Arginine 109 to glutamine mutation in a girl with ornithine carbamoyl transferase deficiency

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Abstract
We studied DNA from 29 families with at least one member with ornithine carbamoyl transferase (OCT) deficiency and have found a mutation in the Taq1 site within exon 5 of the OCT gene in a female presenting at the age of 21 months. Hybridisation with site specific oligonucleotides shows that the mutation is a C to T substitution resulting in a glutamine for arginine substitution at amino acid 109.

Ornithine carbamoyl transferase deficiency is an X linked disorder caused by mutation of the urea cycle enzyme ornithine carbamoyl transferase. Heterozygous females have very variable manifestations, some being virtually asymptomatic while others have severe protein intolerance and recurrent hyperammonaemia. The reason for this variation is not known although it is usually assumed that it is a result of unequal X inactivation. A survey of mutations found in symptomatic females is needed to clarify the variation in clinical symptoms.

A number of mutations have been defined in males with OCT deficiency. These include deletions and point mutations. Of particular interest is the finding of several independent mutations having arisen at codon 109 within a Taq1 restriction enzyme site. Mutations of C to T at this position in the coding strand change arginine to a stop codon and arginine to glutamine when the change occurs on the complementary DNA strand. Cell transfection studies confirmed that this latter change abolishes enzyme activity. The arginine to stop mutation has already been found in a mildly affected female and we now report the other, arginine to glutamine, change arising independently for a second time, in this case in an affected female.

Materials and methods
CLINICAL DETAILS
The female proband is 3½ years of age and presented at the age of 21 months with developmental delay (around the 9 to 10 month level) and poor growth (−3.62 SD). Increased urine glutamine and orotic acid confirmed OCT deficiency. On a low protein diet, sodium benzoate, and arginine she has done well and her growth has caught up somewhat (now being only −1.96 SD). At 3½ years of age her development is now borderline normal. There was no previous family history. Chromosomal analysis was not performed.

We examined DNA from a further 28 families with at least one member, male or female, manifesting OCT deficiency, who had been referred to the Institute of Child Health for DNA testing for carrier detection or prenatal diagnosis.

OLIGODEOXYRIBONUCLEOTIDE SYNTHESIS
Oligonucleotide primers and probes were synthesised on an Applied Biosystems 381A DNA synthesiser.

Taq1 RESTRICTION ENZYME ANALYSIS
DNA was prepared from the nuclei of peripheral leucocytes by guanidinium hydrochloride extraction, digested with Taq1 according to the manufacturer's instructions (NBL), and the fragments separated by electrophoresis on a 0.8% agarose gel. After denaturation of the gel with 0.4 mol/l NaOH, 1.5 mol/l NaCl, the DNA was blotted directly onto Hybond N+ (Amersham International) and fixed by rinsing with 0.1% SDS for 16 hours each. Filters were washed for 3×20 minutes in 3×SSC,

HYBRIDISATION PROBE
The probe used was a 1.5 kb human cDNA insert isolated from Nuseive-GTG agarose or whole plasmid and labelled by random hexanucleotide primer extension with a 32PdCTP. Prehybridisation and hybridisation were carried out in 10×Denhardt's solution, 4×SSC, 50 μg/ml sonicated salmon sperm DNA, and 0.1% SDS for 16 hours each.
0.1% SDS at room temperature followed by a final wash in 0.5 × SSC, 0.1% SDS for 30 minutes at 65°C.

The probe covers exons 1 to 10 and recognises TaqI restriction fragments of 4.6, 3.8, 2.7, 1.9, 1.8, 1.4, 0.9, and 0.8 kb.

POLYMERASE CHAIN REACTION (PCR) ANALYSIS
Exon 5 (for the proband’s family) and exon 1 (for an affected subject or known carrier from each family) were amplified using PCR. Primers for exon 5 have been previously reported.4 Exon 1 primers were, on the 5' side, 5'TCACTGCACTGAACA-CATTCTTTA3' and, on the 3' side, 5'CCTAAATCAAACCCAAGTCTCTGACC3'. The sequences were obtained from Dr M Grompe (personal communication).

Genomic DNA (500 ng) in a total volume of 100 μl was denatured at 94°C for four minutes; 2.5 units of TaqI polymerase (Promega) were added and the mix held at 50°C for five minutes and then 30 cycles of one minute at 72°C, one minute at 94°C and 30 seconds at 50°C for exon 5 and one minute at 72°C, one minute at 94°C, and 30 seconds at 56°C for exon 1.

TaqI RESTRICTION ENZYME ANALYSIS OF PCR PRODUCTS
Twenty microlitres of each amplified product were digested with TaqI according to the manufacturer’s instructions (NBL) and visualised by electrophoresis in 4% Nuseive-GTG-agarose gels and ethidium bromide staining.

ALLELE SPECIFIC DETECTION OF MUTATION
Sixty microlitre samples of amplified exon 5 from the proband’s family were denatured with 540 μl of 0.4 mol/l NaOH for 15 minutes before loading 100 μl into the wells of a MiniFiold Tm filtration manifold (Schleicher and Schuell) containing a Hybond N+ membrane presoaked in 2 × SSC. The wells were washed with 20 × SSC and filters fixed in 0.4 mol/l NaOH.

Three identical blots were prepared. One was hybridised with the wild type oligodeoxyribonucleotide probe as described in Hata et al4 and the other two with the two possible mutant sequences. All probes were end labelled with gamma 32P deoxyadenosine triphosphate. Blots were hybridised for 16 hours at 47°C before washing three times in 3 × SSC, 0.1% SDS at room temperature and once at either 53°C (mutant probe hybridisation) or 57°C (wild type probe hybridisation) for 30 minutes before autoradiography for 16 hours.

Results
TaqI RESTRICTION ENZYME ANALYSIS
A preliminary analysis of 29 patients was carried out using Southern blotting. DNA from seven male and 22 female patients was digested with TaqI and probed with the CDNA probe for OTC. A new band at 3.4 kb was observed in the proband. No such band was observed in her mother. This band would correspond to a band predicted if the TaqI restriction site normally giving rise to bands 1.8 and 1.6 kb within exon 5 were mutated. Mutations of the TaqI sites in exons 3 and 9 would also give altered bands, but none such was observed. Only one other altered band was observed after TaqI digestion. This was found in a mother and affected son but must have arisen as the result of the creation of a new TaqI site and its origin remains to be discovered.

PCR ANALYSIS OF EXON 1
The polymerase chain reaction was used to amplify exon 1 from all 29 patients, as Southern blotting does not readily detect the change in the TaqI site found in exon 1. All amplified bands were digested with TaqI, the resulting products separated on Nuseive agarose gels and visualised by ethidium bromide staining (fig 1). In each case the 217 bp amplification product was digested to a band of 172 bp showing that there were no mutations at the TaqI site in this exon. The other 45 bp band was too small to detect.

PCR ANALYSIS OF EXON 5
Genomic DNA from the proband and her mother was amplified using primers specific for exon 5. TaqI digests of the amplified products showed two bands in the case of the proband (fig 2), the expected digestion product of 121 bp from the unaffected X chromosome and an undigested fragment of 156 bp from the other X chromosome, confirming that the TaqI site has been mutated. Again the other digestion product of 35 bp was too small to detect.

DETECTION OF POINT MUTATION USING ALLELE SPECIFIC OLIGONUCLEOTIDE HYBRIDISATION
The amplified exon 5 products from the proband and her mother were tested for both predicted C to T mutations by hybridisation to radiolabelled oligonucleotides corresponding to the sense strand of the normal sequence, and to the sense strands of both mutations. The proband showed clear hybridisation to the normal oligonucleotide but with reduced intensity, as expected for a heterozygote, and to the oligonucleotide corresponding to the arginine to glutamine change (fig 3). Her mother showed only hybridisation to the normal sequence.
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Discussion
The coding sequence for ornithine transcarbamylase contains four restriction enzyme sites for TaqI. Systematic studies in other diseases, both X linked and autosomal, show that the dinucleotide \(^5'CpG^3\) is particularly prone to mutation to \(^5'TpG^3\). Examples have been found in the analysis of the factor IX gene in haemophilia B\(^10\)\(^11\) and the phenylalanine hydroxylase gene in phenylketonuria.\(^2\) The reason for this is believed to be deamination of methyl C to form T, followed by repair of the resulting G.T mismatch to A.T. As the sequence is symmetrical this can happen in either strand resulting in C to T or G to A. A small proportion of CpG sequences will fall within TaqI sites (recognition site TCGA) and may be directly detectable on Southern blots. The mutation reported here falls within that class. However, the mutation within exon 1 is difficult to detect because of coincidence of band sizes. TaqI digestion of PCR amplified exon products provides a quick and clear screening method.

We have examined 29 X chromosomes giving rise to OCT deficiency at four TaqI sites. We have only

Figure 1  PCR amplification products of exon 1 digested with TaqI. Tracks 1, 3, 5, 9, and 11 undigested product. Tracks 2, 4, 6, 10, and 12 samples digested with TaqI. Tracks 7 and 8 blanks. The band size is reduced from 217 bp to 172 bp when the TaqI site is present. The 45 bp band is too small to detect.

Figure 2  Analysis of amplified DNA from exon 5 in the family of the proband. Each pair of tracks shows the amplified product undigested on the left and digested with TaqI on the right. Tracks 1 and 2 are from the father, tracks 3 and 4 the affected girl, tracks 5 and 6 an unaffected sister, tracks 7 and 8 an unaffected brother, and tracks 9 and 10 the mother.

Figure 3  Detection of point mutation using allele specific oligonucleotide hybridisation. (A) DNA from family members hybridised with wild type oligonucleotide. Duplicate samples were applied in each case except for the brother. (B) DNA from family members hybridised with oligonucleotide detecting Arg-Gln mutation.
found one mutation and have found no deletions in eight male samples analysed. A further 15 CpG sites\textsuperscript{13} are found within the OCT coding region.\textsuperscript{4} They remain to be screened. Five other independent mutations in OCT males have been reported.\textsuperscript{37} They were initially detected either by chemical mismatch cleavage of mRNA\textsuperscript{3} or by denaturing gradient gel electrophoresis.\textsuperscript{7} Two of the five are produced by mutations within CpG dinucleotides.

This report describes the second mutation found in a female patient. So far they both reproduce mutations found in severely affected males. Further reports are awaited in order to deduce whether the severity in females is connected in any way with the mutation or whether the proportion of X inactivation is the only critical factor.

As the patient's mother showed no alteration within her TaqI site in exon 5 it is very likely that the patient represents a new mutation. However, a small possibility of mosaicism within the germ cells of the mother remains\textsuperscript{8} and she could be offered prenatal diagnosis based on PCR amplification of exon 5 and TaqI digestion if required.

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