

**PAPER****CRIMINALISTICS**

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The Probabilistic Genotyping Software STRmix: Utility and Evidence for its Validity*

ABSTRACT: Forensic DNA interpretation is transitioning from manual interpretation based usually on binary decision-making toward computer-based systems that model the probability of the profile given different explanations for it, termed probabilistic genotyping (PG). Decision-making by laboratories to implement probability-based interpretation should be based on scientific principles for validity and information that supports its utility, such as criteria to support admissibility. The principles behind STRmix™ are outlined in this study and include standard mathematics and modeling of peak heights and variability in those heights. All PG methods generate a likelihood ratio (LR) and require the formulation of propositions. Principles underpinning formulations of propositions include the identification of reasonably assumed contributors. Substantial data have been produced that support precision, error rate, and reliability of PG, and in particular, STRmix™. A current issue is access to the code and quality processes used while coding. There are substantial data that describe the performance, strengths, and limitations of STRmix™, one of the available PG software.

KEYWORDS: forensic science, DNA, probabilistic genotyping, validation, STRmix™

A common binary method, dating to the 1990s, for the interpretation of short tandem repeat (STR) typing results from forensic casework analyses, while valid (1–3), had two primary limitations: First, for some specimens, a substantial amount of profile data could not be used for calculating statistical weight,

resulting in more inconclusive results; and second, a number of laboratories faced challenges in interpretation of complex forensic DNA mixtures.

In recent years, more sophisticated approaches to applying the fundamental principles of DNA mixture interpretation have been incorporated into customized software that expands the capabilities of the forensic analyst. These tools bring together refined methods of biological modeling, probability, and computational power that provide more meaningful empirical assignments of evidentiary weight.

Substantial data have been generated and accumulated that demonstrate the utility of probabilistic genotyping (PG). As exemplified herein with one software solution for PG, STRmix™, this document focuses on sound practices for forensic DNA mixture interpretation, the attendant statistical analyses using STRmix™, and other application issues related to admissibility. The topics covered in this effort include:

- An introduction to PG, the likelihood ratio (LR), and setting propositions,
- Validity of STRmix™,
- A discussion about admissibility, peer-review, code disclosure, and independent testing of STRmix™,
- A discussion on assigning the number of contributors, and
- The effect of relatives in mixtures.

This resource should guide the reader in becoming familiar with the salient features of STRmix™, as well as its strengths and limitations.

Introduction to Probabilistic Genotyping

The interpretation of forensic DNA mixture evidence is moving toward PG. The Scientific Working Group on DNA Analysis

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Methods (SWGDM) defines PG as “the use of biological modeling, statistical theory, computer algorithms, and probability distributions to calculate likelihood ratios (LRs) and/or infer genotypes for the DNA typing results of forensic samples...” (4). Biological modeling is based on numerical criteria that can be encoded into the software to aid in interpretation of DNA profile characteristics such as peak height, base pair size, stutter, DNA degradation, allele dropout, and drop-in. The conceptual basis for PG was in place by 2000 (5–9). Advances to this initial concept were made and encoded in the software LoComatioN (10). Workable PG solutions were not implemented into routine forensic casework until about 2009 following advancements and the development of other programs, such as TrueAllele® (11).

Broadly, there are two categories of PG software: semi-continuous and fully continuous. The key difference between them is that semi-continuous models do not consider allele peak heights, while fully continuous methods make direct use of such information. Both the semi- and fully continuous methods assess the probability of observing the mixed DNA profile given proposed genotypes for the contributors. The semi-continuous methods assign a probability to the profile given a genotype combination, and the number is in the continuous interval [0,1]. The mathematical details are not described herein, but all the PG solutions utilize some form of “nuisance parameter” for a factor that must be accounted for in the process. In most semi-continuous applications, this parameter is the assignment of the probability of allele dropout. The semi-quantitative model by Slooten (12) in the software product MixKin removes the nuisance parameter by the preferred method of integration. The programs LRmix (13), LikeLTD (14), or Lab Retriever (15) use plug-in values or the value derived by the method of maximum likelihood estimation (MLE) rather than from the integral. In the case of LRmix, an allele dropout value is assigned, often following a sensitivity analysis; LikeLTD assigns the nuisance parameter by MLE; and in the case of Lab Retriever, this value is assigned using a form of logistic regression that does not account for degradation (see 16; for a review of some logistic regression methods). Lab Retriever contains an additional approximation to a population genetic model introduced for computational convenience (17). This approximation is unlikely to have any large effect.

STRmix™, TrueAllele® (11), and GenoForm Mixture 3 (18) are fully continuous methods that are based on a Markov chain Monte Carlo (MCMC) resampling method (19). The use of MCMC is not novel and has been used to solve many complex problems within chemistry, physics, biology, statistics, and computer science. The continuous model software *Kongoh* (20) utilizes MLE. Other continuous solutions of which we are aware include LikeLTD-ht (21), DNAmixtures (22,23), and EuroForMix (24).

Introduction to Likelihood Ratios (LRs)

The outputs of all PG software are *LRs*. The *LR* is a ratio of the probability, the probability density, or quantities proportional to either probability or density of some specific observations or findings when considering two alternative (i.e., mutually exclusive) propositions. As applied to forensic DNA typing, the ratio, in its simplest form (i.e., a single-source specimen), expresses the probability of the DNA evidence if a person of interest (POI) rather than an unknown individual is the source of the DNA.

Bayes’ theorem follows immediately from the laws of probability and in the current context may be expressed in the following form: posterior odds = $LR \times$ prior odds.

Whatever the odds are on the person of interest (POI) being a contributor without considering the DNA evidence (i.e., the prior odds), this theorem describes that these odds should be increased (or decreased) by *LR* times upon considering the DNA evidence. In practice, it is the *LR* rather than posterior odds that are typically presented in court.

Naming the Propositions

Ian Evett in “What is the probability this blood came from that person?” (25) recognized the work of Dennis Lindley on probability and Bayesian theory. Evett used the terms *C* (contact) and \bar{C} (noncontact) to describe alternate propositions used for the *LR*. Subsequently, H_p and H_d were introduced as the prosecution and defense hypotheses, respectively (26). The prosecution proposition is usually straightforward (i.e., the defendant is the source of the DNA on the evidence), while the defense proposition can vary substantially. Argument can arise about assertions, such as:

- The expert should not assume what the defense proposition may be,
- The defense is entitled to all propositions consistent with exoneration and should not be constrained to one proposition, and
- The defense is not obligated to provide a proposition.

Using the terms prosecution and defense may contribute to some contention when considering propositions. Therefore, as with the earlier *C* and \bar{C} espoused by Evett, alternate propositions without such descriptors might be sensible. H_p and H_d could readily be replaced with, for example, H_1 and H_2 (15,27), H_1 and H_a (where “*a*” stands for alternate), or H_C and $H_{\bar{C}}$ (referring to contributor and noncontributor), to avoid the implications of the “*p*” and “*d*” labels.

The two propositions used for the *LR* must be exclusive, meaning that they cannot both be true at the same time. They should also be exhaustive (cannot both be false at the same time) within the context of the specific case. For example, consider that the prosecution asserts that the defendant is the source of the DNA, and the alternate proposition is that the source of DNA is a random person unrelated to the defendant. In this case, H_p and H_d may both be false, for example, the DNA could be from the defendant’s brother, in which case these propositions are not exhaustive. However, propositions should always be considered in light of the accepted background information (*I*), and if a proposition is very unlikely or impossible given *I*, then it need not be considered by an analyst. In the context of the case discussed above in which *I* is that the defendant has no brother, this possibility need not be considered. We direct the reader to Biedermann et al. (28) for a more in-depth discussion on setting propositions.

Transfer and Persistence of DNA

The concept of hierarchy of propositions is well established (28,29). Gittelson et al. (30) discussed this concept more recently. Propositions are classified into four levels: offence, activity, source, and subsurface.

- Offence level propositions describe the issue for the fact finder which is one of guilt or innocence.
- Activity level propositions describe the activity that deposited the DNA.
- Source level refers to the origin of the body fluid or cell type examined.
- Subsurface level refers to the origin of the DNA (i.e., donor).

Gittelsohn et al. (30) suggested a requirement for interpretation:

Due attention must be paid to the position in the hierarchy of propositions that can be considered. This information must be effectively conveyed to the court to avoid the risk that an evaluation at one level is translated uncritically and without modification to evaluation at a higher level.

They further stated:

We cannot emphasise the importance of this enough. A DNA match may inform decisions about the source of the DNA, but decisions about an activity, say sexual intercourse vs. social contacts, involve additional considerations beyond the DNA profile.

Transfer and persistence of DNA are relevant for an activity level evaluation of the DNA results, whereas PG software and other interpretation and statistics methods evaluate the DNA results at the subsample level. Discussion of transfer and persistence therefore has nothing to do with PG. However, a subsample level evaluation of the DNA results may be necessary for evaluating the findings with regard to a pair of activity level propositions.

Effect of Different Propositions When Using STRmix™

The assignment of propositions should be made from the relevant background information (30,31). The following issues should be considered:

- Which, if any, of the known individuals may be reasonably assumed to be contributors (30–32),
- The number of contributors to the profile (discussed later in this paper) (33–37),
- How to deal with multiple POIs (30,31),
- How to deal with evidential items associated with neither the POI nor the victim (30,31).

The effects of these considerations are summarized here.

Assumption of the Presence of an Individual's DNA in a Mixture

If a genotyped person, say the complainant in a sexual assault, can reasonably be expected to have donated DNA to the sample and the profile suggests his or her presence, then that person should be included under both the prosecution and defense propositions.

There are three principles that could be applied when making this decision.

First, any person should be assumed to be a contributor if the presence of his or her DNA is reasonably expected and the mixture is explained well by his or her inclusion. One reasonable expectation of the person's DNA being present (e.g., Option 1 in Table 1) is if the item of evidence is derived from an intimate sample of this person such as a vaginal swab. This concept can reasonably be extended to other items associated with the person, for example, their clothing. Accordingly, it is important that any such assumption of the presence of the person's DNA be stated/documented.

Second, the contributor proposition should align with the scientific explanation of the evidence informed by any legitimate background information.

Third, reasonable alternate propositions consistent with non-contribution should be considered. For example, in a mixed

TABLE 1—Various options for the propositions H_1 and H_2 . V is for victim, P is the person of interest, and U is an unknown person.

H_1	H_2	
$V + P$	$V + U$	Option 1
	$U + U$	Option 2
	$P + U$	Option 3
$U + P$	$U + U$	Option 4
$U + V$		Option 5

DNA profile, it may be in the interests of the defense to include any person's DNA under both H_1 and H_2 as long as this inclusion is consistent with his or her own noncontribution.

How to Deal with Multiple POIs

Consider a situation where there are two POIs termed P_1 and P_2 . A crime stain is found, and the DNA mixture profile can be explained fully if P_1 and P_2 are the contributors (Table 2). For demonstration purposes, a two-person mixture is assumed (but the three approaches described here can extend to higher order mixtures). U stands for an unknown individual, usually considered to be unrelated to either P_1 or P_2 , although in STRmix™, this assumption may be relaxed to a relative in most, but not all, of the situations illustrated below. Generally, either Approach 1 or 2 (Table 2) is acceptable. Approach 2 has slightly more power to distinguish contributors from non-contributors. It should be used when it aligns with the prosecution allegation.

Approach 3 runs the risk of a major contributor with a high LR (if analyzed separately) "carrying" a noncontributor or a weak/trace contributor with a low LR (if analyzed separately) into the final high LR for $P_1 + P_2$ that could be misleading if reported. This approach should be predicated on separate tests for P_1 and P_2 which both return $LRs > 1$. Approach 3 only should be used in the unlikely event that background information determines that the DNA must originate from both P_1 and P_2 or neither of them.

How to Deal with Evidential Items not Demonstrably Associated (Before DNA Testing) with Either the POI or the Victim

As an example, consider a situation in which a two-person DNA mixture was recovered from somewhere not particularly closely associated with the victim, such as a stain on a bedsheet in a room at a house where a party occurred. The alleged victim, V , states that she was raped in this room by the accused. The stain can be explained as a mixture of the victim and the person of interest, P .

Initially, five options for sets of propositions may be considered (Table 1). The contributor hypothesis may pose H_1 as $V + P$. The noncontributor hypothesis could be Option 1, 2, or 3 for H_2 (or any pair or all three of these). Note that Option 1 is almost always more favorable to the defendant (i.e., a lower LR) than Option 2.

Option 3 suggests the presence of P but not V . It may be difficult for the defense to motivate this option in the context of the case. This option requires that the DNA is from the person of interest and another individual. This proposition asks for a rejection of the victim's statement and an explanation of P 's DNA in the very room where the rape is alleged to have occurred.

TABLE 2—Three approaches to assigning propositions when there are multiple POIs. Propositions are H_1 and H_2 . P_1 and P_2 are POIs, and U is an unknown person.

Approach 1			Approach 2			Approach 3		
H_1	H_2		H_1	H_2		H_1	H_2	
$P_1 + U$ $P_2 + U$	$U + U$	The results are given in the report	$P_1 + U$ $P_2 + U$	$U + U$	The results are in the notes and not the report. They are used in an exploratory manner to inform the inclusion of P_1 and P_2 separately before testing them both together.	$P_1 + U$ $P_2 + U$	$U + U$	The results are in the notes and not the report. They are used in an exploratory manner to inform the inclusion of P_1 and P_2 separately before testing them both together.
$P_1 + P_2$		The result is in the notes and not the report. It is used to check that both P_1 and P_2 may be included together.	$P_1 + P_2$	$P_1 + U$ $P_2 + U$	The results are given in the report	$P_1 + P_2$		The results are given in the report

This leaves H_2 as either $V + U$ or $U + U$ (whichever is considered to be the most reasonable given the case's circumstances). Note that the option $U + U$ will likely lead to higher LR s than the other options. Using the Option 2 set, one could misleadingly produce a very high LR from a major aligned with V and a small scatter of trace alleles consistent with P . As $V + U$ is consistent with noncontribution and in the defendant's interest, we suggest that it should be used.

This approach would lead us to suggest Option 1, in which we have assumed the presence of the complainant, V . This would seem to run contrary to the SWGDAM (32) suggestion that people only be used as conditioning genotypes if their DNA is reasonably expected. One could easily state that no reasonable expectation exists as the item is not strongly associated with either P or V . However, the appearance of V under H_1 comes about because that is indeed the contributor allegation, and under H_2 because it is consistent with noncontribution and is in the defense's interests.

Options 4 or 5 may be used as exploratory investigations to check the inclusion of V and P separately before proceeding to the set actually used for the interpretation, which would be both V and P for H_1 .

In very complex situations, a "search strategy" (akin to proposition sets 4 and 5) may be the only recourse available to the scientist. We note that this approach is suitable in an investigative framework, and some check of the potential for co-contribution should be made. An approach has been proposed for these types of cases in section 4 of Buckleton et al. (31).

Applicability of Probabilistic Genotyping to Forensic DNA Typing Results and Usage Matters

As the use of any tool or technique must be supported both scientifically and for admissibility, we provide some information about PG, and in particular, STRmix™, to support its application for interpretation of DNA profiles derived from forensic evidence.

The General Acceptance Test

General acceptance of the method is one of the Daubert admissibility criteria (38) and is the primary criterion of the Frye standard (39). Most PG software applications are based on established mathematical principles. For example, the MCMC algorithm is not novel (Markov published the first of his papers on this topic in 1906). Other components of MCMC were developed in the middle of the twentieth century; "Monte Carlo

methods were born in Los Alamos, New Mexico during World War II, eventually resulting in the Metropolis algorithm in the early 1950s. . . MCMC was brought closer to statistical practicality by the work of Hastings in the 1970s." (40).

MCMC is a widely used technique and is considered a mainstream statistical tool. It is used in real estate market prediction (41), earthquake and rock fracturing (42), electricity capacity modeling (43), weather prediction (44), betting (45), climate (46), computational biology (47), computational linguistics (48), genetics (49), engineering (50), physics (51), aeronautics (52), stock market prediction (53), and social science (54). The key papers describing the algorithms used within the MCMC are Metropolis et al. (55), with 37,506 cites in Google Scholar (as at May 27, 2018), and Hastings (56), with 12,229 cites providing some measure of their widespread acceptance and use. Searching scientific literature for "Markov chain Monte Carlo" returns more than 512,000 records.

There is substantial interest in PG as evidenced by the number of modern PG software programs that have been developed, or are being developed, by researchers with very strong mathematical or statistical backgrounds (11,13,19,22,57–62). These efforts and recommendations indicate strong support for the general acceptance of PG. While different PG software programs tend to differ in some details, there is a very substantial commonality of principle between these software tools (i.e., they all produce LR s and all model the probability or probability density of the profile given all plausible genotypes). The differences among programs do not indicate a lack of consensus on general acceptance of PG as an analytical/statistical method or of individual software programs. These efforts are strong support for the general acceptance of PG. Both SWGDAM (4) and ISFG (63,64) give recommendations for validation for laboratories that adopt PG for mixture interpretation.

At preparation of this manuscript, STRmix™ is in use in at least 51 laboratories worldwide as their predominant method for the interpretation of DNA profiles in forensic casework. The laboratories using STRmix™ reside in the United States ($n = 36$), Australia ($n = 7$), England ($n = 2$), Scotland ($n = 1$), Republic of Ireland ($n = 1$), Canada ($n = 2$), Finland ($n = 1$), and New Zealand ($n = 1$).

Peer-review, Independent Testing, and Further General Acceptance

Peer-review is another criterion of the Daubert standard that may be considered by the gate keeper. Oxford (<https://en.oxf>

ordictionaries.com/definition/peer_review) defines peer-review as: “Evaluation of scientific, academic, or professional work by others working in the same field.” The scientific concepts that underpin PG software are verified by publication in peer-reviewed journals and wider comments published elsewhere (11,13,16–19,21–23,30,33,34,36,59,61,65–95).

Additional peer-review has been achieved for many PG solutions by those laboratories that performed internal validation studies (11,81,96). Such internal validation studies typically are not published, as journals tend not to find such studies novel. These data, however, are available for review, if desired by the courts, as part of the formal discovery process, and some are available online (97–100).

PG and the PCAST Report

Recently, the President’s Council of Advisors on Science and Technology (PCAST) proffered criticisms about the foundational validity of some forensic disciplines (101). While the 2016 PCAST Report favored the use of PG for forensic DNA mixture interpretation, PCAST neither evaluated the current state-of-knowledge regarding PG at the time of its review nor up to disbandment of the Council in 2017. PCAST considered validity proven for the use of PG for up to three-person DNA mixtures where the minor contributor is greater than 20% of the mixture (amended to the POI being 20% in the Report addendum) and for two-person mixtures where the minor profile is greater than 10%. If taken literally, according to PCAST one cannot reliably interpret mixtures—to include minor as well as major contributors—where the minor contribution is below 10% (footnote 216 of the original report). This statement relative to the major contributor is obviously unfounded. It is likely that PCAST was referring to assessment of the minor and assumed the major could be analyzed. PCAST also incorrectly perceived gaps in proof of validity with high contributor numbers and mixture contributions <20%.

The PCAST Report assessed proof of validity by empirical studies published in the peer-reviewed literature as of 2016 and did not review the totality of available data even at that time. This partial assessment is unfortunate, as publication of all validation studies is difficult as many journals preclude or discourage publication of most internal validation studies and many laboratories do not see it as their role to publish. By taking this limited stand, the PCAST committee members did not avail themselves of the totality of data that was accessible at the time of their review and subsequent report being issued.

Fortunately, validation data exist, peer-reviewed literature is available, and limitations of the applications are described. One example of such work with STRmix™ (81) is summarized in Table 3. This work covers 1–5 person mixtures at much greater

ratios and lower templates than referred to in the PCAST Report.

Recently, the internal validation data from 31 laboratories using or validating STRmix™ were compiled and interpreted (hereafter “internal compilation study” 96) specifically to address the points raised within the PCAST Report. This study concluded that this combined dataset “demonstrates a foundational validity of, at least, the STRmix™ software method for complex, mixed DNA profiles to levels well beyond the complexity and contribution levels suggested by PCAST.” These efforts, representing a substantial resource commitment, are a collation of the validation studies from 31 laboratories and demonstrate that there is support for interpreting a minor contributor much less than 20%, and in fact down to 0% (present but not observed), of the total DNA present in the mixture. As the template tends toward 0, the *LR* tends to approximately 1.

Disclosure of the Algorithms

An issue that has arisen during court proceedings (or during discovery requests) is that there is a need to have access to the source code of PG software to ensure proper peer-review of the validity/reliability of the software. Use of open source software has been advocated (for example, by ISFG <https://www.isfg.org/>) because:

- It allows all parties (including the defense) to have access to software, and
- There is a possibility that review by third parties could improve the code.

Regarding the first point, freeware is accessible without restriction or cost, which may be desirable to some potential users. In contrast, commercially available software comes at a cost but includes continued support and quality control measures. Users need to evaluate these aspects of open source and commercially available software when deciding to implement PG software. Discovery regarding commercial software may involve the code or an executable program but often with cost recovery. STRmix™ comes with a user manual and data on validation associated with the current version. Moreover, STRmix™ code has been and remains available for court purposes under a nondisclosure agreement and supervision, but not to competing software developers. To date, the STRmix™ code has been provided via this process thrice (102).

While code can be made available for legal proceedings, informed empirical testing, which is the basis of validation studies (both developmental and internal), is the best way to assess performance and critically evaluate the results produced by a software tool. Indeed, troubleshooting a PG tool by the developers is typically performed without reference to the source code. STRmix™ has a facility called “extended output” that is

TABLE 3—A summary of the tests undertaken in the internal validation studies by the FBI.

Number of contributors	Input DNA range (per contributor) ng	Contributor ratio range	Number interpreted	Number of true contributors tested	Number of false contributors tested
2	0.006 to 0.9	10:1 to 1:1	105	202	22,504
3	0.021 to 1.0	16:1:1 to 1:1:1	64	192	13,620
4	0.050 to 3.2	16:1:1:1 to 1:1:1:1	84	336	17,808
5	0.016 to 1.25	10:1:1:2:2 to 1:1:1:1:1	24	120	5,256

A selection of 172 one-, two-, and three-person profiles were interpreted as originating from two, three, and four individuals, respectively. The true contributors and 200 noncontributors were tested. These experiments entertain ratios below that espoused as a threshold by the PCAST Report.

available to users as a useful assessment and diagnostic tool. This function outputs the proposed genotypes and other variables and the probability density for each step in the MCMC process. One can then attempt to reproduce these values by independent calculation(s). The code is only accessed to rectify an error, for example, identified during this extended output review, or to modify the program.

In response to the second point, one would expect that free-ware, being more accessible, would enable substantial third-party improvement. There are a few instances to date, however, where programming input to open source PG software has occurred from external parties (e.g., Lab Retriever programmer Kirk Lohmueller made suggestions to the LikeLTD programmers (21) (see pg. 27), while cloning parts of LikeLTD, and David Balding (personal communication) had input from students). All software needs funding for support and development, and noncommercial software may require public funding or donations (see <https://scieg.org/support-our-work/>).

STRmix™ has received input on miscode detection and suggestions for improvement from many sources, such as collaborators, based on empirical testing, or applied usage of the software and/or extended output functions.

Both commercial and open source software should come with additional support beyond an accompanying manual. STRmix™ requires substantial mandatory training provided by experts, as well as making available a user help desk. As one moves to more complex ways of interpreting DNA profiles, proper training is vital to reduce the problems of misuse that may occur. The risks associated with little, improper, or no training and quality control are serious and warrant substantial consideration. This concept was not lost on the Court in a Daubert challenge ruling, stating (103):

As the source code cannot be altered by anyone except the programmers, there is an additional layer of internal controls that govern the STRmix's operation.

Support throughout validation and implementation and participation in a software user group are also critically beneficial to the development and execution of reliable standard operating procedures.

Error Rate

Error and error rate are general scientific issues but also arise in the forensic setting (error rate of the method is one of the Daubert admissibility criteria) (38). Determining the error rate is not always straightforward, largely because error has various meanings; in DNA interpretation, it is determined very much by the sample. With sound quality assurance practices and standard operating procedures derived from judicious validation studies, error may be addressed and reduced.

In the context of forensic cases, concerns surround false associations and false exclusions. A fair justice system would generally favor a false exclusion over a false association. Accordingly, overstating the strength of the evidence when a POI (or victim) cannot be excluded as a potential contributor should be avoided. In forensic science, a tendency to understate the evidential weight is termed conservativeness. The STRmix™ software incorporates key features to drive the *LR* toward a conservative (lower) result.

Most discussions on error rate surround the concepts of a true state and a declared state. For example, one could input

true donors and see if the software outputs a declaration of "true donor." The output of all PG software, however, is not a declaration of true or nondonors but a *LR*, which is a number on a continuous scale. This approach differs from a categorical declaration of inclusion or exclusion in a similar manner to how a probability would differ from a statement of certainty. The result of a true donor indicated as a nondonor could be termed something like evidence supporting H_2 and a false donor indicated as a true donor could be termed something like support for H_1 .

Support for H_1 for a noncontributor is directly related to the strength (or quality) of the DNA profile (76,104). Hence, there will not be one rate of support for H_1 for all DNA profiles examined by PG but a different rate for each sample.

An important distinction to understand is that support for H_1 for a nondonor can be due to the nondonor sharing many alleles with the profile, which is different from an event of "software error." Studies on STRmix™ suggest that the software itself does not contribute to the support for H_1 (76) under this scenario beyond that expected by overlap of the alleles of the nondonor and the profile. Some of the best evidence for this comes from Turing's rule. Turing's rule translated to DNA states that the average *LR* for a large sample of nondonors should be 1. Trials with STRmix™ show the average *LR* to be approximately 1 or, when various levels of conservancy are included, <1 (76,96,104). Considerable research has been undertaken that allows informed statements to be made about the potential uncertainty associated with *LRs* (19,36,67,69). It is very difficult for operator error of the software or false information about a known contributor to cause a false inclusion. There are methods, based on importance sampling (104), that are sufficiently fast and allow massive (up to 10^{30} or even greater) nondonor tests to be run during validation and if needed on a particular case basis. Importance sampling creates a biased sample by drawing from a distribution of importance, in the DNA example, these are profiles likely to produce high *LRs*. As the sample is biased, it is necessary to readjust for the bias after simulation. In a large set of mixtures compiled from 31 laboratories (96), all large (over 10,000) *LRs* for nondonors were investigated; in all instances, the nondonors had high allelic overlap with the profile. This is a correct result. This empirical assessment of nondonor inclusion rate provides additional support on the reliability of STRmix™. To date, there have been no detected instances of a high *LR* using STRmix™ software where there were not also many alleles in common between the donor and the profile.

There are several situations where a false exclusion ($LR < 1$ for a true donor) may occur:

- The contribution of a true donor is low resulting in only a few alleles above the analytical threshold, or for other reasons, the PCR does not generate an optimum profile (i.e., the contributor's alleles are not detected or are poorly represented, or stutter heights appear disproportionately due to stochastic amplification),
- Incorrect typing information for a true contributor(s) is used in the analysis (to include a sample mix-up) (this can be for the tested POI or, for a conditioned analysis, another contributor whose DNA is assumed to be present in a mixture),
- An operator error, notably not removing an artifact before STRmix™ analysis, or
- The number of contributors assigned is too few.

Diagnostics output by STRmix™ align with human judgment and thus allow for a human check of the results (a desirable and recommended feature). In some instances of false exclusions, the

output data may indicate an $LR < 1$ for a single locus, with high LR s at all other loci, suggesting a possible error in the input data. In such situations, or others in which the output from the software and the human operator disagree, the operator should review the input data to determine whether they are correct (e.g., artifacts removed and alleles properly recorded). A retained artifact should be removed or a mislabeled allele corrected, and PG should be carried out again.

Although one might suggest that an error rate should be considered for the software—operator pair, there is in fact no reasonable way to calculate the operator error rate. As the process described here allows for correction, the reported result of such a software—operator pair is not an error for which an appropriate rate could be calculated. Any such corrections should be documented and, together with the output of the software and the original profile(s), are subject to technical review and, if desired, other independent review. Lastly, error rates, at best, are some indirect indication of performance, but are not indicative or predictive of error in a specific case. The important question is whether an error occurred in the case, which can best be addressed by review or by re-calculation, in this situation, of the PG results (105).

Coding Standards and Miscodes

There are no coding standards designed specifically for forensic software. To date, the software has generally been evaluated in systems (i.e., testing the overall process for generating reliable results) and by empirical approaches. Neither the SWGDAM nor ISFG recommendations on validating PG software require accreditation by any software standardization organization, instead seeming to prefer the systems approach for validation. Coding standards in the wider industry, such as the Institute of Electrical and Electronics Engineers (IEEE), might provide guidance when testing the performance of software (106). Third-party assessment, developmental validation studies, internal validation studies, and adherence to gathering community feedback tend to provide more scrutiny than only a single entity assessment.

As all software programs may have coding faults (miscodes), even with the most diligent scrutiny, developers, and maintainers of a software must gather information by continuous testing and from users to identify miscodes. It is important to be transparent and disclose any miscodes discovered that affect the numerical result, so stakeholders are informed and can have confidence that key software is subjected to critical quality review and is continuously improving. Many probabilistic genotyping software such as LRmix Studio (107), Lab Retriever (108), STRmix™ (109), EuroForMix (24), and LikeLTD (21) have disclosed miscodes. The consequences of miscodes should also be investigated and disclosed.

Validation testing selects a broad range of samples but cannot test the myriad possible ways DNA profiles may present in real forensic casework (11,110). However, one can have confidence that a breadth of normal usage is well tested, as exemplified by Bright et al. (96) with more than 2,825 mixtures of DNA from three to six contributors.

Precision of the Output

Inherent in the validation of forensic methods is the assessment of variability. There has been some misuse of terms

TABLE 4—Terms relating to variability.

Repeatability	The degree, within measurement error, to which the same result(s) is obtained for a sample when the assay is repeated by the same operator and/or detection instrument
Reproducibility	The degree, within measurement error, to which the same result(s) is obtained for a sample when the assay is repeated between/among different operators and/or detection instruments
Precision	The degree of mutual agreement among a series of individual measurements, values, and/or results. Precision depends only on the distribution of random errors and does not relate to the true value or specified value
Accuracy	Degree of conformity of a measured quantity to its actual (true) value.
Objective	Little or no judgment required by the analyst
Subjective	Some judgment required by the analyst
Biased	A measurement is systematically above or below the true value

associated with precision. We reprise some definitions in Table 4, following those of (32,111).

It is well known that measurement error is associated with diagnostics, such as DNA typing. For example, the highly reliable generation of STR results using optimum input amounts of DNA is considered repeatable and reproducible. Yet, if a sample was re-amplified or re-injected, no one would expect that precisely the same peak heights would be obtained and that the various peaks of a profile would be in the exact same relative proportions. Precision and accuracy are assessed within some range of measurement that is determined through validation studies. These studies are important for determining the limitations of a methodology, including usage of the STRmix™ software (4).

STRmix™ uses a MCMC resampling method. The MCMC resampling strategy will create run-to-run variability. For STRmix™, the random number generator is run starting from a seed and, in the default setting, that seed is set from the clock. Thus, one should expect a degree of variation in the results. Validation should address the degree of variation as a basis for determining appropriate operating procedures and reporting criteria.

Recognition of variability is a positive aspect of PG, or for that matter any methodology measuring a target of interest. PG results are based on modeling which is the best attempt available at producing a rational and supportable answer that is based on solid mathematics and extensive empirical work. When judgment is exercised in relation to any modeling decision, we favor decisions that tend to reduce the value of the LR , an approach we call conservative. Indeed, introducing a continuous DNA interpretation system helped to better recognize the uncertainty inherent in assigning a LR .

Reliability of PG at Low Template

STRmix™ has now been extensively tested on profiles generated from optimum template levels down to extinction [see in particular (69,75,76,81,96)], as well as across a range of constructed mixture types as encountered in forensic casework with respect to total template amount (i.e., optimal to trace), contributor proportions (i.e., similar to disparate relative contributions within a given mixture), and other features such as allele sharing. The trend is that the LR tends toward 1 for both true donors and nondonors as peak heights of the contributor in question

become lower. This finding is true whether the other contributors are also low or high in average peak height. Trials have been undertaken where the minor contributor is not observable (0%). In such cases, STRmix™ reports a *LR* close to 1, usually between a $\log(LR)$ of -3 to 3 . These results demonstrate that STRmix™ reliably reports that the profile is close to uninformative with respect to whether the POI, at zero template and hence not there, is a contributor or not.

Moretti et al. (81) reported a total of 277 two-, three-, four-, and five person mixtures, prepared using DNA from thirteen contributors with varying individual template amounts (ranging 0.006–3.2 ng) and total template amounts (ranging 0.019–4 ng) tested using STRmix™ (Table 3). Ratios ranged from equal contributions (i.e., 1:1 to 1:1:1:1) to up to a maximum major contribution of approximately 95% (e.g., 20:1), as well as various intermediate proportions. This study also assessed the effect on the *LR* of assigning the number of contributors for STRmix™ analysis as one more than the target number of contributors. The word “target” is used herein to mean the number of contributors input into the mock sample. This usage is differentiated from the “true” number as one or more of the target contributors may be in such a low amount that it is not realistically present at all.

Bright et al. (96) reported a large compilation of the results from 31 different laboratories of internal validation studies of STRmix™. There were 2,825 mixtures generated using eight different STR multiplexes and analyzed on two different types of CE instruments. These mixtures comprised three to six donors, with contributor templates down to extinction and a wide range of mixture proportions. The apparent number of contributors as interpreted by the respective laboratories was assigned as 3, 4, or 5 for PG analysis. Although some trace contributors were not observed in the electropherograms, the assigned *LR*s were appropriate based on the data present for each contributor (i.e., tending to 1 with less information present).

Bright et al. also observed lower *LR*s for true contributors and more *LR*s near 1 for nondonors when the number of assigned

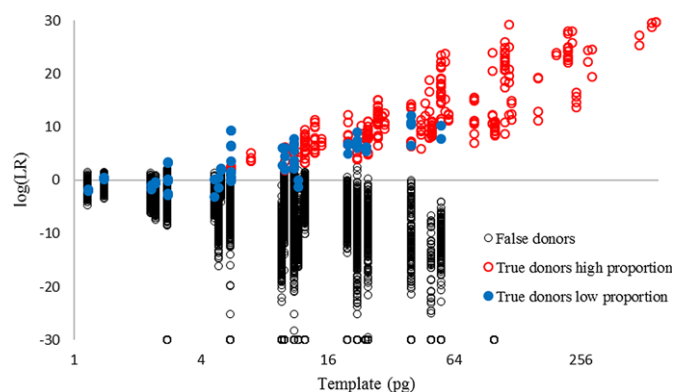


FIG. 1—A plot of $\log_{10}(LR)$ vs. template (pg) for each donor in a four-person mixture, prepared across a range of template amounts and contributor ratios, tested using STRmix™. For the nondonor tests, the template is assigned as the lowest template of the four true donors. For those samples with template above 1 pg, 194 nondonors were tested against the profile. We have also added a fictional contributor, effectively at template 0 pg and tested against 100,000 nondonors. Due to plotting limitations, these samples are represented in this plot at template 0.5 pg. For the true donor tests, the data have been divided into proportion above 10% (high $N = 275$) and those with proportion below 10% and down to 0% (low $N = 72$). As the template diminishes, the *LR*s for both the true and noncontributors tend toward 1 (a $\log(LR)$ of zero is marked with a central horizontal line in the graph). [Color figure can be viewed at wileyonlinelibrary.com]

contributors increased. This finding using STRmix™ also was reported by Taylor (69) and Moretti et al. (81) and demonstrated that mixed DNA profiles containing more contributors are reliably reported as less informative.

These studies also support that the average peak height (APH) of the contributor is a good indicator of information content. It is the low peak heights rather than the extreme ratios that lead to uninformative profiles. For example, testing two low-level contributors with similar APHs (a 1:1 mixture) present more of a challenge to the software than does a 1:20 mixture, as the genotype of the higher contributor has less uncertainty and helps to inform the genotype of the lower contributor. The PCAST position placed significance solely on contributor ratio, ignoring the important component of template amount. Empirical testing (81,96) demonstrates that the positions stated in the PCAST Report are unsupported and the use of complete data should be considered when evaluating performance of PG software. Typical results are shown in Fig. 1. The *LR* tends to approximately, but not exactly, 1 for both true and nondonors as the template is reduced. This is the correct result.

Number of Contributors (NoC)

The number of contributors to a DNA mixture profile from a casework sample is typically unknown and is correctly described as a nuisance variable. A nuisance variable is something needed to do the computation but not available directly from the data. However, depending on the typing results, assigning a NoC to a DNA mixture can range from fairly straightforward to particularly challenging. When the assignment of NoC is more challenging, there are approaches (described herein) available that can provide results that tend to understate the strength of the evidence.

There are two general *LR* approaches in use, often driven by the capabilities of the software. These are as follows:

- Assigning a NoC that is the same under both H_1 and H_2 (termed constrained NoC).
- Allowing the NoC under each proposition to differ (termed unconstrained NoC).

Constrained NoC

Under this process, a NoC is assigned to the profile based on the number of allelic peaks and their heights, often after consideration of artifacts. While most constrained NoC determinations are performed manually, there are software tools available to assist if needed, such as NOCIt (112) based on Monte Carlo methods, PACE (113) based on machine learning, and methods using Bayesian networks (114) and maximum likelihood (115).

Herein, only assignment of NoC by a human operator is discussed (initially drop-in is not considered to simplify the discussion). The DNA profile should be examined and any obvious artifacts such as spikes and pull-up discounted. Peaks in back or forward stutter positions below some preset value derived from internal validation may be considered stutter, or stutter and allelic. If considered solely stutter, such peaks can be discounted in determining NoC. The peaks that remain should be considered potentially allelic. The minimum NoC may be determined based on peak count alone. It is important, then, to consider whether the observed peak heights of the alleles can be supported by this preliminary assignment. If peak imbalances are unrealistic with the preliminary NoC, then at least one additional contributor should be added. After this initial review of the evidentiary

DNA profile, it is desirable to determine which, if any, contributors should be expected under both H_1 and H_2 . This assessment may include, for example, the victim and a consensual partner. The profile of the POI should not be examined at this stage.

Depending on the laboratory's internal validation studies assessing detection sensitivity and allele drop-in, few trace peaks may be discounted for the NoC assignment as potentially being attributed to drop-in. This is usually fewer than a preset number (often only up to two are permitted) and lower than a height established from empirical studies. Although discounted when assigning NoC, these peaks must not be discounted from the analysis of evidential value.

With this general process, a reasonable NoC can be assigned, but there is no guarantee that this estimate is the actual NoC of the sample. Confidence in the assignment varies depending on the complexity of the mixture. Fortunately, any reasonable discrepancy in NoC assignment seems to have a minor effect on the deconvolution or LR (36,67,81,96).

Even for controlled studies, such as those performed during validation, the actual NoC used to generate a mixture (target N) may not be the same as the number *observed* in the mixture (N). Simply stated: low-level contributors may be too low to observe, because a minimum amount of template is needed for any DNA sample to be detectable. Hence, even in mock samples, the number of contributors may not be accurately represented (note that these issues relate to the minor and trace contributors, as major contributors tend to be well represented in mock samples).

The internal validation compilation study (96) described this effect of estimating various NoC to a mixture. Figure 2 (derived from fig. 13 of Bright et al. 96) shows the level of over and underestimation of the apparent NoC (N) determined following the individual laboratories' protocols compared to the target N in their respective studies. Overestimation of N generally led to similar or lower LR s for true contributors. Underestimation of N resulted in exclusions of true contributors, usually affecting the lower/lowest quantity contributor(s).

These data support the view that when assigning N , for false contributors, the risk is overestimation of N , as there is an increase in the number of very low-grade adventitious hits. With respect to the LR for true contributors, when N is either under or overestimated, the result is conservative. Hence, if the LR is large, for example, larger than 1000 and there is uncertainty in N , there is confidence in the LR if N is correct and if not correct,

the LR is more conservative than the already built in buffers [such as the use of a conservative population genetic model (116) and reporting a lower bound on the LR (117)].

Moretti et al. (81) reported the effect on the LR of assuming an incorrect N by both increasing ($N+1$) and decreasing ($N-1$) from the most plausible number. For the $N+1$ tests, 27 total one-, two-, and three-person profiles were interpreted as originating from two, three, and four individuals, respectively. The LR was calculated for both true contributors and 200 noncontributors, which then were converted to lower bounds on the LR . For true contributors (H_1 -true), the majority of lower bounds on the LR s under the assumptions of N and $N+1$ contributors were similar (within one order of magnitude); for 13% of the analyses, the lower bound on the LR decreased by more than one order of magnitude. With regard to noncontributors under the incorrect assumption of an additional contributor, fewer were excluded outright, though overall 94.3% returned lower bounds on the LR <1.

As a means of examining the impact of assuming too few contributors without returning an exclusion outright, Moretti et al. (81) artificially created three mixtures from a two-person mixture (1:5 contributor ratio) by adding a "third" contributor in the range 50–200 rfu, constructed as if it was a child of the two true contributors. The resulting LR s for the major or minor contributor were not affected by the addition of a third contributor at any of the three average peak heights. All noncontributors resulted in exclusions ($LR = 0$). Note that a $LR = 0$ is a practical rounding off, as in theory a LR should not be assigned a value of 0 (81).

Management of the uncertainty of NoC in real casework (usually for complex profiles or low-level contributors) is easily achieved by testing plausible values for NoC. All outcomes of plausible analyses should be retained, and one or a few may be reported.

It may be tempting to revisit the NoC after examination of the POI's profile. For example, it may be possible to sustain the inclusion of the POI by adding a contributor. This approach cannot be entertained with software that uses the constrained NoC approach.

Unconstrained NoC

There is no requirement for the contributor (H_1) and noncontributor (H_2) hypotheses to specify the same number of contributors when calculating LR s. Some PG, for example, LRmix, LikeLTD, and Lab Retriever, can perform calculations with different NoC for each proposition.

Under this approach, when considering NoC under H_1 whether by human judgment or software, the genotype of the POI may be considered. This may lead to the situation where NOC under H_1 is one larger than under H_2 in order to accommodate the POI. We are unaware of any publications addressing the likely effect of this approach.

Software implementing the unconstrained NoC approach can also test an increase in NoC under H_2 while leaving NoC under H_1 at the assigned value. Again we are unaware of a publication outlining the effect of this.

Recently, Slooten and Caliebe (118) published a result that is likely to advance this discussion. If we consider a constrained NoC, then there will be different LR s for each value of the NoC (termed LR_n where n is the NoC). We want the overall LR which we define as the LR where the number of contributors is not known and is treated, correctly, as a nuisance variable.

		Apparent number of contributors				Total samples tested
		3	4	5	6	
Target number of contributors	3	0.98	0.02	0	0	1315
	4	0.23	0.76	0.01	0	1263
	5	0.06	0.58	0.36	0	182
	6	0.03	0.69	0.28	0	65

FIG. 2—Heat map of the fraction of prepared DNA mixtures as interpreted with various differences between the apparent number of contributors (NoC) and the target number (target NoC). The higher numbers are blue and the lower numbers red. [Color figure can be viewed at wileyonlinelibrary.com]

Slooten and Caliebe show that the overall LR is the weighted average of the LR_n values under one reasonable assumption that we discuss later. This is a useful finding.

In their simplest solution (they offer several), the weights for the weighted average are $\Pr(N = n|G_C, G_P, H_2)$ where G_C is the profile of the crime stain and G_P is the profile of the POI. As we consider H_2 , G_P can be removed from the conditioning yielding $\Pr(N = n|G_C, H_2)$. It is likely that only a few values of n need to be considered, maybe often only one or two.

The assumption that leads to this result is that n is equally likely under H_1 and H_2 , specifically $\Pr(N = n|H_1) = \Pr(N = n|H_2)$. Note that the conditioning does not contain G_C or G_S and hence is informed only by whether or not the POI is a donor. This assumption is likely to be true or approximately true in the vast majority of cases.

Subjectivity

Some have suggested that subjectivity equates with bias with attendant negative outcomes. Instead, subjectivity does not automatically imply a bias that will result in an error, and objectivity does not automatically imply an absence of bias that will render an interpretation free from error. The forensic community has begun to appreciate the risks of contextual and conformational bias and is attempting to address these risks and their potential negative effects in a number of ways (see 119 for a discussion). As an example, the use of suspect-driven bias is clearly an unacceptable practice for deciding which loci in a mixture profile may exhibit allele dropout (2,120).

However, the stark view that subjectivity equates to committing error belies the current thinking about cognitive bias. As Jeanguenat et al. (119) recently reminded readers:

Cognitive contamination or bias is inherent in all human beings due to the architecture and operation of the brain. However, it is important to understand that although bias exists it does not always result in an incorrect interpretation, just as enacting bias reduction steps will not guarantee that laboratory results will be error free. Nevertheless, forensic scientists should continue to improve and seek mechanisms to minimize error due to bias.

The idea that subjectivity will inevitably lead to unfair outcomes is incorrect. Indeed, there are methods that have been invoked with intentional bias. For example, some practitioners have generally set stochastic thresholds (STs) rather high to reduce the chances of declaring a false match to a reference sample, at the expense of producing more inconclusive results. SWGDAM defines the ST as “the value above which it is reasonable to assume that allelic dropout has not occurred within a single-source sample” (121). While such a practice does not make use of some potentially useful data, it is “biased” to avoid a more egregious error (i.e., a false inclusion).

The best approach to control the potential negative effects of bias is by providing proper training and education to DNA analysts using these software tools. All humans are susceptible to various biases, and forensic scientists are no exception (119). The more cognizant individuals are to the risks of bias, the better they will be able to develop strategies and procedures to minimize their effects. Thinking that one can overcome bias by force of will is a dangerous misconception.

Many PG programs are designed intentionally to yield a lower bound on the LR that factors in several elements of

uncertainty and thereby “biases” the analysis toward conservatism (i.e., reduced statistical weight). For example, in STRmix™, the highest posterior density (HPD) LR , which if enabled, accounts for the expected amount of run-to-run variation from the Monte Carlo effect and variation in allele probabilities (122). In addition, most PG programs use the population genetic model of Balding and Nichols (123) or close variations. This model has been shown to result in LR s that are conservative (116,124–126) and is used in TrueAllele®, LRmix, STRmix™, EuroForMix, and LikeLTD. A close approximation is used in Lab Retriever. Another key parameter in the population genetic model is the coancestry coefficient θ , which is typically set in a way that tends toward lowering the LR result. It is either set toward the upper end of the plausible range, or a distribution is used based on a diverse set of populations (34). LRmix and LikeLTD utilize the size bias correction described by Balding (127) which produces conservative assignments on average. Balding (128) described a way to combine the contribution of various relatives and unrelated people, which is implemented in STRmix™ (72) and can be set to give an allowance regarding relatedness. Together or separately, all these features are biased toward cautious statements of evidential weight in relation to the contributor proposition. Software has been proposed as a way to eliminate bias by promoting that it is completely objective. This assertion is problematic in that it does not appreciate the effects of bias and dissuades one from embracing the need to consider bias. No one is impervious to bias including those that develop software. While software allows for better repeatability and reproducibility, one should be cognizant that those who develop the software have inserted their own ideas about how the program works best, such as what aspects to weight more so than others. Users of PG software should be informed about the limitations of the software and potential pitfalls and not rely exclusively on software output. Users must apply their training and expertise in DNA profile interpretation and evaluate the software output by visually comparing the PG results with the original DNA profile. For most profiles, the PG results can be assessed to be intuitively correct, or not, by a properly trained DNA analyst. If one were to rely solely and blindly on the PG output, error could occur. For example, a false exclusion of a POI due to incomplete resolution of a TH01 minor 10 allele from the major contributor’s 9.3 allele was shown by Moretti et al. (81) This “false exclusion” was a limitation of CE instrument resolution and not due to the PG software. However, Moretti et al. (81) overcame the limitation with an accompanying manual assessment.

To summarize the issues of bias:

- Bias is an inherent characteristic of human beings.
- Training about cognitive bias is an important aspect of good science, and particularly so for forensic science.
- Software can have inherent bias, some of which is desirable.
- Some PG developers and many users are trained on aspects where bias can impact the decision process.
- Additional review beyond relying solely on software output is recommended as an additional layer to reduce the effects of bias.

PG software removes some decisions or gives substantial support to these decisions. Remaining aspects of subjectivity in STRmix™ include some artifact management of spikes and pull-up at the initial analysis of the electropherogram and an assignment of an exact or approximate number of contributors.

Recently, an interlaboratory study was published (129) that reports the results of 15 different laboratories using LRmix on the same profile. This profile had been constructed from donors with features deliberately chosen to make interpretation difficult (personal communication, MC Márquez). The *LRs* reported vary between 2.6×10^3 and 3.2×10^{14} . This range drew attention in court and suggested some fault in PG. However, the paper clearly describes that the variation arises not from the software but from subjective decisions regarding allele and stutter determinations (see fig. 1 from 129). This finding supports the use of models for stutter as in STRmix™ rather than relying solely on subjective human decisions and may also indicate a need for training that should accompany use of software.

Conclusion

PG had been in gestation for some time from before 2000 (5). The first forensic DNA case of which we are aware that utilized PG methods occurred with TrueAllele® in 2009. Large-scale deployment of PG software, and STRmix™, in particular, to forensic laboratories began in 2012. The efforts to bring PG to fruition, including the initial theoretical development for human identification applications based on STR typing (5–8), span almost two decades, and thus its use today should not be misconstrued as some sudden novel technology. To the contrary, with the maturation of STR typing technologies, great strides in the application of probabilistic solutions to biological phenomena, and the development of several software options, the “coming of age” of PG has been recognized by forensic laboratories. Facilitated by guidance documents from SWGDAM for validating PG systems in 2015 (4), followed by the European Network of Forensic Science Institutes (ENFSI) in 2017 (130), empirical studies performed by many have demonstrated the utility and reliability of PG for mixture analysis and enabled the implementation of reliable procedures for the application of this technology to forensic casework.

Our goal in this study was to provide and address information and issues that relate to PG in general and STRmix™ in particular through our own experience. Our intent is to be informative. By understanding the strengths and limitations of any PG software, users and stakeholders will better understand the system and hopefully use it in a thoughtful manner for the public good.

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