

Introduction to Wright-Fisher Simulations

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Goals

- Simulate the standard neutral model, demographic effects, and natural selection

Hardy-Weinberg Principle



Godfrey H. Hardy:
1877-1947



Wilhelm Weinberg:
1862-1937

● Assumptions:

- Diploid organism
- Sexual reproduction
- Non-overlapping generations
- Only two alleles
- Random mating
- Identical frequencies in males/females
- Infinite population size
- No migration
- No mutation
- No natural selection

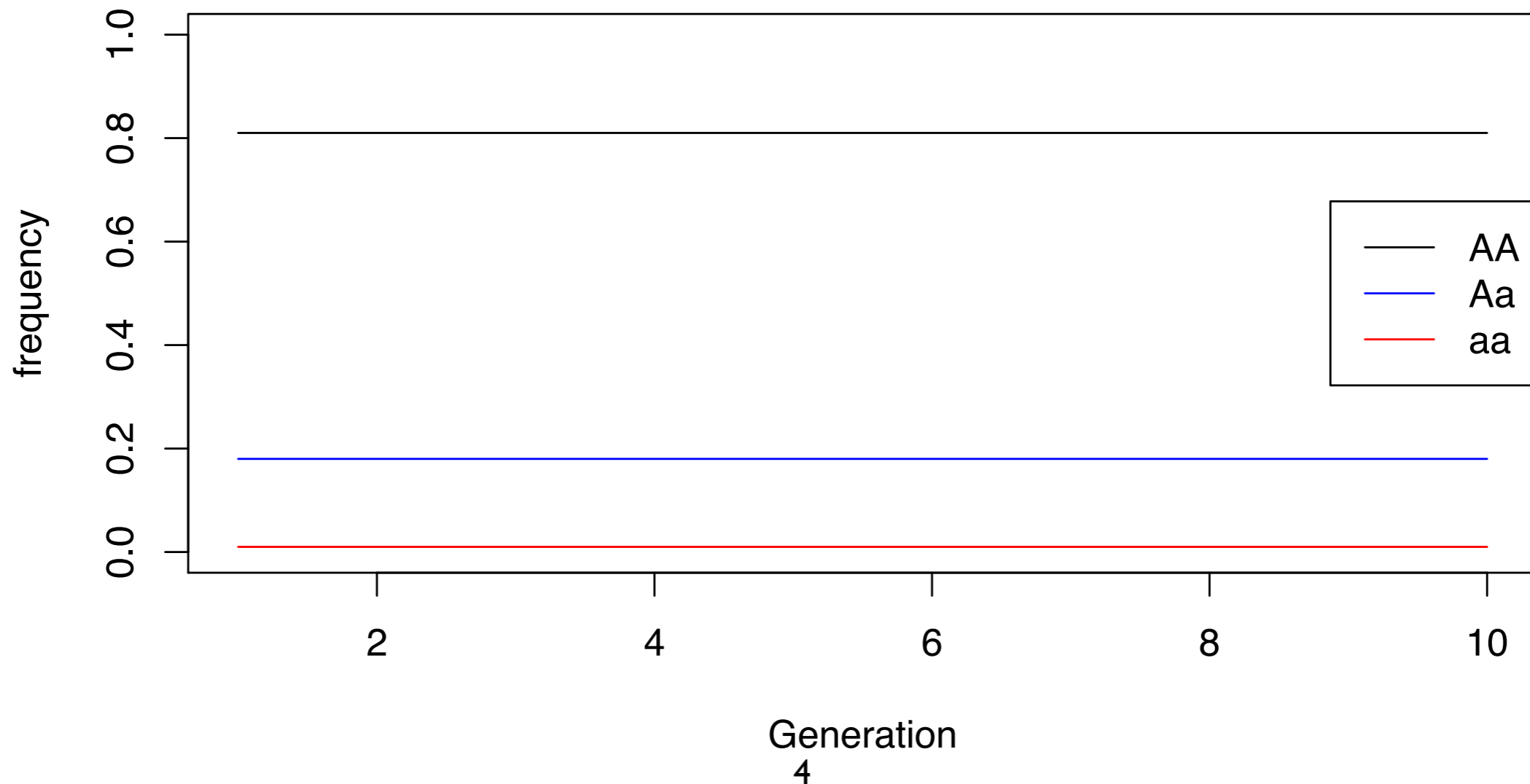
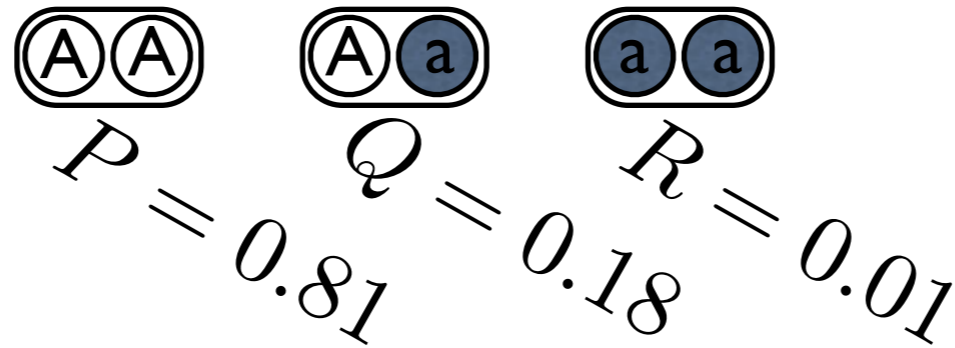
● Conclusion I:

Both allele AND genotype frequencies will remain constant at **HWE** generation after generation... forever!

$$P=p^2$$
$$Q=2p(1-p)$$
$$R=(1-p)^2$$

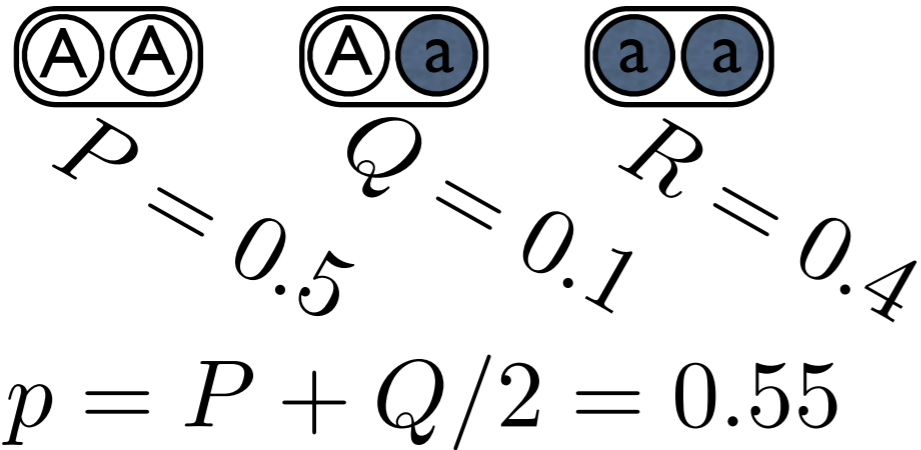
Hardy-Weinberg Principle

- Imagine a population of diploid individuals

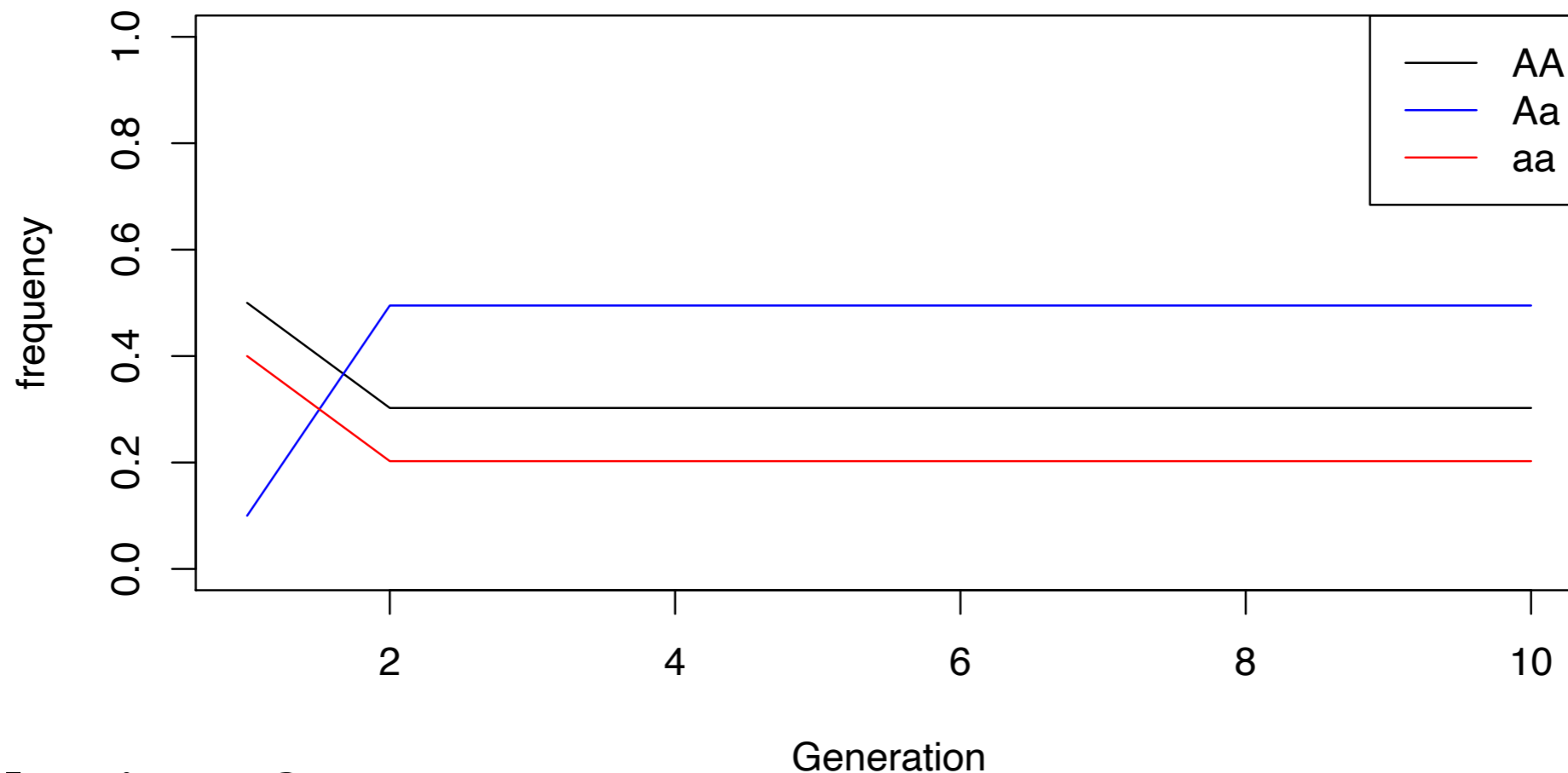


Hardy-Weinberg Principle

- Imagine a population of diploid individuals



$$p^2 = 0.3025$$
$$2p(1 - p) = 0.495$$
$$(1 - p)^2 = 0.2025$$



- Conclusion 2:** A single round of random mating will return the population to HWE frequencies!

Hardy-Weinberg Principle



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1877-1947



Wilhelm Weinberg:
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● Assumptions:

- Diploid organism
- Sexual reproduction
- Non-overlapping generations
- Only two alleles
- Random mating

- Identical frequencies in males/females
- Infinite population size
- No migration
- No mutation
- No natural selection

Genetic Drift

- In finite populations, allele frequencies can and do change over time.
- In fact, EVERY genetic variant will either be lost from the population ($p=0$) or fixed in the population ($p=1$) some time in the future.
- The most common model for finite populations is the **Wright-Fisher model**.
- This model makes explicit use of the *binomial distribution*.

Wright-Fisher Model



Sewall Wright:
1889-1988



Sir Ronald Fisher
1890-1962

- Suppose a population of N individuals.
- Let $X(t)$ be the #chromosomes carrying an allele A in generation t :

$$\begin{aligned} P(X(t+1) = j | X(t) = i) &= \binom{N}{j} p^j (1-p)^{N-j} \\ &= \text{Bin}(j | N, i/N) = \binom{N}{j} \left(\frac{i}{N}\right)^j \left(\frac{N-i}{N}\right)^{N-j} \end{aligned}$$

Wright-Fisher Model

- A simple R function to simulation genetic drift:

```
WF=function(N, p, G){  
  t=array(,dim=G);  
  t[1] = p;  
  for(i in 2:G){  
    t[i] = rbinom(1,N,t[i-1])/N;  
  }  
  return(t);  
}
```

Initial pop size
Starting frequency
number of simulated generations

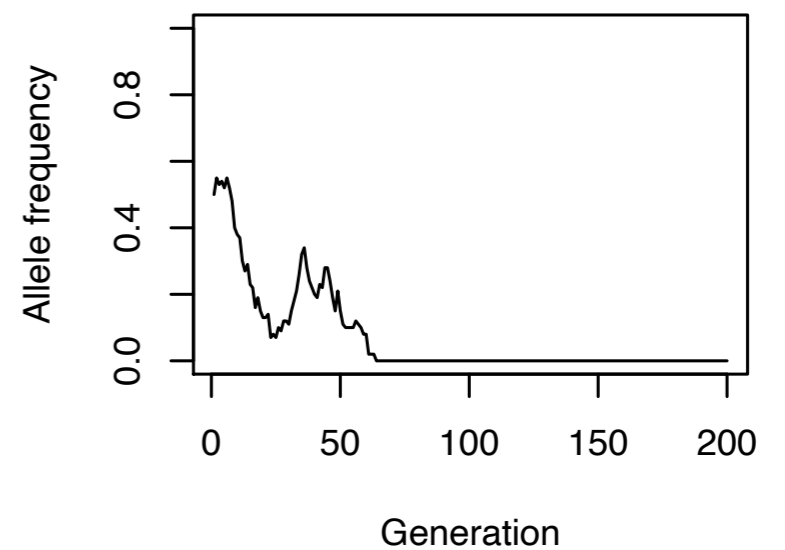
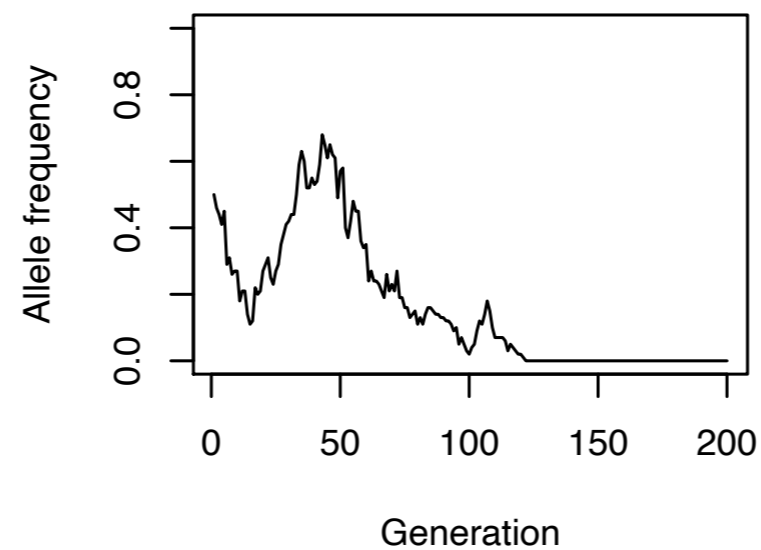
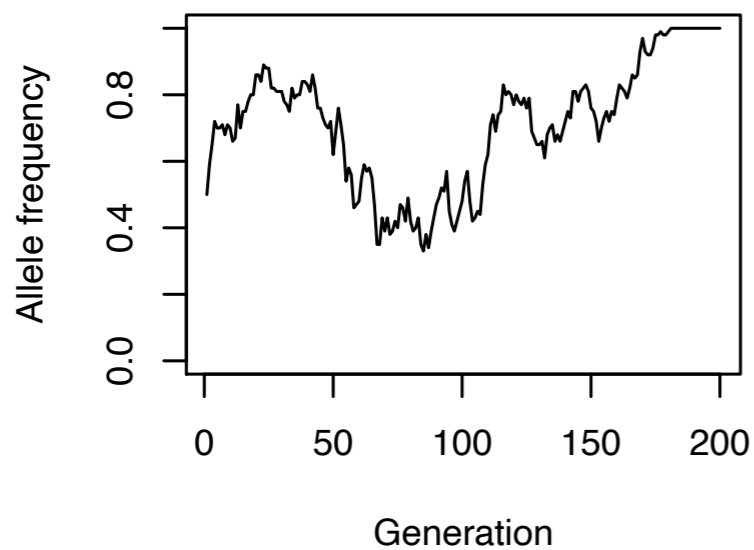
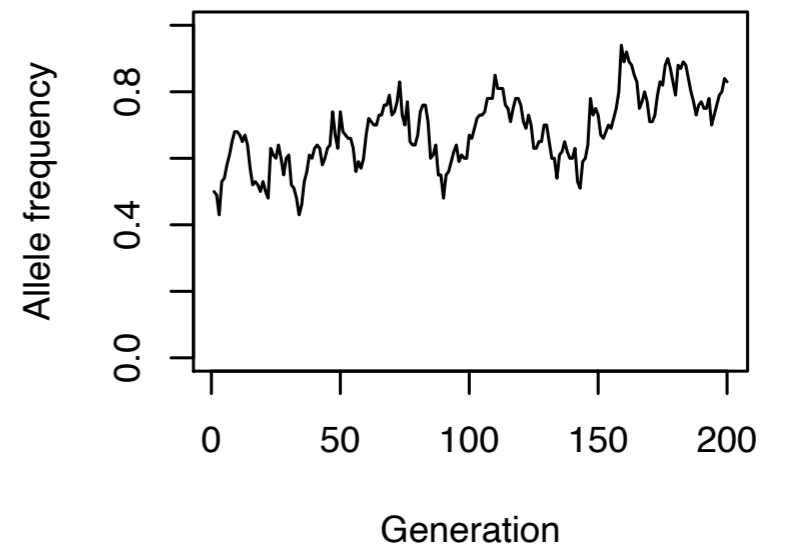
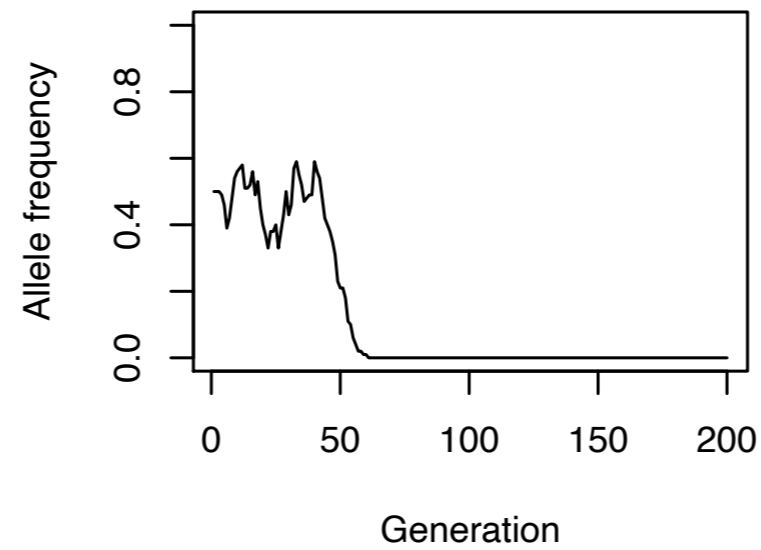
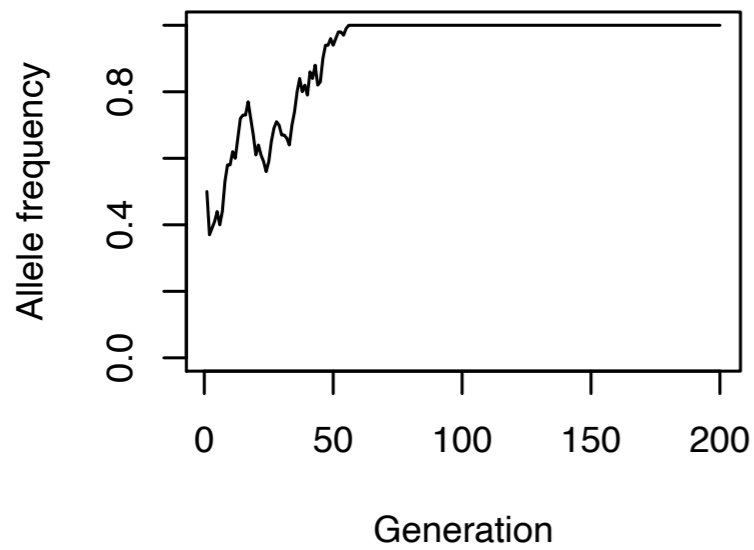
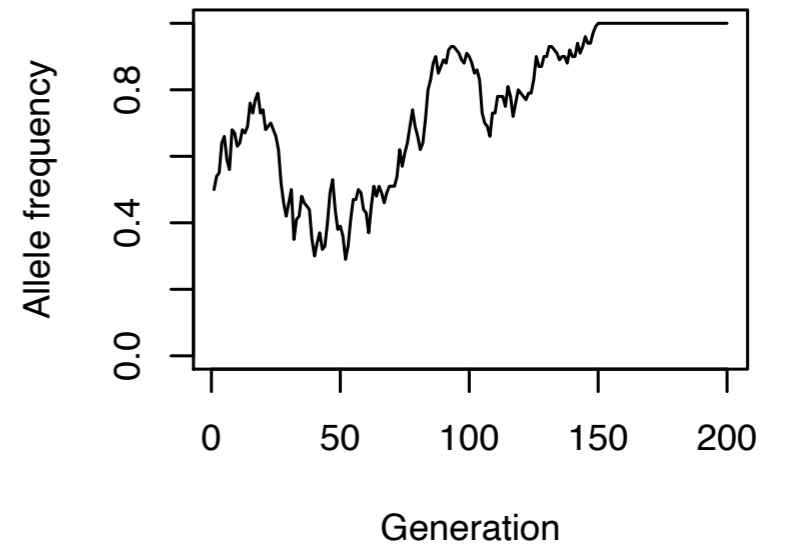
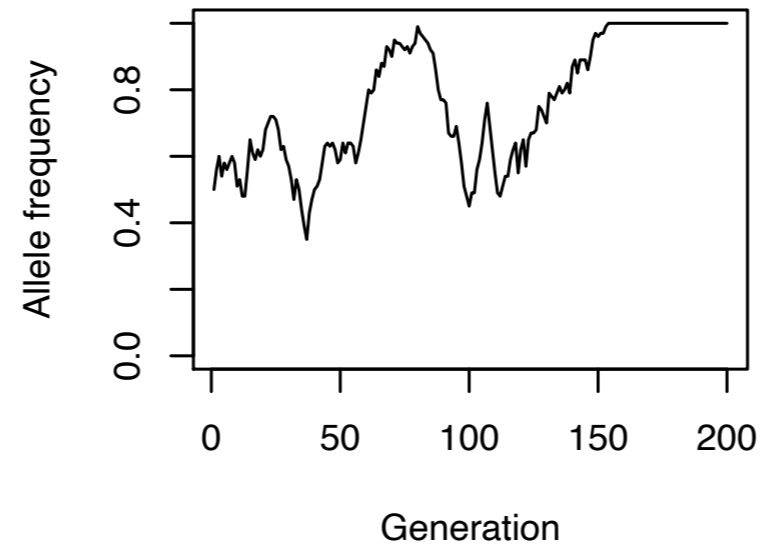
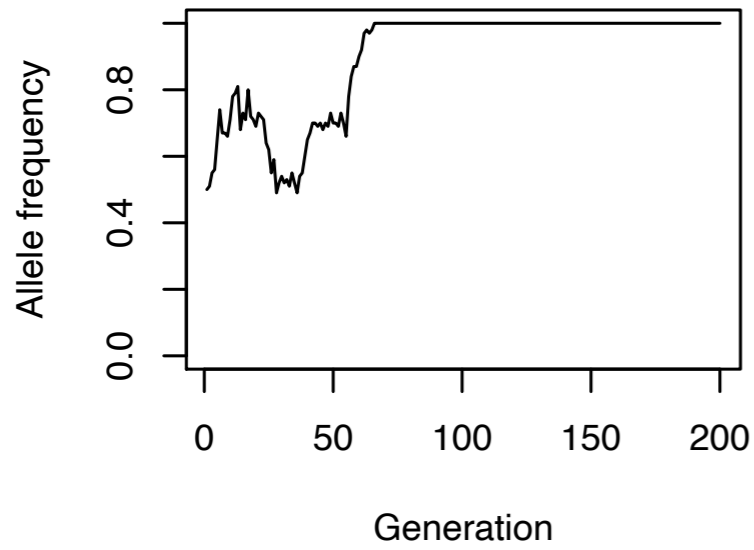
- Run it in R using:

```
f=WF(100, 0.5, 200)  
plot(f)
```

Breakout Groups

- Please work together to code this up and generate the plot.
- Let us know if you have questions, or call for help!
- What happens in your plot?
- Were you able to get any fixations or losses?

Wright-Fisher Model



Demographic Effects

- Population changes size at a given generation

Wright-Fisher Model



Sewall Wright:
1889-1988



Sir Ronald Fisher
1890-1962

- Suppose a population of N individuals.
- Let $X(t)$ be the #chromosomes carrying an allele A in generation t :

$$\begin{aligned} P(X(t+1) = j | X(t) = i) &= \binom{N}{j} p^j (1-p)^{N-j} \\ &= \text{Bin}(j | N, i/N) = \binom{N}{j} \left(\frac{i}{N}\right)^j \left(\frac{N-i}{N}\right)^{N-j} \end{aligned}$$

Wright-Fisher Model

- A simple R function to simulation demographic effects:

```
WFdemog = function(N, p, G, Gd, v){  
  t=array(,dim=G);  
  t[1] = p;  
  for(i in 2:G){  
    if(i == Gd){  
      N = N*v;  
    }  
    t[i] = rbinom(1,N,t[i-1])/N;  
  }  
  return(t);  
}
```

Initial pop size
Starting frequency
Generations to simulate
Gen demographic event happens
Magnitude of size change

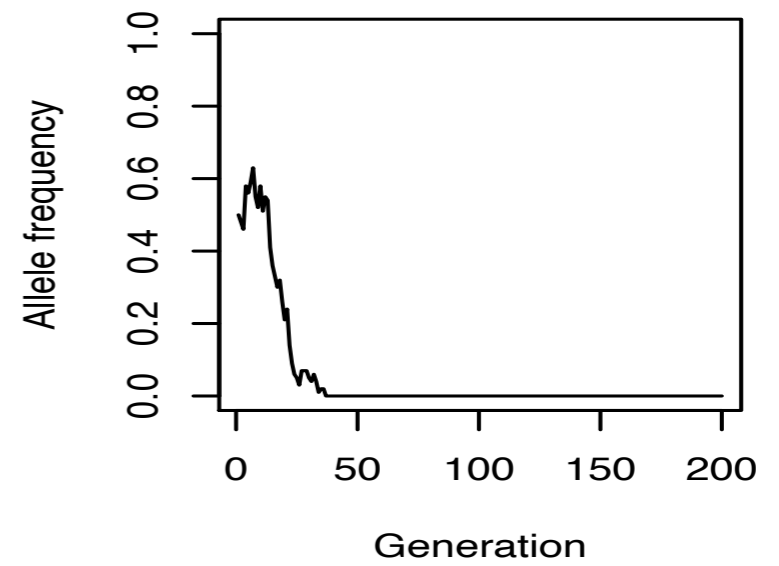
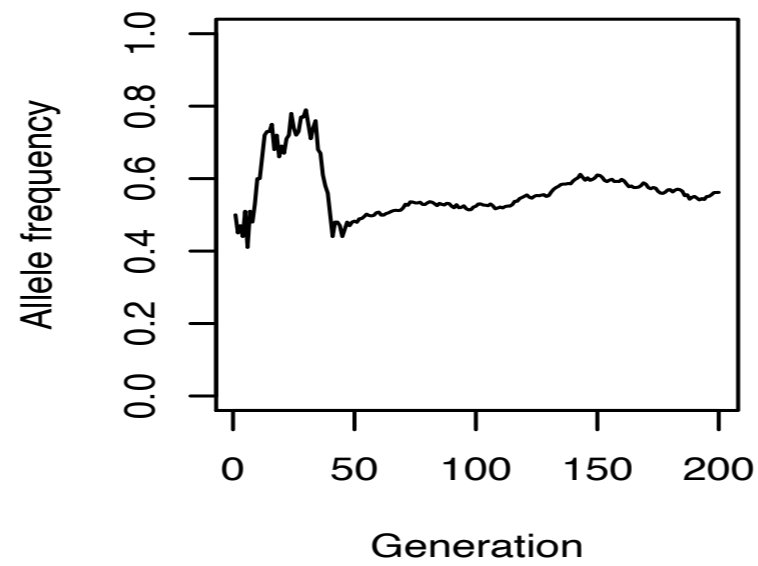
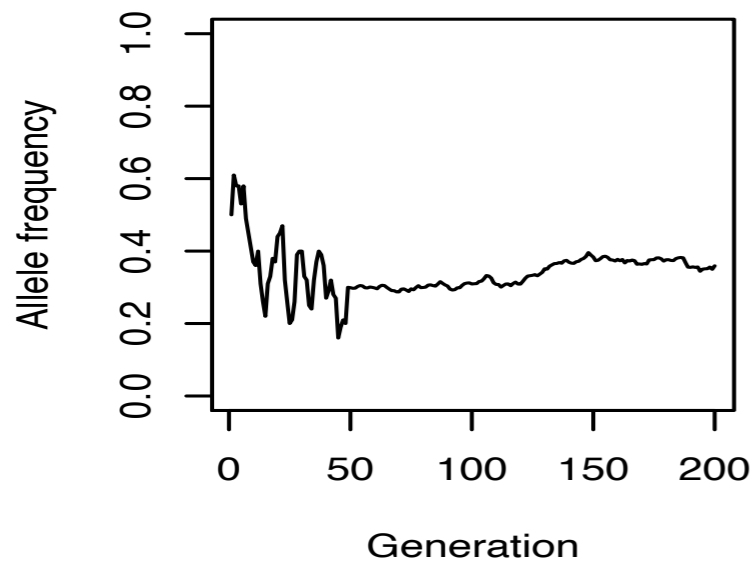
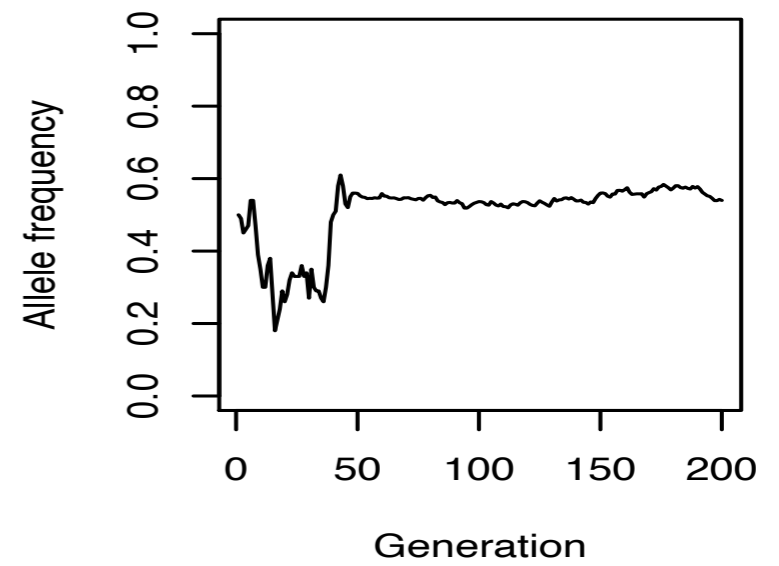
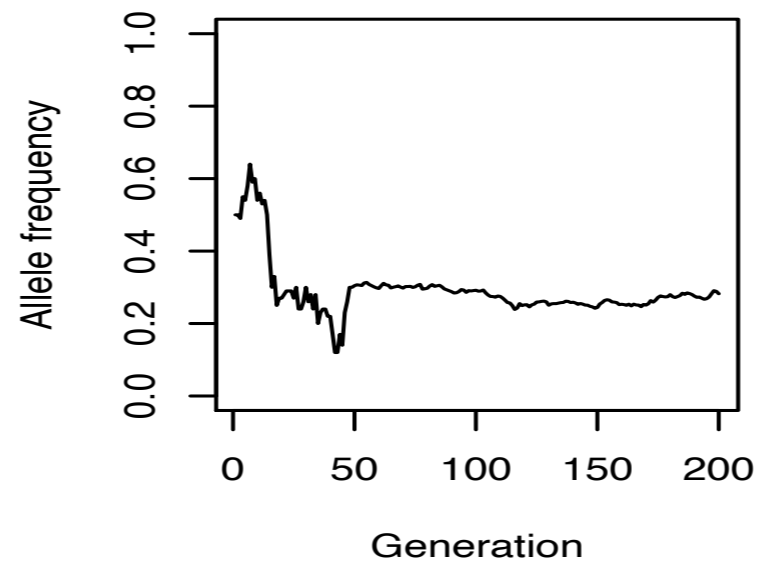
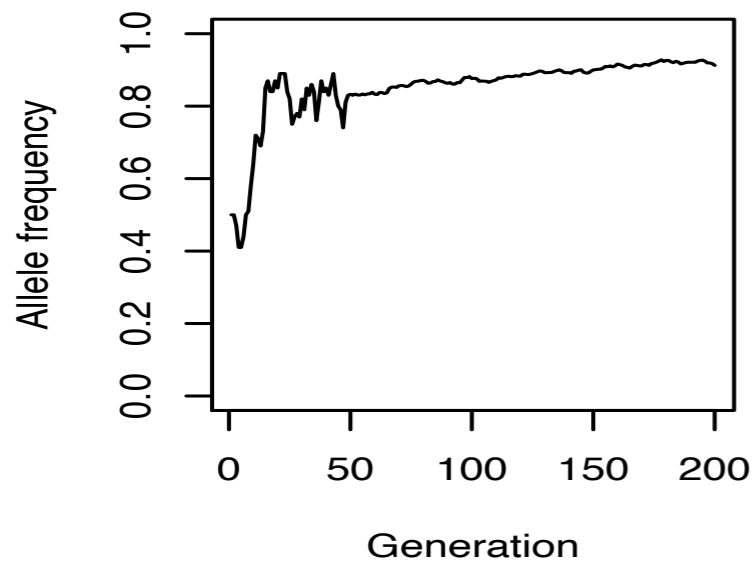
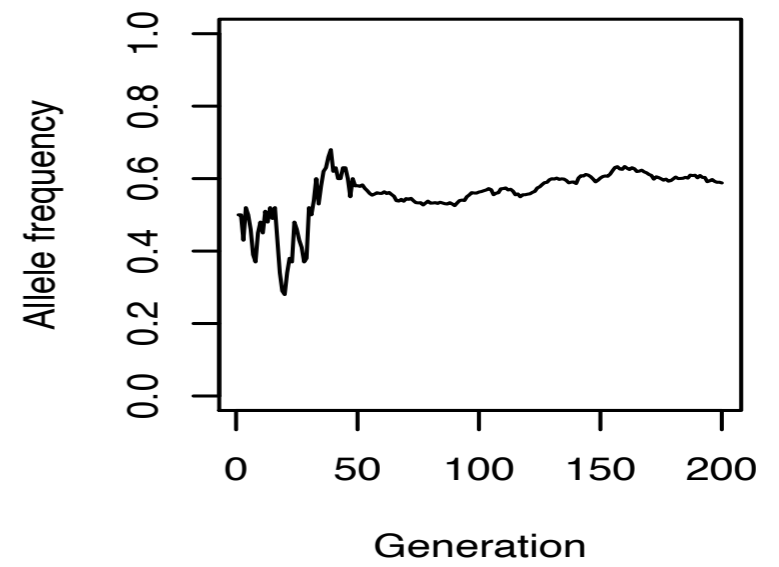
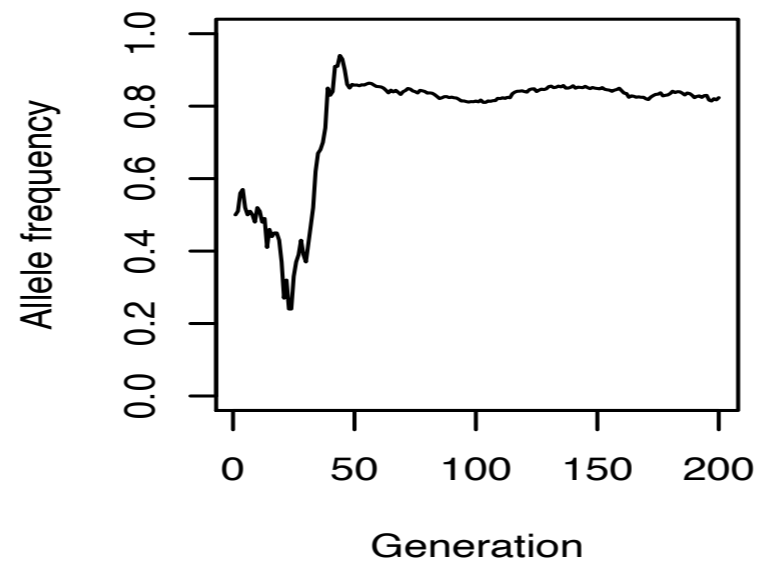
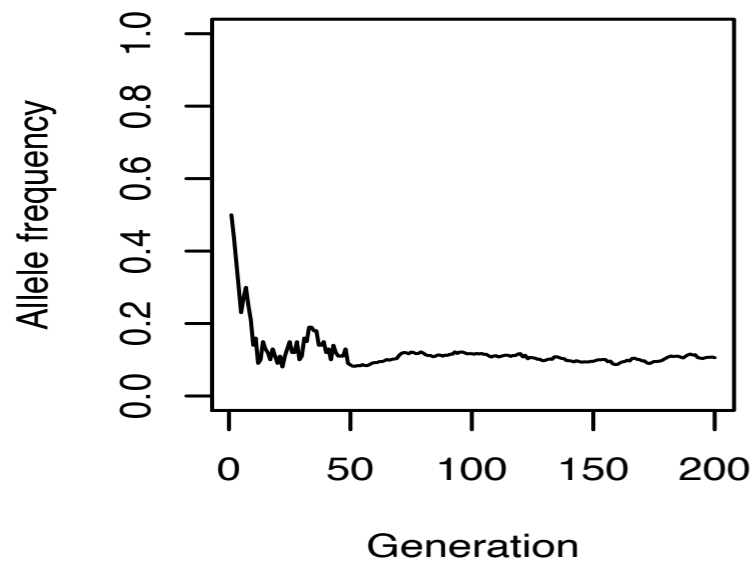
- Run it using:

```
f=WFdemog(100, 0.5, 200, 50, 100)  
plot(f)
```

Breakout Groups

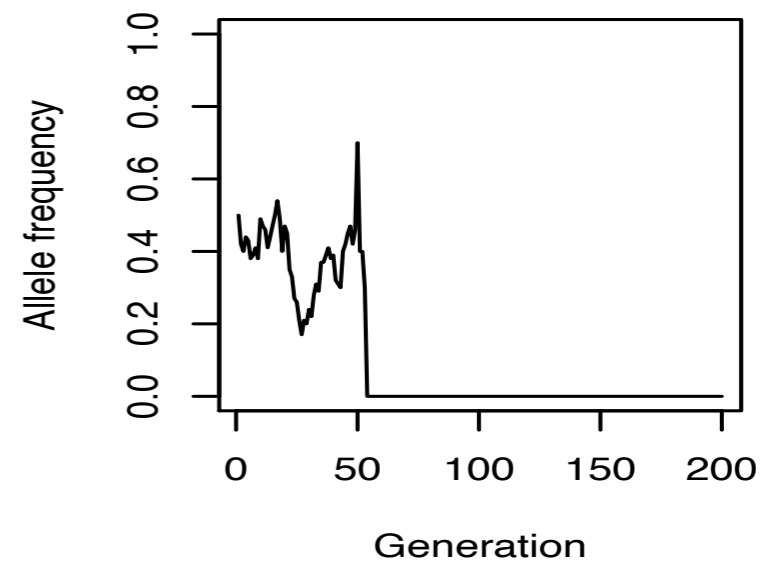
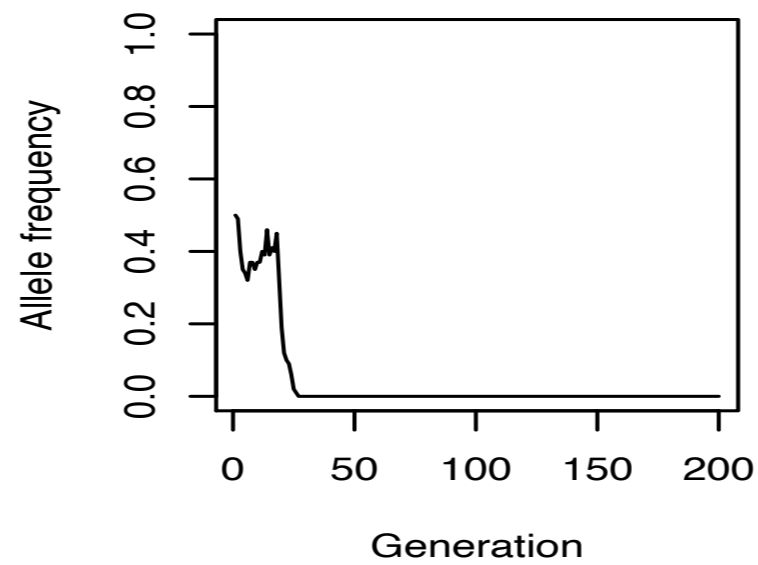
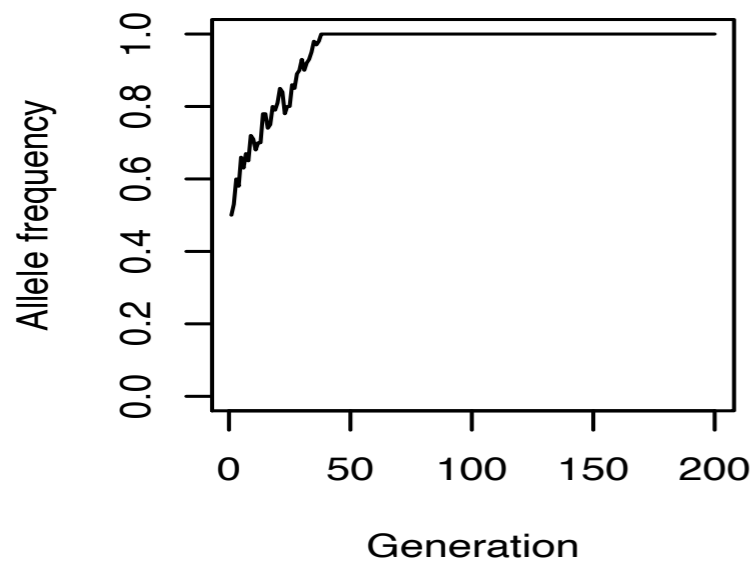
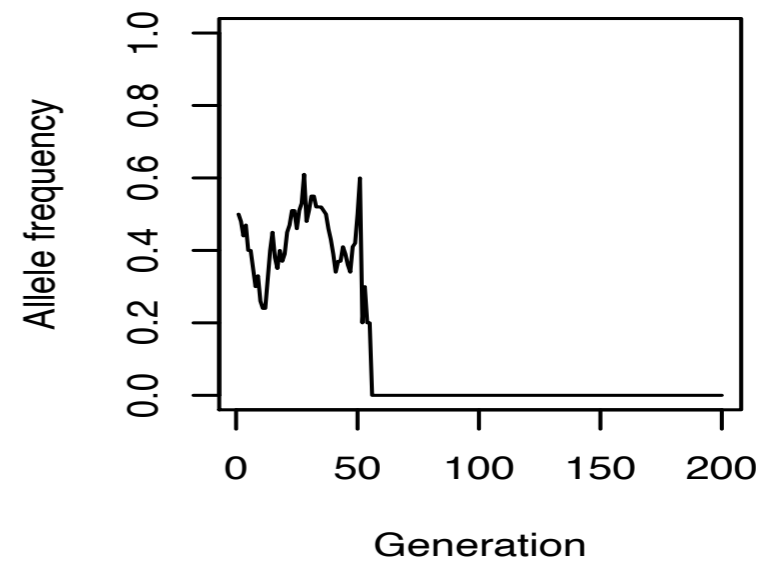
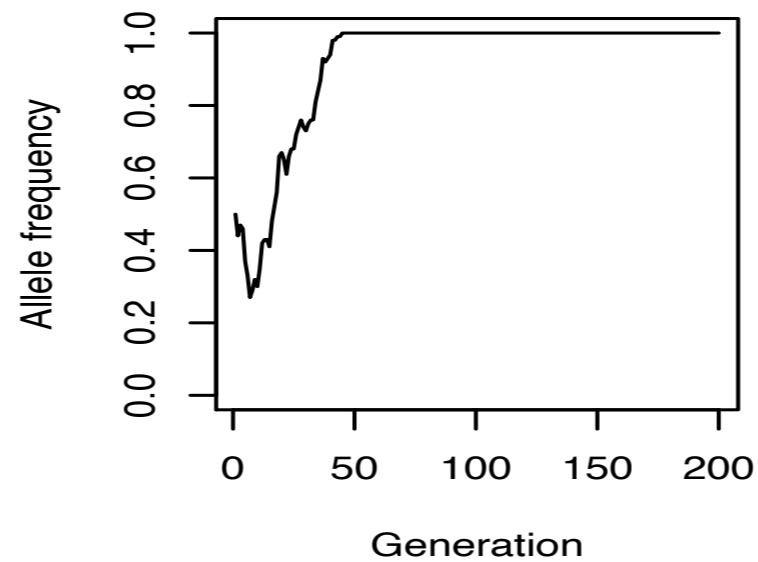
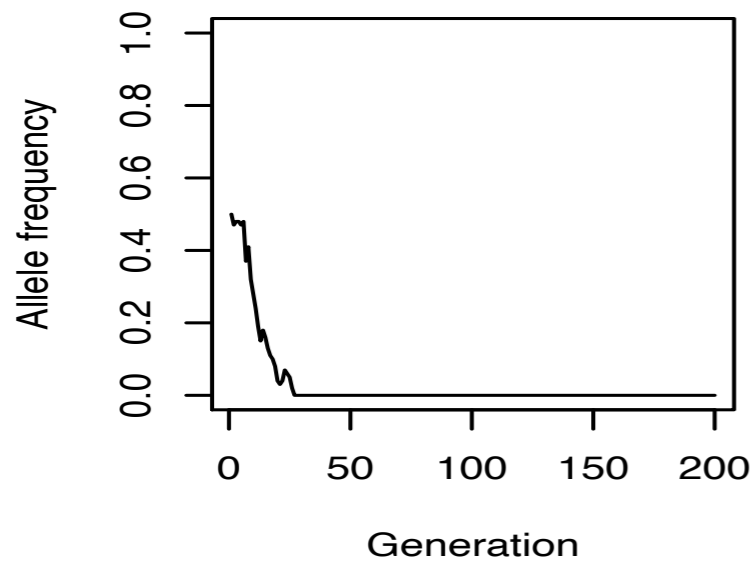
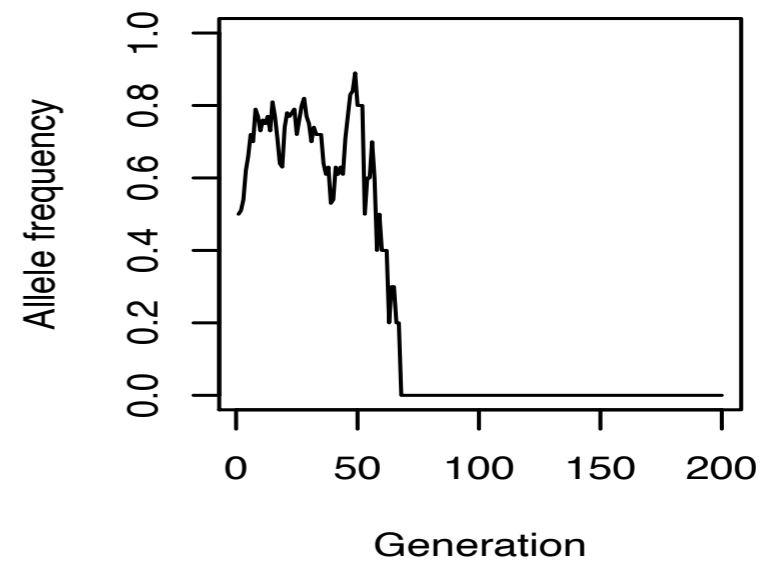
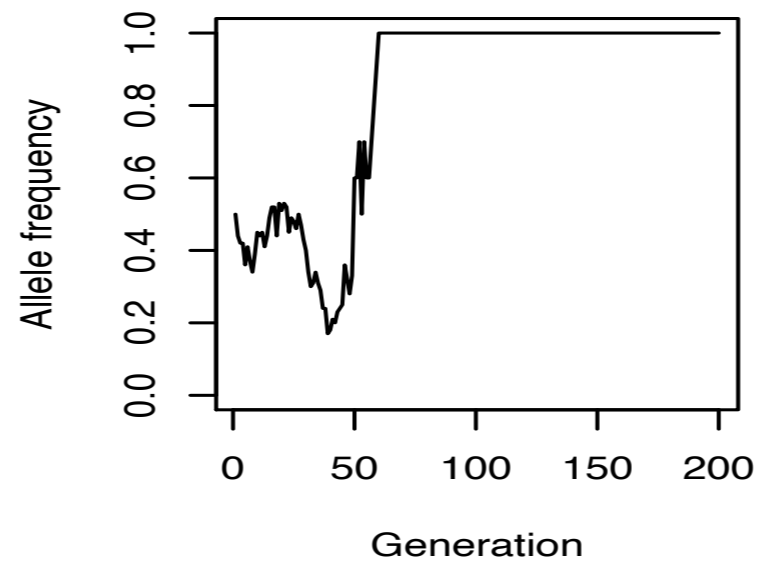
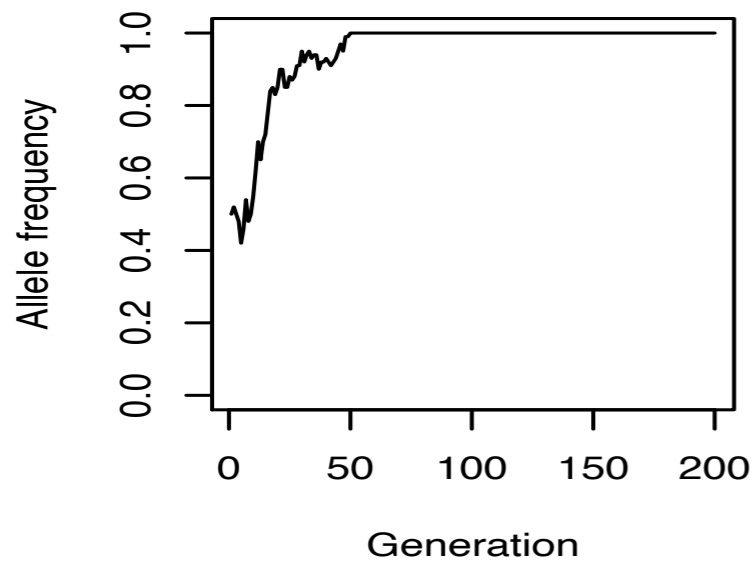
- Please work together to code this up and generate the plot.
- Let us know if you have questions, or call for help!
- What happens in your plot?
 - Were you able to get any fixations or losses?
- Can you simulate a 10-fold contraction?
 - How does it change the trajectory?

Wright-Fisher Model with Expansion



Wright-Fisher Model with Contraction

- Run it using: `WFdemog(100, 0.5, 200, 50, 0.1)`



Hardy-Weinberg Principle

● **Assumptions:**

- Diploid organism
 - Sexual reproduction
 - Non-overlapping generations
 - Only two alleles
 - Random mating
 - Identical frequencies in males/females
 - Infinite population size
 - No migration
 - No mutation
 - No natural selection
- What happens when we allow natural selection to occur?
 - Alleles change frequency!

Natural Selection

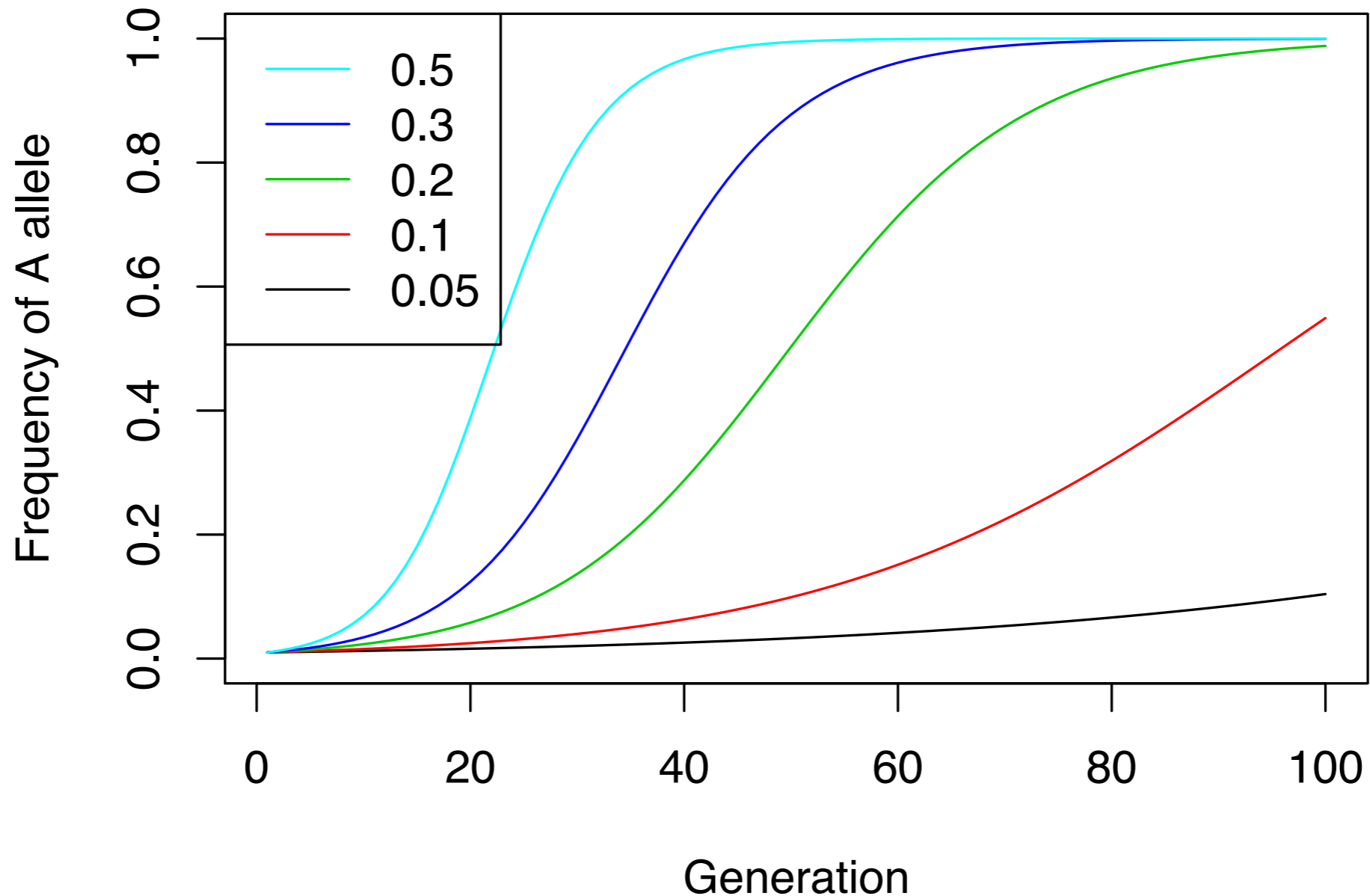
Genotype	AA	Aa	aa
Frequency	p^2	$2pq$	q^2
Fitness	1	$1+hs$	$1+s$

- The *expected frequency* in the next generation (q') is then the density of offspring produced by carriers of the derived allele divided by the population fitness:

$$q' = \frac{q^2(1+s) + pq(1+hs)}{1 + sq(2hp + q)}$$

Natural Selection

- Trajectory of selected allele with various selection coefficients under genic selection ($h=0.5$) in an “infinite” population



Hardy-Weinberg Principle

● Assumptions:

- Diploid organism
- Sexual reproduction
- Non-overlapping generations
- Only two alleles
- Random mating
- Identical frequencies in males/females
- Infinite population size
- No migration
- No mutation
- No natural selection

- What happens with natural selection in a finite population?
 - Directional selection AND drift!

Simulating Natural Selection

- First write an R function for the change in allele frequencies:

```
fitfreq = function(q, h, s){  
  p=1-q;  
  return((q^2*(1+s) + p*q*(1+h*s))/(1 + s*q*(2*h*p+q)));  
}
```

*initial freq
dominance
fitness*

- Now use this in an updated WF simulator:

```
WF.sel=function(N, q, h, s, G){  
  t=array(, dim=G);  
  t[1] = q;  
  for(i in 2:G){  
    t[i] = rbinom(1, N, fitfreq(t[i-1], h, s))/N;  
  }  
  return(t);  
}
```

*pop size
initial freq
dominance
fitness
gens to simulate*

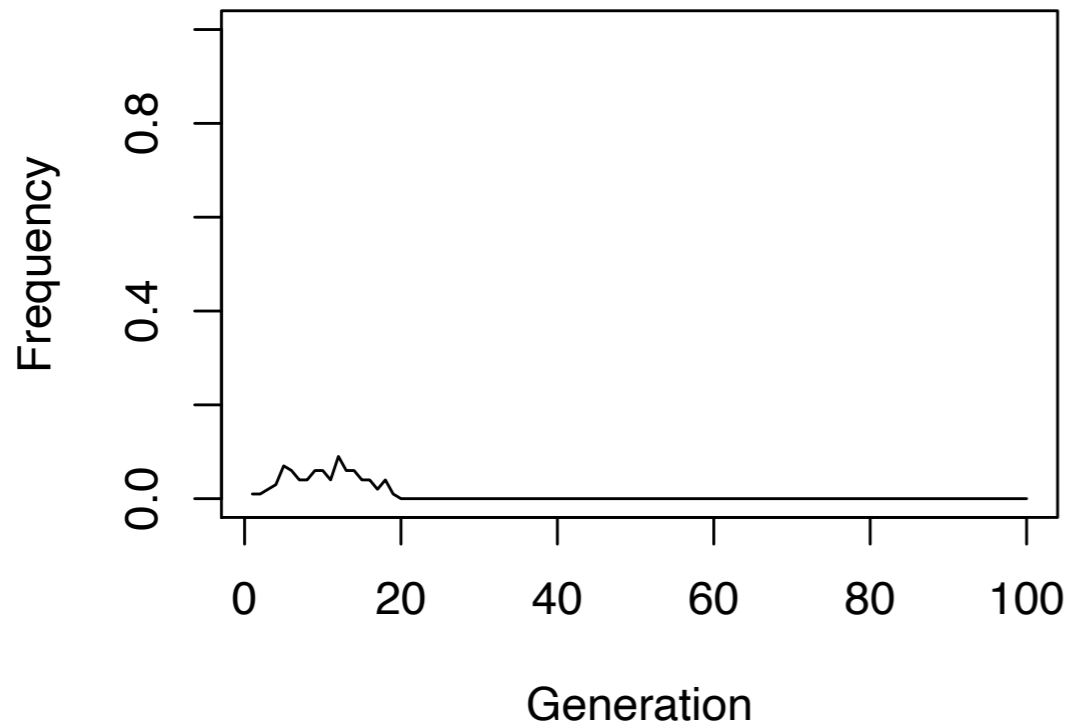
Breakout Groups

- Please work together to code this up.
- Can you simulate a trajectory for 100 generations with these characteristics:
 - Population size = 100
 - Initial frequency is 1%
 - Allele has a 50% fitness advantage
- What happens in your plot?
 - Were you able to get any fixations or losses?

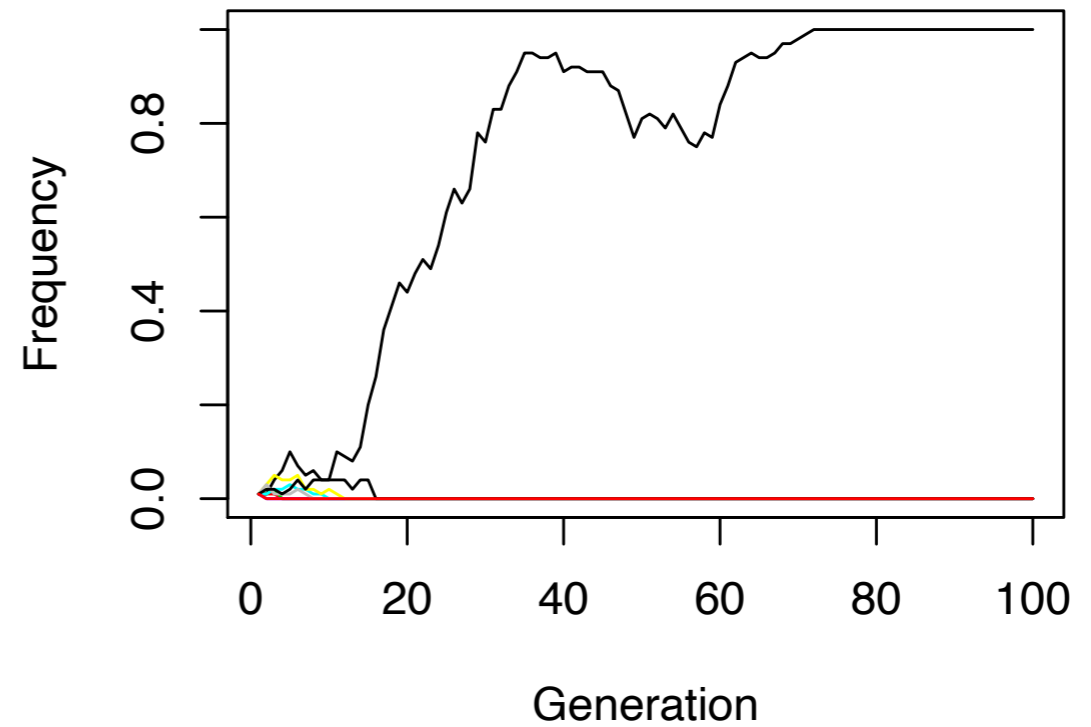
Natural Selection

`WF.sel(100, 0.01, 0.5, 0.1, 100)`

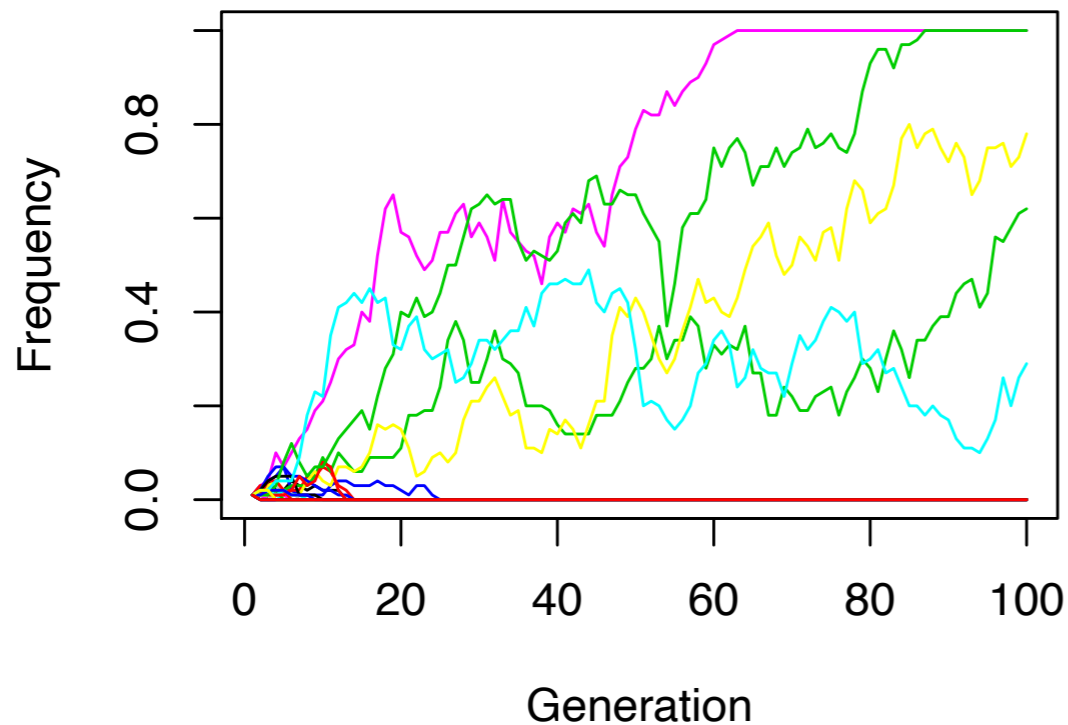
1 simulations



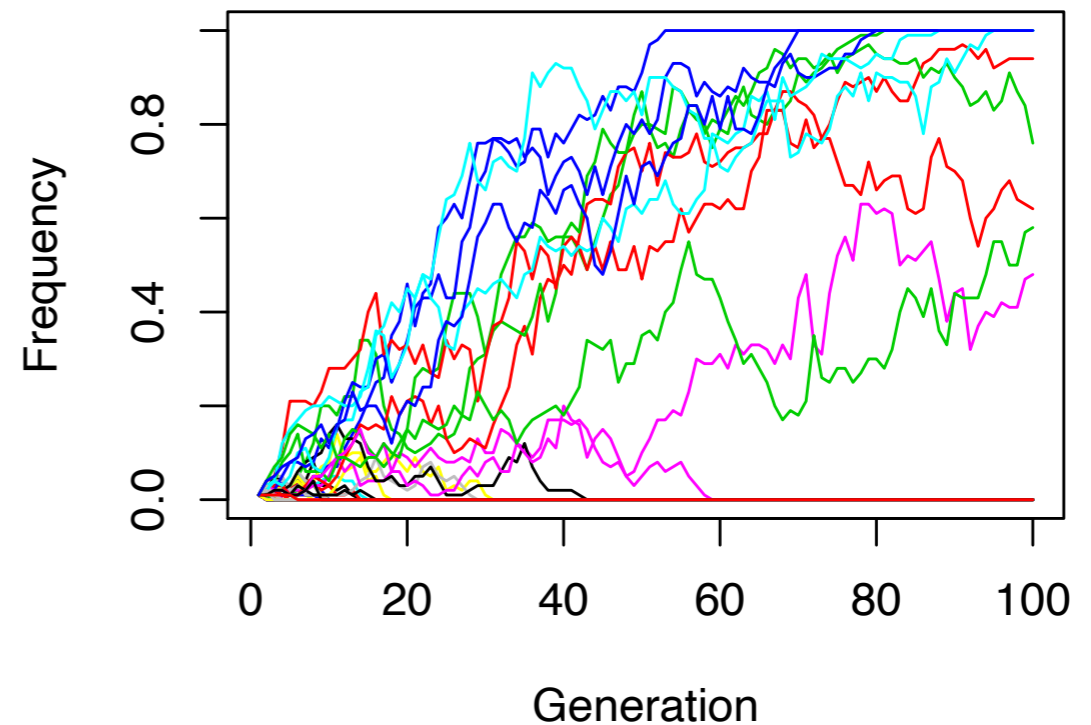
10 simulations



50 simulations



100 simulations



Simulating Natural Selection

- How would you simulate both selection AND demographic effects?

- Now use this in an updated WF simulator:

```
WF.demsel=function(N, q, h, s, G, Gd, v){  
  t=array(,dim=G);  
  t[1] = q;  
  for(i in 2:G){  
    if(i == Gd){  
      N = N*v;  
    }  
    t[i] = rbinom(1, N, fitfreq(t[i-1], h, s))/N;  
  }  
  return(t);  
}
```

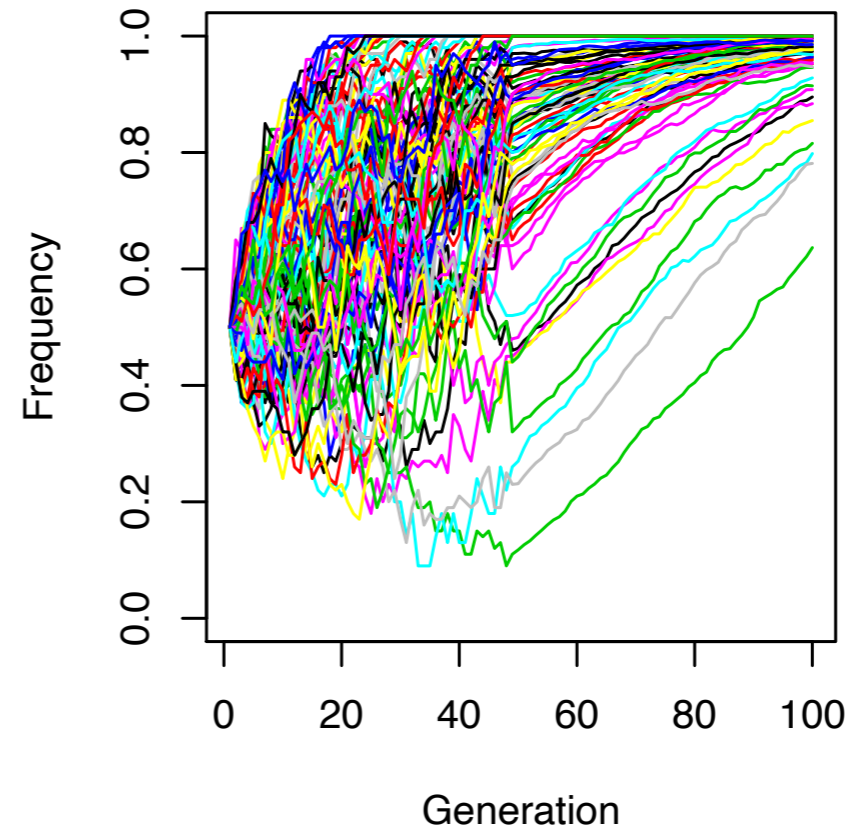
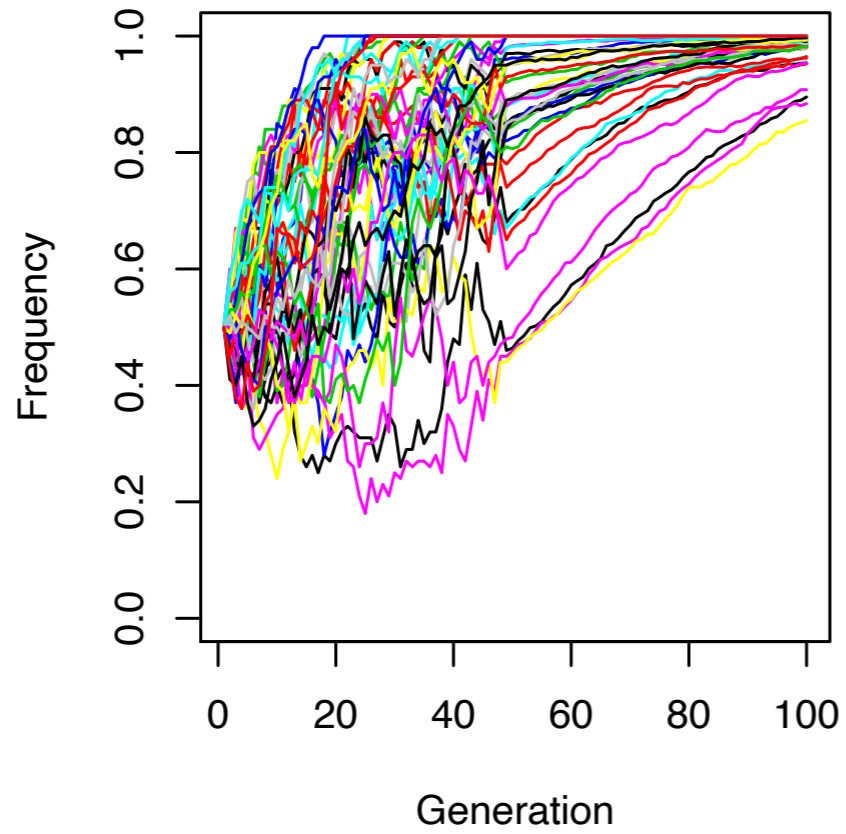
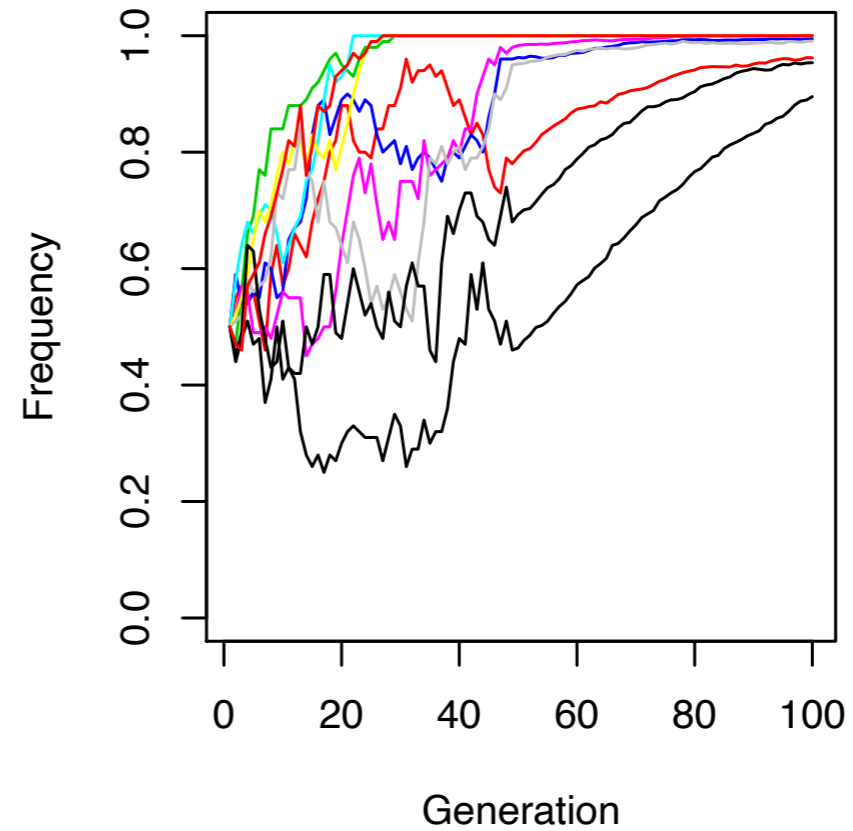
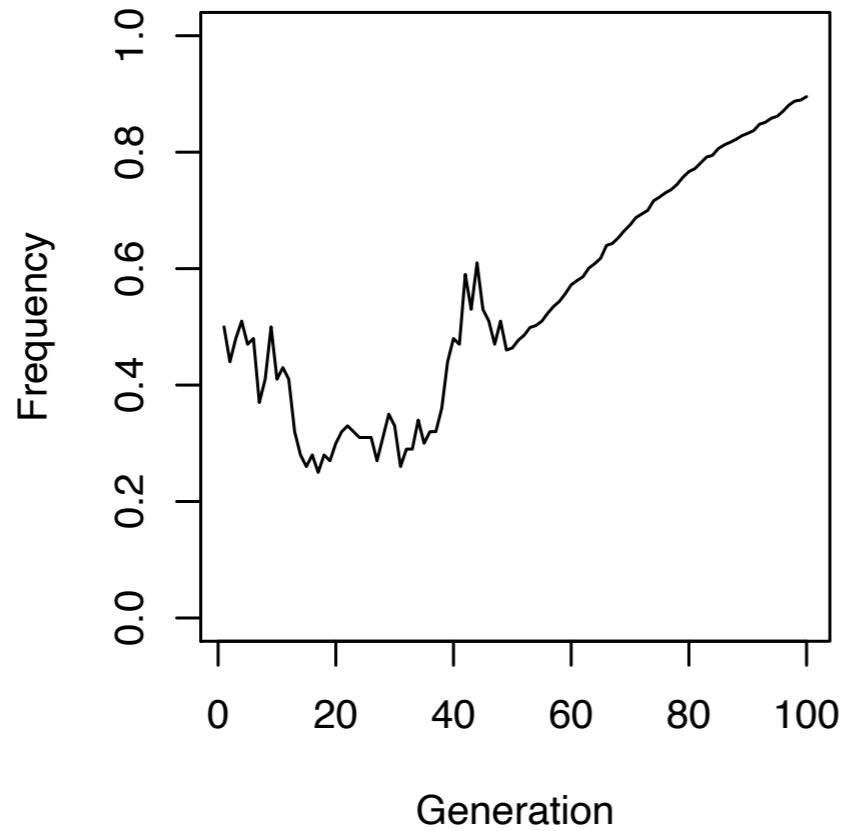
pop size
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Gen demographic event happens
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Breakout Groups

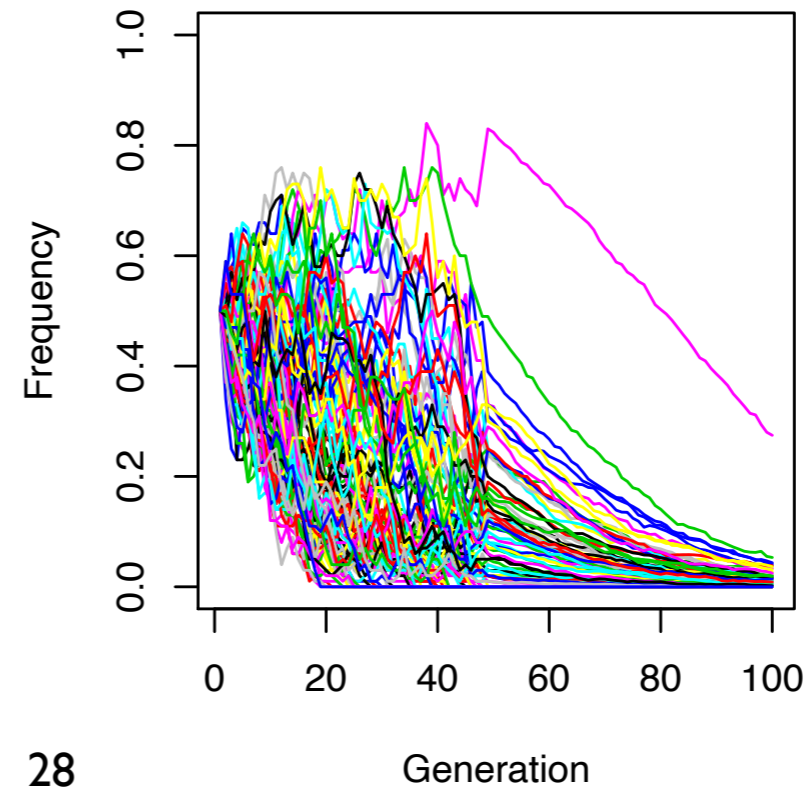
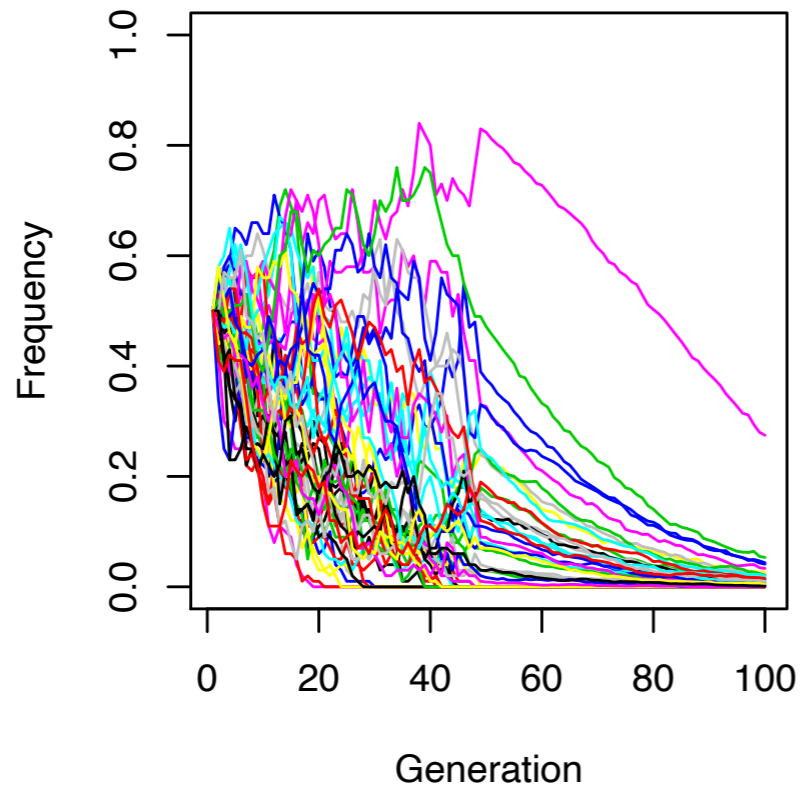
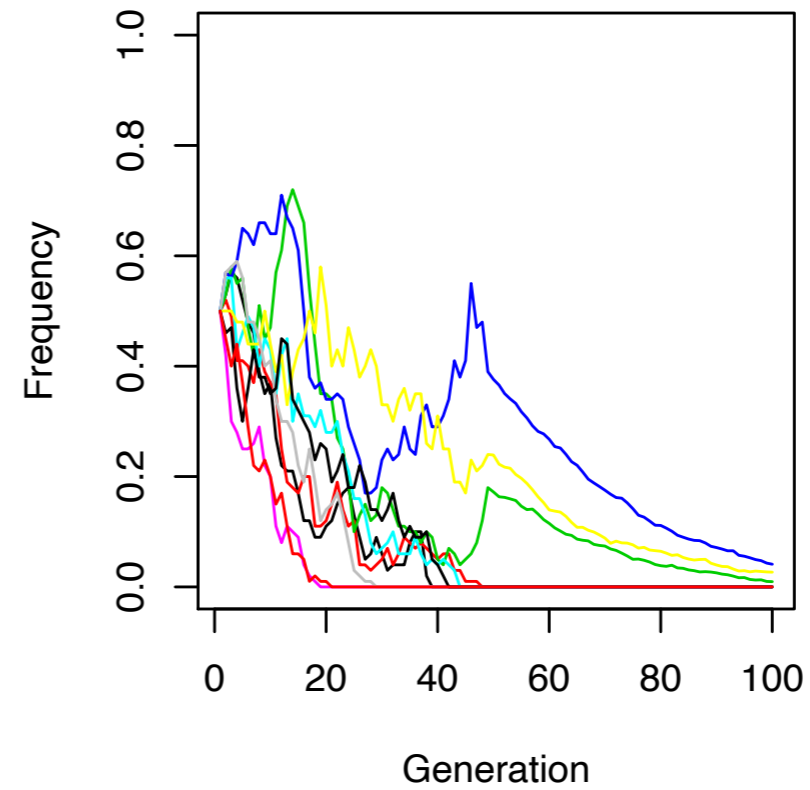
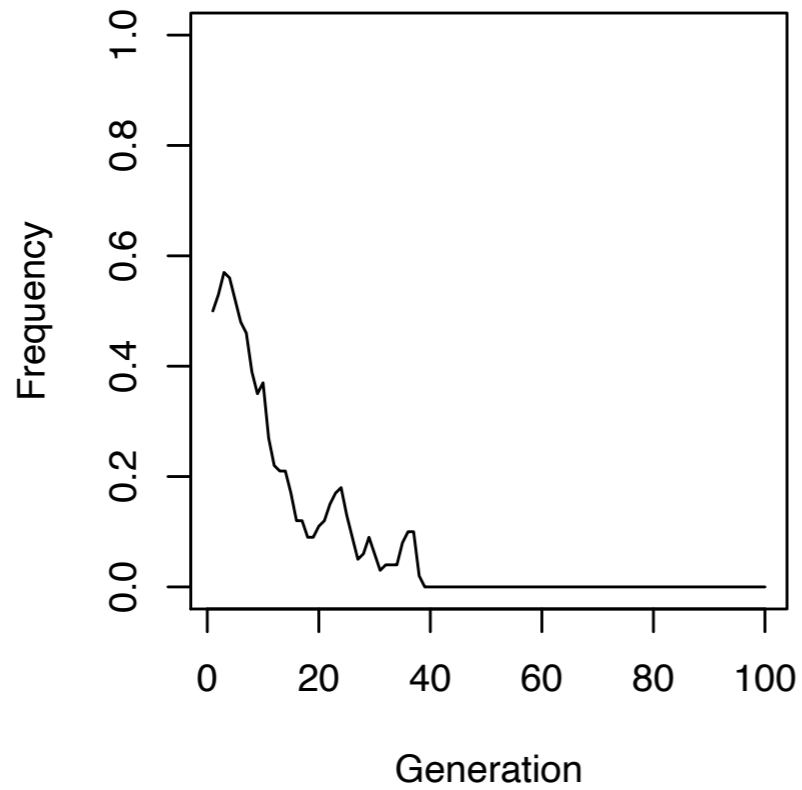
- Please work together to code this up.
- Can you add 100-fold population growth at generation 50 to your previous simulation?
- What happens in your plot?
- What if the initial frequency is 50%?

Wright-Fisher Model with Contraction

- Run it using: `WF.demsel(100,0.5,0.5,0.1,100,50,100)`

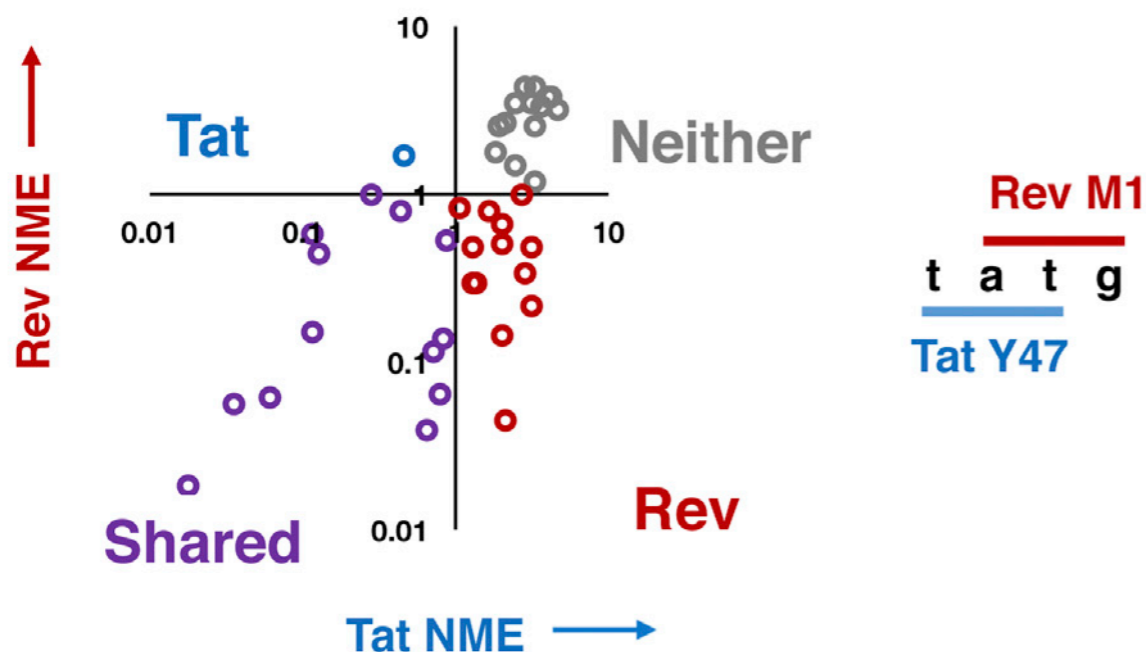
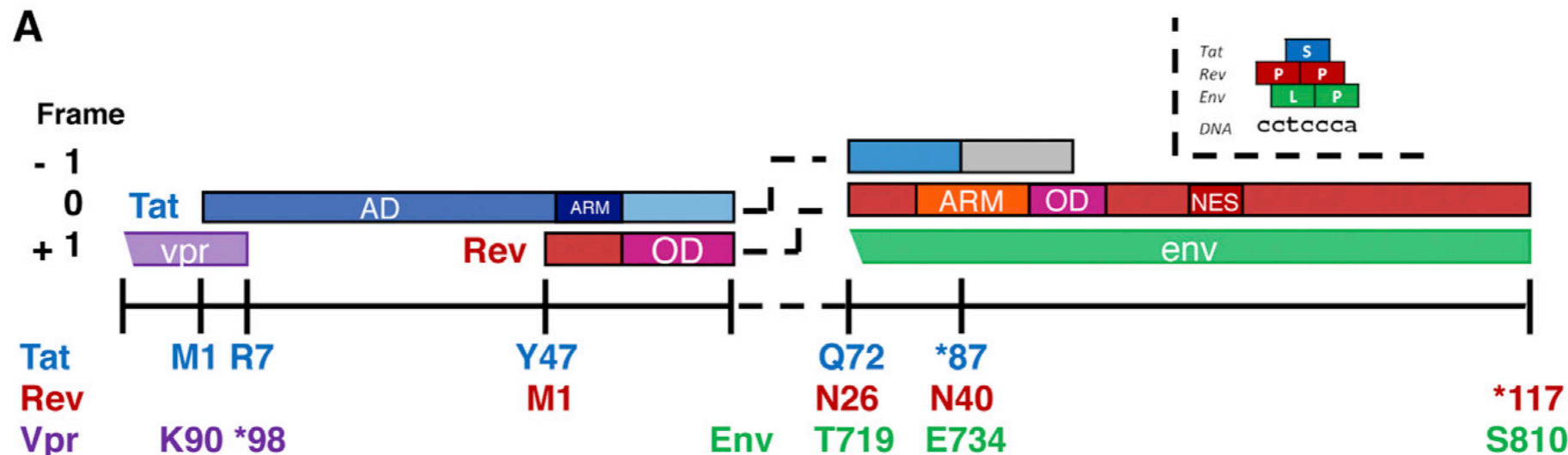


What parameters generated these?



Functional Segregation of Overlapping Genes in HIV

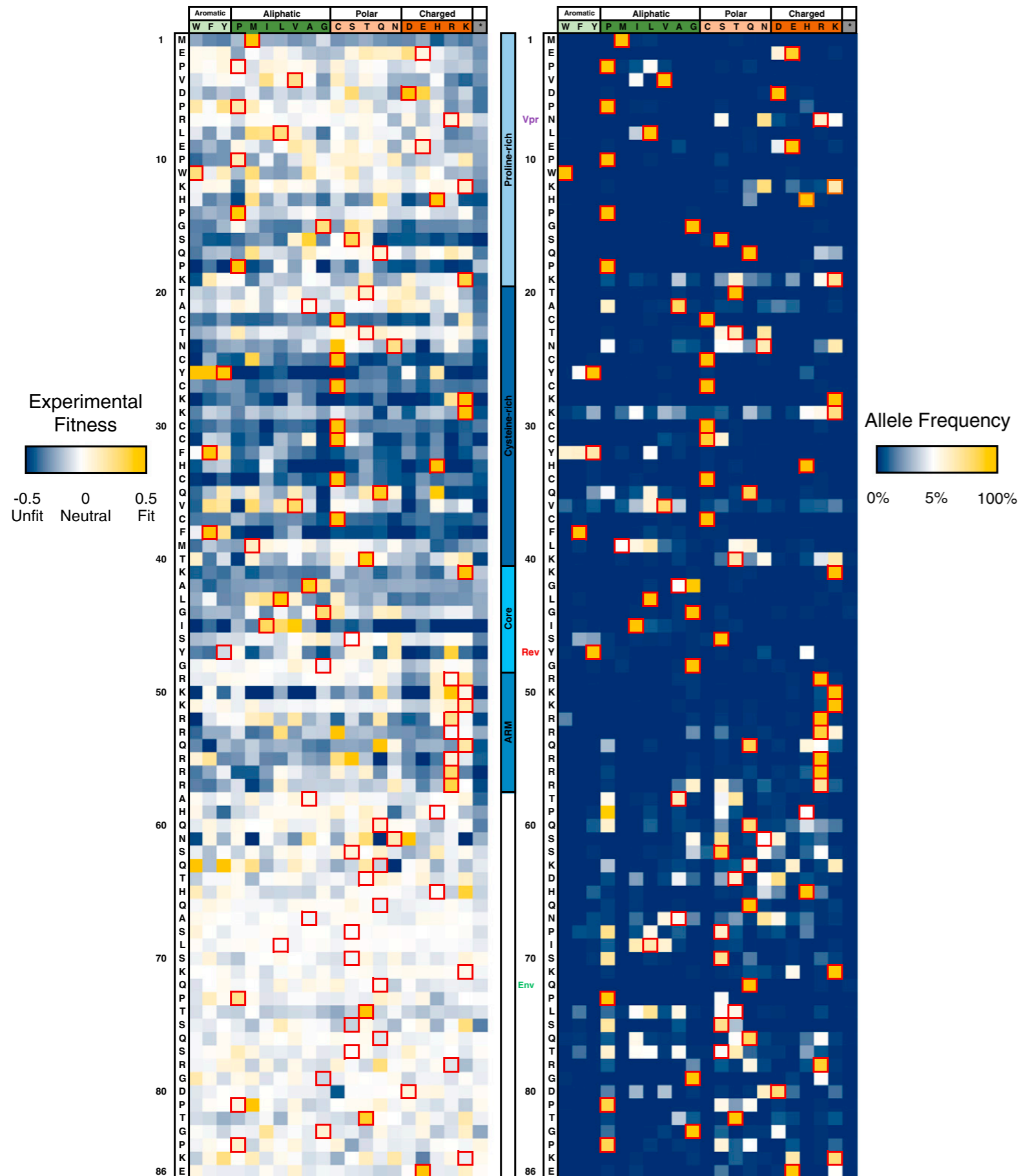
Jason D. Fernandes,^{1,2} Tyler B. Faust,^{1,3} Nicolas B. Strauli,^{4,5} Cynthia Smith,¹ David C. Crosby,¹ Robert L. Nakamura,¹ Ryan D. Hernandez,⁴ and Alan D. Frankel^{1,6,*}



- HIV genes Tat and Rev overlap.
- At protein level, many overlapping sites are conserved in both, but some sites only conserved in Rev.
- Is joint conservation due to dual function or genetic code?

Non-Overlap Deep Mutational Scanning

Overlap Patient Conservation



Functional Segregation of Overlapping Genes in HIV

Jason D. Fernandes,^{1,2} Tyler B. Faust,^{1,3} Nicolas B. Strauli,^{4,5} Cynthia Smith,¹ David C. Crosby,¹ Robert L. Nakamura,¹ Ryan D. Hernandez,⁴ and Alan D. Frankel^{1,6,*}

- In patient data, Tat sites that overlap with Rev are highly conserved.
- HIV can be engineered so that Tat and Rev do not overlap
- Deep mutational scanning in non-overlap context (all possible codons at each position) shows that many sites lack conservation in cell lines.
- Is this due to drift (neutral) or selection?

Functional Segregation of Overlapping Genes in HIV

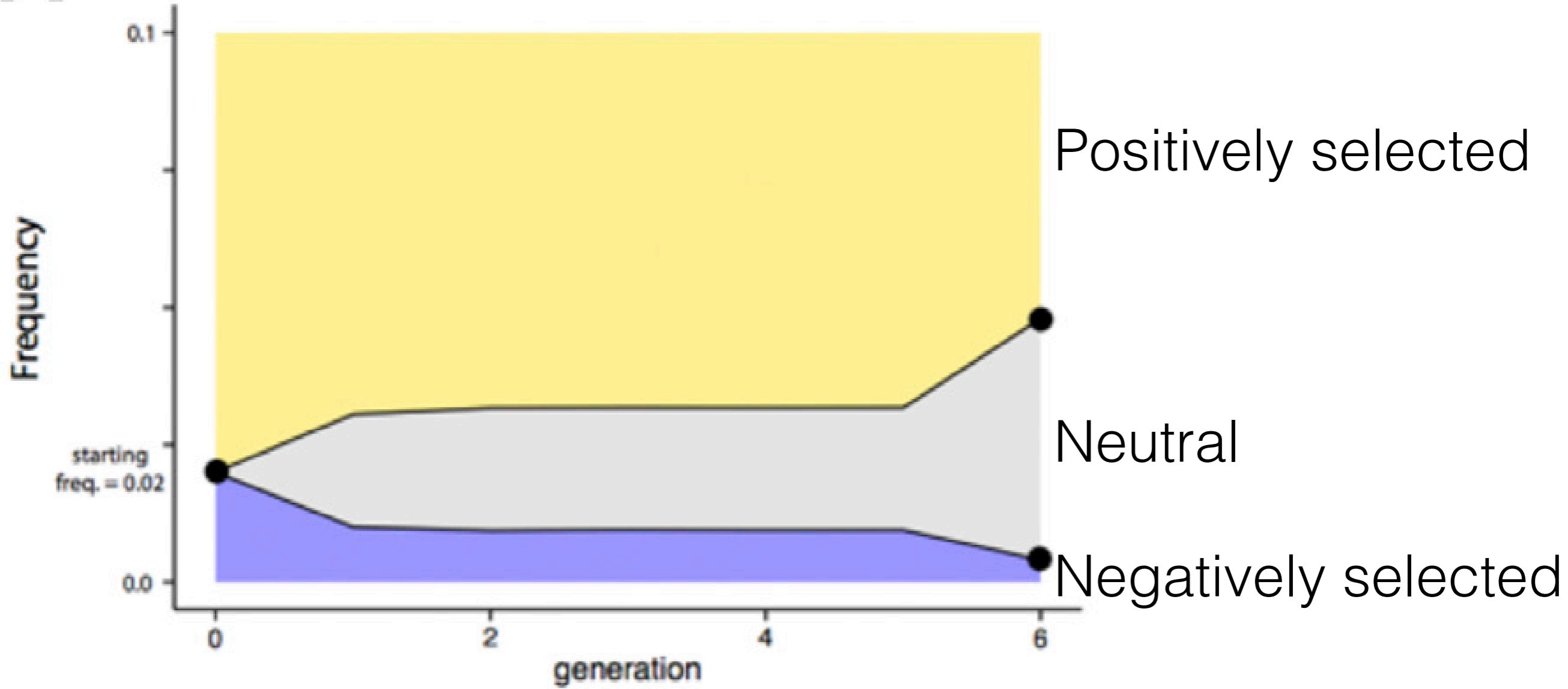
● **Deep mutational scanning:**

- Create exhaustive libraries with all possible codons at all overlapping positions
- Allow population mixture to evolve for **G** generations, then sequence to measure final frequencies of all amino acids
- Simulate to evaluate significance of allele frequency change

● **Factors you might want to include in your simulation:**

- the overall population growth function
- the number of generations
- the starting allele frequency
- the ending read depth for the experiment

A



Natural Selection

Time-course data from artificial selection/ancient DNA

- Let's estimate some selection coefficients!
- Given 2 alleles at a locus with frequencies p_0 and q_0 , and fitnesses w_1 and w_2 (with w the population-wide fitness).
- Expected freq. in next generation is: $p_1 = p' = p_0 w_1 / w$.

- We can then write:

$$\frac{p_1}{q_1} = \frac{p_0 w_1 / w}{q_0 w_2 / w} = \left(\frac{p_0}{q_0} \right) \left(\frac{w_1}{w_2} \right)$$

- Using induction, you could prove for any generation t :

$$\frac{p_t}{q_t} = \frac{p_0 w_1 / w}{q_0 w_2 / w} = \left(\frac{p_0}{q_0} \right) \left(\frac{w_1}{w_2} \right)^t$$

Natural Selection

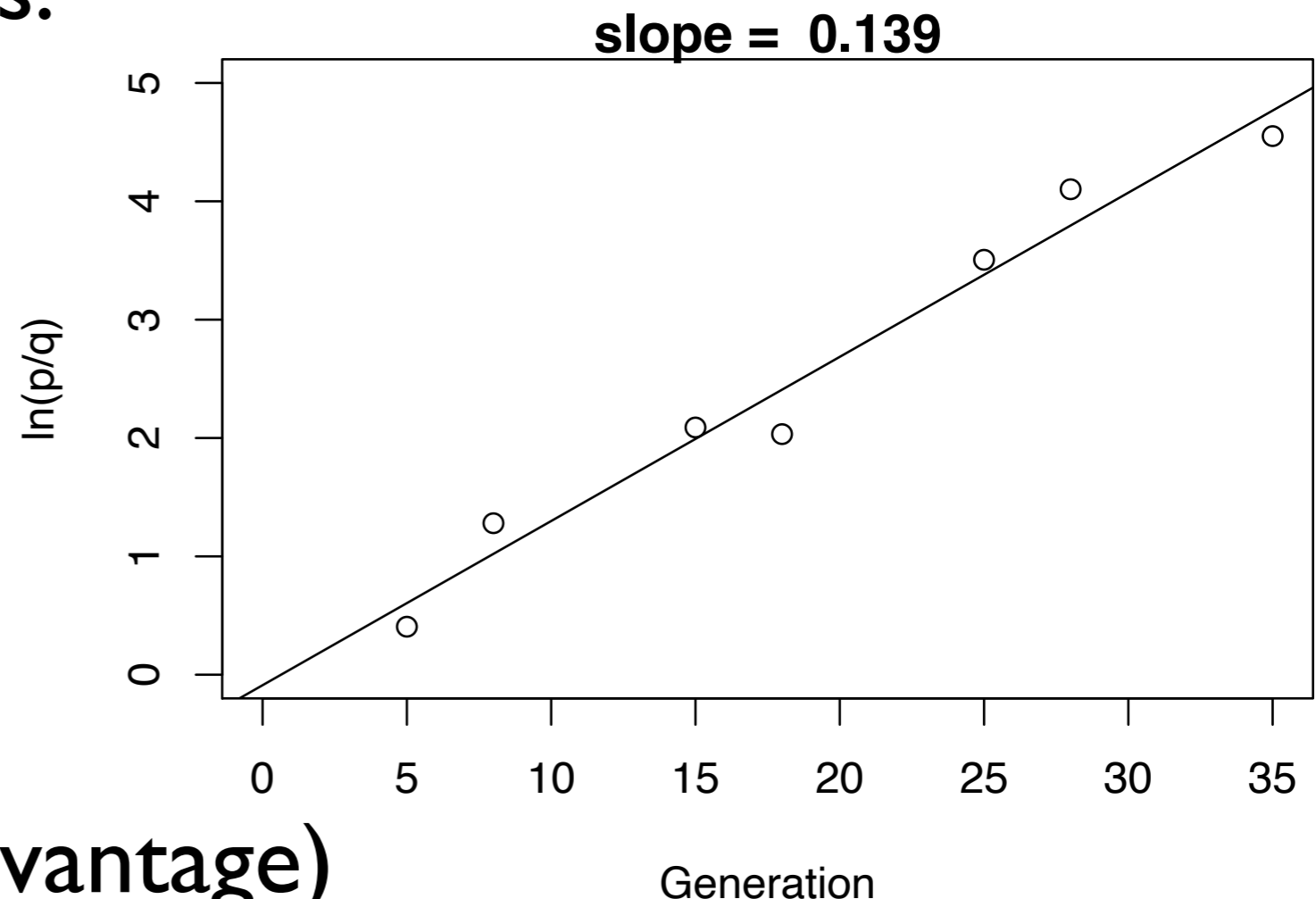
- Taking the natural log of this equation:

$$\log \left(\frac{p_t}{q_t} \right) = \log \left(\frac{w_1}{w_2} \right) t + \log \left(\frac{p_0}{q_0} \right)$$

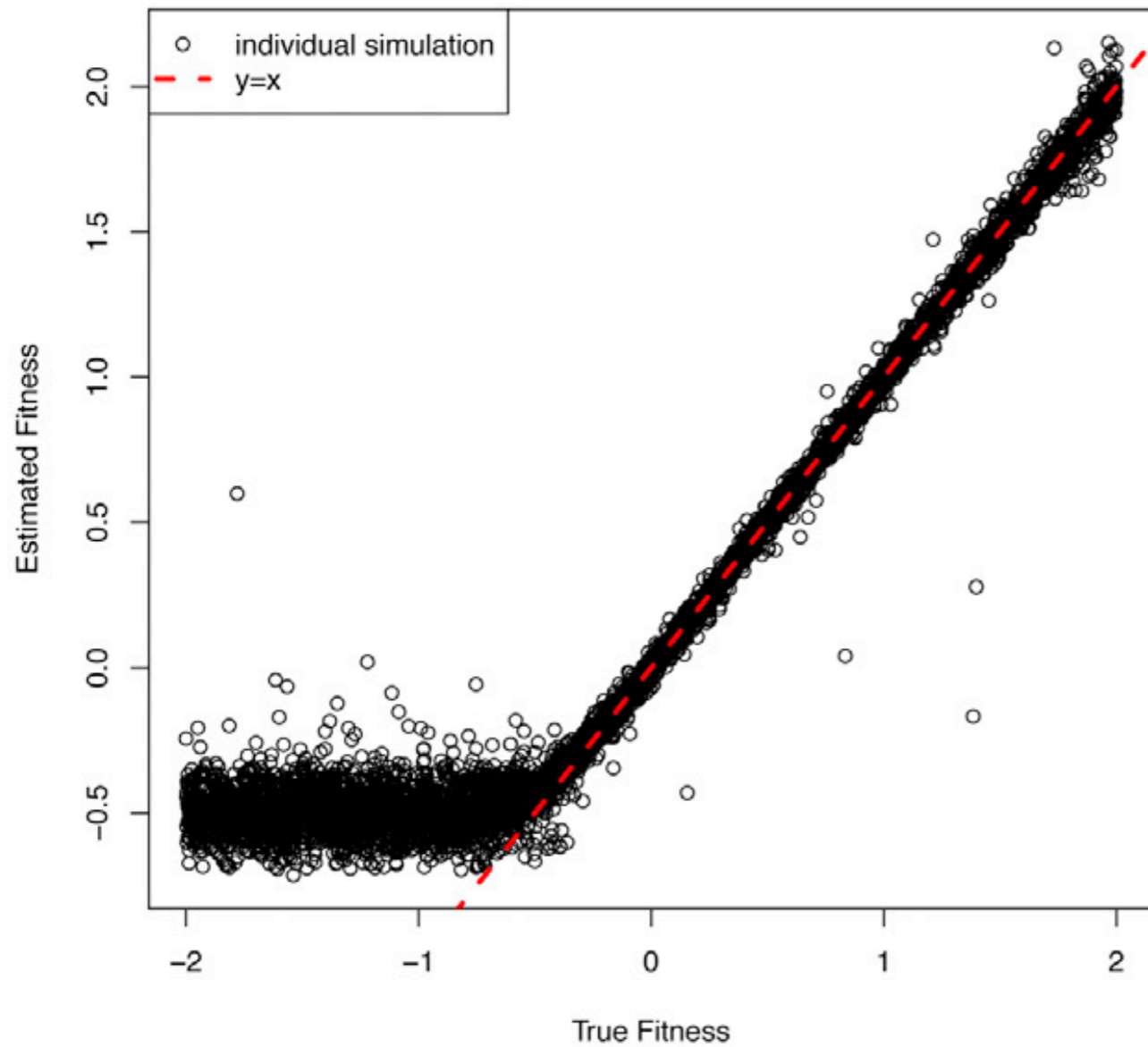
- Which is now a linear function of t , the number of generations.
- Therefore, the ratio of the fitnesses $w_1/w_2 = e^{\text{slope}}$

Natural Selection

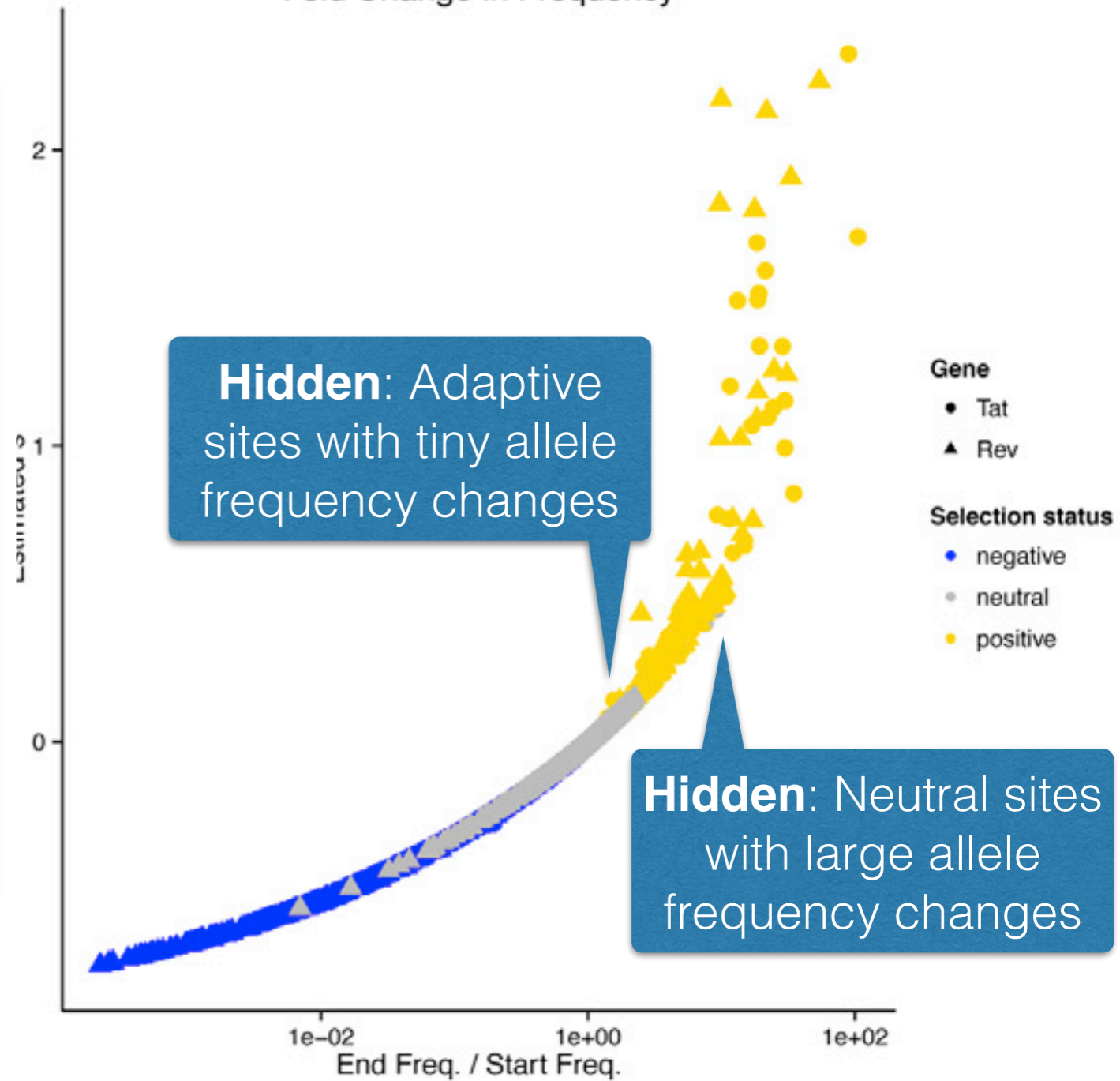
- Experiment: Set up a population of bacteria in a chemostat, and let them reproduce.
- Sample roughly every 5 generations.
- A slope of 0.139 implies:
 $w_1 = e^{0.139} = 1.15$
- Assume $w_2 = 1$.
- Thus, allele p has a 15% fitness advantage over allele q!
- (simulated with 20% advantage)



Accuracy of Fitness Point Estimate



Estimated Selection Coefficient Vs. Fold Change in Frequency



Existing forward simulators

- **SFS_CODE: Hernandez (2008)**
 - Command-line flexibility... shameless plug!
- **FWDPP: Thornton (2014)**
 - C++ library of routines intended to facilitate the development of forward-time simulations under arbitrary mutation and fitness models
- **SLiM 3: Haller & Messer (2019)**
 - Command-line, GUI, and R-like scripting environment that provides control over most aspects of the simulated evolutionary scenarios