

Exact String Matching Algorithms

Introduction

This document is related with the first work proposed at the Bioinformatic lectures. The purpose of this work is to check if the theory explained in class, about the search algorithms for only one pattern, is still true. As it is said in the statement, we have to compare the performance of 4 algorithms: *Brute Force* (BF), *Horspool* (H), *Backward Nondeterministic Dawg Matching* (BNDM) and *Backward Oracle Matching* (BOM). The comparison is made over a DNA alphabet file of at least 2GB, with random patterns of length $k > 0$.

Experiment

First, I downloaded the codes of the algorithms from [?]. Then, I had to modify each of the codes to let them read an external file and accept several parameters. These parameters are the DNA alphabet file and n pairs composed by the size of a pattern and the pattern itself. As an example of how to call one of the algorithms:

```
$ ./algorithm file_path size1 pattern1 size2 pattern2 ... size $n$  pattern $n$ 
```

Moreover, I created a *Python* script to generate a random pattern of length k (input). Finally, I created a *Bash* script that is in charge of generating a string with all the parameters, and then to call each of the algorithms with it. I executed it with $n = 100$, with the size of the patterns from 10 to 1000 with steps of 10.

Results