Basic Principles for Estimating the Rarity of Y-STR Haplotypes Derived from Forensic Evidence

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Introduction
The Y-chromosome STR loci are male-specific and thus typing of these genetic markers enable analysis of previously problematic forensic biological samples, such as mixtures comprised of low quantities of male DNA amidst a large background of female DNA. In addition, some paternal lineage questions may be elucidated with Y STR loci. (1-9). The loci that have been validated for forensic applications include DYS19, DYS385 (counted as two loci), DYS389I, DYS389II, DYS390, DYS391, DYS392, DYS393, DYS437, DYS438, DYS439, DYS456, DYS458, DYS635, DYS448, and Y GATA H4. Their use is promoted by availability of robust and reliable commercial multiplex kits - Powerplex Y (12 locus multiplex kit; Promega Corp., Madison, WI) (10) and Y filer (17 locus multiplex kit; Applied Biosystems, Foster City, CA) (11).

When the evidence and reference sample Y STR profiles are sufficiently similar (i.e., matches or can be a component of a mixed sample) and thus potentially could have originated from the same source, some significance is placed on the observation, typically by estimating the rarity of the evidentiary profile. (12). The interpretation of inclusion or exclusion for Y STR loci profiles is the same as for autosomal STR loci profiles. However, when there is a failure to exclude, the approach for estimating the rarity of a Y DNA profile differs from that applied to autosomal DNA markers, because of the non-recombining features of the Y-STR loci typed. This paper describes the basic aspects of generating reliable statistical inferences of the rarity of Y STR evidentiary profiles. Three concepts are summarized. They are:

1. Use of the counting method and the accompanying upper bound haplotype frequency for estimating a conservative frequency of the observed haplootype

2. Accommodating reasonable possible effects of population substructure

3. Assessing the significance of a mixture
The Weight of Y STR Haplotype Evidence

The forensically important Y STR loci reside in the non-recombining portion of the hemizygous Y chromosome. Thus, the assumption of independence between and among loci does not apply for the Y STR loci. The product rule, even the modified product rule (13), is not appropriate for estimating the rarity of a Y STR haplotype. Essentially, because of linkage, each haplotype is treated as an allele and the total number of possible haplotypes comprises the alleles of a single locus. This is the same for all Y STR systems, no matter how many different loci are used to generate the haplotype profile.

Because the haplotypes are treated as alleles of a single locus, the counting method, which is simple, robust, and conservative, can be applied for assessing the significance of a Y STR haplotype derived from an evidence sample (14,15). The counting method is carried out as follows:

1. The profile of a Y STR haplotype generated from an evidence sample is searched against a reference database(s) of unrelated individuals;

2. The number of times the Y STR haplotype is observed in a database(s) is counted (termed “count”);

3. That count is divided by the number of profiles in the reference database(s);

4. A confidence interval is placed on the proportion of count/total profiles in a database(s).
For the foreseeable future, the size of the available reference databases will be limited such that most haplotypes (certainly the 12-17 loci profiles generated using commercial kits) will not be observed or only observed once in a reference population database. Because of this constraint, an estimate of the frequency of the haplotype in a population is not possible by calculating the proportion of the number of matching profiles divided by the total number of profiles in the database. To address this limitation, a conservative bound is placed on that estimate (count/total) to correct for possible sampling error. The confidence interval allows for a measure of the amount of confidence which can be placed on a value lying between two specified limits (i.e., the interval). One can calculate the upper bound of the confidence interval and this value can be used to convey with a high degree of confidence that the rarity of the evidence Y-haplotype among unrelated individuals in a given population(s) is less than the upper bound of the estimate. The assumption of a Normal distribution may not apply for Y-STR haplotype frequency estimates (16); however, assuming normality will provide a conservative upper bound estimate. Exact methods for constructing confidence intervals are available (17,18).

The counting method has several features that make it desirable for forensic applications. First, it is a conservative approach. Although the counting method treats all loci as physically linked (which they are biologically), not all Y STR loci are in linkage disequilibrium to the same extent. Some pairs of Y-STRs fail to show significant linkage disequilibrium and there are other pairs for which the linkage disequilibrium is incomplete (i.e., less than the maximum possible; 14, 19). Kayser et al. (20), Heyer et al (21), and
Budowle et al (14), have typed confirmed father/son pairs for 12 or more Y-STR loci and have observed mutation rates on the order $10^{-3}$. This rate of mutation will somewhat destabilize the effects of linkage, because of the expansion-contraction (of allele size) nature of mutations. Therefore, the assumption of complete linkage disequilibrium contributes to the conservative nature of the counting method. Second, although Y-STR profiles are not as individualizing as are an equal (and even fewer) number of autosomal STR loci, it is still unlikely to draw two unrelated people at random with the same minimal haplotype. Furthermore, in some cases, where no result can be obtained by autosomal STR typing, Y STR typing may be the only viable approach to obtain a result. Thus, even with their combined lower power, the Y STR loci may be the best tool available in certain cases to exclude those individuals falsely associated with the sample. Although Y STR haplotypes are treated as single locus systems, the forensic Y multiplex systems exhibit greater diversity than any single autosomal STR locus. This observation contributes to exceedingly low Fst values used for correcting for the effects of possible substructure (see below). Third, a similar counting approach has been in use for more than 10 years for estimating the rarity of evidentiary mtDNA haplotypes (22). So the counting method is a well established application in the forensic community and in legal proceedings.

Population Substructure

The effects of population structure are expected to be greater for the Y-chromosome than with the forensically important autosomal STR loci. This is due to: 1) the non-recombining Y STR loci are not shuffled during meiosis as are the autosomal
genetic markers; and 2) Y chromosomes have an effective population size that is 1/4 that of autosomal loci. This linkage and smaller effective population size contribute to population specific distributions, which are affected by genetic drift and geographic differentiation. Indeed, population substructure effects have been shown to be more substantial for Y loci compared with that observed for the autosomal STR loci (see e.g., 23 and their cited references). This would suggest that 1) reference population databases may have to be defined more restrictively (most likely by geography and ethnohistory) than is typically done for autosomal STR loci; or 2) current practices of pooling databases based on major population groups can be employed if corrections are employed for effects of substructure when estimating the rarity of the profile (13). It is incumbent upon forensic scientists to assess the effects of population substructure and employ statistical approaches that address those effects.

As recommended by the NRC II Report (13), \( \theta \) can be used to adjust for possible population substructure. Using the general theory, the unconditional frequency of a haplotype \( (A_i) \), which is the count divided by the sample size, can be modified to obtain the conditional probability

\[
Pr (A_i|A_i) = \frac{p_i^2 + \theta p_i(1-p_i)}{p_i}
\]

\[= p_i + \theta(1-p_i) \]

\[= \theta + p_i(1 - \theta)\]

Based on this formula, the conditional probability always exceeds the quantity of \( \theta \), making \( \theta \) the minimum bound for estimating the rarity of a Y STR haplotype. So the
probability of observing a particular haplotype in an unrelated individual given that it is seen in another male is highly dependent on the value of $\theta$ deemed appropriate for the relevant Y-STR haplotype database.

The value of $\theta$ (computed by either $R_{ST}$ or $F_{ST}$) is dependent indirectly on the number of Y STR loci comprising the haplotype. Generally, as more loci are included in the haplotype, most haplotypes in a data set will become differentiated. Therefore, the greater the number of loci within the haplotypes, the smaller should be the value of $\theta$.

$R_{ST}$ has been used to estimate the value of $\theta$ for forensic calculations (24). However, the analyses based on $R_{ST}$ are better applied to evolutionary biology for studying the phylogenetic history of Y-chromosomes. The $R_{ST}$ values are based on allele size variance, exploiting the extent of difference between different haplotypes. Such an approach, however, typically does not apply to forensic inferences. Forensic applications assess the evidence in terms of match or non-match. In these terms, the $\theta$ values for Y STRs should not be computed on mismatch based approaches (such as AMOVA), but instead by simply treating all haplotypes as different alleles. Thus, $F_{ST}$ (or $G_{ST}$) is a better estimator for $\theta$ relevant for forensic applications, because haplotypes are identified solely by their distinctiveness (i.e., haplotypes are considered simply in terms of identity by state). This generally leads to a more appropriate and much smaller $\theta$ value than estimated by $R_{ST}$.

The $F_{ST}$ values across major populations range from less than 0.002 to less than 0.0008 (depending on the number of loci comprising the haplotype). The population-specific values are smaller, excluding Native Americans (manuscript in preparation). Therefore, with the size of current reference population databases, $F_{ST}$ rarely will
influence the upper bound of the Y STR frequency. Since $F_{ST}$ does not affect the upper bound frequency estimate on any practical level for most population groups (with the current commercially available systems), increasing the size of reference population data sets and including more populations would be more valuable and a better use of resources than adding more loci to the current multiplexes for exploiting the full power of Y STR typing. Dedicating efforts to expanding the battery of Y STR loci will not significantly exploit the power of the assays.

As population comparisons are carried out, significant, but small $F_{ST}$ values at the level of haplotypes for the Y-STRs may and have been observed (24 and manuscript in preparation). This raises the question of the legitimacy of pooling of population samples for forensic calculations, as is done for autosomal STR loci (25). However, as long as the effects of population substructure are addressed adequately, pooling of populations is appropriate. Since most haplotypes are unique in a reference population data set, the conservative counting method already more than compensates for population substructure effects, when pooling geographic samples of populations of broad definitions (e.g., US Caucasian, African American, Hispanic, etc.).

One might attempt to suggest that because of significant differences (albeit small) in population comparisons that no general reference (population specific) database can be used for forensic statistical computations. This position would ignore the fact that regardless of population distinctions, observing an extended Y STR haplotype (12-17 loci) is a rare event in any of the major population groups routinely considered for forensic analyses. For examining the effect of population substructure, haplotype diversity is a very useful indicator. The haplotype diversities are greater than 0.99
(Kayser) and with the extended Y STR haplotypes (with the current commercially available kits) diversities tend to be greater than 0.999 for most populations. Other pragmatic justifications of pooling populations for Y-STR haplotype databases include the unspecified nature of the lineage of paternal (population) ancestry based on self-identification of individuals as member of a specified population.

Mixtures of Y STR Haplotype Profiles

The theory and logic (for use of the exclusion (PE) and/or the likelihood ratio (LR)) for assessing the significance of mixture samples are well-described (26,27 and references within). The PE provides an estimate of the portion of the population that has a genotype composed of at least one allele not observed in the mixed profile. The LR provides the odds ratio of two competing hypotheses, given the mixture evidence. There are two limitations, however, for Y STR mixture analyses that need to be addressed. These are: what frequencies to apply for all possible haplotypes that contribute to the mixture; and 2) user-friendly algorithms of interpretation of Y-STR evidence profiles.

Assume that there 2 alleles observed in a mixed sample at each locus of a 12 locus Y STR profile. Assuming independence across the loci (an obviously incorrect assumption), there can be $2^{12}$ possible haplotypes (4,096) that could possibly contribute to the mixed profile. Most of these possible profiles will not have been seen in a database and some of these possible haplotypes may not exist in the greater population. So for computing the PE (that is for a random male to be a (part) contributor of a DNA mixture) a minimal haplotype frequency (similar to the logic of a minimum allele frequency for autosomal loci (28)) is not practical. However, one can employ the same principles used
for the counting method described above and count the all contributing possible 
 haplotypes that can be part contributors observed in a reference database(s), divide that 
 by the total number of samples in the database(s), calculate the upper bound of the 
 counted proportion and subtract the upper bound estimate from 1.

The PE evaluation does not use any information about the Y-STR profiles of the 
 known persons tested or the combinations of profiles that can explain the observed 
 evidence. Hence, the total strength of observations is not fully captured by the concept of 
 PE. The likelihood ratio is a more informative approach for assessing the significance of 
 a mixed evidentiary profile.

For addressing the LR only a two person mixture will be considered for 
 simplicity. There are four general scenarios for a two contributor sample:

1. S1 and S2 are the source
2. S1 and an unknown male are the source
3. S2 and unknown male are the source
4. two unknown males are source

where S1 and S2 are identified, DNA-typed suspects. Since scenarios 2 and 3 are the 
simpler versions of the same calculations as for 1-PE (or the probability of inclusion) 
described above (with the modification that the unknown male must have alleles that 
explain all of the alleles in the mixture, taken in combination with the alleles in the 
profile of S1 or S2), only scenarios 1 and 4 are described below.
For simplicity, only three loci are shown for the LR. Assume two alleles reside at each of the three loci in a mixture. Two male suspects are identified and the combined Y STR haplotypes of the suspects explain all the alleles observed in the evidence (Table 1).

Table 1. Possible 3 locus haplotypes for fictitious mixed sample DYS319 - 13/15; DYS437 - 14/16; DYS389II - 28/30; S1 has the profile DYS319 - 13; DYS437 - 16; DYS389II - 30; S2 has the profile DYS319 - 15; DYS437 - 14; DYS389II – 28.

<table>
<thead>
<tr>
<th>DYS319</th>
<th>DYS437</th>
<th>DYS389II</th>
<th>Haplotype</th>
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<tbody>
<tr>
<td>13</td>
<td>14</td>
<td>28</td>
<td>haplotype 1</td>
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<tr>
<td>15</td>
<td>14</td>
<td>28</td>
<td>haplotype 2</td>
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<td>13</td>
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<td>haplotype 3</td>
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<td>28</td>
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<tr>
<td>13</td>
<td>16</td>
<td>30</td>
<td>haplotype 7</td>
</tr>
<tr>
<td>15</td>
<td>16</td>
<td>30</td>
<td>haplotype 8</td>
</tr>
</tbody>
</table>

Under the prosecution hypothesis (Hp), the probability is 1 of observing the evidence given the two suspects are the sources of the DNA in the mixed profile. In contrast, under the defense hypothesis (Hd), where two unknown individuals are the source of the evidence, the probability of the evidence is calculated by determining the proportion of the population of pairs of unrelated individuals drawn together at random whose combined profiles are the same as the evidence (note: alternate hypotheses can be generated; only one is described herein for illustration). Essentially, the denominator is the same as the probability of inclusion (1-PE), but with consideration of a prescribed number of contributors. If one considers the haplotypes in Table 1 that could not be excluded as possibly comprising a part of the evidence profile, not all possible pairwise
combinations of the 8 haplotypes could explain all the alleles observed in the mixture. By considering only those pairings the LR (for the 3 loci) is:

\[ LR = \frac{1}{2} [Pr(H_1)Pr(H_8) + Pr(H_2)Pr(H_7) + Pr(H_3)Pr(H_6) + Pr(H_4)Pr(H_5)] \]

The formulation for the LR is correct and has been described by others (see 29) but it is not particularly useful, because no direction is offered for what values (i.e., frequencies to be used for the haplotypes drawn randomly from a population) should be used in the denominator. Again the same principle can be applied as described above for the counting method. All pairwise combinations of profiles in a reference database(s) can be generated and then those pairs that match the evidence can be counted. Then an upper bound of the confidence interval of the proportion of counts over total pairs is calculated.

Conclusion

The Y STR loci currently available through commercial kits are highly informative. For statistical purposes the counting method (and sampling correction) is simple, conservative, and well-established for assessing the rarity of a profile. To correct for possible effects of substructure, gene diversity-based \( F_{ST} \) (treating the haplotypes as alleles of a single composite locus) is the more appropriate measure of \( \theta \) to be used in a conditional probability calculation. Because Y STR haplotype data are treated similarly to that of single locus data and haplotype diversity is high, the \( F_{ST} \) values are very small for most populations. Thus, for current size databases, the upper bound estimate of the count proportion is more than adequate to compensate for uncertainty for most populations (excluding for example Native Americans). Lastly, the basic logic for
calculating Y STR mixtures is the same as for autosomal markers, i.e., the application of
the PE and/or the LR. Appropriate frequencies can be estimated for the unknowns in the
LR by using the same logic as that of the counting method. Software is under
development to facilitate mixture calculations (manuscript in preparation).

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