DNAtools: Tools for empirical testing of DNA match probabilities

James M. Curran University of Auckland Torben Tvedebrink Aalborg University

December 6, 2013

Abstract

DNA evidence is the pre-eminent tool in the modern forensic scientists toolbox. It is widely accepted by the public, scientific and legal communities and it has been instrumental in determining both the innocence and guilt of individuals involved in the legal process. Despite this widespread acceptance there is unease regarding the statistical measures used to evaluate DNA evidence amongst some of members of all these communities. In particular, some people regard the random match probabilities associated with DNA evidence as just too small or basically unsupportable. In this article we discuss what it means for a pair of DNA profiles to match or partially match, and we present an R package that allows a rational examination of the statistical properties of a DNA database.

1 Introduction

In 2001, a poster was presented by a forensic scientist from Arizona (Troyer et al., 2001) at a scient meeting on human identification. This poster reported a nine locus match between two unrelated men, of white and one black (Kaye, 2009). It was not a full match. Both men had been typed at thirteen loci in total and "partially matched" at three of the remaining four loci. These partially or non-matching loci wo have excluded either man as a suspect if the other was the true offender. However, such a match seem to be at odds with the random match probabilities. On one hand, these two men were in a DNA datab which consisted of approximately 65,000 profiles, and on the other hand, the random match probability for the nine locus genotype were "1 in 754 million in Caucasians, 1 in 561 billion in African Americans, at 1 in 113 trillion in Southwest Hispanics", Troyer et al. (2001).

As we will show later this is, in effect, an example of the "birthday problem" and therefore is regarded completely predictable from a statistical perspective. However, most us who have taught a class on birthday problem know that our students are initially skeptical.

1.1 DNA evidence, matches and partial matches

Forensic genetics has its terminology which we briefly explain here. Human DNA consists of 23 pairs chromosomes and those chromosomes are composed of a sequence of nucleotides which are labeled A, G and T after the bases adenine, guanine, cytosine and thymine that are used to form them. Modern Dl typing uses short tandem repeats (STRs). These are regions of DNA which are highly variable, but patterned in that they consist of repeats of a short sequence of DNA bases. The locations at which to information is collected are called loci, and the (length) variations in the patterns observed at each located alleles. We have two alleles at each locus, because humans are a diploid species, meaning the have two copies of each chromosome. One allele comes from our mother, and the other from our fath

A pair of alleles at a locus is called a genotype, and therefore a DNA profile is actually a multi-locus genotype. Modern forensic laboratories genotype DNA evidence using commercial kits, called multiple which consist of 9–17 loci. The multiplex currently used in the United Kingdom (and until recently N Zealand and Denmark) is called AmpFlSTR® SGM Plus $^{\text{TM}}$, or SGM Plus for short, and consists of 10 locus one sex specific locus, Amelogenin. Forensic laboratories in the United States which load profiles in the FBI's Combined DNA Index System (CODIS) collect a core set of thirteen loci, although they are constrained to use one multiplex.

Locus	vWA	D18	TH01	D2	D8	D3	FGA	D16	D21	D19
Alleles	15,18	14,17	6,9.3	17,23	12,15	15,15	19,23	11,12	28,28	13,14

Table 1: A DNA profile from the AmpFlSTR® SGM Plus[™] multiplex

Table 1 shows a DNA profile from the AmpFlSTR® SGM Plus multiplex. There are two numbers each locus representing the two alleles that make up the genotype at that locus. The numbers relate to number of times the pattern or motif that describe the alleles at the locus are repeated. For example, to person's genotype at the locus THO1 is 6,9.3. This means that on one chromosome, the motif for THO TCAT was repeated 6 times, and on the other chromosome it was repeated 9 times, and then followed TCA. The .3 represents the fact that three of the four bases have been repeated.

A pair of profiles is said to (fully) match if every allele at every locus that occurs in one profile occ in the other. A pair of profiles are said to partially match if there are allelic matches at a subset of ke Weir (2004) provided a taxonomy for describing partial matches which depends on the number of furpartial and non-matching loci between a pair of profiles. For any given pair of (full) profiles from the samultiplex there will be: m_2 loci where both alleles match, m_1 loci where only one of the alleles match and m_0 loci where none of the alleles match. For example, the profile that Troyer *et al.* (2001) found wa 9/3/1 partial match - nine fully matching loci, three partially matching loci, and one non-matching loci

1.2 DNA database comparison exercises

The Troyer match came from a database matching exercise. In such an exercise every profile is comparable with every other profile in the database. This type of comparison exercise is absolutely essential at in addition, can provide some interesting information about the statistical properties of the populat under consideration. We say that database comparison is essential in the first instance for the detection duplicates. Duplicates may arise in a number of different ways. For example, an offender may provide a farname or an offender's name may be entered incorrectly. Alternatively, an offender may have an identity win who is already in the DNA database. There are six pairs of identical twins in the New Zealand Natio DNA Database (NZDNADB). Forensic scientists are also interested in 'very close' matches. For example pair of profiles might fully match at nine loci out of ten and partially match at the remaining locus. To may happen either because the donors of the samples are very close relatives. It is more likely, however, the profiles do not match because of allelic dropout, primer binding site mutations, nomenclature changes are very close relatives.

1.3 The birthday problem

or somatic mutation.

Weir (2007) and others (Brenner, 2007; Curran *et al.*, 2007; Mueller, 2008; Kaye, 2009) note that presence of matching profiles in a DNA database is effectively an instance of the well-known "birthed problem" (Wikipedia, 2010) where, in a group of at least 23 randomly chosen people, there is a great

than 50% chance that one pair of them will have the same birthday. Early critics, implicitly calculat the expected number of matches as Np, used the wrong value for N and the wrong value for p. Firstly, number of pairwise matches, not the size of the database, is the relevant quantity. Although the databasize is relatively small, the number of pairwise comparisons is very large. The Arizona database contain of N = 65,493 profiles (Brenner, 2007). Therefore, there are

$$N_{\text{Comparisons}} = \frac{N(N-1)}{2} = 2,144,633,778$$

or approximately two billion, possible pairwise comparisons. Secondly, the random match probability not the probability we need. The random match probability for the pair of profiles in question answers question "What is the probability that someone other than these two men would have this particular n locus profile." The probability we actually want is "What is the probability that two randomly select profiles would match at nine loci, partially match at three loci, and not match at one locus." Weir (200 working on an unrelated case, showed that this probability can be calculated by

$$P_{m_0,m_1,m_2}(\theta) = \sum_{m_{l0},m_{l1},m_{l2}} \prod_{l} P_{l2}(\theta)^{m_{l2}} P_{l1}(\theta)^{m_{l1}} P_{l0}(\theta)^{m_{l0}}$$

where m_{l0} , m_{1l} and m_{l2} are indicator variables that are equal to one if the individuals share zero, or two alleles in common respectively and zero otherwise. The expressions $P_{li}(\theta)$ are the probability sharing i = 0, 1, 2 alleles in common at locus l for a given degree of population substructure θ and given explicitly in Weir (2004, 2007). The coancestry coefficient, θ or F_{ST} models low levels of related between individuals in the same subpopulation, and is typically between 0 and 0.03.

1.4 Modeling the observed data

Weir's original paper (Weir, 2004) contained an informal analysis where the minimum level of θ requited to explain the observed counts was calculated. For example using the FBI Caucasian data (Budowle & Moretti, 1999) a θ value of 0.005 is needed to explain the 679 observed one locus matches (at locus FG. That is, if $\theta > 0.005$, then the expected count at this locus will exceed the observed count. Curran et (2007) formalized and extended this analysis in the following way. We model the expected number of paper of profile which fully match at m_2 loci and partially match at m_1 loci for a given value of θ , $E_{m_2/m_1}(\theta)$.

$$E_{m_2/m_1}(\theta) = \alpha E_{m_2/m_1}^U(\theta) + \beta E_{m_2/m_1}^B(\theta) + \delta E_{m_2/m_1}^C(\theta) + \gamma E_{m_2/m_1}^P(\theta)$$

where $0 \le \alpha, \beta, \delta \le 1$ and $\gamma = 1 - \alpha - \beta - \delta$. The quantities $E_{m_2/m_1}^R(\theta)$, $R \in \{U, B, C, P\}$ are expected number of matching pairs of profiles calculated under four relationship categories: unrelated, siblings (brothers), cousins, and parent/child. These expressions are derived in Curran *et al.* (2007), wit typographical mistake corrected in Curran and Buckleton (2010). Tvedebrink (2010) also derived expression avancular relationships. Curran *et al.* (2007) estimated $\alpha, \beta, \delta, \gamma$ and θ by using a combination of a learnth (across θ) and a Monte Carlo steepest descent method to find the values that minimized several different distance metrics applied to the observed and expected value. Curran *et al.* (2007) recommend minimizing

$$C_3(\theta) = \sum_{i=0}^{L} \sum_{j=0}^{L-m_2} \frac{|E_{i/j}(\theta) - O_{i/j}|}{O_{i/j}}$$

where $O_{i/j}$ is the observed number of pairs of profiles fully matching at $m_2 = i$ and $m_2 = j$ loci us a multiplex consisting of L loci. This metric was chosen because of the belief that it puts emphasis

"explaining" the higher order matches. Further research into this by Tvedebrink (2010); Tvedebrink et (2011) has shown that Mahalanobis distance

$$T_2(\theta,) = \left(\vec{E}(\theta) - \vec{O}\right)^{\top} \Sigma(\theta)^{-} \left(\vec{E}(\theta) - \vec{O}\right)$$

actually provides a "better" fit to the data and does not drive the value of θ to zero. $\Sigma(\theta)^-$ is a pseu inverse because of constraint

$$\sum_{m_2=0}^{L} \sum_{m_1=0}^{L-m_2} P_{m_2,m_1,m_0} = 1$$

2 The DNAtools package

The aim of the **DNAtools** package is to provide statisticians and forensic scientists with access to procedures described in the previous sections. Early implementations by Weir (2004) and then Curret al. (2007) required custom written code for each new database and, in the case of Curran et al. (2007) generation of at least half a dozen precursor files and a significant amount of memory. Tvedebrink (2017) Tvedebrink et al. (2011) reduced the computational effort of Weir (2004) and Curran et al. (2007) deriving recursion formulas for Equation 1, improved the optimization procedures through the use of package **Rsolnp** (Ghalanos and Theussl, 2010), and derived the variances of the probabilities which allow both the computation of Mahalanobis distances and asymptotic confidence intervals. **DNAtools** aims make all of these procedures easier to use in R (R Development Core Team, 2010).

3 Using the package DNAtools

The expected data format of the databases used as input for the functions in **DNAtools** is a data fram which is constituted by a column of DNA profile identifiers (the first column) and two columns per typ DNA marker. An example is given below:

head(get(data(dbExample)))[,1:9]

ıd	D16S539.1	D16S539.2	D18S51.1	D18S51.2	D19S433.1	D19S433.2	D21S11.1	D21S11.2
1	11	11	15	21	14	14	28	29
2	13	12	15	14	16	16	29	28
3	9	9	13	17	14	14	28	27
4	11	12	14	15	15	13	32	29
5	12	12	17	12	15.2	13	31.2	28
6	9	13	17	14	13	14	30.2	28

Budowle and Moretti (1999) published data from six US subpopulations of different ethnicity (Caucasia Hispanics, African Americans, Bahamians, Jamaican and Trinidad). We demonstrate here our packa **DNAtools** using the Caucasian profiles typed at nine forensic STR markers.

(caucasian.summary <- dbCompare(caucasian,hit=5))</pre>

Summary matrix

partial

match 0 1 2 3 4 5 6 7 8 9

0 17 145 628 1531 2416 2516 1822 752 170 26

1 28 178 733 1426 1902 1455 727 211 40

```
2
           121
                  303
                         530
                                492
                                       310
                                              108
                                                       9
     13
                          99
3
      5
            32
                   64
                                 52
                                        23
                                                 6
                            7
                                   2
4
      0
              6
                     6
                                          1
5
              1
                     0
                            1
                                   1
6
       0
              0
                     0
                            0
7
              0
                     0
       0
8
       0
              0
9
```

Profiles with at least 5 matching loci

	ID1	1D2	match	partial
1	10	29	5	4
2	77	116	5	3
3	64	170	5	1

There is a plot method for the returned object. Applying this method to caucasian.summary yields "dropping ball"-picture of Figure 1. The right end of the "distribution" is interesting part, due to the lar number of coinciding loci between profile pairs.

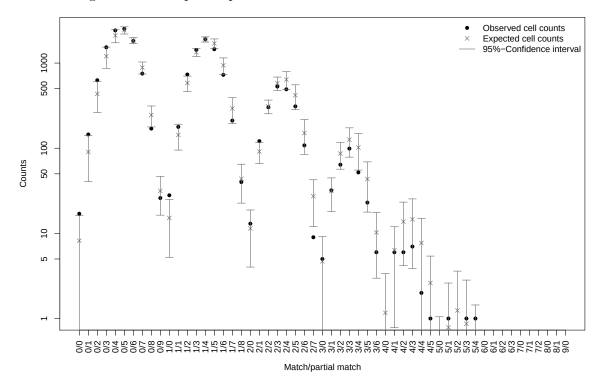


Figure 1: Plot produced by plot(caucasian.summary,pch=16). Superimposed are the expected cou and associated 95%-confidence intervals. The labels of the first axis denote the number of matching a partial-matching loci. The second axis is on a log₁₀-scale.

In Table 2, the estimated parameters are reported using the different object functions implemented in optim.relatedness-function of the **DNAtools**-package. Only for T_2 the estimate of θ is different from

The fitted values can be used to compute $E_{m_2/m_1}(\theta)$. This is done using dbExpect which takes θ and list of locus specific probability vectors as input. The function efficiently computes the expectation usin recursion relation (Tvedebrink *et al.*, 2011). Similarly, the superimposed confidence intervals is Figure 1 w computed using the dbVariance-function, which computes the covariance matrix of the summary statis

	θ	Unrelated	First-Cousins	Avuncular	Parent-child	Full-siblings
C_1	0	9.99e-01	2.32e-13	1.15e-08	5.01e-04	1.57e-04
C_2	0	9.99e-01	6.90e-08	3.99e-08	1.16e-08	3.68e-14
C_3	0	9.99e-01	2.16e-08	1.01e-11	1.08e-04	7.03e-04
T_1	0	9.99e-01	3.97e-09	5.64e-12	4.63e-04	1.58e-05
T_2	0.015	9.99e-01	1.67e-10	4.60e-06	7.47e-04	6.15 e - 06

Table 2: The estimated parameters of the model for the Caucasian subsample.

Availability The R package DNAtools is available at CRAN: DNAtools

4 Conclusion

In this paper we have described an R package which allows statisticians and forensic scientists to ease examine the properties of a forensic DNA database. In particular, our package makes it simple to calculate out a database comparison exercise where every DNA profile in the database is compared to every of database, and compare the resulting numbers of observed pairs of matching and partially matching profit to expectation under a set of population genetic assumptions. There are potential limitations on the of this package in that it may not scale well to extraordinarily large databases (>100,000 profiles), but expect that this will be remedied by further development.

Acknowledgments

We wish to acknowledge Professor Chris Triggs and the Department of Statistics at the University Auckland for financial support for accommodation and travel for T. Tvedebrink. We also wish to thank I John Buckleton for initially introducing us to this problem. T. Tvedebrink would like to thank Poul Sva Eriksen, Aalborg University, for his contributions to the recursion formulas for computing expectations a covariance matrices.

References

Brenner C (2007). "Arizona DNA Database Matches." Accessed 4-January-2010, URL http://dna-viecom/ArizonaMatch.htm.

Budowle B, Moretti TR (1999). "Genotype Profiles for Six Population Groups at the 13 CODIS Short Todem Repeat Core Loci and Other PCR-Based Loci." Forensic Science Communications, 1(2). Access 5-January-2010, URL http://www2.fbi.gov/hq/lab/fsc/backissu/july1999/budowle.htm.

Curran JM, Buckleton JS (2010). "Re: Sign mistake in allele sharing probability formulae of Curran, al." Forensic Science International: Genetics, 4(3), 215–217.

- Curran JM, Walsh SJ, Buckleton J (2007). "Empirical testing of estimated DNA frequencies." Forent Science International: Genetics, 1(3-4), 267–272. ISSN 1878-0326. URL http://www.ncbi.nlm.ncgov/pubmed/19083772.
- Ghalanos A, Theussl S (2010). **Rsolnp**: General Non-linear Optimization. R package version 1.0-7, U http://CRAN.R-project.org/package=Rsolnp.
- Kaye DH (2009). "Trawling DNA Databases for Partial Matches: What Is the FBI Afraid Of?" Cor Journal of Law and Public Policy, 19(1), (in press).
- Mueller LD (2008). "Can simple population genetic models reconcile partial match frequencies observing large forensic databases?" Journal of Genetics, 87(2), 101–108. ISSN 0022-1333. URL http://www.ncbi.nlm.nih.gov/pubmed/18776637.
- R Development Core Team (2010). R: A Language and Environment for Statistical Computing. R Fountion for Statistical Computing, Vienna, Austria. ISBN 3-900051-07-0, URL http://www.R-project.o
- Troyer K, Gilboy T, Koeneman B (2001). "A nine STR locus match between two apparently unrela individuals using AmpFlSTR® Profiler Plus™ and Cofiler™." In Genetic Identity Conference Proceedir 12th International Symposium on Human Identification. Accessed 4-January-2010, URL http://www.promega.com/geneticidproc/ussymp12proc/abstracts/troyer.pdf.
- Tvedebrink T (2010). Statistical Aspects of Forensic Genetics Models for Qualitative and Quantitative STR Data. Ph.D. thesis, Department of Mathematical Sciences, Aalborg University.
- Tvedebrink T, Eriksen PS, Curran JM, Mogensen HS, Morling N (2011). "Analysis of matches and part matches in a Danish DNA reference profile data set." Forensic Science International: Genetics.
- Weir BS (2004). "Matching and partially-matching DNA profiles." Journal of Forensic Sciences, 49(1009–1014. ISSN 0022-1198. URL http://www.ncbi.nlm.nih.gov/pubmed/15461102.
- Weir BS (2007). "The rarity of DNA Profiles." The Annals of Applied Statistics, 1(2), 358-370. IS 1941-7330. URL http://www.ncbi.nlm.nih.gov/pubmed/19030117.
- Wikipedia (2010). "Birthday problem." Accessed 5-January-2010, URL http://en.wikipedia.org/wilibirthday_problem.