Population Frequencies for STR’s

1 Population Frequencies for STR’s

1.1 To interpret the significance of a match between genetically typed samples, it is necessary to know the population distribution of alleles at the loci that were typed. If the STR alleles of the relevant evidence sample are different from the alleles of a subject’s reference sample, then the subject is “excluded,” and cannot be the donor of the biological evidence being tested. An exclusion is independent of the frequency of the alleles in the population.

1.2 If the subject and evidence samples have the same alleles, then the subject is “included”, or is a “match”, and could be the source of the evidence sample. The random match probability, or the probability that another, unrelated, individual would also match the evidence sample, is equal to the frequency of the evidence profile genotypes in the relevant population. Population frequencies are estimated separately for the Asian, Black, Caucasian and Hispanic populations. Additional population frequencies may be used for other population groups. If a source contains more than one frequency for a single population group, then the highest frequency is used for calculations. Allele frequencies are used for all calculations. Profile frequency estimates are calculated according to the National Research Council report entitled The Evaluation of Forensic DNA Evidence (National Academy Press 1996, pp. 4-36 to 4-37).

1.3 Spreadsheets may be used to automate the calculation of the population specific genotype and profile frequency estimates. The spreadsheets, or links to the spreadsheets, are located in Qualtrax.


1.5 If both autosomal and Y-STRs statistics are calculated, the results are reported separately.

2 Random Match Probability for Autosomal STRs

2.1 For a match to an Identifiler evidence profile, enter the evidence profile alleles in the worksheet of the POPSTATS spreadsheet. For Fusion body identification cases (ex. match between a remain to remain or a remain and a reference sample), the CODIS software POPSTATS module is used.

2.2 Off-ladder alleles are entered as decimals (for example, “12.2”) or as “>” or “<” for values above or below the ladder, respectively.

2.3 For loci assigned a “Z” to indicate the possible presence of another allele, only one allele is entered in the calculation. In this manner, the locus is not treated as a true homozygote whose
statistical values are determined by squaring the allele frequency \(p^2\). Rather “Z” loci utilize the probability only of the one assigned allele \(2p\), which allows the second allele to be anything.

2.4 The overall profile frequency estimate for each group is calculated by multiplying the individual locus genotype frequency estimates together.

2.5 In the standard scenario, homozygote genotype frequencies are estimated for each population using the formula \(p^2+p(1-p)\theta\) for \(\theta = 0.03\) and heterozygote genotype frequencies are estimated using the formula \(2p_i p_j\).

2.6 Within the POPSTATS spreadsheet, profile frequencies are also estimated for isolated populations (i.e., “evidence and subject from the same subgroup (isolated village)”) and for relatives using the formulas in the National Research Council Report.

2.7 For each population, the overall profile frequency estimate under the standard scenario of \(\theta = 0.03\) unless there is reason to suspect that the “evidence DNA and subject are from the same subgroup” or a relative of the subject left the biological sample.

2.8 Calculations are retained in the case file for referral at a later date if necessary.

2.9 The most common match probability of the four population groups is reported to three significant figures, truncated. If the match probability is less than 1 in 6.80 trillion, the match probability is reported is 1 in greater than 6.80 trillion people.

3 Frequency for Y STRs

3.1 The frequency for a Y STR haplotype is estimated by counting the number of times the haplotype occurs in each of the population databases and dividing by the total number of individuals in the database.

3.1.1 A haplotype that has not been previously observed in the Asian database, which includes 196 individuals, would be reported as “less than 1 in 196 Asians”.

3.1.2 A haplotype that has been observed once in the Asian database would be reported as “1 in 196 Asians”.

3.1.3 A haplotype that has been observed 5 times in the Asian database is reported as “1 in 39 Asians” (5 in 196 is equal to 1 in 39).

3.1.4 The 95% upper-bound confidence statistic from all ethnic groups is reported, truncated to three significant figures.

3.2 To calculate a statistic for Y-STR haplotypes or Y-STR mixtures, use the Y-Mix Database Filter. Refer to the protocol Usage of the ‘Y-Mix Database Filter’ for instructions.