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Due: 10-Nov-2020, 10 am

Sequence Bioinformatics

WS 2021/22

Assignment 3

In this assignment, we will investigate the idea of using Integer Linear Programming to compute a

>S1 CGTA >S2 CGAT >S3

GATAT

1 Counting (1 point)

How many edges are there between nucleotides that lie in different sequences in the alignment graph? How many simple mixed cycles are there?

Please write a Python program ilp_aligment_YOUR_NAME.py to solve the following tasks:

maximum scoring multiple sequence alignment for the following three sequences:

2 Simple mixed cycles (3 points)

In the following, use Xij_pq to denote the variable that represents the edge connecting the nucleotide $s_i(j)$ in sequence s_i , at position j, with the nucleotide $s_p(q)$ in sequence s_p , at position q.

Generate the list of all simple mixed cycles for the three given sequences and list them in lexicographical order, using the following format (which can be parsed by lp_solve, note that < means "\leq"):

```
X11_21 + X12_21 < 1;
```

3 Objective function (1 point)

Using a match score of 4 and a mis-match score of 1, set up the objective function for the ILP, in the format:

```
max 1*X11_21+4*X12_21+ ...;
```

4 Run the ILP (4 points)

Download the program lp_solve, from https://sourceforge.net/projects/lpsolve/ and install it. Setup the ILP in the format supported by the program, which looks like this:

Run this file using lp_solve.

5 Report the alignment (1 point)

Discuss how to translate the output of lp_solve into an alignment and report the alignment.