Lab 8 – Population Genetics

INTRODUCTION

In 1908, G. H. Hardy and W. Weinberg independently suggested a scheme whereby evolution could be viewed as changes in the frequency of alleles in a population of organisms. In this scheme, if A and a are alleles for a particular gene locus and each diploid individual has two such loci, then p can be designated as the frequency of the A allele and q as the frequency of the a allele. Thus, in a population of 100 individuals (each with two loci) in which 40% of the alleles are A, p would be 0.40. The rest of the alleles (60%) would be a, and q would equal 0.60 (i.e., p + q = 1.0). These are referred to as allele frequencies. The frequency of the possible diploid combinations of these alleles (AA, Aa, aa) is expressed as 

\[ p^2 + 2pq + q^2 = 1.0. \]

Hardy and Weinberg also argued that if five conditions are met, the population’s allele and genotype frequencies will remain constant from generation to generation. These conditions are as follows:

1. The breeding population is large. (The effect of chance on changes in allele frequencies is thereby greatly reduced.)
2. Mating is random. (Individuals show no mating preference for a particular phenotype.)
3. There is no mutation of the alleles. (No alteration in the DNA sequence of alleles.)
4. No differential migration occurs. (No immigration or emigration.)
5. There is no selection. (All genotypes have an equal chance of surviving and reproducing.)

The Hardy-Weinberg equation describes an existing situation. If the five conditions are met, then no change will occur in either allele or genotype frequencies in the population. Of what value is such a rule? It provides a yardstick by which changes in allele frequency, and therefore evolution, can be measured. One can look at a population and ask: is evolution occurring with respect to a particular gene locus? Since evolution is difficult (if not impossible) to observe in most natural populations, we will model the evolutionary process using the class as a simulated population. The purpose of this simulation is to provide an opportunity to test some of the basic tenets of population genetics and evolutionary biology.

EXERCISE 8A: Estimating Allele Frequencies for a Specific Trait within a Sample Population

Using the class as a sample population, the allele frequency of a gene controlling the ability to taste the chemical PTC (phenylthiocarbamide) could be estimated. A bitter-taste reaction to PTC is evidence of the presence of a dominant allele in either the homozygous condition (AA) or the heterozygous condition (Aa). The inability to taste the chemical at all depends on the presence of homozygous recessive alleles (aa).

To estimate the frequency of the PTC-tasting allele in the population, one must find p. To find p, one must first determine q (the frequency of the non-tasting PTC allele), because only the genotype of the homozygous recessive individuals is known for sure (those with the dominant trait could be AA or Aa).

Procedure

1. Using the PTC taste-test papers provided, tear off a short strip and press it to your tongue tip. PTC tasters will sense a bitter taste. For the purposes of this exercise these individuals are considered to be tasters.
2. A decimal number representing the frequency of tasters \(p^2 + 2pq\) should be calculated by dividing the number of tasters in the class by the total number of students in the class. A decimal number representing the frequency of non-tasters \(q^2\) can be obtained by dividing the number of non-tasters by the total number of students. You should then record these numbers in Table 8.1.
3. Use the Hardy-Weinberg equation to determine the frequencies \(p\) and \(q\) of the two alleles. The frequency \(q\) can be calculated by taking the square root of \(q^2\). Once \(q\) has been determined, \(p\) can be determined because \(1 - q = p\). Record these values in Table 8.1 for the class and also calculate and record values of \(p\) and \(q\) for the North American population.
Table 8.1: Phenotypic Proportions of Tasters and Non-tasters and Frequencies of the Determining Alleles

<table>
<thead>
<tr>
<th>Phenotypes</th>
<th>Allele frequency based on Hardy Weinberg equilibrium</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Tasters ( (p^2 + 2pq) )</td>
</tr>
<tr>
<td>Class Population</td>
<td>#</td>
</tr>
<tr>
<td>North American Population</td>
<td>0.45</td>
</tr>
</tbody>
</table>

**Topics for Discussion**

1. What is the percentage of heterozygous tasters \((2pq)\) in your class? 

2. What percentage of the North American population is heterozygous for the taster trait?

**EXERCISE 8B: Case Studies**

**CASE I – A Test of an Ideal Hardy-Weinberg Population**

The entire class will represent a breeding population, so find a large open space for this simulation. In order to ensure random mating, choose another student at random. In this simulation, we will assume that gender and genotype are irrelevant to mate selection.

The class will simulate a population of randomly mating heterozygous individuals with an initial gene frequency of 0.5 for the dominant allele \( A \) and the recessive allele \( a \) and genotype frequencies of 0.25 \( AA \), 0.50 \( Aa \), and 0.25 \( aa \). Your initial genotype is \( Aa \). Record this on the Data Page. Each member of the class will receive four cards. Two cards will have \( A \) written on them and two cards will have \( a \). The four cards represent the products of meiosis. Each "parent" contributes a haploid set of chromosomes to the next generation.

**Procedure**

1. Turn the four cards over so that the letters do not show, shuffle them, and take the card on top to contribute to the production of the first offspring. Your partner should do the same. Put the two cards together. The two cards represent the alleles of the first offspring. One of you should record the genotype of this offspring in the Case I section on the Data Page. Each student pair must produce two offspring, so all four cards must be reshuffled and the process repeated to produce a second offspring.

2. The other partner should then record the genotype of the second offspring on the Data Page. The very short reproductive career of this generation is over. You and your partner now become the next generation by assuming the genotypes of the two offspring. That is, Student 1 assumes the genotype of the first offspring and Student 2 assumes the genotype of the second offspring.

3. Each student should obtain, if necessary, new cards representing the alleles in his or her respective gametes after the process of meiosis. For example, Student 1 becomes genotype \( Aa \) and obtains cards \( A, A, a, a \); Student 2 becomes \( aa \) and obtains cards \( a, a, a, a \). Each participant should randomly seek out another person with whom to mate in order to produce the offspring of the next generation. Remember, the sex of your mate does not matter, nor does the genotype. You should follow the same mating procedures as you did for the first generation, being sure to record your new genotype after each generation. Class data should be collected after each generation for five generations. At the end of each generation, remember to record the genotype that you have assumed. Your teacher will collect class data after each generation by asking you to raise your hand to report your genotype.
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4. **Allele Frequency:** The allele frequencies, $p$ and $q$, should be calculated for the population after five generations of simulated random mating.

**Number of $A$ alleles present at the fifth generation**

Number of offspring with genotype $AA$ _____ x 2 = ______ $A$ alleles

Number of offspring with genotype $Aa$ _____ x 1 = ______ $A$ alleles

Total = ______ $A$ alleles

\[ p = \frac{\text{TOTAL number of } A \text{ alleles}}{\text{TOTAL number of alleles in the population} \times (\text{number of students} \times 2)} \]

In this case, the total number of alleles in the population is equal to the number of students in the class x 2.

**Number of $a$ alleles present at the fifth generation**

Number of offspring with genotype $aa$ _____ x 2 = ______ $a$ alleles

Number of offspring with genotype $Aa$ _____ x 1 = ______ $a$ alleles

Total = ______ $a$ alleles

\[ q = \frac{\text{TOTAL number of } a \text{ alleles}}{\text{TOTAL number of alleles in the population} \times (\text{number of students} \times 2)} \]

**Questions**

1. What does the Hardy-Weinberg equation predict for the new $p$ and $q$?

2. Do the results you obtained in this simulation agree?

3. Based on your answer to #2, what major assumption(s) were not strictly followed in this simulation?

**CASE II – Selection**

In this Case you will modify the simulation to make it more realistic. In the natural environment, not all genotypes have the same rate of survival; that is, the environment might favor some genotypes while selecting against others. An example is the human condition of sickle-cell anemia. This is a disease caused by a mutation on one allele, and individuals who are homozygous recessive often do not survive to reach reproductive maturity. For this simulation you will assume that the homozygous recessive individuals never survive (100% selection against), and that heterozygous and homozygous dominant individuals survive 100% of the time.

**Procedure**

The procedure is similar to that for Case I.

1. Start again with your initial genotype and produce your "offspring" as you did for Case I. This time, however, there is one important difference. Every time your "offspring" is $aa$, it does not reproduce. Since we want to maintain a constant population size, the same two parents must try again until they
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produce two surviving offspring. You may need to get new "allele" cards from the pool, allowing each individual to complete the activity.

2. Proceed through five generations, selecting against the homozygous recessive offspring 100% of the time. Then add up the genotype frequencies that exist in the population and calculate the new \( p \) and \( q \) frequencies in the same way you did for Case I.

**Questions**

1. How do the new frequencies of \( p \) and \( q \) compare to the initial frequencies in Case I?

2. What major assumption(s) were not strictly followed in this simulation?

3. Predict what would happen to the frequencies of \( p \) and \( q \) if you simulated another five generations.

4. In a large population would it be possible to completely eliminate a deleterious recessive allele? Explain.

**CASE III – Heterozygote Advantage**

From Case II it is easy to see what happens to the lethal recessive allele in the population. However, data from many human populations show an unexpectedly high frequency of the sickle-cell allele in some populations. Thus, our simulation does not accurately reflect the real situation; this is because individuals who are heterozygous are slightly more resistant to a deadly form of malaria than homozygous dominant individuals. In other words, there is a slight selection against homozygous dominant individuals as compared to heterozygotes. This fact is easily incorporated into our simulation.

**Procedure**

1. In this round keep everything the same as it was in Case II, except that if your offspring is \( AA \), flip a coin. If the coin lands heads up, the individual does not survive; if tails, the individual does survive.

2. Simulate five generations, starting again with the initial genotype from Case I. The genotype \( aa \) never survives, and homozygous dominant individuals only survive if the coin toss comes up tails. Since we want to maintain a constant population size, the same two parents must try again until they produce two surviving offspring. Get new "allele" cards from the pool as needed. Total the class genotypes and calculate the \( p \) and \( q \) frequencies.

3. Starting with the \( F_5 \) genotype, go through five more generations, and again total the genotypes and calculate the frequencies of \( p \) and \( q \) as done in Case I.

4. If time permits, the results from another five generations would be extremely informative.

**Questions**

1. How do the changes in \( p \) and \( q \) frequencies in Case II compare with Case I and Case III?

2. Do you think the recessive allele will be completely eliminated in either Case II or Case III?

3. What is the importance of heterozygotes (the heterozygote advantage) in maintaining genetic variation in populations?
Case IV – Genetic Drift

It is possible to use our simulation to look at the phenomenon of genetic drift in detail.

Procedure
1. Divide the lab into several smaller "populations" (for example, a class of 30 could be divided into three populations of ten each) so that individuals from one isolated "population" do not interact with individuals from another population.
2. Now go through five generations as you did for Case I. Record the new genotypic frequencies and calculate the new frequencies of $p$ and $q$ for each population.

Questions
1. Explain how the initial genotypic frequencies of the populations compare.
2. What do your results indicate about the importance of population size as an evolutionary force?

Hardy-Weinberg Problems
1. In *Drosophila*, the allele for normal-length wings is dominant over the allele for vestigial wings (vestigial wings are stubby little curls that cannot be used for flight). In a population of 1,200 individuals, 360 show the recessive phenotype. How many individuals would you expect to be homozygous dominant? How many would be heterozygous for this trait?
2. The allele for unattached earlobes is dominant over the allele for attached earlobes. In a population of 700 individuals, 15% show the recessive phenotype. How many individuals would you expect to be homozygous dominant? How many would be heterozygous for this trait?
3. The allele for the hair pattern called "widow's peak" is dominant over the allele for no "widow's peak." In a population of 3,000 individuals, 720 show the dominant phenotype. How many individuals would you expect for each of the possible three genotypes for this trait?
4. In the United States about 15% of the population is Rh negative. The allele for Rh negative is recessive to the allele for Rh positive. If the student population of South is 1,900, how many students would you expect for each of the three possible genotypes?
5. In certain African countries 3% of the newborn babies have sickle-cell anemia, which is a recessive trait. Out of a random population of 1,000 newborn babies, how many would you expect for each of the three possible genotypes?
6. In a certain population, the dominant phenotype of a certain trait occurs 89% of the time. What is the frequency of the dominant allele?
CASE I
Hardy-Weinberg Equilibrium
Initial Class Frequencies:

\[ \begin{array}{ccc}
AA & Aa & aa \\
p & q & \\
\end{array} \]

My Initial Genotype: ___________

\[ \begin{array}{ccc}
F_1 & F_2 & F_3 \\
& & \\
\end{array} \]

Final Class Numbers:

\[ \begin{array}{ccc}
AA & Aa & aa \\
p & q & \\
\end{array} \]

CASE III
Heterozygote Advantage
Initial Class Frequencies:

\[ \begin{array}{ccc}
AA & Aa & aa \\
p & q & \\
\end{array} \]

My Initial Genotype: ___________

\[ \begin{array}{ccc}
F_1 & F_2 & F_3 \\
& & \\
\end{array} \]

Final Class Numbers:

\[ \begin{array}{ccc}
AA & Aa & aa \\
p & q & \\
\end{array} \]

CASE II
Selection
Initial Class Frequencies:

\[ \begin{array}{ccc}
AA & Aa & aa \\
p & q & \\
\end{array} \]

My Initial Genotype: ___________

\[ \begin{array}{ccc}
F_1 & F_2 & F_3 \\
& & \\
\end{array} \]

Final Class Numbers:

\[ \begin{array}{ccc}
AA & Aa & aa \\
p & q & \\
\end{array} \]

CASE IV
Genetic Drift
Initial Group Frequencies:

\[ \begin{array}{ccc}
AA & Aa & aa \\
p & q & \\
\end{array} \]

My Initial Genotype: ___________

\[ \begin{array}{ccc}
F_1 & F_2 & F_3 \\
& & \\
\end{array} \]

Final Group Numbers:

\[ \begin{array}{ccc}
AA & Aa & aa \\
p & q & \\
\end{array} \]