

Through the Years: A Quantitative Biology Curriculum

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INTRODUCTION

QUANTITATIVE REASONING is a skill that must be mastered by biology students before they graduate (NRC, 2003; AAAS, 2011; Hurney et al., 2011). Yet, experience suggests that biology students do not always see the need for this skill (“I just want to work with plants.”) or fear analytical problems (“I am not good at math.”). Student aversion to quantitative analysis poses pedagogical problems for teaching the process of science.

To address these problems, some biology faculty at Southeast have coordinated their courses to teach skills early that are built upon later. For example, the first biology course for majors uses Microsoft Excel extensively to teach fundamental graphing and math functions that students use for data analysis. Later courses require students to employ those early skills and to learn more advanced skills. Some courses introduce new software for simulation and analysis. Here, I highlight exercises from three courses that span the curriculum of a typical organismal biology student at Southeast.

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COLOPHON

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YEAR 1: EVOLUTION AND ECOLOGY

EVOLUTION AND ECOLOGY is a new lecture and laboratory course for freshmen biology majors. The lab exercises help students develop skills with Microsoft Excel and emphasize graphical analyses of data.

HARDY-WEINBERG EQUILIBRIUM

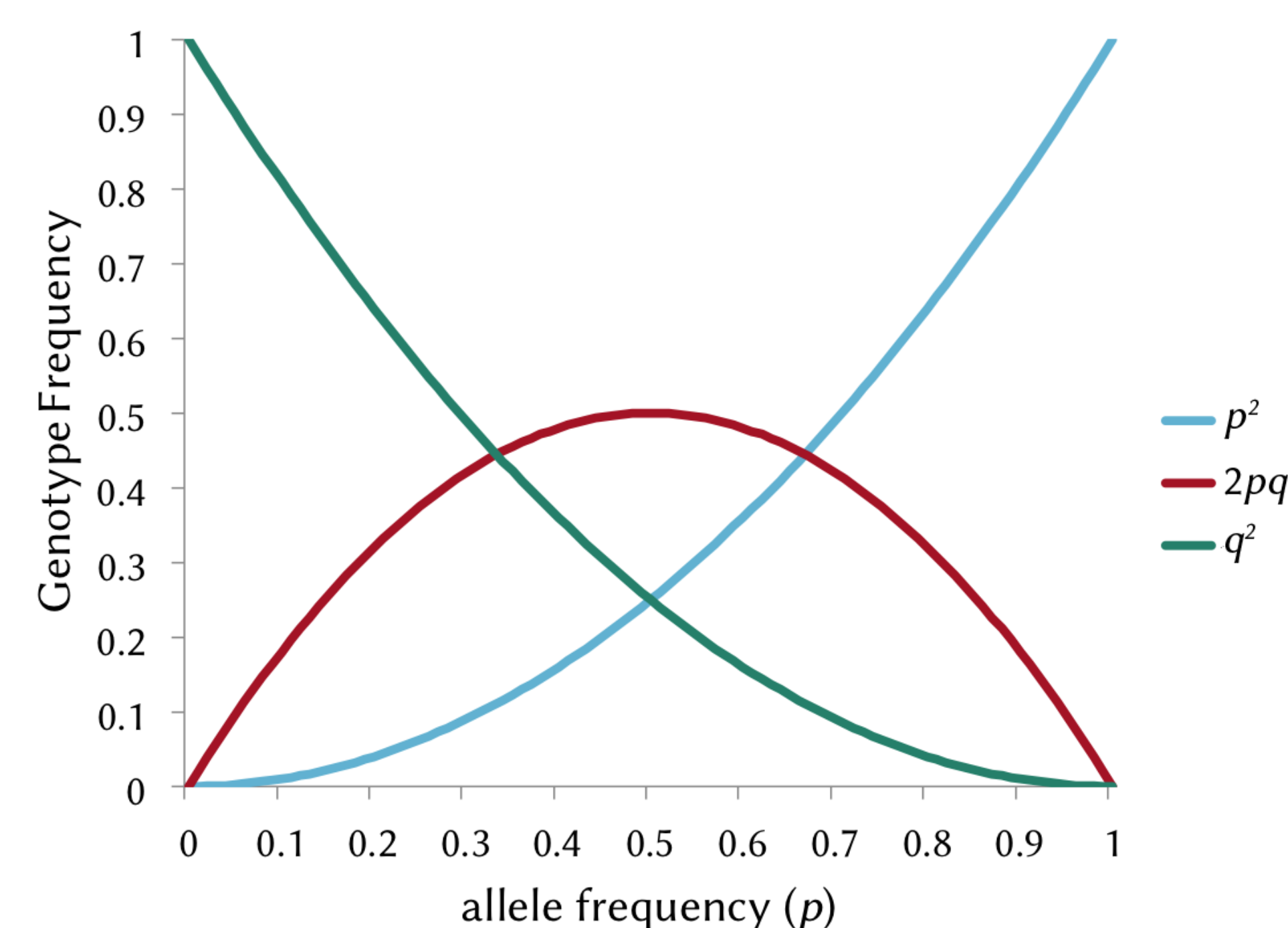
Students use two equations to model allele and genotype frequencies in a population of organisms. The equation for the allele frequencies is

$$p + q = 1,$$

where p and q are the frequencies of two alleles. The formula for the three possible genotypes is

$$p^2 + 2pq + q^2 = 1.$$

Students build formulas in Excel to generate values for the five variables from $p = 0.01$ to 1.0, and then plot the three genotype frequencies against the p allele frequency. The graph is a visual representation of *expected* genotype frequencies for a population in Hardy-Weinberg equilibrium.



Students next apply “beanbag” biology (Jungck et al., 2010) by sampling “alleles” from a “population” with known allele frequencies, and then apply a χ^2 test to compare observed to expected genotype frequencies. Students also explore how migration and natural selection affect allele frequencies in a population.

Representative questions from the migration component are

- Predict how the frequency of each allele will change for each population due to gene flow.
- Use the results of your migration simulation to explain why gene flow between populations should prevent them from becoming genetically different.

YEARS 2–3: EVOLUTIONARY BIOLOGY

INTRODUCTION TO EVOLUTIONARY BIOLOGY is taken by sophomores and juniors. The course is now lecture-only but is being redesigned to include a once-per-week lab that requires to students to reinforce lecture concepts by analysis of real data.

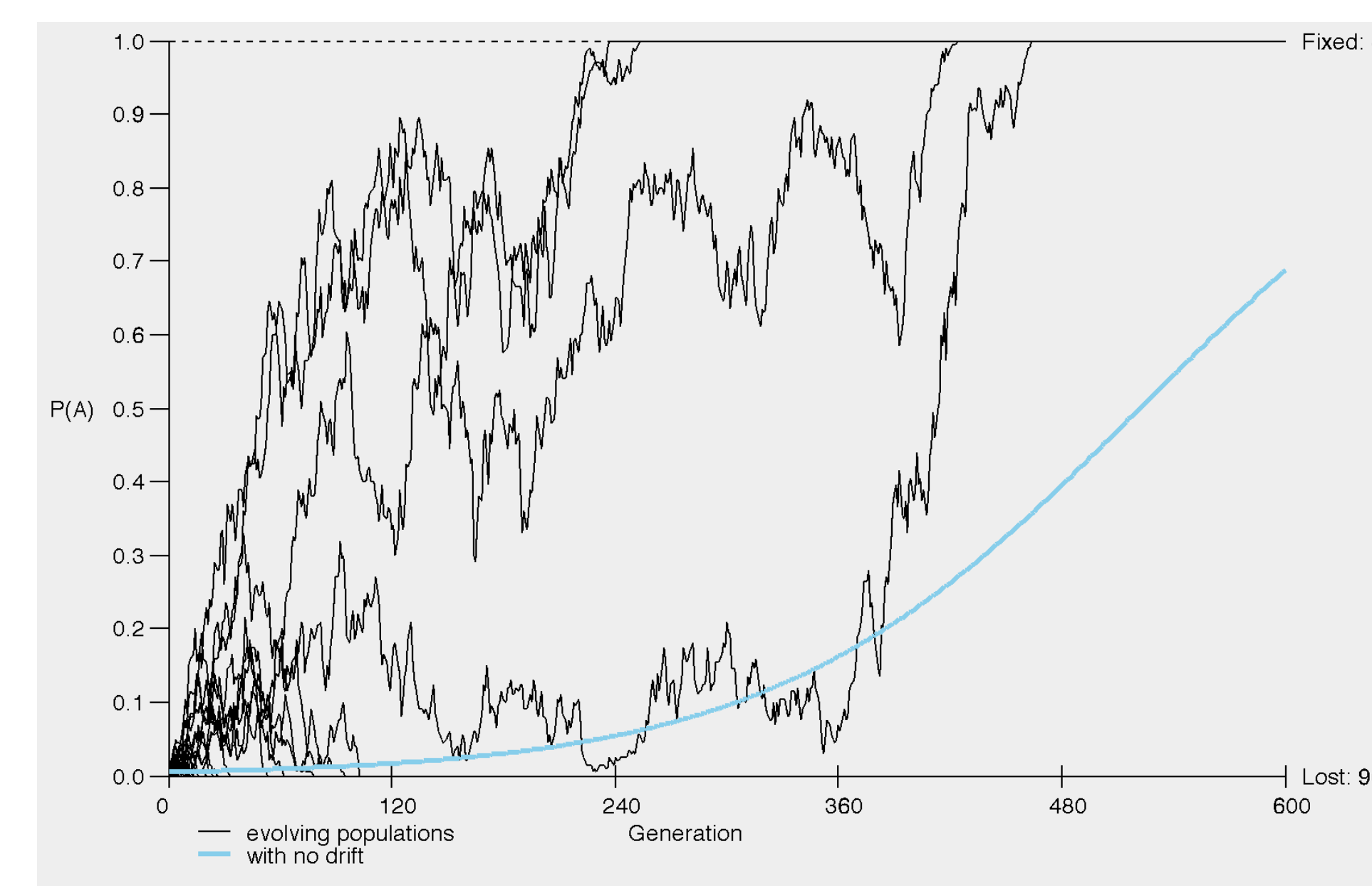
POPULATION GENETICS

Students use PopG software (Felsenstein, 2016) to simulate allele frequencies in a population subject to different evolutionary scenarios such as natural selection and genetic drift. With PopG, students change population size, strength of selection, and migration and mutation rates to learn how these affect allele frequencies over time.

In one exercise, students explore the interaction between the strength of selection (s) and the effective population size (N_e):

$$2N_e s \approx 1.$$

If $2N_e s \gg 1$ then selection has a better chance of overcoming drift to cause a beneficial mutation to become fixed in the population. In a series of simulations where $2N_e s = 0$ and $2N_e s = 2$, students see that a beneficial allele has about a $4\times$ greater chance of becoming fixed in the population. The figure below shows the results of one simulation run on 100 populations of $N = 100$ each and $s = 0.01$.



Representative questions from this exercise are

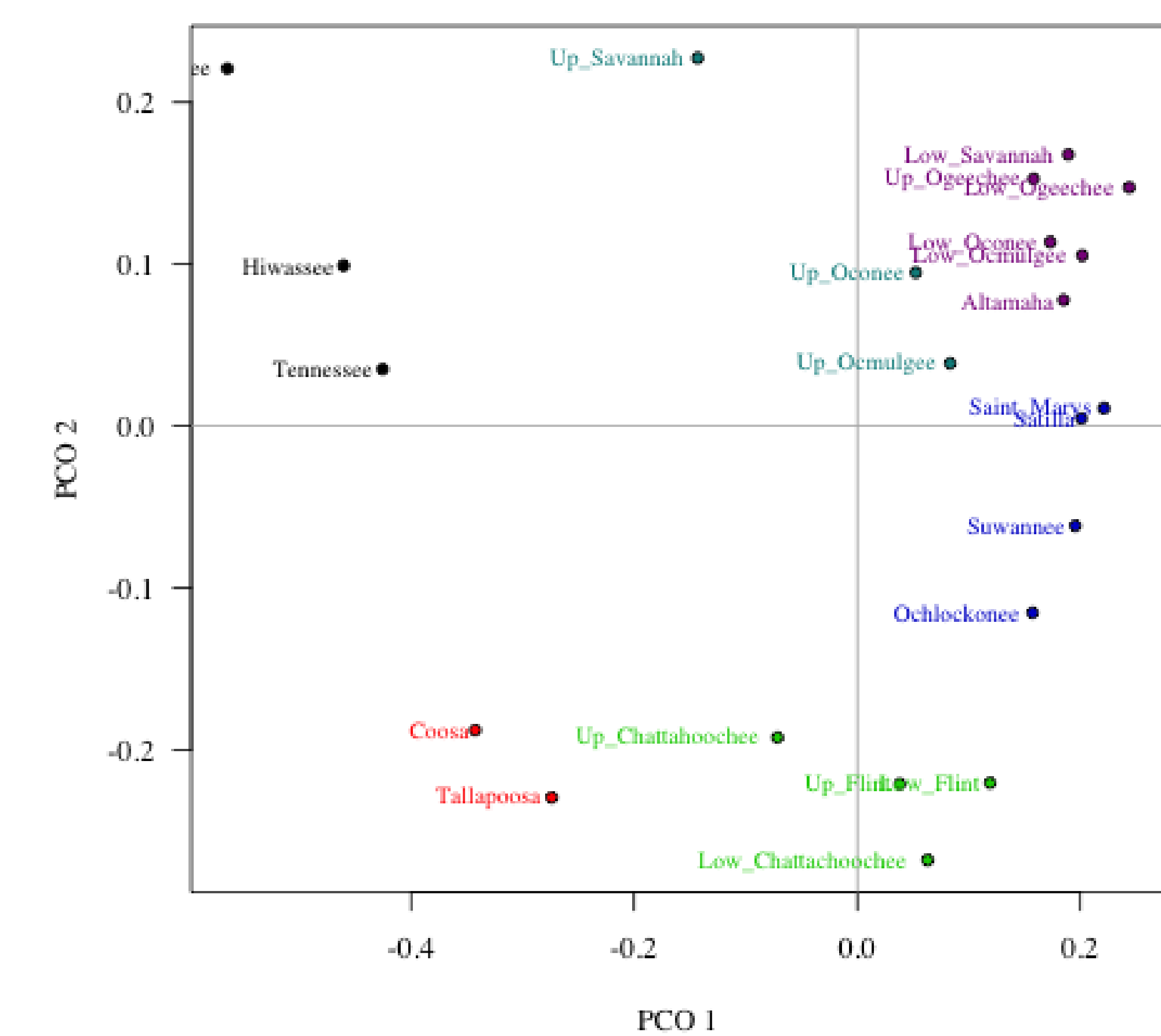
- Predict how N_e affects the probability of fixation for an allele under weak positive selection ($s = 0.01$). Justify your predictions for populations of $N = 1000$ and $N = 10,000$.
- What should be the strength of selection in a population of $N_e = 1000$ to overcome the effects of genetic drift? If $N_e = 10,000$ or $50,000$? Show your work.

YEAR 4: BIOGEOGRAPHY

BIOGEOGRAPHY is a senior and graduate level course. Students use the R Statistical Package (R Core Team, 2016) to analyze real-world data to discover and report biogeographic patterns.

BIOGEOGRAPHIC PROVINCES

Students compare distribution patterns of freshwater fishes and crayfishes by performing a principal coordinates ordination (PCO) for each taxonomic group. PCO groups watersheds based on shared species, as shown here for Georgia fishes.



Students then compare the similarities of the fish and crayfish ordinations with a Procrustes analysis. A significant result indicate that the watersheds have the same spatial relationship in each ordination, suggesting that fishes and crayfishes share a common biogeographic history.

