presented by a population of size one. For each individual of a population, the allelic composition at all loci defined in the map must be given by two arbitrarily chosen characters that represent the two alleles, for example, aa, bb, CC, 11, or 12. Background loci carry at the start of the simulation the same allele as the tightest linked locus whose genotype is known.

**Step 3: Description of the Breeding Program**

Subsequently the breeding program has to be described by a list of commands for reproduction and selection. Commands for crossing, selfing, and outbreeding are available. Furthermore, varying modes of reproduction and various breeding designs are implemented, such as single-seed descent, double haploids, factorial, and top-cross mating designs.

Selection can be carried out for a certain genotype by specifying a list of loci and the respective allelic composition, or with respect to a selection index. A selection index is constructed by summing the number of loci of a certain class that have a specified allelic composition. For example, in a backcross program the following indices are possible. Selection of individuals (1) heterozygous for the donor allele at all target loci; (2) heterozygous for the donor allele at the maximum number of foreground selection markers; (3) homozygous for the recurrent parent allele at the background selection markers flanking a target gene; or (4) homozygous for the recurrent parent allele at a maximum number of background selection markers. An arbitrary number of selection steps can be carried out in sequence, this allows flexibility in the design and investigation of various selection strategies. In addition to reproduction and selection commands, population manipulations such as random sampling, taking subsets of populations, and merging of populations can be performed.

Because of the stochastic nature of a simulation study, it must be repeated to obtain reproducible results. PLABSIM implements the option to execute the input file in total or in part repeatedly for a number of repetitions specified by the user. The populations that are generated during the repetitions of a simulation can be stored in order to evaluate the results together after finishing all repetitions. Storage of the data can be done in an internal format for evaluation with PLABSIM or in an export format for analysis with statistical software.

**Step 4: Analysis of the Simulated Data**

The simulated data can be evaluated with PLABSIM for gene frequencies in three ways: (1) The frequency of an allele that occurs at a defined locus can be calculated. (2) The frequency of the alleles originating from one ancestor can be calculated for a class of loci. This can be used, for example, to estimate the proportion of the recurrent parent alleles at marker loci in backcrossing. (3) The frequency of the alleles originating from one ancestor can be calculated for the whole genome. A possible application is the estimation of the recurrent parent genome proportion in backcrossing.

Genotype frequencies can be calculated either by specifying for each locus on the map an allelic composition, or by specifying for an arbitrary subset of the map an allelic composition. The frequency of the combination of two alleles can be determined for a class of loci. This is useful for estimation of homozygosity or heterozygosity. Furthermore, the distribution of the length of chromosome segments that originate from one ancestor can be calculated.

In addition to estimation of genetic parameters, the number of marker data points required in the selection steps of the breeding program can be estimated.

**Availability**

Compiled versions for varying operating systems such as AIX, LINUX, UNIX, or WindowsNT are available. Noncommercial users can obtain the PC version of PLABSIM via e-mail for a nominal charge. Please contact the corresponding author.

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**WHICHRUN (version 3.2): A Computer Program for Population Assignment of Individuals Based on Multilocus Genotype Data**

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Microsatellite DNA provides essentially limitless, highly varied information within species. That this provides a means for distinguishing not only among populations but also individuals has not escaped current theoretic interest (Smouse and Chevillon 1998; Waser and Strobeck 1998).

Here we present a C++ computer program, WHICHRUN, that uses multilocus genotypic data to allocate individuals to their most likely source population. This program runs on Windows95, 98, or NT (including Macintosh emulations of these operating systems) and has no specific hardware requirements. WHICHRUN differs from a similar individual-based population assignment program “the assignment test” (Paetkau et al. 1995; Waser and Strobeck 1998) in that it provides a variety of methods for evaluating population assignments including maximum likelihood, jackknife, and critical population routines. WHICHRUN also provides resources for converting data into formats required for the population-based Statistical Package for Analysis of Mixtures (SPAM) available from L. Seeb, Alaska Department of Fish and Game.
WHICHRUN requires baseline genotype data for all potential source populations, as well as genotype data for candidate individuals for which population origin is to be determined. Data should be provided in simple ASCII format as required for GE-NEPOP (Raymond and Rousset 1995). The download available at the site described below includes sample input files.

**Theory and Program Outline**

It is assumed that each baseline population \((B_1, \ldots, B_n)\) has Hardy–Weinberg–Castle (HWC) genotype frequencies and that genetic loci employed are independent. The likelihood that an individual sample \((s_1, \ldots, s_n)\) may come from each of the source populations \((B_1, \ldots, B_n)\) is presumed to be equal to the HWC frequency of its specific genotype at each locus in each respective source population. Thus for homozygotes the likelihood that a sample \((s_1)\) is an element \((\Sigma)\) of baseline population \(B_i\) is \(p_i^2\), [the square of its allele frequency \((p)\) in population \(B_i\)]. For heterozygotes, \(s \neq B_i = 2p_i q_i\) \((q_i\) being the frequency of an alternate allele in population \(B_i\)), and the likelihood that \(s \neq B_i = p_i^2\) or \(2p_i q_i\). Likelihood values for each locus are multiplied to give a series of multilocus likelihood functions for assignment to each of the source populations. Alternate hypotheses that individual samples in question may come from each source population are considered in three ways:

1. Multilocus likelihood functions may be grouped to form ratios considering all possible pairs of baseline populations under consideration. If the ratio of the most likely allocation grouped with the second most likely allocation approaches one, there is ambiguity in the assignment of the particular sample under study. Conversely, samples for which this ratio yields a large result in comparison to all other ratios can be assigned to a single population with more confidence. For the two populations considered in the ratio, the chance of error is equal to the inverse of this ratio. Stringency for population allocation can be applied by defining a selection criterion for the log10 of this ratio. For example, by selecting only assignments that have a log of odds (LOD) ratio of at least 2, all results will have a 1/100 chance of error or less.

2. Multilocus likelihood functions may be grouped in a maximum likelihood format according to the equation \(L(\eta) = L(\eta, s)/L(\max)\). This yields a series of ratios between 1 (most likely) and close to 0 (least likely). Analysis of variance of log transformed data followed by a Tukey’s multiple comparison enables evaluation of statistical significance in the classical sense.

3. Jackknife iterations provide an empirical means for evaluating baseline data and the chances of correct allocation. Iterations sample individuals from the baseline one at a time, recalculating allele frequencies in the absence of each individual genotype sampled before determining the most likely population origin for that individual. Experimenting with alternate loci and populations enables one to determine which population comparisons and loci combinations enable reliable population reallocation.

**Reporting Options and Special Cases**

Sample ID, genotypic data, and multilocus likelihoods for population allocation can be displayed for verification. A critical population routine allows one to select a target population for calculation of LOD scores. All scores are then calculated with the critical population as the numerator in the ratio. A special case where test samples may have an allele or pair of alleles not observed in one or all of the baseline populations is treated as follows. For source populations in which the allele is not observed, an estimated allele frequency of \(1/(2N + 1)\) is applied. This hypothesis that the nonobservation of the allele in question is due to sampling error and that the allele in question would have been observed in the baseline population if one more allele had been sampled. Note that this estimation may introduce substantial bias if baseline population size \((N)\) is small, as would be likely for any allele frequency estimation given small \(N\), particularly when dealing with highly polymorphic marker types. The program implements a warning describing this consideration when small baseline population sizes \((N < 30)\) are encountered. Alternatively, if sampling error is low, an unknown sample allele not observed in a baseline population may constitute strong evidence that the sample in question may indeed not originate from the particular baseline population under consideration. Any alleles for which the \(1/(2N + 1)\) estimation is necessary are noted on the genotype output.

It is obvious that a technique such as WHICHRUN will only be effective if there is reasonable reproductive isolation among populations under study. Three other considerations are also important. First, the rate of accumulation of variance for molecular loci employed should be closely matched with estimated divergence times among populations under study. For example, highly polymorphic microsatellites prone to homoplasy would not be suitable for diagnosis among populations that have diverged over substantial evolutionary time. However, highly polymorphic microsatellites are likely one of a few molecular marker types that have sufficient information to resolve diagnosis among recently diverged populations such as the global radiation of *Drosophila melanogaster*, which is estimated to have occurred within the last 10,000–15,000 years (Bénassi and Veuille 1995; David and Capy 1988). Second, the accuracy of determination is crucially dependent upon the lack of differential sampling error among baseline allele frequencies. While this problem is partially addressed through ensuring that sample size is equal for all populations, highly polymorphic marker types such as microsatellites require substantial sampling. Third, for population origin diagnoses where source populations are recently diverged, there will be a number of loci that have not accumulated differences in the time since divergence. As a result, simply increasing the number of loci employed may not necessarily increase the power of diagnosis. For closely related populations, additional loci that have marked differences in allele frequency profiles among populations will be necessary to achieve increased power.

WHICHRUN may be downloaded from http://www-bml.ucdavis.edu/whichrun.htm.

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