# Creating an Evolution Simulator for Protein Low Complexity Regions and Attempting to Utilize it for an Approximate Bayesian Computation

Alexander Turco

April 5, 2023



## Overview

Background Information				
Simulating LCR Evolution				
Applications to ABC-MCMC				
Conclusions				
Acknowledgements				

# What are Low Complexity Regions?

#### Saccharomyces cerevisiae SRP40 Protein LCRs

>CAA82171.1(298-316) complexity=2.18 (15/1.90/2.20) tpassnestpsasssssan

# LCRs Present in Unique Ways

#### Homorepeats

Consecutive iterations of a single residue



#### Direpeats

Consecutive iterations of two ordered, different residues

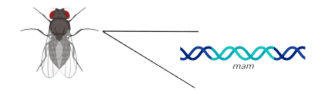


#### Imperfect Repeats

Regions in which the repeat units are not the same



# LCRs are Hypermutable

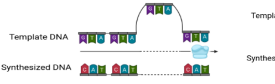


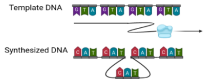
mam domain	Size (bp)	Amino Acid Substitutions	Amino Acid/ Total Substitutions
Unique	933	26	0.15
Repetitive	810	47	0.42

<sup>&</sup>lt;sup>1</sup>Newfeld, Smoller, and Yedvobnick, 1991

## Proposed Mechanisms of LCR Evolution

#### 1. Polymerase Slippage/Slipped Strand Mispairing





**Polymerase Slips Forward** 

Polymerase Slips Backwards

<sup>&</sup>lt;sup>2</sup>Levinson and Gutman, 1987; Sehn, 2015

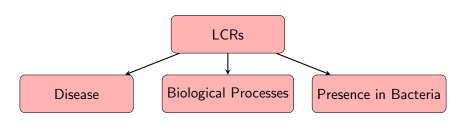
## Proposed Mechanisms of LCR Evolution

#### 2. Unequal Recombination



<sup>&</sup>lt;sup>3</sup>Mirkin, 2007

# Why Care about LCRs and their Evolution?









Huntington's Disease

Genetic Recombination

Neisseria meningitidis

## What we Did in this Study

- ► Utilized C++ to build an evolution simulator which altered protein sequences via point mutations, insertions, and deletions
- ► Tested the simulator with various insertion/deletion rates and mutation rates
- ▶ Attempted to program an ABC-MCMC in C++ using the evolution simulator as an important step in the algorithm, in order to estimate parameters like mutation and indel rates.

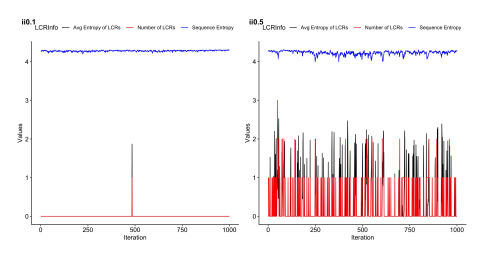
• mutation rate = 0.14 indel rate = 0.14 Random Protein Sequence GGAGGGAQ

- $\begin{tabular}{ll} \textbf{0.14} & \textbf{mutation rate} = 0.14 \\ \textbf{Random Protein Sequence} \\ \textbf{GGAGGGAQ} \\ \end{tabular}$
- ② Assign Exponential Deviates mutation deviates = (0.83, 1.82, 2.35, 0.54, 0.98, 0.76, 1.53, 2.34) indel deviates = (0.21, 1.21, 1.49, 0.86, 0.97, 1.13, 0.53, 0.35) exp $(\beta)$ , where  $\beta$  = mutation rate exp $(\beta)$ , where  $\beta$  = length of repeat \* indel rate

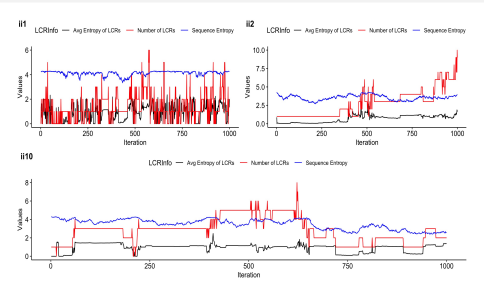
- $\begin{tabular}{ll} \textbf{0.14} & \textbf{mutation rate} = 0.14 \\ \textbf{Random Protein Sequence} \\ \textbf{GGAGGGAQ} \\ \end{tabular}$
- ② Assign Exponential Deviates mutation deviates = (0.83, 1.82, 2.35, 0.54, 0.98, 0.76, 1.53, 2.34) indel deviates = (0.21, 1.21, 1.49, 0.86, 0.97, 1.13, 0.53, 0.35) exp $(\beta)$ , where  $\beta$  = mutation rate exp $(\beta)$ , where  $\beta$  = length of repeat \* indel rate
- Point Mutation, Insertion, or Deletion
  Lowest value deviate = Residue that mutates fastest

- mutation rate = 0.14 indel rate = 0.14 Random Protein Sequence GGAGGGAQ
- ② Assign Exponential Deviates mutation deviates = (0.83, 1.82, 2.35, 0.54, 0.98, 0.76, 1.53, 2.34) indel deviates = (0.21, 1.21, 1.49, 0.86, 0.97, 1.13, 0.53, 0.35) exp $(\beta)$ , where  $\beta$  = mutation rate exp $(\beta)$ , where  $\beta$  = length of repeat \* indel rate
- Opint Mutation, Insertion, or Deletion Lowest value deviate = Residue that mutates fastest
- Upon point mutation, insertion, or deletion, scan the sequence again to see if the landscape of the sequence was affected, assign new deviates to affected amino acids only.

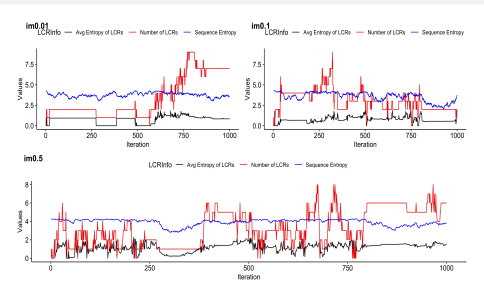
### Lower Indel Rates Produce Less LCRs



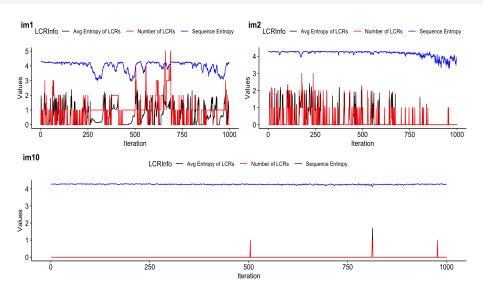
# Higher Indel Rates Produce More LCRs



### Lower Mutation Rates Produce More LCRs



## Large Mutation Rates Prevent the Formation of LCRs



## Bayesian Statistics Overview

#### Bayesian Statistics: Model-based statistical inference

$$p(D|\theta) \tag{1}$$

Likelihood

## Bayesian Statistics Overview

#### Bayesian Statistics: Model-based statistical inference

$$p(D|\theta) \tag{1}$$

Likelihood

$$p(\theta|D) \tag{2}$$

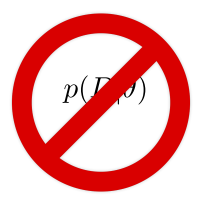
Posterior

## Why use an ABC-MCMC

► The increasing complexity and magnitude of available data can make the likelihood difficult to calculate

$$p(D|\theta)$$

## Why use an ABC-MCMC



Calculation of the likelihood is replaced with a simulation step

### MCMC for ABC

- **①** Propose a move from  $\theta$  to  $\theta'$  according to a transition kernel  $q(\theta, \theta')$ .
- ② Generate simulated dataset D' using  $\theta'$  and calculate S'.
- **3** If  $\rho(S', S) \leq \epsilon$  continue to 4, otherwise remain at  $\theta$  and go to 1.
- Calculate

$$\alpha(\theta, \theta') = min(1, \frac{\pi(\theta')q(\theta', \theta)}{\pi(\theta)q(\theta, \theta')})$$

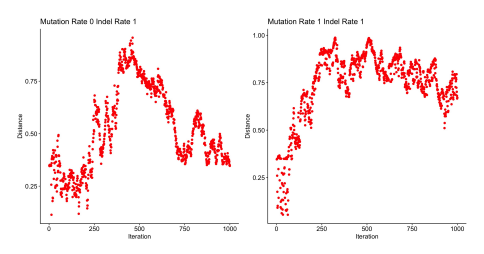
- **5** Accept  $\theta'$  with probability  $\alpha$ , otherwise stay at  $\theta$ .
- Return to 1.

<sup>&</sup>lt;sup>4</sup>Marjoram et al., 2003

# MCMC for ABC: Modified Algorithm

- $\bullet$  Propose a move from  $\theta$  to  $\theta'$  according to the normal distribution N(0.0,1.0)
- ② Create and mutate a random protein sequence using  $\theta'$  to generate the simulated Dataset D' do this 1000 times per newly proposed parameter value.
- ullet Calculate summary statistics for simulated dataset D' (average of all 1000 vectors of summary statistics)
- If  $d(S',S) < previous\_distance$ , go to next step, otherwise employ a one-sample t-test to assess the probability of accepting a larger distance. If the newly proposed distance is close to the previous distance, there is a higher chance we accept the value, otherwise we reject.
- lacktriangle Accept  $\theta'$
- Return to step 1

# ABC MCMC Preliminary Results



### Conclusions

- Created and tested a program to simulate the evolution of low complexity regions based off of two evolutionary parameters, mutation and indel rates
- ► The simulation program is compatible in a program written for an ABC-MCMC
- Struggling with creating posterior distribution, potentially due to selection of poor summary statistics or a lack of weight placed on each statistic

# Acknowledgements

- Dr. Brian Golding
- Sam Long
- Zachery Dickson
- ▶ Johanna Enright