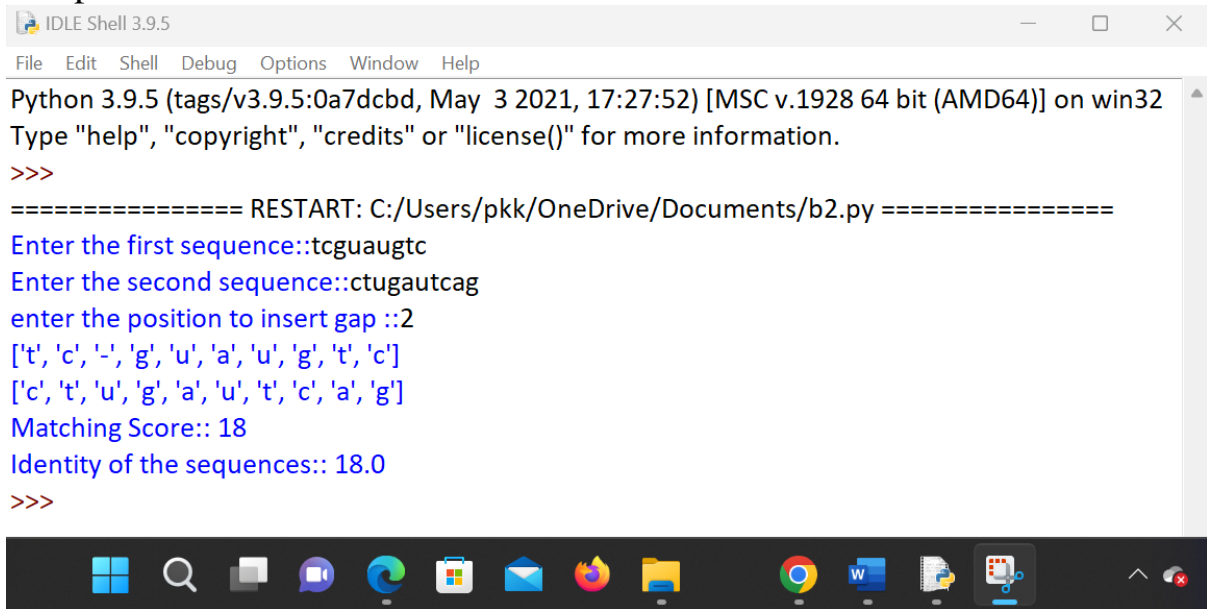
Aim: Write a Python/Java code to find the identity value of a given sequences. Take the sequence from user.

Code:

```
se1=input("Enter the first sequence::")
se2=input("Enter the second sequence::")

seq1=list(se1)
seq2=list(se2)
def find_identity(a,b):
    gap(a,b)
    print(a)
    print(b)
    score=0
    length=len(a)
    total_elements=len(a)*len(b)
    for i in range(0,length):
        for j in range(0,length):
            if(a[i]==b[j]):
                score=score+1
    identity=(score/total_elements)*100
    print("Matching Score::",score)
    print("Identity of the sequences::",identity)
def gap(a,b):
    if(len(a)==len(b)):
        print()
    else:
        k=int(input("enter the position to insert gap ::"))
        if (len(a)<len(b)):
            a.insert(k,'-')
        else:
            b.insert(k,'-')
    return(a,b)
find_identity(seq1,seq2)
```

Output:



```
IDLE Shell 3.9.5
File Edit Shell Debug Options Window Help
Python 3.9.5 (tags/v3.9.5:0a7dcdb, May 3 2021, 17:27:52) [MSC v.1928 64 bit (AMD64)] on win32
Type "help", "copyright", "credits" or "license()" for more information.
>>>
===== RESTART: C:/Users/pkk/OneDrive/Documents/b2.py =====
Enter the first sequence::tcguaugtc
Enter the second sequence::ctugautcag
enter the position to insert gap ::2
['t', 'c', '-', 'g', 'u', 'a', 'u', 'g', 't', 'c']
['c', 't', 'u', 'g', 'a', 'u', 't', 'c', 'a', 'g']
Matching Score:: 18
Identity of the sequences:: 18.0
>>>
```

Practical No: 3

Aim: Write a Python/Java code to find the Similarity value of a given sequences. Take the sequence from user.

Code:

```
sequence_one=input("Enter the first sequence: ")
sequence_two=input("Enter the second sequence: ")
how_many=int(input("How many elements for similarity condition?"))
similarities=[]
for i in range(0,how_many):
    a=input("Enter an element: ")
    c=int(input("How many elements is it similar to? "))
    similarities.append([])
    similarities[i].append(a)

    for j in range(0,c):
        b=input("What is it similar to? ")

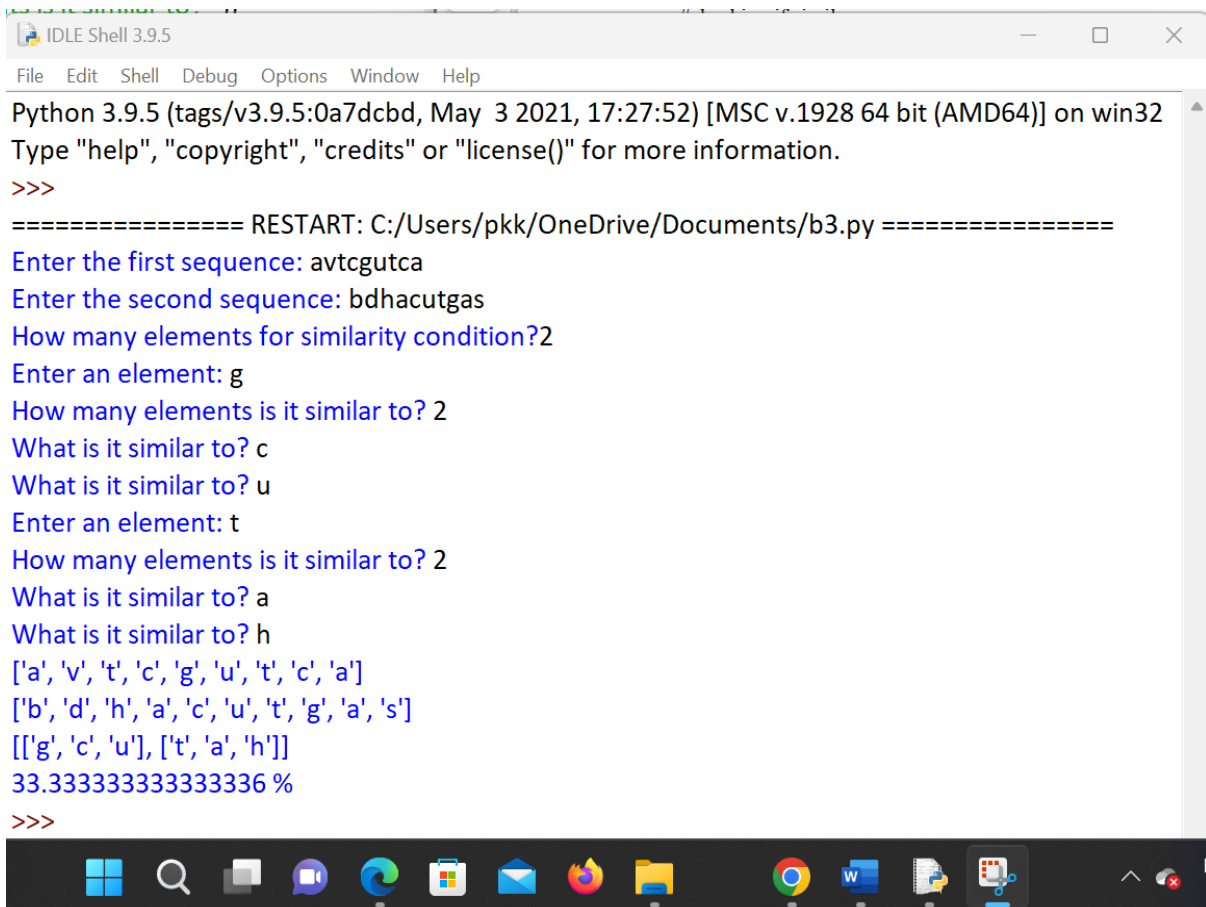
        similarities[i].append(b)

def compare(o,t,s):
    print(o)
    print(t)
    print(s)
    #checking if similar
    score=0
    for i in range(len(o)):
        for j in range(len(s)):
```

```
        if o[i] in s[j] and t[i] in s[j] and o[i] != t[i]:
            score+=1
        #calculating similarity
        similarity= (score*100)/len(o)
        return similarity

print(compare(list(sequence_one),list(sequence_two),similarities),"% ")
```

Output:



```
IDLE Shell 3.9.5
File Edit Shell Debug Options Window Help
Python 3.9.5 (tags/v3.9.5:0a7dcdb, May 3 2021, 17:27:52) [MSC v.1928 64 bit (AMD64)] on win32
Type "help", "copyright", "credits" or "license()" for more information.
>>>
===== RESTART: C:/Users/pkk/OneDrive/Documents/b3.py =====
Enter the first sequence: avtcgutca
Enter the second sequence: bdhacutgas
How many elements for similarity condition?2
Enter an element: g
How many elements is it similar to? 2
What is it similar to? c
What is it similar to? u
Enter an element: t
How many elements is it similar to? 2
What is it similar to? a
What is it similar to? h
['a', 'v', 't', 'c', 'g', 'u', 't', 'c', 'a']
['b', 'd', 'h', 'a', 'c', 'u', 't', 'g', 'a', 's']
[['g', 'c', 'u'], ['t', 'a', 'h']]
33.333333333333336 %
>>>
```

Practical No: 4

Aim: Enter genome of five different organism and write a python/java program to find consensus sequence using Multiple Sequence Alignment (MSA) technique.

Code:

```
import java.io.*;
import java.util.*;

public class Consensus
{
    public static void main(String str[]) throws IOException
    {
        int n, i,j,k,count;
        String seq[],cons[];
        ArrayList<Integer> a = new ArrayList<Integer>();
        ArrayList s = new ArrayList();
        BufferedReader br=new BufferedReader(new InputStreamReader(System.in));
        System.out.println("Enter the no of Sequences");
        n=Integer.parseInt(br.readLine());
        seq=new String[n];
        System.out.println("Enter sequences");
        for(i=0;i<n;i++)
```

```
seq[i]=br.readLine();

    cons=new String[seq[0].length()];
    for(j=0;j<seq[0].length();j++)
    cons[j]=" ";
    for(j=0;j<seq[0].length();j++)
    {
        a.clear();
        s.clear();
        for(i=0;i<n;i++)
        {
            count=1;
            for(k=i+1;k<n;k++)

                if(seq[i].charAt(j)==seq[k].charAt(j))
                    count++;

        }
        System.out.println("count="+count);
        a.add(count);
        s.add(seq[i].charAt(j));
    }

    /**Updated Snippet 1 **/

    Set<String> set = new HashSet<>(s);
    ArrayList setlist = new ArrayList(set);
    Collections.sort(setlist);

    if (setlist.contains('-') && setlist.size()==2){
        cons[j]+="-"+setlist.get(1);
    }
    else if (setlist.size()==1){
        cons[j]+="-"+setlist.get(0);
    }
    else{
        int m = Collections.max(a);
```

```
int index=a.indexOf(m);

System.out.println("Max="+m);

cons[j]+=s.get(index);

System.out.println("index="+index);

for(i=index+1;i<a.size();i++)

{

if(a.get(i)==m)

cons[j]+="/" +s.get(i);

}

}

}

System.out.println("Consensus=");

for(j=0;j<seq[0].length();j++){

    /**Updated Snippet 2**/

if(cons[j].length()==2)

System.out.print(cons[j].toLowerCase());

else if(cons[j].length()==3)

System.out.print(cons[j].replace("-", ""));

else

System.out.print(cons[j]);

}

}

}
```

Output:


```
Terminal
Enter the no of Sequences:
3
Enter sequences:
TAGC
CGAT
ATCG
count=1count=1count=1
Max=1index=0
count=1count=1
count=1
Max=1
index=0
count=1count=1
count=1
Max=1index=0
count=1
count=1
count=1Max=1
index=0Consensus=T/C/A A/G/T G/A/C C/T/G
```



Practical No: 5

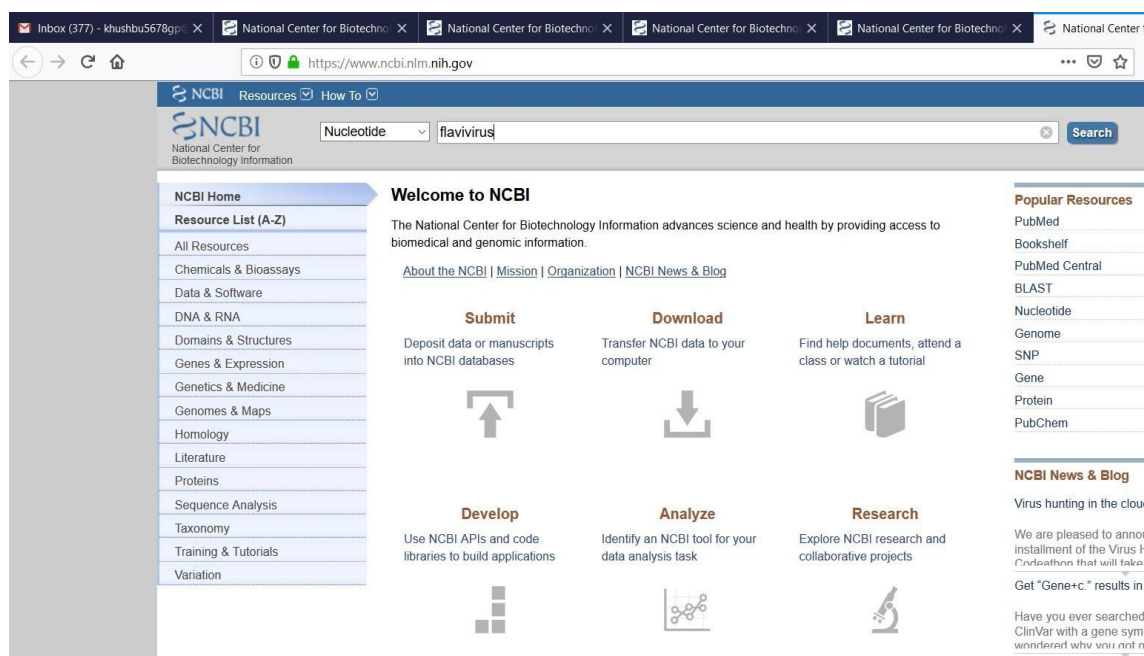
Aim: Perform a BLAST search on a specific gene sequence of a specify organism.

Steps:

Go to the National Center for Biotechnology Information Site

<https://www.ncbi.nlm.nih.gov/>

Select Nucleotide from All Databases and find any organism in a search bar.



Name: Anjalee Pagare.
RollNo:525

NCBI Resources How To

Nucleotide

Create alert Advanced

Species Summary 20 per page Sort by Default order Send to: Filters: Manage Filters

Animals (307)
Plants (12)
Fungi (24)
Protists (20)
Bacteria (793)
Archaea (7)
Viruses (62,020)
Customize ...

Molecule types
genomic DNA/RNA (65,124)
mRNA (1,298)
Customize ...

Source databases
INSDC (GenBank) (67,508)
RefSeq (191)
Customize ...

Sequence Type
Nucleotide (67,735)

Genetic compartments
Mitochondrion (163)
Plasmid (4)

Sequence length
Custom range...

Items: 1 to 20 of 67735

1. ☐ [Tembusu virus flavivirus polyprotein \(flavivirus polyprotein gene\) gene, complete cds](#)
10,990 bp linear RNA
Accession: NC_015843.2 GI: 381333920
[Assembly](#) [BioProject](#) [Protein](#) [PubMed](#) [Taxonomy](#)
[GenBank](#) [FASTA](#) [Graphics](#)

2. ☐ [Cacipacore virus flavivirus polyprotein \(flavivirus polyprotein gene\) and truncated polyprotein \(flavivirus polyprotein gene\) genes, complete cds](#)
10,284 bp linear RNA
Accession: NC_026623.1 GI: 765702599
[Assembly](#) [BioProject](#) [Protein](#) [PubMed](#) [Taxonomy](#)
[GenBank](#) [FASTA](#) [Graphics](#)

3. ☐ [Duck flavivirus KPS54A61/THA, complete genome](#)
10,990 bp linear RNA
Accession: KF573582.1 GI: 673459548
[Protein](#) [Taxonomy](#)
[GenBank](#) [FASTA](#) [Graphics](#)

☐ [Duck flavivirus TA, complete genome](#)

Results by taxon

Top Organisms [Tree]
Dengue virus (28466)
West Nile virus (21550)
Japanese encephalitis virus
Tick-borne encephalitis virus
Zika virus (1694) [viruses]
All other taxa (10744)
More...

Find related data
Database:

Search details
"Flavivirus"[Organism]
flavivirus[All Fields]

NCBI Resources How To

Nucleotide

Create alert Advanced

GenBank Send to: Change region shown

Customize view

Analyze this sequence
Run BLAST
Pick Primers
Highlight Sequence Features
Find in this Sequence

Related information
Assembly
BioProject
Protein
PubMed
Taxonomy
Full text in PMC
Functional Class

Tembusu virus flavivirus polyprotein (flavivirus polyprotein gene) gene, complete cds

NCBI Reference Sequence: NC_015843.2
[FASTA](#) [Graphics](#)

Go to:

LOCUS NC_015843 10990 bp ss-RNA linear VRL 13-AUG-2018
DEFINITION Tembusu virus flavivirus polyprotein (flavivirus polyprotein gene) gene, complete cds.
ACCESSION NC_015843 NC_016958 NC_018670
VERSION NC_015843.2
DBLINK BioProject: [PRJNA485481](#)
KEYWORDS RefSeq.
SOURCE Tembusu virus (TMUV)
ORGANISM [Tembusu virus](#)
Viruses; Riboviria; Flaviviridae; Flavivirus.
REFERENCE 1 (bases 1 to 10990)
AUTHORS Han,K., Huang,X., Li,Y., Zhao,D., Liu,Y., Zhou,X., You,Y. and Xie,X.
TITLE Complete genome sequence of goose tembusu virus, isolated from jiangnan white geese in jiangsu, china
JOURNAL Genome Announc 1 (2), E0023612 (2013)
PUBMED 23516233
REMARK Publication Status: Online-Only
DESCRIPTION 2 (bases 1 to 10990)

Run BLAST option we have to select

Align two or more sequences

Choose Search Set

Database: ☐ Human genomic + transcript ☐ Mouse genomic + transcript ☒ Others (nr etc.):
Nucleotide collection (nr/nt)

Organism: Enter organism name or id—completions will be suggested ☐ exclude

Exclude: ☐ Models (XM/XP) ☐ Uncultured/environmental sample sequences

Limit to: ☐ Sequences from type material

Entrez Query: Enter an Entrez query to limit search

Program Selection

Optimize for: ☒ Highly similar sequences (megablast) ☐ More dissimilar sequences (discontiguous megablast) ☐ Somewhat similar sequences (blastn)

Choose a BLAST algorithm

BLAST Search database Nucleotide collection (nr/nt) using Megablast (Optimize for highly similar sequences)

☐ Show results in a new window

Algorithm parameters

BLAST is a registered trademark of the National Library of Medicine

BLAST

Sequences producing significant alignments

Download Manage Columns Show

select all 99 sequences selected

	Description	Max Score	Total Score	Query Cover	E value
<input checked="" type="checkbox"/>	Tembusu virus strain JS804, complete genome	20064	20064	99%	0.0 9
<input checked="" type="checkbox"/>	Tembusu virus strain JS/2010, complete genome	20064	20064	99%	0.0 9
<input checked="" type="checkbox"/>	Duck egg-drop syndrome virus strain byd1, complete genome	20048	20048	99%	0.0 9
<input checked="" type="checkbox"/>	Tembusu virus isolate Tembusu virus strain, complete genome	20026	20026	99%	0.0 9
<input checked="" type="checkbox"/>	Duck Tembusu virus isolate df-2, complete genome	20020	20020	99%	0.0 9
<input checked="" type="checkbox"/>	Duck egg-drop syndrome virus strain JXSP, complete genome	20020	20020	99%	0.0 9
<input checked="" type="checkbox"/>	Tembusu virus isolate YY5, complete genome	20015	20015	99%	0.0 9
<input checked="" type="checkbox"/>	Tembusu virus isolate SDMS, complete genome	20009	20009	99%	0.0 9
<input checked="" type="checkbox"/>	Tembusu virus isolate ZJ-6, complete genome	20009	20009	99%	0.0 9
<input checked="" type="checkbox"/>	Tembusu virus strain AH-F10 from China, complete genome	20004	20004	99%	0.0 9
<input checked="" type="checkbox"/>	Duck egg-drop syndrome virus strain pigeon, complete genome	20004	20004	99%	0.0 9
<input checked="" type="checkbox"/>	Tembusu virus genomic RNA, complete genome, strain TMUV-YY1Du	19998	19998	99%	0.0 9
<input checked="" type="checkbox"/>	Duck Tembusu virus strain BZ_2010, complete genome	19998	19998	99%	0.0 9
<input checked="" type="checkbox"/>	Duck egg-drop syndrome virus strain duan, complete genome	19998	19998	99%	0.0 9
<input checked="" type="checkbox"/>	Duck Tembusu virus strain GDLH01, complete genome	19989	19989	99%	0.0 9

Here the result will be display

Inbox (377) - khushbu5678gp

National Center for Biotechno

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NCBI Blastref|NC

https://blast.ncbi.nlm.nih.gov/Blast.cgi?alnHdr_381283018

Download

GenBank

Graphics

Next

Previous

Tembusu virus strain JS804, complete genome

Sequence ID: [JF895923.2](#) Length: 10990 Number of Matches: 1

Range 1: 1 to 10990

[GenBank](#)

[Graphics](#)

Next Match

Previous Match

Score	Expect	Identities	Gaps	Strand
20295 bits(10990)	0.0	10990/10990(100%)	0/10990(0%)	Plus/Plus

Query

1

AGAAGTTCGCCTGTGTGAACCTATTCCAAACAGCTTTTGGAGTAGTGC

60

Sbjct

1

AGAAGTTCGCCTGTGTGAACCTATTCCAAACAGCTTTTGGAGTAGTGC

60

Query

61

ACACAGTTTGAACGTTTTTGGATAGAGACAACTATGTCTAACAAAAACCAGGAAGACC

120

Sbjct

61

ACACAGTTTGAACGTTTTTGGATAGAGACAACTATGTCTAACAAAAACCAGGAAGACC

120

Query

121

CGGCTCAGGCCGGGTTGTCAATATGCTAAAGCGCGGAACGTC

180

Sbjct

121

CGGCTCAGGCCGGGTTGTCAATATGCTAAAGCGCGGAACGTC

180

Query

181

GCGGATAAAGAGGACGATTGATGGGTCCTGAGAGGAGCAGGACCCATAAGGTTTGTGCT

240

Sbjct

181

GCGGATAAAGAGGACGATTGATGGGTCCTGAGAGGAGCAGGACCCATAAGGTTTGTGCT

240

Query

241

GGCTCTACTGACTTTCTTCAAGTTTACAGCCCTGAGGCCAACCATTTGGAATGCTGAAGAG

300

Sbjct

241

GGCTCTACTGACTTTCTTCAAGTTTACAGCCCTGAGGCCAACCATTTGGAATGCTGAAGAG

300

Query

301

ATGGAAGCTGGTTGGAGTTAATGAGGCGACCAACATCTGAAAAGCTTCAAGCGTGACAT

360

Sbjct

301

ATGGAAGCTGGTTGGAGTTAATGAGGCGACCAACATCTGAAAAGCTTCAAGCGTGACAT

360

Query

361

TGGACAGATGCTCGACGGACTGAATAAGCGGAAGCGGAAACCTCgggggggggAGTTGCTC

420

Sbjct

361

TGGACAGATGCTCGACGGACTGAATAAGCGGAAGCGGAAACCTCGGGGGGGGAGTTGCTC

420

Related Inform

[Gene](#) - associated

Practical No: 6

Aim: Write a Python/Java code to find motif in a given sequence.

Code:

```
import random

l=int(input("Enter the length of motif"))

file=open("mot.txt","r")

r=file.read()

print("Sequence",r)

size=len(r)

print("Size of the sequence",size)

pos=random.randint(0,len(r)-5)

#pos=1

print("Position",pos)

motif=r[pos:pos+l]

print("Motif",motif)

i=pos+1

while(i<=size-1):

    if(motif==r[i:i+l]):

        str1=r[i:i+l]

        print("Match motif",str1)

        file1=open("motoutput.txt","a")

        file1.write(str1+" ")

    i+=1
```

Output:

Enter the length of motif4

Sequence AGAAGTTCGAGAAGCCGTAGT

Size of the sequence 21

Position 0

Motif AGAA

```
import random

l=int(input("Enter the length of motif"))

file=open("mot.txt","r")

r=file.read()

print("Sequence",r)

size=len(r)

print("Size of the sequence",size)

pos=random.randint(0,len(r)-5)

#pos=1

print("Position",pos)

motif=r[pos:pos+l]

print("Motif",motif)

i=pos+1

while(i<=size-1):

    if(motif==r[i:i+l]):

        str1=r[i:i+l]

        print("Match motif",str1)

        file1=open("motoutput.txt","a")

        file1.write(str1+" ")

    i+=1
```

Output:

```
Enter the length of motif4

Sequence AGAAGTTCGAGAAGCCGTAGT

Size of the sequence 21

Position 0

Motif AGAA
```

Aim: Perform a BLAST search on any genes sequence and write a java/python code to count the no of repetition of each nucleotide in the sequence.

Code:

```
file=open("genes.txt","r")
r=file.read()
size=len(r)
score_A=0
score_C=0
score_T=0
score_G=0
for i in range(size):
    if(r[i]=='A'):
        score_A+=1
    elif (r[i]=='C'):
        score_C+=1
    elif (r[i]=='T'):
        score_T+=1
    elif (r[i]=='G'):
        score_G+=1
print("score of A is ",score_A)
print("score of C is ",score_C)
print("score of T is ",score_T)
print("score of G is ",score_G)
```

Output:

score of A is 6

score of C is 4

score of T is 7

score of G is 6

Practical No: 8

Aim: Generate a regular expression enter three protein sequence of three different organism. Write Python/Java code to find regular expression for this sequences.

Code:

```
def gen_reg_exp(seq_list, no_of_col):  
    final_list=[]  
    for colnum in range(no_of_col):  
        collist=[]  
        for colseq in seq_list:  
            collist.append(colseq[colnum])  
        if len(set(collist))==len(collist):  
            #print(final_list)  
            final_list.append('x')  
        else:  
            if len(set(collist))==1:  
                final_list.append(collist[0])  
            else:  
                final_list.append(".".join(set(collist)))  
    display_output(final_list)
```

```
def display_output(final_list):  
    print(*final_list, sep='-')  
  
no_of_seq=int(input("Enter the number of sequence: "))  
print("Enter all the sequences")  
seq_list=[]  
for _ in range(no_of_seq):  
    seq_list.append(list(map(str, input("").split())))  
gen_reg_exp(seq_list, len(seq_list[0]))
```

Output:

```
Enter the number of sequence: 4  
Enter all the sequences  
A D L G A V F A L C D R Y F Q  
S D V G P R S C F C E R F Y Q  
A D L G R T Q L R C D R Y Y Q  
A D I G Q P H S L C E R Y F Q  
SA-D-IVL-G-x-x-x-x-FRL-C-ED-R-YF-YF-Q
```

Practical No: 9

Aim: Enter six protein sequence of different organism and write a program to find a fingerprint of sequence.

Code:

```
def solve_fingerprint(seq_list, no_of_col):  
    seq_dict=dict()  
    for colnum in range(no_of_col):  
        counta,countc,countt,countg=0,0,0,0  
        for colseq in seq_list:
```

```
        if colseq[colnum]=='A':
            counta+=1
        elif colseq[colnum]=='T':
            countt+=1
        elif colseq[colnum]=='C':
            countc+=1
        elif colseq[colnum]=='G':
            countg+=1

    seq_dict[colnum]=[counta,countc,countt,countg]

display_results(seq_dict)

def display_results(seq_dict):
    print("\tA \tC \tT \tG")
    for key in seq_dict:
        print("\n",*seq_dict[key],sep="\t")

no_of_seq=int(input("Enter the number of sequence: "))
print("Enter all the sequences")
seq_list=[]
for _ in range(no_of_seq):
    seq_list.append(list(map(str, input("").split()))))
solve_fingerprint(seq_list,len(seq_list[0]))
```

Output:

Enter the number of sequence: 4

Enter all the sequences

A C T G A T G

A T C A G A A

A T A A G C A

A G T T A G C

A	C	T	G
---	---	---	---

4	0	0	0
---	---	---	---

0	1	2	1
---	---	---	---

1	1	2	0
---	---	---	---

2	0	1	1
---	---	---	---

2	0	0	2
---	---	---	---

1	1	1	1
---	---	---	---

2	1	0	1
---	---	---	---