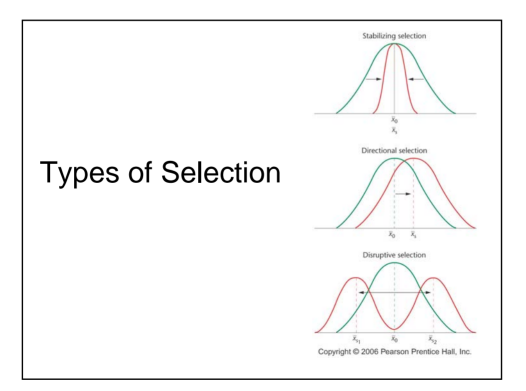
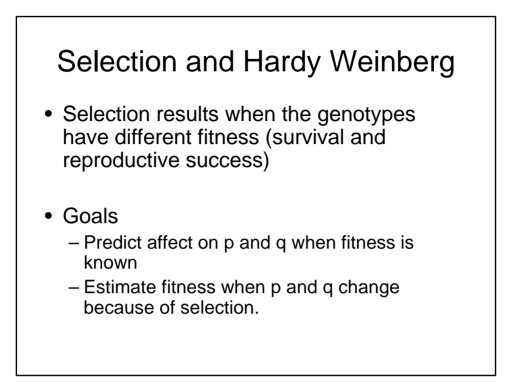


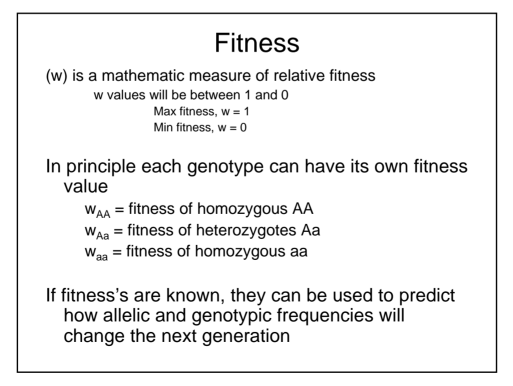
Effect Exception Assumption 1. Individuals must randomly mate Inbreeding Decrease or /Outbreeding increase in heterozygosity 2. No migration Migration Homogenize different Populations 3. No mutation Source of all Increase Variation Heterozygosity Disrupts return to equilibrium Genes are autosomal Sex Linkage 4 5. No selection **Directional Selection** Reduce variation **Disruptive Selection Increase Variation Stabilizing Selection** Increase Variation 6. Population must be very large Genetic Drift **Reduce Variation** (tend to loose rare alleles)

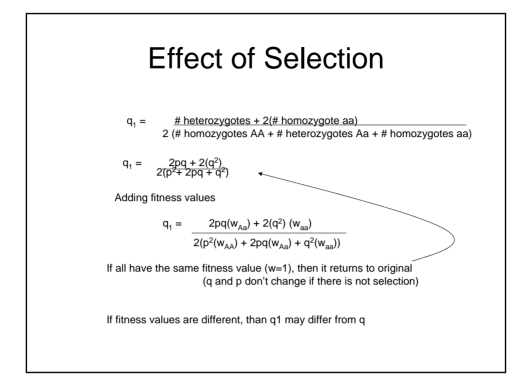


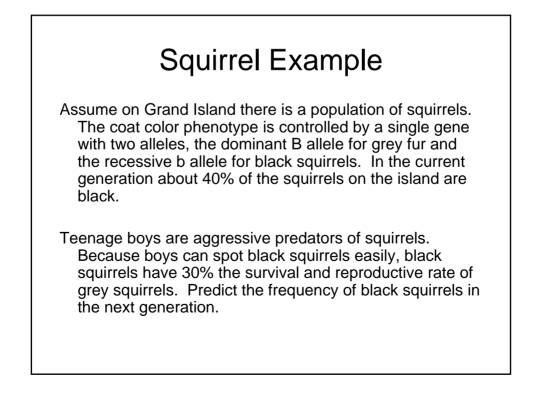
Value of Hardy-Weinberg Model

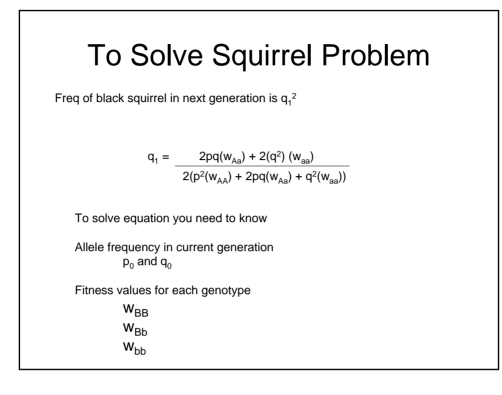
3. Evaluate factors that cause populations to evolve

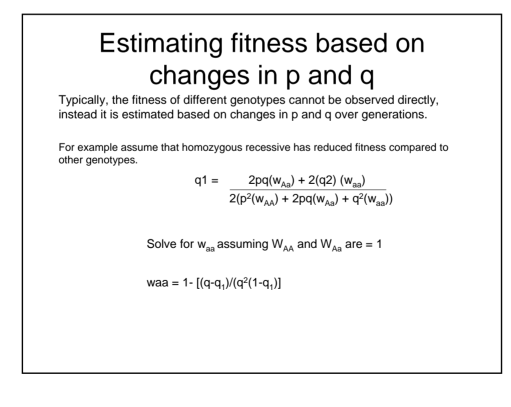












Example

The phenotypic frequency of black squirrels in 25% in one generation, but after heavy predation by boys it decreases to 21% in the next generation. What is the relative fitness of black squirrels?

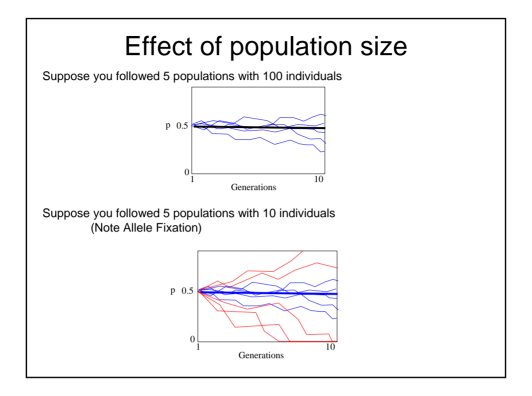
Need q and q₁

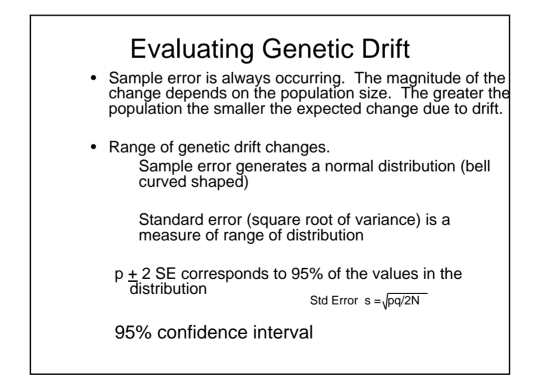
Genetic Drift

• Definition - Changes in allele frequency due to sampling error of gametes during fertilization.

Consider this example.

- Suppose you had a population of 100 in which p and q were both equal to 0.5. That population would generate 1000's of gametes, 50% of the gametes would have an A allele and 50% would have an a allele.
- To generate the next generation of 100, you would need 200 gametes to fuse during fertilization. If there was no sampling error than you would expect 100 of these gametes would have an A allele and 100 of these gametes would have an a allele. Without sampling error, p and q would remain at 0.5.
- However, because of sampling error, the chances that the 200 gametes would be exactly 50% A and 50% a is not likely. For example it would be unlikely that if gametes are randomly sampled 105 might have the A allele and 95 might have the a allele. With this level of sampling error p = 52.5 and q = 47.5.





Example

 Suppose that a population of 64 individuals had a p value of 0.6 in one generation but a p value of 0.5 the next generation. Would it be reasonable to believe that genetic drift could account for this change in p?