Prediction of consanguinity using human DNA fingerprints

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SUMMARY DNA fingerprinting was performed to verify the pedigree structure of a family under investigation for an unusual case of β thalassaemia. A higher than expected proportion of hypervariable bands was shared by the proband and his mother, leading to suspicion that the child had been the product of a consanguineous mating. Further analysis of the mother’s brother showed that he was almost certainly the proband’s father.

Accurate assessment of pedigree structure is of considerable importance in genetic studies. Traditionally blood group, protein, and tissue antigen markers have been used for this purpose. These approaches are limited by virtue of their low individual informativeness and their major application is in the exclusion of paternity.

Over the last few years many sequence variants of human DNA have been described which are detectable as restriction endonuclease fragment length polymorphisms (RFLPs). The advantage of DNA polymorphisms is their virtual limitless number. Recently, a class of DNA probes capable of detecting multiple sequences of highly polymorphic length, known as hypervariable regions (HVRs), has been isolated. Under appropriate hybridisation conditions, multiple HVRs can be detected simultaneously in human DNA to produce a DNA fingerprint that is completely individual specific and shows somatic and germline stability. The component alleles of these DNA fingerprints are inherited in a Mendelian fashion and are derived from a large number of dispersed autosomal loci. Hence, DNA fingerprints are highly effective for individual indentification and have been successfully used to resolve disputes arising from lack of proof of family relationships. As another example of this application, we report the use of DNA fingerprinting analysis in the prediction of consanguinity in a family.

Materials and methods

CASE REPORT
A five year old Asian Indian boy was referred with anaemia and poor growth. He was born in Africa and had been previously admitted to several hospitals with jaundice and anaemia, but had never been transfused. On examination he was pale, mildly icteric, and rather small for his age; he had bifid thumbs, webbed toes, and a palpable spleen. Haematological investigations showed Hb 6-5 g/dl, MCV 85 fl, MCH 27 pg, and 3-2% reticulocytes. Hb electrophoresis showed Hb A2 0-5% with non-Hb A, the rest of the haemoglobin being Hb F. Globin chain synthesis studies confirmed the absence of β chains with an α:non-α globin chain ratio of 0-33. Cytogenetic studies showed a balanced reciprocal translocation between chromosomes 15 and 20. A provisional diagnosis of thalassaemia intermedia with an unassociated chromosomal anomaly was made. The child was thought to be homozygous for βthalassaemia or a compound heterozygote for two different β thalassaemia mutations. Haematological investigation of the mother showed Hb 12-0 g/dl, MCV 57-5 fl, MCH 18-9 pg, Hb A2 6%, and Hb F 2-6%. The father’s indices were Hb 16-1 g/dl, MCV 86-6 fl, MCH 30-6 pg, Hb A2 2-2%, and Hb F 0-1%. Thus, the mother was heterozygous for β thalassaemia but the father was apparently normal and this was confirmed by a β/α globin chain synthesis ratio of 0-99. Non-paternity was denied.

In order to obtain a more definitive diagnosis, particularly for genetic counselling purposes, DNA fingerprinting was carried out on all available family members.

DNA ANALYSIS Genomic DNA was extracted from peripheral leucocytes by standard techniques. From each subject 5 µg DNA was digested with HinfI and
electrophoresed through a 20 cm 1-0% agarose gel at 2 Vcm\textsuperscript{-1} for 46 hours, at which time the 2 kb molecular weight marker had reached the bottom of the gel. The DNA was transferred to a nylon membrane (Hybond-N, Amersham) and incubated at 37°C for three hours in heparin prehybridisation buffer (3\times SSC, 50 \mu g/ml heparin, 0-2% SDS, 50% formamide). Minisatellite probes 33-15 and 33-6 were prepared from single stranded template, as described by Jeffreys \textit{et al},\textsuperscript{1} and the filter was hybridised to each probe in turn by incubation at 37°C overnight in heparin hybridisation buffer (3\times SSC, 200 \mu g/ml heparin, 0-2% SDS, 50% formamide, 5% dextran sulphate) at a concentration of 2\times 10\textsuperscript{6} cpm/ml. After hybridisation, the membrane was washed at 65°C in 1\times SSC/0-1% SDS for 45 minutes, then autoradiographed. Between the two hybridisations, the probe was eluted from the filter by incubating at 45°C in 0-4 mol/l NaOH for 30 minutes and then in 0-2 mol/l Tris pH 7-5, 0-1\times SSC, 0-1% SDS for 30 minutes.

Results

The confirmation or exclusion by DNA fingerprinting of a reported family relationship is accomplished by comparing the patterns of hypervariable fragments of the child and its putative parents. The figure shows the DNA fingerprints generated by probes 33-15 and 33-6 for the family described.

The mean probability, \( x \), that a band present in a subject’s DNA fingerprint will occur in the fingerprint of an unrelated subject has been estimated in population studies to be \( x = 0.2 \).\textsuperscript{2} Thus, a child whose parents are not related to one another will, on average, inherit one half of the bands appearing in its mother’s fingerprint as well as half of the one-fifth of his father’s bands which are shared by chance with the mother, giving a total average band sharing frequency of 0-6 between parent and offspring.

This frequency, \( x \), of band sharing in the population is related to the allele frequency, \( q \), by the relation \( x = 2q - q^2 \). Where \( q \) is small, as is the case for the hypervariable alleles of DNA fingerprints, \( x \approx 2q \), and so for \( x = 0.2, q \approx 0.1 \).

Under Hardy-Weinberg equilibrium, the probability that an allele present in a subject is shared by that subject’s sib is given by the expression \((4+5q-6q^2+q^3)/(4(2-q))\). Using the approximation \( q \approx 0.1 \), this gives a frequency of band sharing between sibs of 0.69. Thus, the expected frequency of fingerprint band sharing between second degree relatives (for example, between aunt and nephew) can be calculated as \((0.69 \times 0.5) + (0.2 \times 0.5)\) = 0.44. We have used these expected frequencies of band sharing to analyse the fingerprint data for the family under study in this report (table). The fingerprints of S1 (proband) and S2 (his brother) comprise 49 and 50 resolvable bands, respectively. Of the 51 bands present in the fingerprint of M, the putative mother, 39 are shared by S1 and 27 by S2. Assuming a binomial distribution, the likelihood that S1 is not related to M is \( L(p=0.2) = (\binom{51}{39}) \times 0.2^{39} \times 0.8^{12} = 6.0 \times 10^{-18} \), and the likelihood of S1 and M being first degree relatives is \( L(p=0.6) = (\binom{51}{39}) \times 0.6^{39} \times 0.4^{12} = 5.9 \times 10^{-3} \), giving odds of 9.9 \times 10^{14} to one in favour of a first degree relationship. Similarly, for S2, the odds are 4.5 \times 10^7 to one in favour of a first degree relationship with M. Hence, it is highly likely that M is the mother of S1 and S2.

To establish the paternity of S1 and S2 we compared the patterns of DNA fingerprints in S1 and S2 with H, M’s husband. All the 23 non-
maternal bands in S2 are present in H. The probability that H is not the father of S2 and possesses all these 23 bands by chance is \( (0.2)^{23} = 8.4 \times 10^{-17} \). Therefore, S2 is almost certainly the child of H and M. However, of the 49 bands present in S1, 10 cannot be traced to either M or H and thus the reported family relationship is false. Of the S1 bands that can be resolved in M's fingerprint, 27 are shared with S2. This degree of band sharing is consistent with a first \( (\chi^2 = 1.06, 1 \text{ df}, p > 0.02) \) or a second \( (\chi^2 = 1.68, 1 \text{ df}, p > 0.02) \) degree relationship, but not with unrelatedness \( (\chi^2 = 17.3, 1 \text{ df}, p < 0.005) \). S1 shares 39 of the S1 bands present in M's fingerprint. The expected number of shared bands if S1 is a first degree relative of M is 30-6, so the number of shared bands between S1 and M is significantly higher than would be expected even for a first degree relationship \( (\chi^2 = 5.76, 1 \text{ df}, p < 0.02) \), and suggests that the father of S1 may be related to M. Examination of the DNA fingerprints showed that all the 10 non-maternal bands in S1’s fingerprint were present in M’s brother, B. The probability that B is unrelated to S1 but happens to share these bands is \( (0.2)^{10} = 1 \times 10^{-7} \). If, however, some other male relative of M shares these 10 bands with B, then that relative could equally be the father of S1. The probability of a sib of B sharing these 10 bands is \( (0.69)^{10} = 0.025 \). We therefore conclude that the true parents of S1 are M and B. Haematological investigations showed that B is heterozygous for β-thalassaemia which would explain the phenotype of homozygous β-thalassaemia in S1.

After completion of the studies, M volunteered that her brother, B, was the true father of her son, S1.

Discussion

DNA fingerprinting is a robust technique which allows positive determination of paternity with a high degree of certainty. In this family it has permitted the detection of consanguinity within a nuclear family. This use of DNA fingerprinting is unlikely, however, to be universally applicable in the detection of incest because of the variance of parent-offspring band sharing, which will sometimes obscure consanguinity. In this case, S1’s unusual clinical phenotype, combined with his very high degree of band sharing with his mother, M, led us to suspect incest. Had the band sharing between M and her brother, B, been slightly lower, and had S1 shared 37 of M’s 51 bands instead of 39, then the observations would have been within the bounds of what might be expected for a first degree relationship \( (\chi^2 = 3.35, 1 \text{ df}, p > 0.05) \). In such a case, and in the absence of clinical clues, the possibility of consanguinity might go unsuspected. Thus, this case must be regarded as anecdotal, and we do not wish to suggest that DNA fingerprinting with these two probes alone will detect consanguinity in all cases.

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References


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