DNA Fingerprinting for Forensic Identification: Potential Effects on Data Interpretation of Subpopulation Heterogeneity and Band Number Variability

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Summary
Some methods of statistical analysis of data on DNA fingerprinting suffer serious weaknesses. Unlinked Mendelizing loci that are at linkage equilibrium in subpopulations may be statistically associated, not statistically independent, in the population as a whole if there is heterogeneity in gene frequencies between subpopulations. In the populations where DNA fingerprinting is used for forensic applications, the assumption that DNA fragments occur statistically independently for different probes, different loci, or different fragment size classes lacks supporting data so far; there is some contrary evidence. Statistical association of alleles may cause estimates based on the assumption of statistical independence to understate the true matching probabilities by many orders of magnitude. The assumptions that DNA fragments occur independently and with constant frequency within a size class appear to be contradicted by the available data on the mean and variance of the number of fragments per person. The mistaken use of the geometric mean instead of the arithmetic mean to compute the probability that every DNA fragment of a randomly chosen person is present among the DNA fragments of a specimen may substantially understate the probability of a match between blots, even if other assumptions involved in the calculations are taken as correct. The conclusion is that some astronomically small probabilities of matching by chance, which have been claimed in forensic applications of DNA fingerprinting, presently lack substantial empirical and theoretical support.

Introduction
Minisatellites are regions of the genome in which a DNA sequence is repeated tandemly for a variable number of times. Minisatellites are dispersed in the genomes of humans and other mammals. A subset of the repeated DNA sequences contains a short, shared “core” sequence (Jeffreys et al. 1985a). Different subsets of minisatellites share different core sequences. Minisatellites that contain a given core sequence can be detected by a hybridization probe that consists of the core sequence repeated in tandem.

When human DNA is digested by a restriction enzyme and hybridized with a specific core probe, the resulting fragments can be distributed according to size by a Southern blot. This procedure reveals the sizes of fragments that contain a specific core. The pattern of fragment sizes is highly variable or polymorphic from one individual to another (except for identical twins) but is apparently highly stable or conserved both from one tissue to another and over time within a given individual (Jeffreys et al. 1985a). The combination of variability among individuals and stability within an individual suggests that the patterns of fragment sizes revealed by specific probes might serve as useful individual identifiers, or “DNA fingerprints” (Jeffreys et al. 1985b).

In forensic practice, the patterns of DNA fragment sizes are usually determined by one of two distinct procedures. One procedure, common in the United States,
uses several single-locus probes serially. The other procedure, common in the United Kingdom, uses a single probe that detects alleles at multiple loci.

In forensic applications of DNA fingerprints, a specimen's pattern of DNA fragment sizes is compared with a collection of such patterns from some set of people. A measure of similarity between any two such patterns is defined. Then a probability of observing, by chance alone, a given level of similarity is computed. If the specimen's pattern has a priori a very low chance of being similar to the pattern of a randomly chosen person, and if, nevertheless, the specimen's pattern is observed to be very similar to the pattern of some particular person, then it is concluded that the individual whose pattern matches that of the specimen is the source of the specimen, with high probability.

The scope of the present paper is limited to statistical problems in the analysis of data on matches between DNA fragments of various sizes for the purpose of identifying an individual by means of a specimen. Some published procedures appear to lack adequate empirical or theoretical support. The following sections address problems raised by (1) heterogeneity in populations, (2) calculating the probability that a given match between two fingerprints would arise at random, and (3) calculating the average power of a fingerprint. As the present paper is intended to be constructive as well as critical, some procedures, experiments, and analyses are suggested which could provide a firmer foundation for the use of DNA fingerprinting for forensic identification.

The present paper does not deal with uncertainties and ambiguities in the underlying biochemical procedures and data, such as the problems of collecting uncontaminated specimens at the scene of a crime, degradation of materials prior to analysis, use of internal controls and mixture experiments in electrophoretic gels, the necessity for "blind" judgments and probabilistic assessment of a match between bands in different lanes of a gel or on different gels, and others (e.g., see Lander 1989; Lewin 1989; Sensabaugh and Witkowski 1989). The present paper does not deal with questions of laboratory protocol, such as the chain of custody of samples and quality assurance (Sensabaugh and Witkowski 1989). Although recent journalistic accounts suggest that routine laboratory error rates for this kind of test may be as high as 1%-5% (according to a referee), procedures for reducing these error rates are in principle well understood and could be applied in sensitive cases. The present paper does not deal with legal questions, such as the admissibility or presentation of DNA fingerprinting data in court or pretrial hearings and the determination of a quantitative threshold for a probability that is "beyond a reasonable doubt" (e.g., see Tribe 1971; Fienberg and Schervish 1986). The present paper does not deal in detail with the use of DNA fingerprinting to establish genetic relationships among people—e.g., in paternity testing—although it will be apparent that many of the problems discussed below arise also in paternity testing. In genealogical applications, a high spontaneous mutation rate to minisatellites of different lengths has been noted (Jeffreys et al. 1988).

### Population Heterogeneity and Statistical Dependence of Alleles

The methods that have been used to infer probabilities of identification by using DNA fingerprinting appear to overlook population heterogeneity in gene frequencies. A simplified example will show that such heterogeneity may render invalid a key assumption made in applications of DNA fingerprinting. This example contains no new genetics or statistics, but the issues it raises have not received adequate attention. These issues apply equally to fingerprinting procedures based on single-locus probes and to those based on multi-locus probes.

Consider a large human population (e.g., Britain) in which one probe detects allelic variation in restriction-fragment length at one autosomal locus and in which another probe detects allelic variation in restriction-fragment length at another autosomal locus. Suppose there are just two alleles at each locus. Call the alleles at the first locus A and a, and call the alleles at the second locus B and b. No dominance between alleles is implied: A, a, B, and b each are assumed to correspond to well-defined distinct bands, so that the genotype of an individual (at these two loci) can be unequivocally determined from inspection of a gel. Suppose also that the two loci are located either on different autosomes or far enough apart on the same autosome so that the recombination fraction between loci is 1/2. Suppose also that inheritance at each locus is strictly Mendelian. So far this is a textbook model of two loci with two alleles (e.g., see Crow and Kimura 1970).

Now suppose that Britain contains two subpopulations. Call the two subpopulations F and G. Suppose that within each subpopulation the two loci are in linkage equilibrium and that there is random mating and no selection with respect to both loci. Let $p(A,F)$ denote the gene frequency of allele A in subpopulation
More generally, let \( p(i, H) \) denote the gene frequency of allele \( i \) in subpopulation \( H \), where \( i = A, a, B, b \) and \( H = E, F \). Thus \( p(A, F) + p(a, F) = 1 \), \( p(B, F) + p(b, F) = 1 \), and similarly with \( F \) replaced by \( G \).

Under the preceding assumptions, within each subpopulation each band (allele) is statistically independent: the genotype frequencies within a subpopulation are simply the products of the appropriate gene frequencies within that subpopulation. For example, if \( f(AABB, F) \) denotes the relative frequency of the AA homozygote at the first locus and of the Bb heterozygote at the second locus in subpopulation \( F \), then \( f(AABB, F) = p(A, F)^2p(B, F)p(b, F) \).

If the gene frequencies are different in the two subpopulations, it is not in general true that each band (allele) is statistically independent in the population as a whole. On the contrary, in the population, different bands may be positively or negatively associated, depending on the proportions of people in the different subpopulations and on the differences of the allele frequencies at each locus. Statisticians have known for a long time (Yule 1903) that attributes (allele frequencies, in this case) may be positively or negatively associated in a population as a result of pooling the frequency of the attributes in subpopulations in which the attributes are independent. This phenomenon, still of active interest (Good and Mittal 1987), appears to have been overlooked in forensic applications of DNA fingerprinting.

A numerical example, using completely arbitrary figures, illustrates the phenomenon. Suppose the fractions of the whole population that belong to subpopulations \( F \) and \( G \) are given by \( \pi(F) = .9 \) and \( \pi(G) = .1 \). Suppose that \( p(A, F) = .3 \), \( p(A, G) = .6 \), \( p(B, F) = .4 \), and \( p(B, G) = .8 \). The overall relative frequency of the A allele in the population is \( p(A) = p(A, F)\pi(F) + p(A, G)\pi(G) = .33 \), whence \( p(a) = 1 - p(A) = .67 \) is the overall relative frequency of the a allele. Similarly, \( p(B) = .44 \) and \( p(b) = 1 - p(B) = .56 \). The actual relative frequency of the AABB genotype in the population is \( f(AABB) = p(A, F)^2p(B, F)^2\pi(F) + p(A, G)^2p(B, G)^2\pi(G) = .36 \). However, if the relative frequency of the AABB genotype in the population is calculated assuming independence between alleles (bands), the estimate is \( f'(AABB) = p(A)^2p(B)^2 = .021 \).

To make the example slightly more realistic, assume that, within each subpopulation, the alleles \( A_1, A_2, \ldots, A_{10} \) have the same allele frequency as \( A \) (so that \( a_1, a_2, \ldots, a_{10} \) have the same allele frequency as \( a \)) and that the alleles \( B_1, B_2, \ldots, B_{10} \) have the same allele frequency as \( B \) (so that \( b_1, b_2, \ldots, b_{10} \) have the same allele frequency as \( b \)) and that the different alleles are statistically independent. The actual relative frequency of the homozygous genotype \( A_1A_1A_2A_2 \ldots A_{10}A_{10}B_1B_1B_2B_2 \ldots B_{10}B_{10} \) in the whole population is \( [p(A, F)^2p(B, F)^2]^{10}\pi(F) + [p(A, G)^2p(B, G)^2]^{10}\pi(G) = 4.2 \times 10^{-8} \). However, if the relative frequency of the genotype in the population is calculated assuming independence between alleles (bands), the estimated relative frequency of the homozygous genotype \( A_1A_1A_2A_2 \ldots A_{10}A_{10}B_1B_1B_2B_2 \ldots B_{10}B_{10} \) is \( p(A)^{20}p(B)^{20} = 1.7 \times 10^{-17} \). Now suppose a forensic specimen is determined to have the homozygous genotype \( A_1A_1A_2A_2 \ldots A_{10}A_{10}B_1B_1B_2B_2 \ldots B_{10}B_{10} \) by DNA fingerprinting, and suppose a suspect individual is identified whose genotype exactly matches that of the specimen. In this case, the estimated probability of a match, when independence of alleles is assumed, is lower than the true probability of a match when one allows for the heterogeneity of subpopulations, by a factor of more than \( 10^{-9} \). The estimated probability, being lower than the true probability, exaggerates the significance of a match and unnecessarily incriminates the suspect. The numerical values in this example were chosen in advance for simplicity, rather than being selected to illustrate a worst case.

This hypothetical example resembles reality in that there is likely to be significant genetic heterogeneity in real populations. The allele frequencies of genes of medical interest differ from one human subpopulation to another. Minisatellite regions and other RFLPs that serve as markers of disease-related genes (Gusella et al. 1983; Jeffreys et al. 1986) are likely to share that heterogeneity. The DNA probes used for forensic identification detect alleles that have heterogeneous allele frequencies: Lander (1989, p. 504), using Wahlund’s formula, found excess homozygosity (relative to Hardy-Weinberg equilibrium) at loci identified by DNA probes in the Hispanic population used as the reference population in a murder trial, demonstrating “the presence of genetically distinct subgroups within the Hispanic sample.”

The hypothetical example given above differs from reality in that neither the assumption of just two subpopulations nor the particular allele frequencies and subpopulation frequencies assumed are likely to be realistic. The actual effect of subpopulation heterogeneity could be larger or smaller than that in this example.

The example demonstrates three points. First, the population used to obtain estimates of allele frequencies is crucial for subsequent applications of match probabilities to individual cases. Future studies should care-
fully define a reference population (what statisticians call a sampling universe) which is to be studied, and they should make explicit the procedure (such as systematic sampling or random sampling) that is used to sample from this population. Options and procedures for proper sampling have been clearly described elsewhere (e.g., see Snedecor and Cochran 1980, chap. 21).

There are good practical and scientific reasons for giving serious attention to sampling. In practice, if a study that attempts to derive matching probabilities for DNA fingerprinting is based on an ill-specified sample, the resulting probabilities can be challenged in court on the grounds that the study sample is not the sample most appropriate to the accused individual. If the proponents of DNA fingerprinting wish to claim that the probabilities of matching at random are astronomically low for virtually all populations, they are obliged to demonstrate the claim for at least several well-defined populations. Scientifically, DNA fingerprinting provides a means of assessing the genetic heterogeneity of populations. Studies of well-defined samples offer an opportunity to compare the genetic variability of different populations and could be of potential interest to students of human evolution (Lander 1989, p. 504).

Second, alleles (bands) may be significantly statistically associated in a population if there is heterogeneity between subpopulations in the allele frequencies, even though the loci involved may be strictly Mendelian, unlinked, and at linkage equilibrium within each subpopulation. Wherever subpopulations are heterogeneous, true random samples of populations are required to measure directly whether any statistical association of minisatellite alleles results from pooling across subpopulations.

Third, the statistical association of alleles, though undetectable either in terms of chromosomal mechanisms or within homogeneous subpopulations, may induce significant errors in estimates of match probabilities if the estimates ignore the statistical association.

Among statisticians, the gratuitous assumption of independence is well known as a source of superficially persuasive arguments for the existence of miracles (Kruskal 1988), which correspond in the present situation to extravagantly small alleged probabilities of obtaining a match at random.

Calculating the Probability of a Match at Random

Given a match between the DNA fingerprint of a specimen and the DNA fingerprint of an individual, how can one calculate the probability that this match could have arisen at random? An obvious possibility is to record the complete genetic pattern (not broken into discrete bands) of each individual in a population survey and then count how many complete patterns match that of the specimen. If S individuals are surveyed, this approach cannot yield a probability of match lower than 1/S (given that one individual's pattern matches the specimen's pattern). To obtain, by this approach, the extremely low probabilities of a match quoted in many forensic applications of DNA fingerprinting would require surveying more people than are alive. Since this would be difficult, the quoted low probabilities are obtained from calculations based on simplifying assumptions about the occurrence of bands in different regions of Southern blots. The results of the calculations may be in error if the assumptions are not justified.

Among the major assumptions sometimes made are the following:

1. The probability of a match for a given probe and fragment size class has been estimated by random sampling of a well-defined, genetically homogeneous reference population.
2. Matching of DNA fragments identified with one probe or at one genetic locus is independent of matching fragments identified with any other probe or at any other genetically unlinked locus.
3. For a given probe, the fragments identified may be categorized into size classes, and matching of DNA fragments in one size class is independent of matching in any other size class.
4. Within a size class of DNA fragments identified by a given probe, the probability of a match is constant for all fragments in the size class, and matching is independent for any two different fragments within the class, and there is no variability in the number of fragments per specimen or per person in the size class.

There is considerable evidence against these assumptions in some applications of DNA fingerprinting.

Regarding the first assumption, Jeffreys et al. (1985b) report data based on “a random sample of 20 unrelated British caucasians” who were, in fact, “20 volunteer white students from our university [University of Leicester]” (A. J. Jeffreys, personal communication, August 28, 1988). Similarly, Gill et al. (1987) analyze, among other materials, whole blood samples from 41 individuals of unspecified origin and characteristics. Such subjects do not represent a genuine random sample, in the technical sense (Snedecor and Cochran 1980,
chap. 21), of the entire "British caucasian" population. Such a sample requires a list or sampling frame from which a sample is randomly selected. The special population from which the subjects were drawn should be remembered before using the data for inferences about any other population.

Blood taken from each individual in the sample of Jeffreys et al. (1985b) was digested with a restriction enzyme and Southern blot hybridized with minisatellite probes 33.6 and 33.15. "Each DNA fingerprint (individual A) was compared with the pattern in the adjacent gel track (individual B), and the number of bands in A which were clearly absent from B, plus those which had a co-migrating counterpart of roughly similar autoradiographic intensity in B, were scored. The data shown are averages for all pairwise comparisons" (Jeffreys et al. 1985b, p. 76).

Table 1 reproduces table 1 of Jeffreys et al. (1985b). The data are based on approximately 15 resolved fragments (or bands) in the 4–20-kb size range. Jeffreys et al. (1985b, p. 76 fn.) describe the procedure used to compute probability estimates, as follows: "Probability estimates: the mean probability that all fragments detected by probe 33.15 in individual A are also present in B is 0.082.9 \times 0.205.1 \times 0.276.7 = 3 \times 10^{-11}." In their text, they observe that "the probability that the fingerprints of A and B are identical, that is, that all fragments less than 4 kb also match and that B does not possess any additional 4–20-kb fragments, is therefore << 3 \times 10^{-11}. Similarly, the probability that A and B have identical fingerprints for both probes 33.15 and 33.6 is << 5 \times 10^{-19}.

Retracing their steps, I confirm their arithmetic, except that the product of the separate probabilities for the two probes should be 5.0 \times 10^{-21}, not 5 \times 10^{-19} (a simple arithmetic error [A. J. Jeffreys, personal communication, August 28, 1988], which, however, is conservative, since 5 \times 10^{-19} > 5.0 \times 10^{-21}).

At the last step, when the probabilities for the different probes are multiplied, Jeffreys et al. (1985b) implicitly assume that matching (more precisely, the inclusion of A's bands among B's) for probe 33.15 is independent of matching for probe 33.6. This is the second assumption listed above. Jeffreys (personal communication, August 28, 1988) considers this assumption of independence to be supported by a subsequent study of a large sibship of 11 English individuals. Jeffreys et al. (1986, pp. 15–18) found that "none of the polymorphic DNA fragments scored using probe 33.15 were present in the set of fragments detected by 33.6; any such fragment that hybridized to both probes would have been detected as bands of equal size that were transmitted from the same parent to the same children (i.e., 'linked'). These two probes therefore hybridize to essentially completely different subsets of human mini-satellites. In addition, no bands detected by probe 33.6 were allelic to, or linked with, any fragments hybridizing to 33.15." However, as explained in the previous section, even if alleles segregate independently within a family, as the example of Jeffreys et al. (1986) suggests, the same alleles are not necessarily statistically independent in a population. To the best of my knowledge, Jeffreys et al. (1985b, 1986, and elsewhere; Wong et al. 1987) offer no direct statistical data, based on random samples, that different probes or loci match independently in a population. As another example of the unexamined assumption that different loci are independent in a population, Collaborative Research Inc. (1989) advertised 11 DNA probes for identification and parentage testing. The ad-

![Table 1](image-url)

**Table 1**

<table>
<thead>
<tr>
<th>Probe, and DNA Fragment Size (in kb)</th>
<th>No. of Fragments/ Individual Mean ± SD</th>
<th>Probability x That Fragment in A Is Present in B</th>
<th>Maximum Mean Allelic Frequency/ Homozygosity</th>
</tr>
</thead>
<tbody>
<tr>
<td>33.6:</td>
<td>2.8 ± 1.0</td>
<td>.11</td>
<td>.06</td>
</tr>
<tr>
<td>33.15:</td>
<td>2.9 ± 1.0</td>
<td>.08</td>
<td>.04</td>
</tr>
<tr>
<td></td>
<td>5.1 ± 1.3</td>
<td>.18</td>
<td>.09</td>
</tr>
<tr>
<td></td>
<td>5.9 ± 1.6</td>
<td>.28</td>
<td>.14</td>
</tr>
<tr>
<td></td>
<td>6.7 ± 1.2</td>
<td>.27</td>
<td>.14</td>
</tr>
</tbody>
</table>

Source.—Jeffreys et al. (1985b, p. 76).
vertiment listed a “probability of identity” for each probe, followed by a “probability of identity” for “all loci combined” of $4.96 \times 10^{-15}$. According to Stanley D. Rose, Director of DNA Products and Services for Collaborative Research Inc. (personal communication, July 18, 1989), two of the probes recognize different polymorphisms at the same locus; one of these probes was excluded in calculating the probability of identity for all loci combined. After that exclusion, according to Rose, “the cumulative average probability of identity is based on the assumption that all loci segregate independently.” Rose notes that, “although these probes provide an extremely powerful tool for identification of individuals, it should be pointed out that we (collectively speaking) know relatively little about the characteristics of DNA polymorphisms in large populations, or how the frequency distributions of specific alleles may differ in sub-populations.”

At the penultimate step of their calculation, when the match probabilities for different size classes of fragments (identified by a single probe) are being multiplied, Jeffreys et al. (1985b) implicitly assume that there is no statistical association between any two minisatellite regions in different size classes, so that matching in one size class of fragments is statistically independent of matching in any other size class. This is the third assumption listed above.

Later evidence argues against the assumption of statistical independence among fragments. One dog family displayed clear associations (positive and negative) among fragments of different lengths generated by DNA fingerprinting (Jeffreys and Morton 1987, pp. 8, 11). A large sibship of English humans displayed several allelic pairs of both paternal and maternal fragments identified by both probes; there was a linked pair of fragments in this and another pedigree (Jeffreys et al. 1986, p. 15). Moreover, alleles at the same locus span different categories of fragment length: “large differences in minisatellite allele lengths must exist, arising presumably by unequal exchange in these tandem repetitive regions; several allelic pairs identified . . . do indeed show substantial length differences” (Jeffreys et al. 1986, p. 18). These examples of statistical association (allelism or linkage) of fragments in different length categories within families could be amplified at the population level, as mentioned above, by heterogeneity among subpopulations.

At the first step of their calculation, in computing the match probability for DNA fragments within each size class (identified by a single probe), Jeffreys et al. (1985b) implicitly make three major approximations (as in the fourth assumption listed above): (1) that the probability (denoted by $x$ in table 1) that a fragment in A is also present in B is the same for all fragments in the size class, (2) that different fragments within a size class match or fail to match independently, and (3) that there is no variability in the number of fragments, either per specimen or per person, in the size class. Under these assumptions, it is correct to compute the probability of a match as Jeffreys et al. (1985b) have done: if $x$ is the probability that a fragment in A is present in B, and if $n$ is the (putatively constant) number of fragments per person in the size class, then the probability that all the fragments in A in the given size class are also present in B is precisely $x^n$.

In the remainder of this section, I examine the first two approximations: constancy of match probabilities within a size class and independence of matching. The third approximation, constancy in the number of fragments in the size class per specimen or per person, will be examined in the following section, because it relates to the major issue of calculating the average power of a match.

The assumption that the match probabilities are constant within a size class seems implausible, given the data in table 1, because the fragment match probability rises as the DNA fragment size falls, for both probes. It seems unlikely that the match probabilities change abruptly by discrete steps, precisely at the boundaries of the arbitrarily selected fragment-size classes. More likely, the match probabilities rise smoothly with falling fragment size. However, using a fixed average $x$ is a conservative approximation, in that it overstates the probability of a match (as may easily be shown by constructing a numerical example). Reanalysis of the raw data on which table 1 is based, by using finer size categories, or analysis of more extensive other data, could show whether the match probabilities are indeed constant within the given size classes.

Gill et al. (1987, table 2) also assume a constant probability of matching for fragments of different sizes in their analysis of the frequency distribution of the number of matches between individuals. They use two measures of matching: inclusion of the bands of individual A among those of individual B (like Jeffreys et al. (1985b)) and identity between the bands of individuals A and B (equivalent to inclusion in both directions). They use the binomial distribution to compute the probability of each possible number of matches under each definition of matching, and they find a reasonable agreement with the observed frequency distribution of number of matches. This calculation assumes explicitly that
the probability of any one band matching is .22, even though their own table 1 shows that x varies from .07 for the 10–20-kb fragments to .26 for the 4–6-kb fragments. The agreement between the expected and observed frequency distribution of number of matches is an insensitive test of the hypotheses underlying the binomial distribution, because the expected and observed counts are small; only two of 12 observed cells have more than 10 counts, and six observed cells have zero counts. Given the heterogeneity in x demonstrated in both table 1 of Gill et al. (1987) and table 1 of Jeffreys et al. (1985b), the binomial distribution is not the expected distribution of the number of matching bands, under either measure of matching used by Gill et al. (1987).

Returning to the data of Jeffreys et al. (1985b), we find that the assumption that the match probabilities are constant within a size class is not consistent with their data in table 1 under the assumption of independence between fragments. To test the combined assumptions of constant fragment match probabilities within a size class and independence between fragments, observe that, since different individuals are assumed to be unrelated and therefore independent, the presence of a fragment of a particular size in A has no effect on the presence of a fragment of that size in B, so x (as defined by Jeffreys et al. [1985b] and above) equals the probability that a fragment in the size class will be present. From the assumed independence of different fragments, it follows that the number of fragments present per individual should be binomially distributed in a population of unrelated individuals with a probability x of having a fragment present in the size class. Of course, for any particular individual, the presence of a fragment is determined by whether the individual inherited the corresponding region from his or her parents.) If N is the (unknown) total number of alleles or distinguishable bands or fragments in the size class, then the average or mean number of observed fragments per individual in the size class must be Nx and the variance (the square of SD) must be Nx(1–x); these formulas are well-known properties of the binomial distribution (e.g., see Snedecor and Cochran 1980, chap. 7). Therefore the variance Nx(1–x) divided by the mean Nx should approximate 1 – x.

When (a) the sample variance (obtained by squaring the SD given in table 1) is divided by the sample mean for each size class of fragment measured by probe 33.15 in table 1 and (b) the quotient is subtracted from 1, the resulting estimates of x differ substantially from the reported values of x. From largest to smallest fragments, the estimates of x derived from the binomial distribution are .66, .76, and .79; the corresponding reported values are .08, .20, and .27. The discrepancies are similar for the data from probe 33.6. Using an estimate of N that is not available to me, A. J. Jeffreys (personal communication, August 28, 1988) computed the binomially expected SD [Nx(1–x)]½ for probe 33.15 as 1.6, 2.0, and 2.2, compared with the reported SDs of 1.0, 1.1, and 1.2 from largest to smallest fragments, and similarly for probe 33.6; the binomially predicted SDs according to Jeffreys were larger, by approximately 30%–80%, than those reported in all cases in table 1. I conclude that the combined assumptions that x is constant and that fragments are independent are not compatible with the reported data.

This conclusion does not determine whether it is the constancy of x or the independence of fragments (or both) that must be modified. Nonconstancy of x within a size class could explain at least part of the reduction in the reported variance, compared with the predicted variance. Nonindependence of fragments might have orthodox biological origins; for example, "a locus with many alleles ranging exclusively from 6–10 kb (such loci do exist) will show a low band sharing but will always show 2 bands of 6–10 kb (i.e., S.D. for this locus in 6–10 kb range = 0)" (A. J. Jeffreys, personal communication, August 28, 1988).

Without appropriate data, no amount of statistical theory can say whether different fragments are statistically independent or are statistically associated in a population. In principle, log-linear models for multidimensional contingency tables (e.g., see Haberman 1974; Bishop et al. 1975; Fienberg 1980) could be used to analyze blot data in order to determine appropriate models for the possible independence or dependence of DNA fragments. For a given probe and a given class of fragment sizes, suppose that N fragments can be distinguished and matched reliably. Each individual’s blot may be coded by a vector (or list) of N 0’s or 1’s; thus individual A’s blot may be summarized as A = (a1, a2, . . . , aN), where aj = 0 if fragment j is absent from A’s blot and where aj = 1 if fragment j is present in A’s blot; j = 1, 2, . . . , N. The blots of a sample of M individuals may be represented by M such vectors, each containing N 0’s or 1’s. The frequencies of each possible combination of bands may be organized into a contingency table with N dimensions. Each dimension corresponds to one size of fragment, and each dimension is divided into two cells (fragment absent or fragment present). The frequencies can be analyzed for consistency with the assumption of independence.
between fragments. If independence fails, the family of log-linear models provides a hierarchy of alternative descriptions of the data.

The number \(2^N\) of cells in the contingency table becomes enormous for realistic numbers of fragments \(N\) ranging from 20 to 60. Hence alternative approaches are required, such as methods for the analysis of large sparse contingency tables (e.g., see Koehler 1986) or of contingency tables with incompletely classified data (e.g., see Chen and Fienberg 1976), tests for pairwise independence (e.g., see Haber 1986), and methods of reducing the dimension of the table (e.g., see Bishop et al. 1975).

A stepwise approach to the problem of independence of fragments seems reasonable for practical purposes. Depending on the available data, one could begin with tests of pairwise independence for all fragments or of complete independence among small numbers of fragments and then extend to complete independence of larger numbers of fragments if the data permit.

### Calculating the Average Power of a Match

To estimate the average power of a match, Jeffreys et al. (1985b) assume implicitly that every person has the same number of fragments in the given size class. This is false (obviously) because the SDs of \(n\) are positive. The level of variability appears to be lower in cats and dogs than in humans (Jeffreys and Morton 1987, p. 6). I now analyze how ignoring the variability in the number of fragments affects the calculated average power of a match, under the temporary assumptions of (a) constant matching probability within a size class and (b) independent matching.

The conclusion of the following analysis is that the procedure used by Jeffreys et al. (1985b) underestimates the true probability of a match between a randomly chosen person and a given specimen or a given gel, because their procedure uses the geometric mean instead of the arithmetic mean. This is a straightforward mathematical error.

Consider a fixed probe and a fixed size class of fragments identified by that probe. Let \(f_n\) denote the fraction or proportion of all individuals who have exactly \(n\) DNA fragments; \(n = 0, 1, 2, \ldots, N\). (The finite resolution of Southern blots imposes an upper bound \(N\) on the number of possible fragments that can be distinguished within any size class.) Then \(f_0 + f_1 + f_2 + \ldots + f_N = 1\). The average number of fragments, averaging over all individuals in the population, is \(\bar{n}/f_N\), which I denote by \(E(n)\), as is usual. The \(E\) in \(E(n)\) stands for “expected number” or “expectation” of \(n\).

Given a randomly chosen individual \(A\), who may have varying numbers of DNA fragments, and a specimen \(B\), the probability \(P\) that every fragment of \(A\) also belongs to \(B\) (the match probability) is a weighted average of the match probabilities for those individuals \(A\) with each different possible number of fragments. Let \(x\) be the probability that a fragment in any randomly chosen individual \(A\) is also present in \(B\), as before. If \(A\) has exactly \(n\) fragments, the probability of a match is then \(x^n\). This match probability must be weighted by the fraction \(f_n\) of the population that has exactly \(n\) fragments. Thus the probability \(P\) of a match in the population is \(P = f_0x^0 + f_1x^1 + f_2x^2 + f_3x^3 + \ldots + f_Nx^N\). Using the same notation Jeffreys et al. (1985b) calculate the quantity \(Q = x^{0f_0}x^{1f_1}x^{2f_2}x^{3f_3} \cdots x^{Nf_N} = x^{E(n)}\). The inequality of arithmetic and geometric means guarantees that \(Q < P\) as long as there is at least some variation in the number of fragments per person, i.e., so long as there is no \(n\) with \(f_n = 1\). (If there is an \(n\) with \(f_n = 1\), then \(Q = P\).)

To get a quantitative idea of the effect of incorrectly using the geometric mean \(Q\) instead of the arithmetic mean \(P\), I invented several artificial frequency distributions \(f_0, f_1, \ldots, f_N\) to match the means and SDs given in table 1 for probe 33.15. These invented data are presented in table 2, along with their means, SDs, and other statistics. For example, to simulate the distribution of fragment numbers in the size class from 10 to 20 kb, suppose that among every 40 people there were exactly one with \(n = 0\) fragments, one with \(n = 1\) fragment, 11 with \(n = 2\) fragments, 15 with \(n = 3\) fragments, 11 with \(n = 4\) fragments, and one with \(n = 5\) fragments. Thus \(f_0 = 1/40, f_1 = 1/40, f_2 = 11/40, \ldots, f_N = 1/40\). The mean number of fragments would be \(E(n) = 2.929\) and the SD of the number of fragments would be 1.0, which are close to the observed mean and SD (table 1). Taking \(x = .08\) from table 1, I find \(Q = .08^{2.929} = 6.1 \times 10^{-4}\) and \(P = .029\). A second set of artificial data (table 2), with \(E(n) = 2.925\) and SD 0.9710, gives \(Q\) as before and \(P = .0061\). The difference between \(P\) and \(Q\) is at least an order of magnitude, for these two sets of artificial data.

Using the second of these two sets of artificial data (in order to give a low rather than high estimate of the difference between \(P\) and \(Q\)) for the 10–20-kb class and using other sets of artificial data for the 6–10-kb and 4–6-kb size ranges (table 2), together with the assumptions of independence already mentioned, gives an overall match probability, for probe 33.15, of \(1 \times 10^{-7}\),
which is substantially larger than the estimate of $3 \times 10^{-11}$ given by Jeffreys et al. (1985b).

The reported "probability that the same pattern will occur in a randomly chosen individual," in tables 1 and 3 of Gill et al. (1987), has the same problems as $Q$ in the work of Jeffreys et al. (1985b).

In response to these hypothetical calculations, A. J. Jeffreys (personal communication, August 28, 1988) computed $P$ by using the original data of Jeffreys et al. (1985b). Assuming independence between probes, he obtains (table 3) an overall match probability of $8.4 \times 10^{-17}$, approximately four orders of magnitude larger than the original estimate of $3.4 \times 10^{-21}$ (though still a very small probability). Jeffreys correctly observed that $P$ is heavily influenced by the proportions $f_0$ and $f_1$ of individuals with no or only one fragment or band.

Miscalculation of the match probability by using the geometric mean rather than the arithmetic mean can make a substantial difference—in this case, four orders of magnitude. However, this criticism applies only when the geometric mean is used to summarize the effectiveness of DNA fingerprinting for a randomly chosen person, as Jeffreys (personal communication, August 28, 1988) has pointed out. When a specific person is the subject of matching, the number of fragments is fixed, not a random variable, and then the same answer is obtained by either the geometric or the arithmetic mean. However, it seems doubtful that, in forensic applications, a specific person should be considered as the only

### Table 2

**Artificial Frequency Distribution of the Number of DNA Fragments/Individual, Chosen to Match the Mean and SD Reported for the Corresponding Size Classes in Table 1, for Probe 33.15**

<table>
<thead>
<tr>
<th>No. of Fragments</th>
<th>10–20-kb</th>
<th>10–20-kb</th>
<th>6–10-kb</th>
<th>4–6-kb</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sample 1</td>
<td>100</td>
<td>120</td>
<td>10</td>
<td>10</td>
</tr>
<tr>
<td>Sample 2</td>
<td>120</td>
<td>100</td>
<td>10</td>
<td>10</td>
</tr>
</tbody>
</table>

Mean no. of fragments: 2.9
SD of no. of fragments: 1.0

### Table 3

**Geometric Mean Probability ($Q$) and Arithmetic Mean Probability ($P$) of a Match When Using the Original Data of Jeffreys et al. (1985b), According to Calculations of A. J. Jeffreys**

<table>
<thead>
<tr>
<th>Probe</th>
<th>$P$</th>
<th>$Q$</th>
</tr>
</thead>
<tbody>
<tr>
<td>33.6</td>
<td>$6 \times 10^{-9}$</td>
<td>$1.8 \times 10^{-10}$</td>
</tr>
<tr>
<td>33.15</td>
<td>$1.4 \times 10^{-8}$</td>
<td>$3 \times 10^{-11}$</td>
</tr>
<tr>
<td>Both probes jointly, when independence between them is assumed</td>
<td>$8.4 \times 10^{-17}$</td>
<td>$5.4 \times 10^{-21}$</td>
</tr>
</tbody>
</table>

Source.—A. J. Jeffreys (personal communication, August 28, 1988).
possible candidate for matching. If a given suspect is exonerated because his DNA fingerprint does not match that of a specimen, the police will not, in general, simply stop looking for the criminal but will seek another person; when they do seek another suspect, the number of fragments in the person being compared to the specimen is indeed a random variable, and the arithmetic mean, not the geometric mean, is appropriate.

**Conclusion**

Scientific data and statistical analyses play increasing roles in the courtroom (DeGroot et al. 1986; Black 1988; Marx 1988). Increasingly, judges are requiring that the details of the evidence and the analysis be explicit and well founded (Black 1988). It is the responsibility of scientists and statisticians to provide measurements, analyses, and conclusions that justify lawyers' faith in "relatively simple and well-defined techniques like electrophoresis or polygraph lie detection" (Black 1988, p. 1509). When such faith is not justified, it is scientists' responsibility to provide clear warning labels to the contrary. Since human lives and liberty are at stake in uses of DNA fingerprinting for forensic identification, it is important that there be little room for doubt about the assumptions underlying the analysis and interpretation of DNA fingerprinting data, including their statistical analysis and statistical interpretation.

The difficulty in establishing the statistical basis of DNA fingerprinting for forensic identification lies in assuring that the assumptions implicit in the calculations are justified by evidence or theory and that any simplifying approximations made give conservative estimates (i.e., overstatements) of match probabilities.

Some methods of statistical analysis of data on DNA fingerprinting suffer serious weaknesses. Unlinked Mendelizing loci that are at linkage equilibrium in subpopulations may be statistically associated, not statistically independent, in the population as a whole if there is heterogeneity in gene frequencies between subpopulations. In the populations where DNA fingerprinting is used for forensic applications, the assumption that DNA fragments occur statistically independently for different probes, different loci, or different fragment size classes lacks supporting data so far; there is some contrary evidence. Statistical association of alleles may cause estimates based on the assumption of statistical independence to underestimate the true matching probabilities by many orders of magnitude. The assumptions that DNA fragments occur independently and with constant frequency within a size class appear to be contradicted by the available data on the mean and variance of the number of fragments per person. The mistaken use of the geometric mean instead of the arithmetic mean to compute the probability that every DNA fragment of a randomly chosen person is present among the DNA fragments of a specimen may substantially underestimate the probability of a match between blots, even if other assumptions involved in the calculations are taken as correct. The conclusion is that some astronomically small probabilities of matching by chance, which have been claimed in forensic applications of DNA fingerprinting, presently lack substantial empirical and theoretical support.

Many of the above issues apply to paternity testing through DNA fingerprinting as well as to forensic identification of unrelated individuals. Because of the possible relatedness of individuals in paternity testing, the genetic formulas are more complicated than the formulas used in identifying unrelated individuals. However, most of the underlying hypotheses are the same, and most of the same caveats apply.

Future experiments and analyses could provide a firmer foundation for DNA fingerprinting by giving careful attention to both sampling and possible statistical dependence among fragments. DNA fingerprinting can be the basis of a useful method of identifying individuals, provided that claims for it are not exaggerated.

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**References**