

Detecting Selection in Experimental Evolution Experiment

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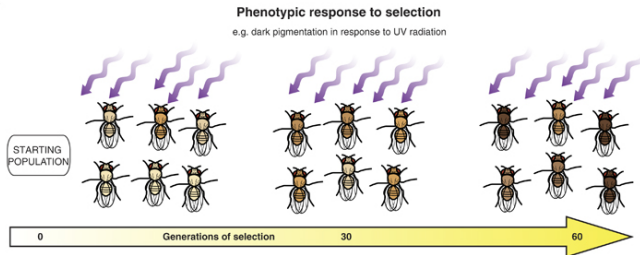
Introduction

- We are interested in (**quickly**) identifying genetic adaptations, in different organisms in find genes (or alleles) that are beneficial w.r.t. a **selective pressure**.
- Examples of interesting selective pressures:
 - harsh environmental conditions
 - antibiotics
 - chemotherapy
 - anti-viruses
 - pesticides
 - etc.

- **Classical Static Datasets** Given a single observation of Allele Frequency Spectrum (AFS) or Haplotypes of a region (say 50Kbp) a whole library of tests of selection is available.
- **Modern Dynamic Datasets**, multiple observations are available.

Experimental Evolution

a



b

Genotypic response to selection

e.g. causative variant increases in frequency



c

Pool-Seq

e.g. base and selected population



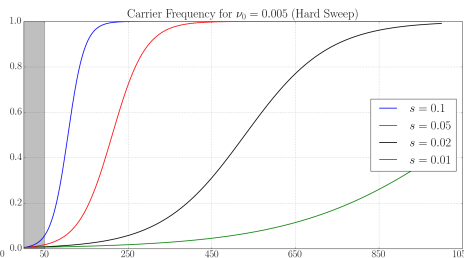
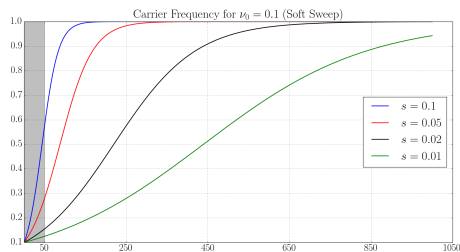
Goals

- Given the genome time series data (allele frequencies) we are interested in
 - I Detecting selection
 - II Locating the favored mutation
 - III Estimating strength of selection

Simulations

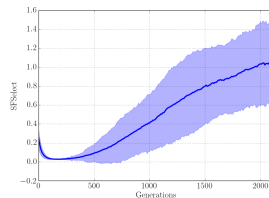
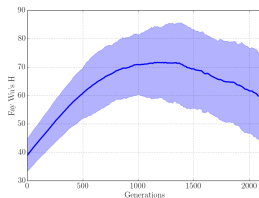
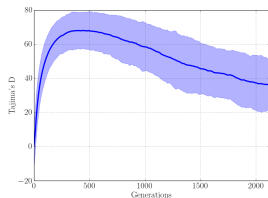
- (i) For each simulation, population of $F = 200$ founder lines is created in `msms` program with parameters
 - window size $L = 50\text{Kbp}$.
 - scaled mutation rate $\theta = 200$. (≈ 1100 SNPs)
- (ii) Using $F = 200$ founder lines a population diploid is created.
- (iii) Using forward simulator population is evolved and AF is sampled every 10 generation for 50 generations.
- (iv) This procedure is repeated 100 times for each $s = \{0.1, 0.05, 0.02, 0.01, 0\}$.

Single Locus Logistic Model: Carrier Frequency



Bottleneck

- side-effect of not tracking new mutations and restricting population to founder-lines.



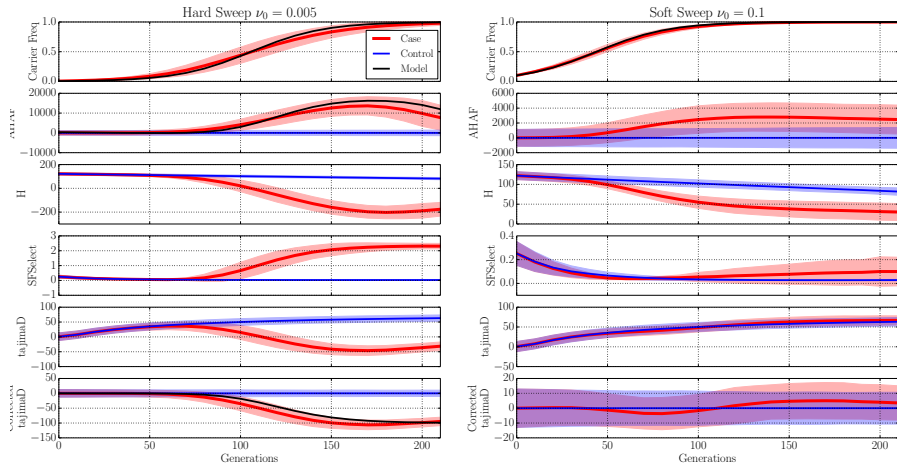


Figure: Mean and 95% CI of 1000 simulations for strong selection.

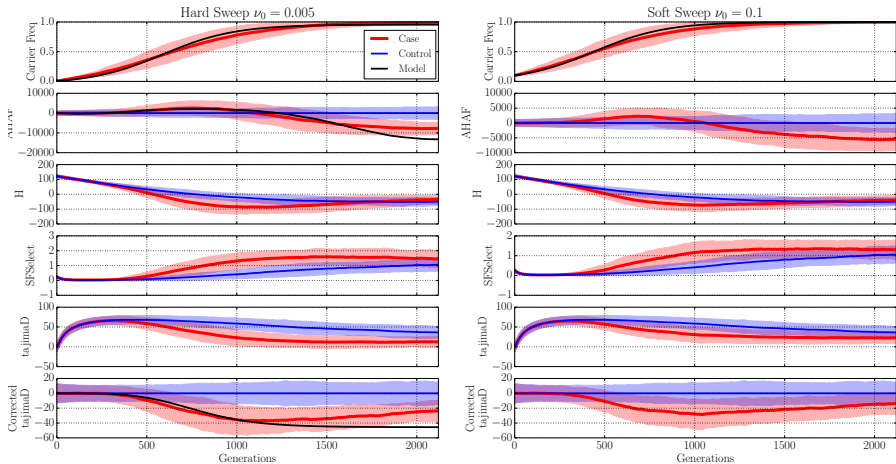
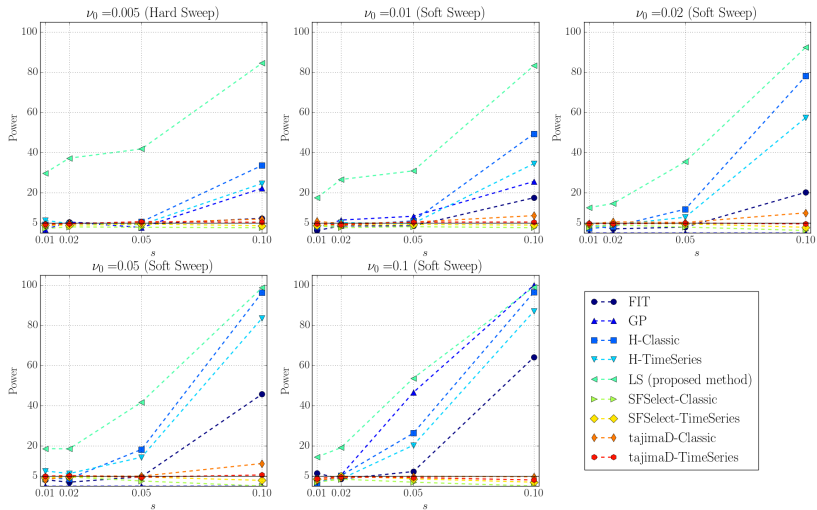


Figure: Mean and 95% CI of 1000 simulations for weak selection.

Experiments

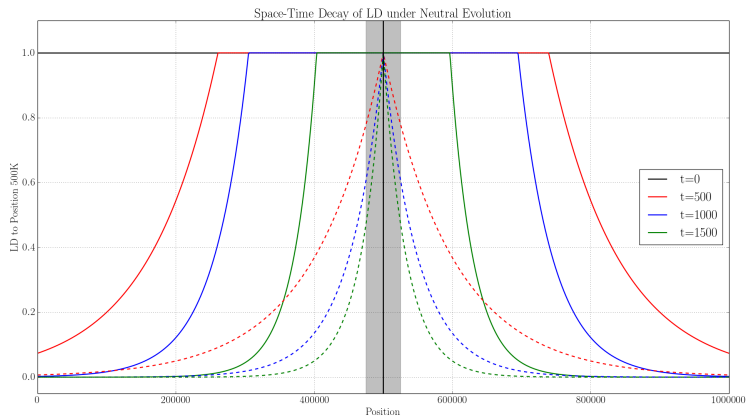
- for each setting we performs 200 simulations, (100 neutral and 100 selection), and predictive performance of all the methods in detecting selection.
- We computed ROC curve and defined power of a method as area under ROC curve when False-Positive rate is less than 0.1.

Power of detection

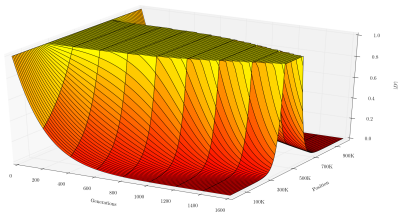
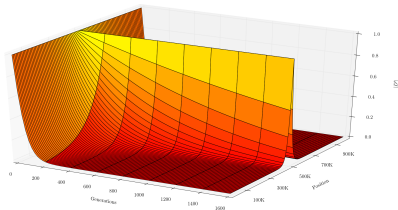


Locating selection I

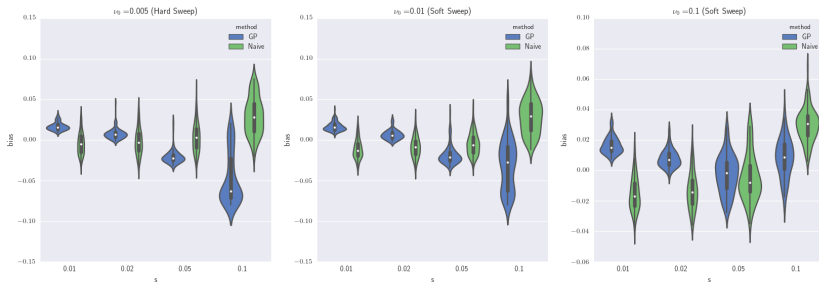
Strong LD in a window



Locating selection II



Power of detection



Summary

- Behaviour of different population statistics in time is studied.
- A method based on single locus AF is proposed and shown to have superior performance than multi-locus and traditional methods.
- Analysis of real data and locating genes in the genome is our next step.

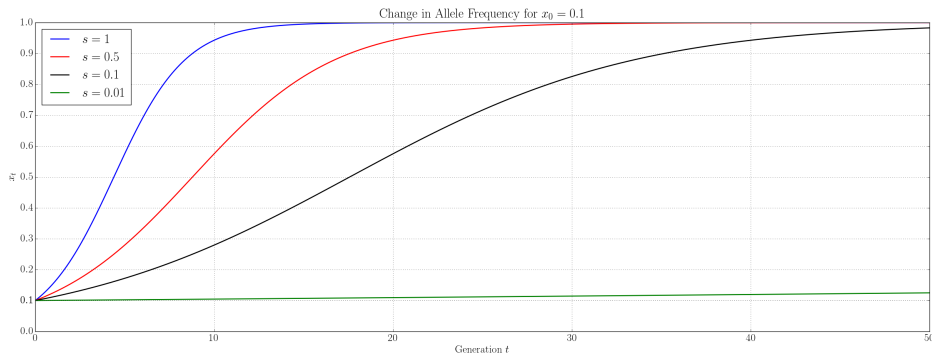
Thanks!

Single Locus Model: Carrier Frequency

- By differentiating update equations ($x_{t+1} = x_t + \frac{sx_t(1-x_t)}{2+2sx_t}$ w.r.t. t and solving differential equation, we have

$$\nu_t = \sigma(st/2 + \eta(\nu_0))$$

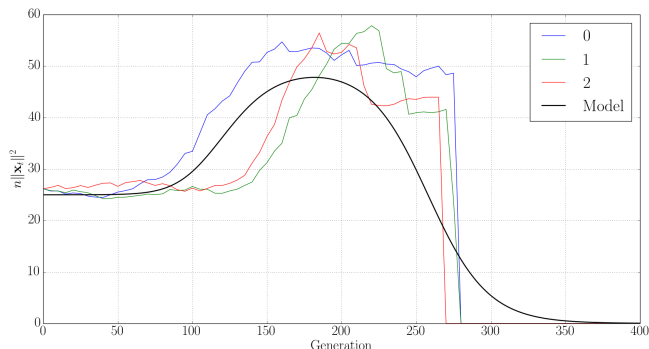
where $\sigma(\cdot)$ is logistic function and $\eta = \sigma^{-1}$ is the logit function and ν_t is the frequency of the carrier at time t .



Multi Locus Model: Average Haplotype Allele Frequency

$$\mathbb{E}[1\text{-HAF}(t)] = \|\mathbf{x}_t\|^2 \approx \theta \nu_t \left(\frac{\nu_t + 1}{2} - \frac{1}{(1 - \nu_t)n + 1} \right) + \theta(1 - \nu_t) \left(\frac{n + 1}{2n} - \frac{1}{(1 - \nu_t)n + 1} \right)$$

where \mathbf{x}_t is vector of AF at time t and $\nu_t = \sigma(st/2 + \eta(\nu_0))$.



- Its too complex, likely to overfit with small number of iid replicates.
- Although the likelihood model is based on different parameters, in practice, it can learn only one parameter at a time.
- not tractable, its time complexity is quartic!
- worse, each iteration requires **maxGeneration** recursion which makes it very hard to analyse late epochs of sweeps.
- In addition to PoolSeq data, it requires initial population haplotypes.
- Despite its elegant theory, it has not compared with classical methods.
- In practice, single locus scan is performed and multi-locus (with 3-7 seg. sites.) model is fitted at regions of interests.