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Identification and Separation of DNA Mixtures using Peak Area Information

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November 26, 2004

Abstract

We show how probabilistic expert systems can be used to analyse forensic identification problems involving DNA mixture traces using quantitative peak area information. Peak area is modelled with conditional Gaussian distributions. The expert system can be used for ascertaining whether individuals, whose profiles have been measured, have contributed to the mixture, but also to predict DNA profiles of unknown contributors by separating the mixture into its individual components. The potential of our methodology is illustrated on case data examples and compared with alternative approaches. The advantages are that identification and separation issues can be handled in a unified way within a single network model and the uncertainty associated with the analysis is quantified.

Some key words and phrases: Bayesian network, conditional Gaussian distributions, DNA mixture, DNA profile, forensic identification, mixture separation, probabilistic expert system, peak weight.

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1 Introduction

Recent work has demonstrated the potential of using probabilistic expert systems (PES) for evaluating DNA evidence (Dawid *et al.* 2002). This article is concerned with the analysis of *mixed traces* where several individuals may have contributed to a DNA sample left on a scene of crime. Mortera *et al.* (2003) showed how to construct a PES using information about which alleles were present in the mixture, and we refer to this article for a general description of the problem and for genetic background information.

However the results of a DNA analysis are usually represented as an *electropherogram* (EPG) measuring responses in *relative fluorescence units* (RFU) and the alleles in the mixture correspond to peaks with a given height and area around each allele, see Figure 1. The band intensity around each allele in the relative fluorescence units represented, for example, through their *peak areas*, contains important information about the composition of the mixture. This information was ignored by Mortera *et al.* (2003) but exploited by e.g. Evett *et al.* (1998) who emphasized and discussed its rôle in evidential calculations. Perlin and Szabady (2001) and Wang *et al.* (2002) used numerical methods known as *Linear Mixture Analysis* (LMA) and *Least Square Deconvolution* (LSD) for separating mixture profiles using peak area information.

In this article we build a PES for mixture traces based on conditional Gaussian distributions for the peak areas, given the composition of the true DNA mixtures; see Chapter 7 of Cowell *et al.* (1999) as well as Lauritzen and Jensen (2001). Such a network enables us to perform evidential calculation and separation of DNA mixtures in a unified way, yielding a natural quantification of any possible uncertainty associated with the analysis.

For the sake of simplicity we only consider the case where the mixed trace contains DNA from exactly two contributors, but we emphasize that the flexibility and modularity of the PES approach enables extension and modification of our network to include complications such as an unknown number of contributors, indirect evidence, etc. along the lines given in Mortera *et al.* (2003). An analysis of a mixed trace can have different purposes, several of which can be relevant simultaneously, making a unified approach particularly suitable. However, for the sake of exposition we consider the issues separately. The first focus of our analysis will be that of *evidential calculation*, detailed in § 4. Here a *suspect* with known genotype is held and we want to determine the likelihood ratio for the hypothesis that the suspect has contributed to the mixture vs. the hypothesis that the contributor is a randomly chosen individual. We distinguish two cases: the other contributor could be a *victim* with a known genotype or a *contaminator* with an

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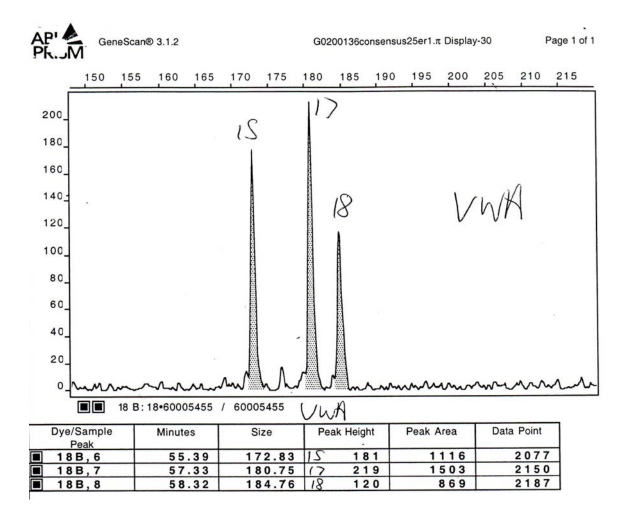


Figure 1: An electropherogram (EPG) of marker VWA from a mixture. Peaks represent alleles at 15, 17 and 18 and the areas and height of the peaks express the quantities of each. Since the peak around 17 is high, this indicates two alleles with repeat number 17. This image is supplied courtesy of LGC Limited, 2004.

unknown genotype, possibly without a direct relation to the crime. In § 5, we show how to use our network for *separation of profiles*, i.e. identifying the genotype of each of the possibly unknown contributors to the mixture, the evidential calculation playing a secondary role. Finally, in § 6 we will contrast our approach with existing alternatives and discuss perspectives for further developments.

2 Basic model assumptions

We assume the usual Mendelian genetic model for the allele composition of the mixture traces with gene frequencies of single STR alleles being those reported in Evett *et al.* (1998) and Butler *et al.* (2003) for U.S. Caucasians, the latter being used for data taken from Perlin and Szabady (2001)¹ and Wang *et al.* (2002). The peak area information is included in the analysis through the relative peak weight. The (absolute) *peak weight* w_a of an allele with *repeat number a* is defined by scaling the peak area with the repeat number as

$$w_a = a\alpha_a,$$

where α_a is the peak area around allele *a*. Multiplying the area with the repeat number is a crude way of correcting for the fact that alleles with a high repeat number tend to be less amplified than alleles with a low repeat number. The *relative peak weight* r_a is obtained by scaling with the total weight

$$r_a = w_a/w_+, \quad w_+ = \sum_a w_a.$$

Our simple model for the observed peak weight, R_a , uses a normal distribution

$$R_a \sim \mathcal{N}(\mu_a, \tau^2), \quad \mu_a = \{\theta n_a^1 + (1-\theta)n_a^2\}/2,$$
 (1)

where θ is the fraction of DNA in the mixture originating from the first contributor, n_a^i is the number of alleles with repeat number *a* possessed by person *i*, and $\tau^2 = \sigma^2 \mu_a (1 - \mu_a) + \omega^2$, where σ^2 and ω^2 are variance factors for the contributions to the variation from the amplification and measurement processes. Unless stated otherwise, we have used $\sigma^2 = 0.01$ and $\omega^2 = 0.001$, corresponding approximately to a standard deviation for the observed relative weight of about

$$\sqrt{0.01/4 + 0.001} = 0.06$$

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¹This dataset has an observed allele 25.2 of the marker FGA. As none of the 302 subjects in Butler *et al.* (2003) had this allele, we chose to use 1/604=0.00166 as the allele frequency.

for $\mu_a = 0.5$. This is about the variability in peak imbalance that has been reported in the literature when amplifying DNA from one heterozygous individual, for which $\mu_a = 0.5$.

Our model ignores the obvious correlation between weights due to the fact that they must add up to unity. The model can be seen as a second order approximation to a more sophisticated model based on gamma distributions for the absolute scaled peak weights to be discussed elsewhere. The simple model above seems in any case sufficiently accurate and adequate for the purposes of the present paper. In general the variance factors may depend on the marker and on the amount of DNA analysed, but here we use the values above.

3 Bayesian networks for DNA mixtures with peak weights

3.1 Object-Oriented Networks

Object-oriented Bayesian networks (Koller and Pfeffer 1997; Dawid 2003) have a hierarchical structure where any node itself can represent a (objectoriented) network containing several *instances* of other generic *classes* of networks. This framework is particularly suited for an application area such as the present because we can exploit the similarity between elements of the networks in a modular and flexible construction, making the networks more and more complex by simply adding new objects which perform different tasks.

Instances have interface *input* and *output* nodes as well as ordinary nodes. Instances of a particular class have identical conditional probability tables for non-input nodes. Instances are connected by arrows from output nodes to input nodes. These arrows represent identity links whereas arrows between ordinary nodes represent probabilistic dependence (Cowell *et al.* 1999). Implementation of object-oriented Bayesian networks is supported by the program HUGIN 6^2 .

3.2 Description of the network classes

Below we describe the different objects used in our construction of the network. In what follows, **bold** will indicate a network class, and **teletype** will indicate a node. In the figures, instances of a certain class are represented by

 $^{^2 \}mathrm{See}$ www.hugin.com

a rounded rectangle, discrete nodes have a single outline, whereas continuous nodes have a double outline. Interface nodes are represented with a grey ring; input nodes having a dotted outline and output nodes having a solid outline. Also, dark grey nodes will indicate where possible evidence might be inserted and black nodes are target nodes or nodes of interest where results will be read.

3.2.1 The founder class

The network class **founder** of Figure 2 contains a single node **founder** with the relevant repertory of alleles as its states, and an associated probability table describing their frequencies.

For illustration, we show marker FGA having observed alleles coded A to C and the aggregation of all unobserved alleles coded as x. The probability table is shown in Table 1.

Table 1: Gene frequencies for marker FGA as reported in Evett et al. (1998).

Allele	A	В	C	x
Frequency	0.187	0.165	0.139	0.509

3.2.2 The genotype class

The network class **gt** in Figure 3 represents an individual's genotype **gt**, which is given by the unordered pair of paternal and maternal genes, {**pg**, **mg**}. Input nodes **pg** and **mg** are copies of node **founder** of class **founder**. The paternal and maternal genes, **pg** and **mg**, are chosen independently from the same population whose allele frequencies are assumed known. Output node **gt** is the logical combination of **pg** and **mg**.

3.2.3 The query class

The network class **whichgt** of Figure 4 describes the selection between two genotypes.

If the Boolean node query? has the value *true*, then the output node, outgt, will have identical genotype to the input node, ingt; otherwise it will be identical to input node othergt. This is written in the HUGIN expression language as: outgt := if(query? == true, ingt, othergt).³

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³The function if (A, x, y) takes the value x if condition A is satisfied, otherwise y.



Figure 2: Network **founder** for founder gene

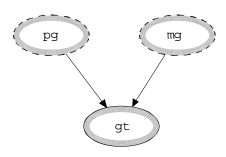


Figure 3: Network **gt** for genotype

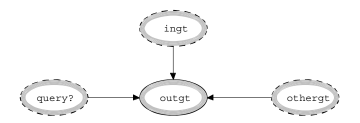


Figure 4: Network **whichgt** for selecting a genotype

3.2.4 The joint genotype

The network class **jointgt** of Figure 5 represents the combined genotype of two individuals, p1 and p2. Node p1gt&p2gt is simply the logical combination of the two input genotypes in p1gt and p2gt.

3.2.5 The number of alleles

The network class **nalleles** shown in Figure 6 counts the number, varying from 0 to 2, of a certain allelic type in a genotype. Thus, for allele A say, nA := if(gt == AA, 2, if (or (gt == AB, gt == AC, gt == Ax), 1, 0)).

3.2.6 The weight of an allele in the mixture

The network class **alleleinmix** shown in Figure 7 computes whether a certain allelic type is in the mixture and its mean contribution to the peak area of the mixture.

Input nodes p1gt and p2gt, the genotypes of the two people, p1 and p2, contributing to the mixture, have identity links to the input node gt in the two instances of class nalleles, n1A and n2A. The Boolean node Ainmix? is *true* if at least one of the two contributors has allele A. Thus, Ainmix? := if(and (n1A_nA == 0, n2A_nA == 0), *false*, *true*), where n1A_nA and n2A_nA refer to the output nodes of the two instances of class nalleles, n1A and n2A.

Input node frac represents the fraction of DNA contributed by p1, denoted by θ in §2. The states of node frac are on a discrete scale ranging from [0,5] for convenience. The scale of node frac can easily be modified to a finer grid. Output node meanA:= n1A_nA * frac + n2A_nA * (5 - frac). This differs from the expression for the mean in (1) by a scale factor of 10 which is appropriately corrected for throughout.

3.2.7 The peak weight

The network class **peakweight** shown in Figure 8 represents the observable peak weight.

The input node mean is identified, for example, with output node meanA of class alleleinmix. The intermediate continuous node weight, with discrete parent mean, has a mean given by the values of node mean and variance equal to 0.01 * mean * (10 - mean), representing variations in the amplification process. The observed peak weight is modelled by the continuous node weightobs to allow for additional measurement error of the true weight, in node weight. Thus this network class is representing the Gaussian part of the model (1).

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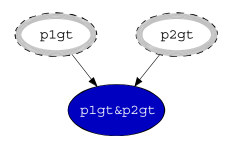


Figure 5: Network **jointgt** for genotype pairs

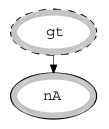


Figure 6: Network **nalleles** for counting the number of alleles

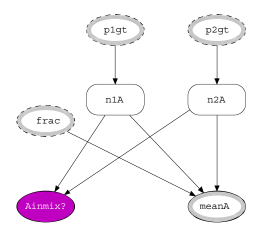


Figure 7: Network **alleleinmix** for alleles in mixture

3.2.8 The target class

The network class **target** shown in Figure 9 has two Boolean output nodes p1=s? (p2=v?) with *true*, *false* states, representing whether contributor p1 (p2) is the *suspect* (*victim*) or not. The black **target** node is the logical conjunction of the nodes p1=s? and p2=v?. These nodes are given a uniform prior distribution so that **target** node has a uniform prior distribution over its states. This enables the computation of the likelihood ratio as described in Mortera *et al.* (2003).

3.2.9 The marker class

The network class **marker** in Figure 10 is an upper level network containing several instances of the classes defined above. This class is made to represent information related to a particular marker. Here it is illustrated for a marker having three alleles represented in the mixture. Input nodes spg, smg, u1pg, u1mg, vpg, vmg, u2pg and u2mg are all copies of node founder of class founder; u1 and u2 are two unspecified individuals. Input nodes p1=s? and p2=v? are identified with the corresponding output nodes of class target. The nodes sgt, u1gt, vgt and u2gt are all instances of class gt; p1gt and p2gt are instances of whichgt and when p1=s? is true (false), p1gt will be identical to sgt (u1gt). A similar relationship holds between nodes p2=v?, p2gt, vgt and u2gt. The node jointgt is an instance of jointgt; Amean, Bmean, Cmean and xmean are instances of class peakweight. Input node frac is identified with the corresponding node in the master network described below.

3.2.10 The master network

Figure 11 gives the master network used for both identification and separation of DNA mixtures from two contributors. It refers to the data from Evett et al. (1998) shown in Table 2.

D8, D18, FGA, and THO1 are all instances of network class marker; D21 and VWA are instances of a simple modification of network class marker and the other network classes it calls, in order to account for 4 observed alleles. D8, D18, FGA, THO1, D21 and VWA each have 8 founder instances with their appropriate gene frequencies as input to the 8 input nodes of class marker. The frac node is connected to all the markers showing their dependence via this quantity. target, an instance of class target is connected to each marker via its output nodes p1=s? and p2=v?. Once constructed, the master network can be used to insert and propagate case evidence in the appropriate

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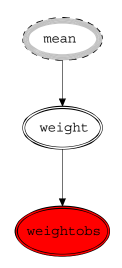


Figure 8: Network **peakweight** for peak weight

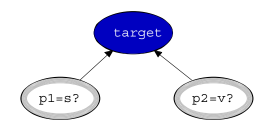


Figure 9: Network **target**

Marker	Alleles	Peak area	Relative weight	Suspect
D8	10	6416	0.4347	10
	11	383	0.0285	
	14	5659	0.5368	14
D18	13	38985	0.8871	13
	16	1914	0.0536	
	17	1991	0.0592	
D21	59	1226	0.0525	
	65	1434	0.0676	
	67	8816	0.4284	67
	70	8894	0.4515	70
FGA	21	16099	0.5699	21
	22	10538	0.3908	22
	23	1014	0.0393	
THO1	8	17441	0.4015	8
	9.3	22368	0.5985	9.3
VWA	16	4669	0.4170	16
	17	931	0.0884	
	18	4724	0.4747	18
	19	188	0.0199	

Table 2: *Evett* data showing mixture composition, peak areas and relative weights from a 10:1 mixture of two individuals, with suspect's genotype specified.

internal nodes, and the marginal posterior probabilities of the quantities of interest can be read from the corresponding nodes.

3.2.11 Amelogenin marker

In the analysis of DNA mixtures the determination of the sex of the two contributors, based on the *amelogenin* marker, is extremely important. To build a network for amelogenin one needs to make the following changes to the previous classes. No founder class is needed and the genotype class has a single output node gt with states XX for female and XY for male, with equal prior probabilities. The **query** and **jointgt** classes only need trivial modifications to reduce their state spaces. The allele counting class **nalleles**, for a male contributor, gt==XY, (for a female contributor, gt==XX) has nX==1 (2) and nY==1 (0). The network class that gives the weight of an allele in the mixture is similar to **alleleinmix** of § 3.2.6, except Xinmix? is always set to *true*. All other network classes remain unchanged.

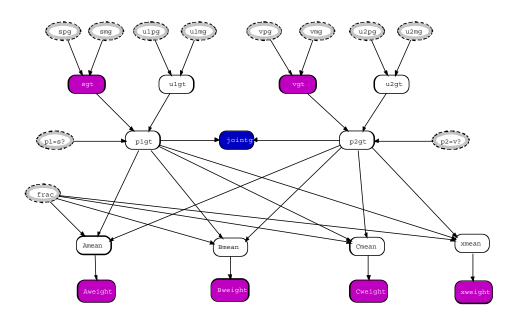


Figure 10: Network **marker**

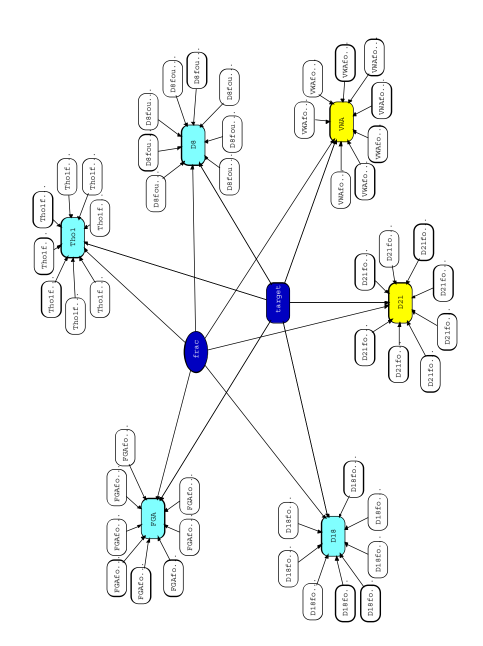


Figure 11: Master network for identification and separation of mixtures

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3.3 MAIES: An expert system for analysing mixed traces

In parallel to the development of the object-oriented networks described in § 3.2 an alternative computer program MAIES— Mixture Analysis In Expert Systems — was developed to provide an independent check of the calculations as well as a flexible environment for specification and input or output of data to allow for experimentation with different methods.

In MAIES, information concerning allele frequencies of markers, and genotypic information (if available) about the potential contributors, is entered via simple popup dialog boxes activated by menu selections. The main program interface displays three user-selectable tabbed windows. One window is used for entering the peak area measurements obtained from the EPG. A second tabbed window displays the graphical model generated on the basis of the marker data, which can be used for entering or retracting evidence on individual nodes (if desired) and for showing the marginal distributions on nodes as bar charts or density curves. The third window is a simple text area that is used to display the marginal distributions of the nodes in the network - the contents of this window may be saved to a file, or cut-and-pasted to other applications, if desired.

After entering peak area information and available genetic profiles on people, the user constructs — by mouse clicking — a single Bayesian network on which the probability calculations are performed. In constructing the Bayesian network the user may specify the scale σ^2 of the amplification error variance, the measurement error variance ω^2 , and the granularity in the discrete states of a node that represents the true fraction of DNA originating from individual 1 (again through a popup dialog box). Sensitivity analysis may be performed in a simple, straightforward manner by varying these three inputs. Peak areas are automatically converted to normalized weights by the program, and entered as evidence in the relevant nodes.

The user can temporarily retract or reinstate evidence on the two potential contributors to the mixture by use of menu selections, thus allowing an evidential calculation to be converted to one of deconvoluting a mixture arising from two unknown contributors, or vice versa.

The Bayesian network generated by MAIES may be considered equivalent to an "unfolded" version of the object-oriented networks described earlier \S 3.2. An example of a network generated for a single marker with two alleles observed in the mixture is shown in Figure 12. The structure is similar to the network shown in Figure 10, and like the object-oriented network described earlier there are several distinct modules of repetition that can be seen in the figure: indeed it is this repetitive structure that makes it possible for MAIES

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to create the much larger Bayesian networks required to analyse mixtures on several markers. We now describe these various structures and how they interrelate.

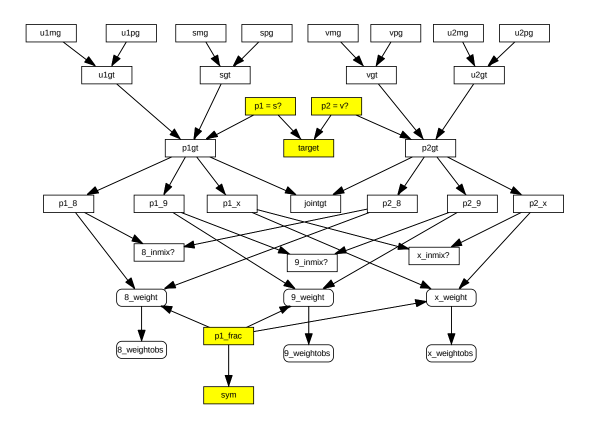


Figure 12: The structure of a Bayesian network generated by MAIES for a single marker, in which two allele peaks (8 and 9) were observed.

3.3.1 Founding people

MAIES currently assumes that DNA from two individuals are in the mixture. Thus it sets up nodes for four founding individuals who are paired up, prefixed by **s** (for suspect), **v** (for victim), and **u1** and **u2** representing two unspecified persons from the population. Corresponding to each of these individuals is a triple of nodes representing their genotype on the marker, and the individuals' paternal and maternal genes. They are joined up as in Figure 3 and their function is the same. The probability tables associated with the maternal and paternal genes contain the allele frequencies of the observed alleles, whilst

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the conditional probability table associated with the genotype node is the logical combination of the maternal and paternal gene.

3.3.2 Actual contributors to the mixture

The genotypes on the marker of the two individuals p1 and p2 whose DNA is in the mixture are the nodes labelled p1gt and p2gt. Node p1gt has incoming arrows from nodes u1gt, sgt and a (yes,no) valued binary node labelled p1 = s?. The function of this latter node is similar to the query? node of Figure 4, namely to set the genotype of node p1gt to be that of sgt if p1 = s? takes the value yes, otherwise set the genotype of node p1gt to be that of u1gt. An equivalent relationship holds between the genotype nodes p2gt, vgt, u2gt and p2 = v?.

The node labelled target represents the four possible combinations of values of the two nodes p1 = s? and p2 = v? as in Figure 9 and described in § 3.2.8.

The network also has a node representing the joint genotypes of individuals p1 and p2, which is labelled jointgt, with incoming arrows from p1gt and p2gt; the function of this part of the network is equivalent to the object shown in Figure 5.

3.3.3 Allele counting nodes

On the level below the genotype nodes for p1 and p2 is a set of nodes representing the number of alleles (taking the value of 0, 1 or 2) of a certain type in each individual. Thus, for example, the node p1_8 counts the number of alleles of repeat number 8 in the genotype of individual p1 for the given marker: this value only depends upon the genotype of the individual p1 and hence there is an arrow from p1gt to p1_8. These nodes model the n_a^i variables introduced in (1).

3.3.4 Repeat number nodes

On the level below the allele counting nodes are the repeat number nodes, labelled 8_inmix?, 9_inmix? and x_inmix?. These are (yes,no) binary valued nodes representing whether or not the particular alleles are present in the mixture: thus for example allele 8 is present in the mixture if either of the allele counting nodes p1_8 or p2_8 takes a non-zero value. For the node x_inmix? the x refers to all of the alleles in the marker that are not observed. When using repeat number information as evidence in calculations, this will be given the value no, the other repeat number nodes will be given the value yes.

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3.3.5 True and observed weight nodes

These nodes are represented by the rounded rectangle shapes. The nodes 8-weight, 9-weight and x-weight represent the true relative peak weights r_8 , r_9 and r_x respectively of the alleles 8, 9 and x in the amplified DNA sample; the nodes 8-weightobs, 9-weightobs and x-weightobs represent the measured weights. The observed weight is given a conditional-Gaussian distribution with mean the true weight, and variance ω^2 . Each true-weight node is given a conditional-Gaussian distribution with mean $\mu_a = \{\theta n_a^1 + (1 - \theta)n_a^2\}/2$, where the fraction θ of DNA from p1 in the mixture is modelled in the network by a discrete distribution in the node labelled p1-frac. The variance is taken to be $\sigma^2 \mu_a (1 - \mu_a)$, as specified in §2. The sym node is only used for separating a mixture of two unknown contributors, and will be described later in §5.2.

3.3.6 Networks with more than one marker

The network displayed in Figure 12 generated by MAIES is for a single marker; for mixture problems involving several markers the structure is similar but more complex because the number of nodes grows with the number of markers (in the *Graham* example, see § 4.1 below, there are 325 nodes). In such a network the nodes shaded in Figure 12 occur only once. The unshaded nodes are replicated once for each marker, with each node having text in their labels to identify the marker that the allele or genotype nodes refer to. There will also be extra repeat number, allele counting and allele weight nodes in each marker having more than two observed alleles in the mixture, extending the pattern for the one-marker network in the obvious manner.

4 Evidence calculations

All evidence calculations reported have been made using MAIES, combined with strategic independent checks using HUGIN. The variable describing the mixture ratio has been discretized to having 101 states $0, 0.01, 0.02, \ldots, 0.99, 1$, but experiments indicate very low sensitivity to the discretization as long as it is not far too rough and 10-20 states would probably be fully appropriate.

4.1 Genotype of suspect and victim available

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This example is taken from Wang *et al.* (2002), stating P. Graham of the Texas Department of Public Safety as the data source. Table 3 displays the measured peak area, the relative weight, and DNA mixture composition on

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9 markers, together with the genotypes of two potential contributors, here named suspect and victim. We will in the following refer to this data as the *Graham* data.

The evidence in this table is now entered into the network and the information propagated. Taking appropriate ratios in the posterior probabilities associated with the target node yields the likelihood ratio in favour of the hypothesis that the victim and suspect vs. that of the victim and a random individual being the contributors to the mixture. Table 4, column "Areas" displays the logarithm of this likelihood ratio, and column "Alleles" the corresponding ratio when only the evidence on the repeat number of the alleles is used. The last columns show the log-likelihood ratio when the mixture ratio θ is assumed known at given values.

The posterior distribution of the mixture ratio θ is displayed in Figure 13. Note that the likelihood ratio is essentially constant in the region $0.3 < \theta < 0.4$ which is the plausible region in the light of the data. Note also that this posterior distribution has its maximum around 0.34, close to the value reported in Wang *et al.* (2002).

The inclusion of area information is indeed strengthening the evidence against the suspect, increasing the logarithm of the likelihood ratio from 12.93 to 14.48, approximately corresponding to a factor 36. This is a modest increase and reflects the fact that when information about the genotype of the victim is available, peak area does not make much difference to the likelihood ratio as genotypes themselves are very informative.

4.2 Only genotype of suspect available

Our next example is taken from Evett *et al.* (1998) and has only information of the genotype from one potential contributor, here named the *suspect*, whereas the other contributor is a *contaminator*. The data is displayed in Table 2 and is henceforth referred to as the *Evett* data. Table 5 displays the logarithm of this likelihood ratio together with the corresponding ratio when peak weights are ignored, and the ratios when the mixture ratio θ is assumed known at given values.

Note that the strengthening of evidence against the suspect is more dramatic when information on the contaminator is absent: the logarithm of the likelihood ratio changes from 4.4 to 8.23, corresponding to an additional factor around 6000, as compared to a factor 36 above.

Also here the likelihood ratio is essentially constant over a region which completely covers the plausible $0.85 < \theta < 0.95$. The posterior distribution of the mixture ratio θ is displayed in Figure 14. The maximum occurs around the value 0.89 which is a little off the true 10:1 mixture proportion.

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Marker	Alleles	Peak area	Relative weight	Suspect	Victim
D3	15	1242	0.3361		15
	16	657	0.1897	16	
	17	1546	0.4742	17	17
D5	7	486	0.0999	7	
	12	512	0.1804	12	
	13	1886	0.7198		13
D7	10	614	0.3232	10	
	11	1169	0.6768		11
D8	12	1842	0.6166		12
	13	490	0.1777	13	
	16	461	0.2057	16	
D13	8	734	0.3128		8
	9	1068	0.5120	9	9
	11	299	0.1752	11	
D18	12	440	0.1724	12	
	13	1503	0.6380		13
	15	387	0.1896	15	
D21	30	842	0.3087		30
	30.2	490	0.1808	30.2	
	31.2	509	0.1941	31.2	
	32.2	804	0.3164		32.2
FGA	22	850	0.3483		22
	23	468	0.2005	23	
	24	681	0.3045		24
	25	315	0.1467	25	
VWA	16	616	0.1900	16	
	17	2021	0.6625		17
	18	425	0.1475	18	

Table 3: *Graham* data showing mixture composition, peak areas, relative weights, suspect's and victim's profiles.

Table 4: Logarithm of the likelihood ratios of s&v vs. u&v for the Graham data.

	Areas	Alleles	Assumed known mixture ratio								
θ			0.1	0.1 0.2 0.3 0.4 0.5 0.6 0.7 0.8 0.9						0.9	
$\rm Log_{10}LR$	14.48	12.93	11.07	13.65	14.48	14.47	11.10	4.5	-5.51	-22.84	-59.52

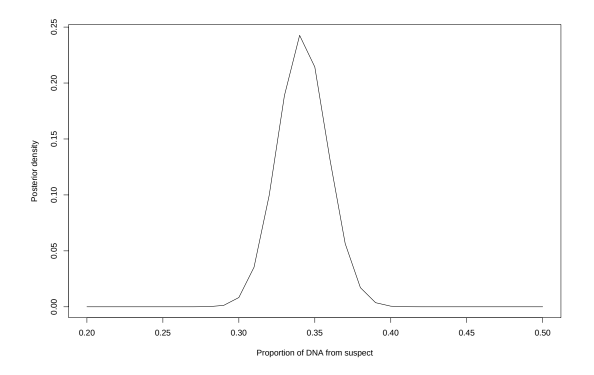


Figure 13: Posterior distribution of the mixture ratio for the *Graham* data using both the suspect's and the victim's genotype.

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Table 5:	Logarithm	of the	likelihood	ratios of	s&u vs.	2u for t	the <i>Evett</i> data.

	Areas	Alleles		Assumed known mixture ratio							
θ			0.1	0.1 0.2 0.3 0.4 0.5 0.6 0.7 0.8 0.9							
$\mathrm{Log}_{10}\mathrm{LR}$	8.23	4.40	-272.78	-167.52	-97.42	-41.88	5.07	8.03	8.53	8.53	8.53

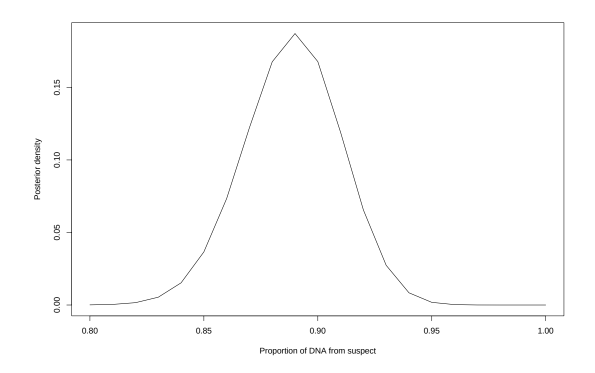


Figure 14: Posterior distribution of the mixture ratio for the *Evett* data using the suspect's genotype.

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The absolute value of the likelihood ratios are slightly different from those given by Evett *et al.* (1998), who report a logarithm of the likelihood ratio of 7.3. This discrepancy is most likely due to slight differences between our model and the model used by Evett *et al.* (1998). On the other hand, they report a likelihood ratio based on allele presence alone of 5800, whereas we find a ratio around 25000, and this appears to be somewhat strange, as we have used the gene frequencies reported in their paper.

5 Separation of mixtures

Deconvolution of mixtures or separating a mixed DNA profile into its components has been studied by Perlin and Szabady (2001) and Wang *et al.* (2002), among others. A mixed DNA profile has been collected and the genotypes of one or more unknown individuals who have contributed to the mixture is desired, for example with the purpose of searching for a potential perpetrator among an existing database of DNA profiles.

The easiest case to consider is clearly that of separation of a single unknown profile, i.e. when the genotype of one of the contributors to the mixture is known. The case when both contributors are unknown is more difficult. In the latter situation this is only possible to a reasonable accuracy when the contributions to the DNA mixture has taken place in very different proportions.

We have chosen to show two alternative methods for predicting the genotype of the unknown contributor(s). In the first method we report the most probable genotype (or pair of genotypes) of the unknown contributor(s) for each marker separately. This result is obtained directly from the standard propagation method in the probabilistic expert system, known as sum-propagation. Note that this genotype is not necessarily the jointly most probable across markers. We therefore also report the joint probability of the genotypes chosen in this way. If this happens to be larger than 0.5, the most probable genotype has clearly been identified.

The second method calculates, by a method termed *semimax*-propagation, the most likely joint configuration of all unobserved discrete nodes, given the evidence available, and reports the genotypes of the unknown contributor(s) associated with this configuration. The semimax propagation first integrates over all unobserved continuous variables and then performs max-propagation as described in Cowell *et al.* (1999), Section 6.4.1, to identify the most probable configuration. Note again that this may not be the most probable genotype across markers. There is no general efficient method for calculating the latter, but identifying the two configurations above and reporting their joint

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probabilities would be fully satisfactory for most purposes as they are most interesting when their joint probability is high.

The two methods generally give results that agree quite closely, the difference largely being due to correlations between the markers originating from the fact the fraction of DNA supplied by each contributor is unknown. When this fraction is well determined by the evidence, the markers are close to being independent. In such cases the two methods tend to given identical results. It then also holds that the joint posterior probability of the genotypes of the unknown contributors is approximately equal to the product of those probabilities for each marker separately.

It would generally seem appropriate to report a list of probable genotypes for the unknown contributor(s), with their associated probabilities, but this would demand a slightly more sophisticated calculation and is beyond the scope of this particular paper.

5.1 Separating a single unknown profile

Our next example is using data from Perlin and Szabady (2001), henceforth referred to as the *Perlin* data, displayed in Table 6.

The two individuals contributing to the mixture are here named *suspect* and *victim* and Table 7 displays the predicted genotype of the suspect, using information from the victim alone.

As in Perlin and Szabady (2001) the genotype of the unknown contributor is essentially determined exactly and the posterior distribution of the mixture ratio concentrates around the true value of 0.7, as displayed in Figure 15. For comparison we have also made a similar calculation for the other two examples. The results are displayed in Table 8 and Table 9.

Here the situation for the *Graham* data is similar to the *Perlin* data: all markers are correctly identified, with probabilities very close to 1 in all cases. Analysis of the *Evett* data also yield probabilities close to 1 on all markers, but not so close as the for *Perlin* and *Graham* data. Evett *et al.* (1998) does not contain the genotype of the second contributor so we do not know whether there are classification errors for this example. Figure 14 and Figure 16 display the posterior distribution of the mixture ratio for these two cases.

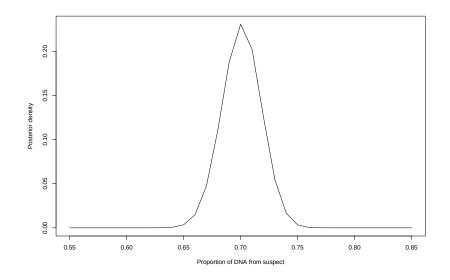


Figure 15: Posterior distribution of the mixture ratio for the *Perlin* data, using genotypic information on the victim only.

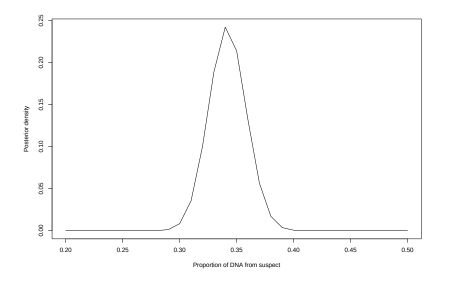


Figure 16: Posterior distribution of mixture ratio for *Graham* data using genotypic information from the victim only.

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Table 6: *Perlin* data showing mixture composition, peak areas, relative weights, suspect's and victim's genotypes from a 7:3 mixture of the two individuals.

Marker	Alleles	Peak area	Relative Weight	Suspect	Victim
D2	16	0.3190	0.1339		16
	18	0.6339	0.2992	18	
	20	0.3713	0.1947		20
	21	0.6758	0.3722	21	
D3	14	1.0365	0.5010	14	14
	15	0.9635	0.4990	15	15
D8	9	0.7279	0.2832	9	
	12	0.2749	0.1426		12
	13	0.6813	0.3829	13	
	14	0.3160	0.1913		14
D16	11	1.4452	0.6801	11	
	13	0.2889	0.1607		13
	14	0.2660	0.1593		14
D18	12	0.3443	0.1504		12
	13	0.6952	0.3290	13	
	14	0.6755	0.3443	14	
	17	0.2850	0.1764		17
D19	12.2	0.6991	0.3109	12.2	
	14	0.6060	0.3092		14
	15	0.6949	0.3799	15	
D21	27	0.2787	0.1289		27
	29	0.7876	0.3913	29	
	30	0.9337	0.4798	30	30
FGA	19	1.0580	0.4621	19	19
	24	0.2830	0.1561		24
	25.2	0.6589	0.3817	25.2	
THO1	6	0.3178	0.1268		6
	7	1.0074	0.4691	7	7
	9	0.6749	0.4041	9	
VWA	17	1.4755	0.7265	17	
	18	0.5245	0.2735		18

Table 7: Predicted genotype of suspect for the *Perlin* data, using genotype information for victim only. All markers are correctly identified by both sum and semi-max propagation.

Marker	Genotype	Probability
D2	18 21	1
D3	14 15	1
D8	9 13	1
D16	11 11	1
D18	$13 \ 14$	1
D19	12.2 15	1
D21	29 30	1
FGA	$19\ 25.2$	1
TH01	7 9	1
VWA	17 17	1

Table 8: Predicted genotype of suspect for *Graham* data, using genotype for victim only. All markers are correctly identified by both sum and semi-max propagation.

Marker	Genotype	Probability
D3	$16\ 17$	0.997638
D5	7 12	1
D7	10 10	0.999464
D8	$13 \ 16$	1
D13	9 11	0.999607
D18	12 15	1
D21	$30.2 \ 31.2$	1
FGA	$23 \ 25$	1
VWA	16 18	1

Table 9: Predicted genotype of contaminator for *Evett* data, using information from suspect. Identical results are obtained using sum and semi-max propagation. The number in brackets is the product of individual marker probabilities.

Marker	Genotype	Probability
D8	11 14	0.904607
D18	$16\ 17$	1
D21	5965	1
FGA	21 23	0.922646
TH01	$9.3 \ 9.3$	0.926062
VWA	$17 \ 19$	1
joint	0.7757173	(0.772921)

5.2 Separating two unknown profiles

We now turn to the problem of separating a mixture into two components, using peak area and repeat number information but no information regarding the two contributors to the mixture. Using only this information will lead to an identifiability problem in assigning genotype combinations to each person, because of the symmetry between the individuals p1 and p2 in the network of Figure 12 or in the equivalent object-oriented network Figure 10.

To remove this problem it is sufficient to enter evidence that the proportion of DNA in the sample from individual p1 is at least one half of the total DNA in the sample. (The alternative, that individual p1 contributes at most half of the DNA to the mixture sample could as equally well be used to break the symmetry.) Using HUGIN this symmetry breaking may be achieved by entering likelihood evidence directly into the fraction node; in MAIES direct entering of likelihood evidence is not possible, so instead this is achieved by entering evidence on the sym node mentioned in § 3.3.5. The node sym has two possible states, $\theta \geq 0.5$ and $\theta \leq 0.5$. Selecting one state as evidence breaks the symmetry (the user does this via a menu selection).

Our first example uses the *Evett* data, ignoring the information on the suspect. The posterior distribution of the mixture ratio θ is displayed as the solid curve in Figure 17. The distribution is similar in shape to that in Figure 14, which uses the suspect genotype information. The broken curve in Figure 17 shows the posterior using the larger variance factor $\sigma^2 = 0.1$. We note that this change of variance has a notable effect on the posterior distribution of mixture ratio.

The predicted genotypes of the two contributors are shown in Table 10, with the suspect's profile being predicted correctly for both choices of variance even though the probability of the chosen genotype is strongly reduced.

Our next example uses the *Perlin* data. The posterior distribution for θ is shown as the solid curve in Figure 18, with the mode at 0.69 very close to the value reported of 0.7. The predicted genotypes of the two contributors is shown in Table 11, with all classifications correct. Still, the joint probability of the chosen genotype indicates that other plausible explanations are available, essentially due to uncertainty about the genotype for marker VWA.

Increasing σ^2 by a factor of 10 to $\sigma^2 = 0.1$ yields the posterior distribution shown by the broken line Figure 18. In this case the effect of choosing an inflated variance factor is dramatic, also yielding reduced genotype probabilities and several classification errors as shown in Table 12. Note also that here there is a large discrepancy between probability of the joint genotype and the product of the probabilities for each marker.

Similar behaviour occurs in our final example that uses the Graham data.

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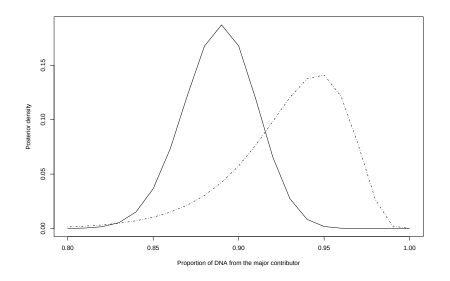


Figure 17: Posterior distribution of mixture ratio from *Evett* data using no genotypic information: solid line $\sigma^2 = 0.01$, broken line $\sigma^2 = 0.1$.

Table 10: Predicted genotypes of both contributors for *Evett* data with $\sigma^2 = 0.01$ and $\sigma^2 = 0.1$. Identical results are obtained using sum and semi-max propagation, with suspect (p1) correct on every marker. The number in brackets is the product of individual marker probabilities.

		$\sigma^2=0.01$			$\sigma^2=0.1$	
Marker	Genotype p1	Genotype p 2	Probability	Genotype p1	Genotype p2	Probability
D8	10 14	11 14	0.904607	10 14	11 14	0.669688
D18	$13 \ 13$	$16\ 17$	1	$13 \ 13$	$16\ 17$	0.995388
D21	67 70	5965	1	67 70	5965	0.999967
FGA	$21 \ 22$	$21 \ 23$	0.922646	$21 \ 22$	$21 \ 23$	0.517756
TH01	8 9.3	9.3 9.3	0.926062	8 9.3	$9.3 \ 9.3$	0.606364
VWA	16 18	17 19	1	16 18	17 19	0.999964
joint	0.775	57173	(0.772921)	0.21	0368	(0.209264)

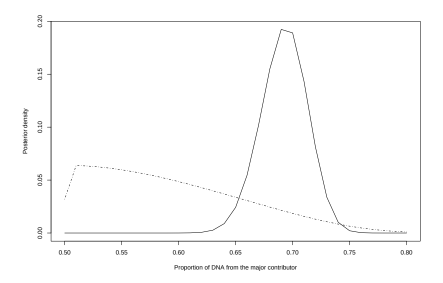


Figure 18: Posterior distribution of mixture ratio from *Perlin* data using no genotypic information: solid line $\sigma^2 = 0.01$, broken line $\sigma^2 = 0.1$.

Table 11: Predicted genotypes of both contributors for *Perlin* data with $\sigma^2 = 0.01$. The number in brackets is the product of individual marker probabilities. All classifications are correct but the marker VWA is uncertain.

		sum prop			semi-max	
Marker	Genotype p1	Genotype p2	Probability	Genotype p1	Genotype p2	Probability
D2	18 21	16 20	0.999072	18 21	16 20	0.999072
D3	14 15	14 15	0.998342	14 15	14 15	0.998342
D8	9 13	12 14	0.997656	913	12 14	0.997656
D16	11 11	13 14	0.999805	11 11	13 14	0.999805
D18	$13 \ 14$	12 17	0.999928	$13 \ 14$	12 17	0.999928
D19	12.2 15	14 14	0.828718	12.2 15	14 14	0.828718
D21	29 30	27 30	0.987463	29 30	27 30	0.987463
FGA	$19\ 25.2$	19 24	0.990340	$19\ 25.2$	19 24	0.990340
TH01	$7 \ 9$	67	0.969079	7 9	67	0.969079
VWA	17 17	18 18	0.537648	17 17	18 18	0.537648
joint	0.434	2251	(0.420059)	0.434	2251	(0.420059)

Table 12: Predicted genotypes of both contributors for *Perlin* data with $\sigma^2 = 0.1$. The number in brackets is the product of individual marker probabilities. There are classification errors in markers D3, FGA and VWA (italicized).

	sum prop			semi-max			
Marker	Genotype p1	Genotype p2	Probability	Genotype p1	Genotype p2	Probability	
D2	18 21	16 20	0.349986	18 21	16 20	0.349986	
D3	14 15	14 15	0.601826	14 15	15 15	0.149452	
D8	913	$12\ 14$	0.339805	$9\ 13$	$12\ 14$	0.339805	
D16	11 11	13 14	0.371007	11 11	13 14	0.371007	
D18	$13 \ 14$	12 17	0.361439	$13 \ 14$	12 17	0.361439	
D19	12.2 15	14 14	0.214513	12.2 15	14 14	0.214513	
D21	29 30	27 30	0.438172	29 30	27 30	0.438172	
FGA	$19\ 25.2$	19 24	0.444306	$19\ 25.2$	24 24	0.114614	
TH01	7 9	$6\ 7$	0.413837	7 9	$6\ 7$	0.413837	
VWA	17 18	17 17	0.432392	17 17	18 18	0.0819453	
joint	0.000510		(7.1723e-05)	0.00	0107	(8.7074e-07)	

The posterior distribution of θ is shown as the solid curve in Figure 19, with a maximum around 0.65; the predicted profiles are shown in Table 13, with one classification error. However note for this classification error (in D7, using sum-propagation) the probability assigned to the genotype pair is around 0.68, with the correct classification (picked out out by the semimax method) has a probability of around 0.32. Note that the two chosen genotypes together account for essentially all of the probability mass.

Increasing the variance factor σ^2 to 0.1 yields more classification errors and also probabilities much lower, as shown in Table 14. The corresponding posterior distribution of θ is plotted as the broken line in Figure 19.

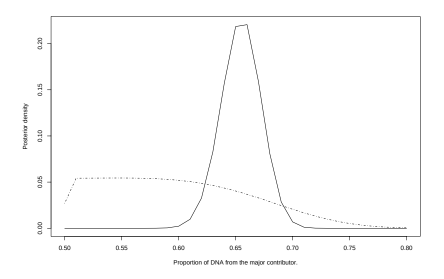


Figure 19: Posterior distribution of mixture ratio from Graham data using no genotypic information: solid line $\sigma^2 = 0.01$, broken line $\sigma^2 = 0.1$.

6 Discussion

In the previous sections we have demonstrated how a probabilistic expert system can be used for analysing DNA mixtures using peak area information, yielding a coherent way of predicting genotypes of unknown contributors and assessing evidence for particular individuals having contributed to the mixture. We emphasize the advantages of using a model-based approach as ours over numerical separation techniques such as Linear Mixture Analysis (LMA) (Perlin and Szabady 2001) and Least Square Deconvolution (LSD)

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Table 13: Prediction of two unknown genotypes for *Graham* data, with $\sigma^2 = 0.01$. The number in brackets is the product of individual marker probabilities. There is a classification error in marker D7 (italicized).

	sum prop			semi-max			
Marker	Genotype p1	Genotype p2	Probability	Genotype p1	Genotype p2	Probability	
D3	16 17	15 17	0.993063	16 17	15 17	0.993063	
D5	7 12	$13 \ 13$	0.999836	712	13 13	0.999836	
D7	11 11	10 11	0.677898	10 10	11 11	0.321722	
D8	$13 \ 16$	12 12	0.952498	$13 \ 16$	12 12	0.952498	
D13	9 11	89	0.997447	9 11	89	0.997447	
D18	12 15	$13 \ 13$	0.990162	12 15	13 13	0.990162	
D21	$30.2 \ 31.2$	$30 \ 32.2$	0.994523	$30.2 \ 31.2$	$30 \ 32.2$	0.994523	
FGA	$23 \ 25$	$22 \ 24$	0.995630	$23 \ 25$	$22 \ 24$	0.995630	
VWA	16 18	17 17	0.997152	16 18	17 17	0.997152	
joint	0.6242217		(0.6251788)	0.297	79028	(0.2967021)	

Table 14: Prediction of two unknown genotypes for *Graham* data, using $\sigma^2 = 0.1$. There are now classification errors in three markers (italicized).

	sum prop			semi-max			
Marker	Genotype p1	Genotype p2	Probability	Genotype p1	Genotype p2 Probabili		
D3	16 17	15 17	0.339087	16 17	15 17	0.339087	
D5	712	13 13	0.425175	712	13 13	0.425175	
D7	11 11	10 11	0.375857	10 10	11 11	0.155175	
D8	12 13	12 16	0.327032	$13 \ 16$	12 12	0.279082	
D13	911	89	0.317089	11 11	89	0.107251	
D18	12 15	13 13	0.310149	12 15	13 13	0.310149	
D21	$30.2 \ 31.2$	$30 \ 32.2$	0.309763	$30.2 \ 31.2$	$30 \ 32.2$	0.309763	
FGA	$23 \ 25$	$22 \ 24$	0.340326	$23 \ 25$	$22 \ 24$	0.340326	
VWA	16 18	17 17	0.350301	16 18	17 17	0.350301	
joint	1.74778e-04		(6.4358e-05)	7.249	1e-05	(7.6696e-06)	

(Wang *et al.* 2002). Since the results are model based there is a natural and directly interpretable quantification of all uncertainties associated with the analysis, and the analysis is extendable to similar but different situations using the modularity and flexibility of the PES approach.

None of the data used as examples had information on the amelogenin marker, but this marker can clearly be very informative concering the unknown fraction of DNA for each contributor when these are of different sex, in particular when genotypes (including sex) is known for one of the contributors.

The examples considered have also demonstrated that there are issues which need further consideration. In particular it appears that the performance of the system is sensitive to the scaling factors used to model the variation in the amplification and measurement processes. This is a serious problem which needs attention. Preliminary investigations seem to indicate that this factor depends critically on the total *amount* of DNA available for analysis. As this necessarily is varying from case to case, a calibration study should be performed to take this properly into account. In any case we find it comforting that the system itself would warn against trusting an uncertain prediction, by yielding an associated low classification probability.

Methods for diagnostic checking and validation of the model should be developed based upon comparing observed weights to those predicted when genotypes are assumed correct. Such methods could also be useful for calibrating the variance parameters σ^2 and ω^2 . To indicate a possible way ahead we note that the network can itself be used for predicting peak weight given a hypothesised composition of the mixture and of the two contributors. Table 15 gives the predicted peak weights for the *Perlin* data based on the repeat numbers in the mixture composition, the true mixture composition, and on the suspect's and victim's genotype. The last two columns show the limits of the 95% predictive interval $[\mu_a - 1.96\tau, \mu_a + 1.96\tau]$ for the weight. For a 95% predictive interval we might expect about one of the areas of the table to lie outside of its predicted interval, as 21 of the 31 intervals are independent (the weights for each marker must necessarily add to one); all expected areas are within their intervals, indicating that the variance at least is not too small.

The predicted peak weights are also useful for identifying measurement errors. For example, if the predicted weight is of the same order of magnitude as the cut-off threshold, the peak is likely to be missed.

Another issue to be further investigated is the possibility of using a model based on gamma distributed absolute peak weights, thereby treating the correlation between individual peak areas in a proper way and avoiding the somewhat unfortunate fact that Gaussian distributions can take negative

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Marker	Allele	Relative Weight	Predicted relative weight		
			$\mu_a - 1.96\tau$ $\mu_a + 1.96\tau$		
D2	16	0.1339	0.0565	0.2435	
	18	0.2992	0.2378	0.4622	
	20	0.1947	0.0565	0.2435	
	21	0.3722	0.2378	0.4622	
D3	14	0.5010	0.3840	0.6160	
	15	0.4990	0.3840	0.6160	
D8	9	0.2832	0.2378	0.4622	
	12	0.1426	0.0565	0.2435	
	13	0.3829	0.2378	0.4622	
	14	0.1913	0.0565	0.2435	
D16	11	0.6801	0.5909	0.8091	
	13	0.1607	0.0565	0.2435	
	14	0.1593	0.0565	0.2435	
D18	12	0.1504	0.0565	0.2435	
	13	0.3290	0.2378	0.4622	
	14	0.3443	0.2378	0.4622	
	17	0.1764	0.0565	0.2435	
D19	12.2	0.3109	0.2378	0.4622	
	14	0.3092	0.1909 0.4091		
	15	0.3799	0.2378	0.4622	
D21	27	0.1289	0.0565	0.2435	
	29	0.3913	0.2378	0.4622	
	30	0.4798	0.3840	0.6160	
FGA	19	0.4621	0.3840	0.6160	
	24	0.1561	0.0565	0.2435	
	25.2	0.3817	0.2378	0.4622	
THO1	6	0.1268	0.0565	0.2435	
	7	0.4691	0.3840	0.6160	
	9	0.4041	0.2378	0.4622	
VWA	17	0.7265	0.5909	0.8091	
	18	0.2735	0.1909	0.4091	

Table 15: Prediction of relative peak weight for *Perlin* data, using the mixture, the suspect's and the victim's DNA composition.

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values. Ideally the method should be generalised to deal with higher complexity such as the simultaneous analysis of several traces, an unknown but large number of contributors, etc., and we have not yet made a proper investigation of the computational complexity issues associated. Finally we emphasize that for the moment we have not dealt with incorporating artifacts such as stutter, pull-up, allelic dropout, etc., but we hope to pursue this and other aspects in the future.

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