Computer Simulation #5: F-Statistics of Population Structure

We have used the **F** statistic to describe several kinds of population structures, based on a difference between the observed and expected numbers of heterozygous genotypes within or among populations. When examination of a population for expected Hardy-Weinberg proportions shows a deficiency of heterozygotes, that deficiency can be quantified as $\mathbf{F} = (\mathbf{H}_{obs} - \mathbf{H}_{exp}) / \mathbf{H}_{exp}$. We called this the **Wahlund Effect**, where the analysis of *two* (or more) populations as a *single* population always results in a deficiency of heterozygotes, and as such led to rejection of the null hypothesis of *no* population structure.

We also used **F** as the **inbreeding coefficient**, where inbreeding is defined as the expectation of drawing two alleles from a population that are <u>identical by descent</u>, that is, genetic copies of an allele existing in a common ancestor. This expectation applies to both (1) **gamete formation** by two alleles **I** by **D**, and (2) **random** '*draw*' of two alleles **I** by **D** from different individuals in a population because of common ancestry. We saw that inbreeding necessarily occurs in any *closed finite* population, because over time all individuals become more or less closely related. **F** can then be used to estimate the <u>effective size (Ne)</u> of different populations, and also to detect mating patterns where mating between more or less closely related individuals is variously avoided, encouraged, or occurs randomly.

F as a measure of breeding structure *within* populations can be extended to **measures of structure** *among* **sub-populations within more inclusive populations units**, which may be hypothesized geographic or ecological units. **F** becomes a *hierarchal* statistic, including as components the structure of **individuals within sub-populations** (F_{IS}), individuals with respect to the total structure (F_{IT}), and perhaps most importantly structure of sub-populations with respect to the total (F_{ST}). F_{ST} and its extensions is one of the most widely used tools in evolutionary population genetics.

The **Excel Workbook** for **F-Statistics** provide spreadsheets for calculation of **F-statistics** in the special case of *three* sub-populations, where sample size **N** in all sub-populations is *identical*. This avoids the complications of weighting the contributions of sub-populations of different size to the various calculations, which obscures the logic of what is really a simple hierarchal procedure.

This lab exercise will develop a sense for recognition of population structures as inferred F-statistics, for hypothetical and real population structures. The data are (1) examples of simple population structures, including hybrid zones between two isolated populations, and (2) Real data from MN blood types used to investigate population structure among Philippine islanders (Arcellana et al., 2011; Carr, 2021). The Philippine archipelago comprises more than 7,600 islands and more than 180 ethinic groups, many of which are more or less isolated by distance, ethnicity, and language. The data allows tests of (a) *intra*-island differentiation among populations on Luzon in the north, and on Mindanao in the south, (b) *inter*-island differentiation, including isolation by distance and two alternative samplings of Luzon, Mindanao, and the island of Palawan in the west.

References:

AES Arcellana, RMS Guzman, and IKC Fontanilla (2011). Distribution of MN blood group types in local populations in Philippines. *Journal of Genetics* **90**, e90–e93.

SM Carr (2021) Corrigendum to Arcellana et al. 2011, J Genet 90, e90-93. Journal of Genetics 100:

Instructions:

- 1. The data are **3x3** blocks of **counts** of **AA**, **Aa**, & **aa** for each of **three** sub-populations selected from the tests of hypotheses below. The worksheet will then calculate (1) **Local F** for each of the three sub-populations, and (2) **F**_{is}, **F**_{st}, & **F**_{it} for the combination of sub-populations entered.
 - a. Arrange data blocks on the spreadsheet; Copy & paste Values Only into the Grey Block
 - b. For all exercises below, **record** blocks (1) and (2) for the sets of sub-populations analyzed.
- 2. Examine the series of simple models of population structure, including
 - a. All sub-populations identical for allele frequencies in Hardy-Weinberg proportions (HWP)
 - b. All sub-populations identical for allele frequencies *not* in HWP
 - c. Sub-populations with different allele frequencies, in HWP
 - d. Sub-populations with different allele frequencies, not in HWP
 - e. Any other simple models that occur to you
- 3. The data below are **MN blood group counts** from populations in the **Philippine Islands** (Arcellana *et al.*, 2011; corrected by Carr, 2021). Counts have been normalized to 100@.

 Examine the population structure of the Philippine archipelago, with the following *a priori* hypotheses:
 - a. Structure occurs according to distance
 - b. Structure occurs within individual islands
 - Structure occurs among different island
 Calculate Chi-Square tests of whether the F-statistics show significant departure from HWE.

