Group Project Report: siRNA Generator

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1 Overview

I will first discuss the motivation for our work and our group's broad objective.

1.1 Motivation

Our primary motivation for our project was the 2014 Ebola outbreak. There are five identified species of the Ebolavirus: Bundibugyo, Taï, Sudan, and Zaire [3]. The last, the Reston virus, does not harm humans. The Zaire ebolavirus is the most deadly and the one responsible for the current outbreak, the most deadly to date.

Symptoms of Ebola virus disease (EVD) include: fever, chills, fatigue, weakness, muscle/join pain, vomiting, reddened eyes, and hemorrhaging [6] [8]. These symptoms appear within eight to ten days, on average [3].

This outbreak primarily affected people in Guinea, Liberia, and Sierra Leone, with additional cases reported in Nigeria, Mali, and Senegal (Figure 1) [3]. Thus far, there have been an estimated 25,907 infected in 2014 outbreak, and nearly 11,000 people have succumbed to the disease [3]. Two cases of Ebola were imported to the United States and was spread to two nurses. This, plus cases Spain and the United Kingdom sparked a large level of concern worldwide [4] [9].

Ebola can be fought, but the most effective method is prevention. Ebola does not spread easily when compared to other viruses such as influenza. Infection requires contact with the bodily fluids of a infected person who is displaying symptoms of Ebola virus disease [8] [6]. Thus, the most effective way currently available is to quarantine the infected individuals and inform those who have had contact with the infected [6].

1.2 Our approach

We chose using siRNA since a couple of group members were involved in active research on siRNA and were very familiar with it.

shRNA is precursor to siRNA Binds to mRNA in virus in order to render it ineffective.



Figure 1: Map of total cases of the 2014 Zaire Ebolavirus outbreak. Source: Center for Disease Control [3].

2 Contributions

2.1 Assigned Tasks

Our group separated into three rough groups to tackle each of the three tasks we identified.

The first group was tasked with creating a script to identify the conserved region within the species of virus. These conserved regions would then be compared against the human genome using BLAST [1]. Any sequence that was shared with the human genome would be filtered out of output for the next group, since we targt that sequence with an siRNA, it could potentially target that sequence in humans.¹

The second group takes the conserved regions that did not show up in the human genome and uses those to design siRNA to bind with the virus's mRNA. The resulting siRNA needs to then be compared against the human genome, for the same reasons as the first check. This would output a fasta file containing the siRNA sequences that would target the genus of virus. These potential siRNA should be further analyzed for any consequences. A member of the group also implemented an algorithm to estimate the amount of mutations it would take for the virus to render the siRNA ineffective.

I was the third group and my responsibilities were quite broad.

- I needed to write a web application to accept a FASTA file from the user. This file would then be passed the beginning of the first group's code. I could accomplish this by spinning up a thread that called the first function needed to run.
- I needed to ensure that the publicly accessible web application did not

¹This is generally considered a bad idea

have write capabilities on the server other than writing the file.²

- I was to help setup the webserver we would use for our application: apollo.³
- I needed to ensure the rest of the group had the tools they needed to run their python programs.
- I made myself available to each of our group members to aid with debugging.

In summary, I needed to do some web programming and act as the IT guy.

2.2 Implementation

I worked with Brendan to install Ubuntu 14.04 on the server. This was a fairly trivial task.

I then had to acquire the various pieces of software we needed

- BLAST [1]
- clustalw [7]
- biopython [2]
- RNAfold [5]
- human genome

I will talk specifically about the goal of each part of the application and my implementation of it.

2.2.1 Application

The first part of the application is the part the user sees. pic:

2.2.2 Scheduler

Once fasta file is retrieved, the scheduler spins up a new thread start the task. The thread is not a daemon, os it will live on.

²And I just thought of an attack! I could upload a very, very, large file. I'm not sure what this would do, but it would probably not be good.

³Highly germane, since Apollo's portfolio includes plagues and medicine.

2.2.3 Email

3 Evaluation

3.1 Test Case

We used a fasta file of the five species of the Ebolavirus as our initial input and test case for the program.

3.2 Deliverables

A deadline for combining the code was set for April 19th and not met. There are a number of reasons for this, but I do not believe that the problem was too difficult or large.

I observed a number bad coding practices during the project. One group member was using Python 3.4 despite the rest of the group working in 2.7. Many members hardcoded file names into commands and wrote duplicate components.

None of these alone were an issue that completely impeded the implementation, or even together, but were symptoms of the true issues: a lack of communication and miscommunication. It seemed like every member assumed what the other member was doing without consulting anyone else. As a result, each member made their own assumptions about how the components their program were connecting to would work, even though we had determined that Fasta was the common input and output between files.

I do not believe that our final product in completely unusable, but

References

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