Vitreoretinopathy with phalangeal epiphyseal dysplasia, a type II collagenopathy resulting from a novel mutation in the C-propeptide region of the molecule

A J Richards, J Morgan, P W P Bearcroft, E Pickering, M J Owen, P Holmans, N Williams, C Tysoe, F M Pope, M P Snead, H Hughes

SUBJECTS, METHODS, AND RESULTS

 Patients in the extended family were identified following independent referral of two family members to the genetics department of the University Hospital of Wales. Written consent to examination and DNA analysis was obtained in all cases as part of their clinical management. Affected subjects ranged in age from 22 to 60 years and presented with both ophthalmic and skeletal phenotypes. The vitreoretinal changes were characteristic with extensive lattice retinopathy, covering approximately two to three clock hours in each quadrant of some patients. Although the vitreous did not exhibit the congenital membranous anomaly characteristic of Stickler syndrome type 1, the architecture was strikingly abnormal, with absence of the usual lamellar array. Affected subjects had a spherical mean refractive error of –1.46 dioptres (SD 1.5), which was not significantly greater than that in unaffected subjects (mean refractive error –0.71, SD 0.99, p=0.13, Mann-Whitney test). The axial length was slightly greater in affected eyes (mean 24.66 mm, SD 0.73) compared with unaffected eyes (mean 23.8 mm, SD 1.1, p=0.008, t test). A single affected subject, whose axial length had increased to 33.5 mm, following retinal detachment surgery, distorted any apparent variation in myopia between affected and unaffected subjects and is not included in the analysis of ametropia or axial lengths. In one case (III.8) bilateral retinal detachment occurred at 13 years of age resulting in blindness. All affected subjects are at risk of rhegmatogenous retinal detachment (retinal detachment secondary to holes or tears in the retina). This is common in SEDC, Kniest dysplasia, SEMD, and Stickler syndrome.

All were considered to be of normal stature, so not all were formally measured. Of those that were, the shortest was 1.55 m (female) and tallest 1.8 m (male); there was no disproportionate stature. The skeletal changes were present in varying degrees in all but one of the affected subjects with retinopathy. The most common feature was changes to their hands. As all affected subjects were adults, it was difficult to assess the development of this feature. It was reported by the patients that at around the time of puberty they had pain and swelling of the joints in their hands, followed by disturbances in the growth of the fingers. However, the feet were normal in all cases. In the most severely affected, the fingers were recalled as being short from early childhood. The changes in the hands were highly variable with one affected subject (III.8) having a normal clinical and radiological examination (aged 34). Another (III.4) had only mild shortening of the terminal phalanges. The majority had shortening of all phalanges (figs 1 and 2), while in severe instances, involvement of the metacarpal and carpal bones was apparent together with the distal epiphyses of the radius and ulna (fig 2C). Epiphysial changes in the hips were also variable. At its most severe, premature osteoarthritis requiring the need for hip replacement surgery was evident and occurred in III.4 at 22 years of age. In this person, there was no radiological evidence of a slipped upper femoral epiphysis, or of avascular necrosis, and the shape of the proximal femurs and bony pelvis was normal (fig 3A). Radiographs of the spine from two older subjects (fig 3B-D), who had both undergone hip replacement, were essentially normal with only minor degenerative changes compatible with age seen in one case (fig 3D). The facies appeared normal without the midfacial hypoplasia usually associated with Stickler syndrome and the more severe type II collagenopathies. None of the affected subjects had a cleft palate.
Because of the involvement of both the skeletal and ocular systems, the gene for type II collagen, the major component of both cartilage and vitreous, was considered to be a good candidate for the mutant locus. Polymorphic dinucleotide repeat markers on either side of the \textit{COL2A1} gene were analysed by amplification using fluorescently labelled primers and electrophoresis through a polyacrylamide gel using an ABI 377 machine and the Genescan Analysis software. Primer sequences for markers D12S85, D12S1701, D12S1661, and D12S361 were obtained from the Genethon Linkage Map. Lod scores were calculated using the LIPEd program. Positive lod scores were obtained for markers on either side of \textit{COL2A1}, the highest being 5.4 at zero recombination with marker D12S361.

The \textit{COL2A1} gene was analysed for mutations by using amplified DNA to sequence all 54 exons as previously described. A heterozygous single base (G-A) change was seen in exon 52, which converted a codon for glycine (G1105D) to aspartic acid (fig 4). In our experience this change was novel and so, using DNA from family members, its segregation with the disease was investigated. The single base change removed a \textit{Bse}I restriction enzyme site and this was used to test the family (fig 5). All affected family members were heterozygous for the change, which was absent in all unaffected subjects. Further tests on 50 unrelated controls (100 chromosomes) were negative for the change. These were of the same ethnicity as the affected family (white); 45 were British, three were Dutch, one was Anglo-American, and another Greek. In addition, 35 samples from white British subjects with Stickler syndrome, SEDC, or Kniest dysplasia were negative for the change as determined by sequencing.

Analysis of illegitimate \textit{COL2A1} transcripts using cultured dermal fibroblasts from an affected subject showed that both alleles were amplified from the mRNA of these cells (fig 4). Amplification of transcripts was achieved essentially as previously described. An antisense primer in the 3' untranslated region (tggaagcactggtgctttg) was used for the reverse transcription reaction. Primers in exon 36 (ccagctggtgctaacg-gcagtttttgg) and 53 (tgaacctgctattgccctctgccc) were used in the primary PCR. Primers for the secondary PCR were nested in exons 39 (agctgtggtcgtacg-ggagcttctg) and 53 (tggaagcactggtgctttg) were used in the primary PCR. Primers for the secondary PCR were nested in exons 39 (agctgtggtcgtacg-ggagcttctg) and 53 (tggaagcactggtgctttg) were used in the primary PCR. Primers for the secondary PCR were nested in exons 39 (agctgtggtcgtacg-ggagcttctg) and 53 (tggaagcactggtgctttg) were used in the primary PCR. Amplification products were sequenced with a primer in exon 51 (ccagctggtgctttg). The observed heterozygosity in amplified cDNA supported the observation of a missense mutation, rather than haploinsufficiency, through nonsense mediated decay, causing the disorder.

Amino acid sequences for the fibrillar collagens were determined from the published cDNA sequences and aligned using the computer program PILEUP. By comparing the sequences for all nine human fibrillar collagens, it was seen that the altered amino acid was within a highly conserved region of the C-propeptide (fig 6). The usual convention for numbering amino acids in collagen molecules is to assign the first glycine of the triple helical domain as 1. Using this system the mutation is G1105D. However, different numbering systems have been used when describing alterations to propeptide regions. For instance, Unger \textit{et al} numbered amino acids from the initiating methionine, but excluded amino acids encoded by
exon 2, which is not expressed by mature chondrocytes. Exon assignment in some published reports is also confusing. Some reports number the 54 exons 1-54, while others use the numbering system of Chu and Prockop, where exons 1-5 are numbered 1, 1A, 2, 3, and 3B. Consequently, exons 6-54 are numbered 4-52. For clarity, the different missense mutations in the C-propeptide of type II collagen are listed in table 1, amino acids are numbered from the start of the C-propeptide (DQAAG), and exons numbered 1-54.

DISCUSSION

The family we have described here have an unusual phenotype with involvement of the hands. As they have normal stature they cannot be classified as SEDC, SEMD, or Kniest dysplasia, yet they are not typical of Stickler syndrome, the most common type II collagenopathy, where stature is normal. Like Stickler syndrome this family presented with dominantly inherited retinal detachment, lattice retinopathy, and premature osteoarthritis. Stickler syndrome is highly variable, and it is not unusual for an affected subject to present with minimal features of the disorder. Indeed, some mutations result in a predominantly ocular form of Stickler syndrome. It is, however, unusual for all affected members of a large family, with both retinal detachment and premature arthropathy, as described here, to have no instance of high myopia, cleft palate, or midfacial hypoplasia. Cleft palate, for instance, is commonly associated with mutations in COL2A1, The family described here also lacked the congenital membraneous vitreous anomaly usually present in cases of Stickler syndrome, instead exhibiting highly disorganised vitreous lamellae. This presumably reflected the different nature of the mutation which if expressed would result in mutant proα1(II) collagen chains. Unfortunately, we did not have cartilage tissue available to confirm expression of mutant protein. However, like other examples of missense mutations in COL2A1, heterozygosity of the mutation was observed in illegitimate transcripts, whereas premature termination mutations exhibit nonsense mediated decay and a homozygous normal cDNA sequence.

The skeletal phenotype varied, but the most striking trait in this family was the phalangeal epiphyseal dysplasia in the hands, which in most cases the patients reported appeared in late childhood. Brachydactyly has been noted in other cases of COL2A1 mutations. A family with an Arg704Cys mutation was described as “Stickler-like”. They had short stubby fingers along with myopia, a flattened midface, and mild generalised epiphyseal dysplasia. A 5 bp duplication in exon 53 resulted in a premature termination codon in the same exon and a phenotype described as spondyloperipheral dysplasia. In this case, where there was short stature and myopia, both hands and feet were affected with shortened digits. Again the family described here is different from these two cases. Two other COL2A1 missense mutations have previously been described in the C-propeptide. A threonine to asparagine at position 149 resulted in hypochondrogenesis, while a threonine to methionine at position 198 caused SEDC. Like the mutation described here, these were both close to cysteines (fig 6). Certain cysteines, in the C-propeptides of type I and type III collagens, form intrachain disulphide bonds before chain association. In type II collagen, the corresponding cysteines are at positions 82-244 and 152-197. Other cysteines
form interchain disulfide bonds after chain association. So, unlike the mutation described here which is adjacent to Cys65 that forms an interchain disulfide bond, T149N and T198M are close to cysteines that form a single intrachain disulfide bond. It is tempting to speculate that these three mutations are close to cysteines that form a single intrachain disulfide bond, possibly delaying disulfide bond formation, leading to excess post-translational modifications. In particular, we have not yet determined if mutant G64D α chains are capable of assembly and if mutant molecules are modified and secreted normally.

It should also be remembered that the COL2A1 gene also encodes proα1(XI) collagen and so these mutations will affect type XI collagen as well. Mutations in the C-propeptide region of type I collagen have also been described, in particular a mutation which substituted a cysteine in the C-propeptide region of proα1(1) collagen, disrupted an intrachain disulfide bond, and resulted in a mild form of osteogenesis imperfecta.

In summary, the disorder in the family described here was linked to COL2A1 and a novel glycine to aspartate substitution was detected. Analysis of cDNAs suggested that both alleles were expressed, as nonsense mediated decay was not apparent. Sequencing of all 54 exons showed no other amino acid changes. Family analysis showed that the aspartate containing allele segregated with the affected subjects; this change was also absent in a panel of control chromosomes. The mutated amino acid is highly conserved in all fibrillar collagen molecules. These observations strongly indicate that this is the causative mutation in this family. Some instances of mutations in COL2A1 can result in unusual phenotypes (that is, spondyloepiphyseal dysplasia) that do not easily fit into the more established criteria for other type II collagenopathies, such as SEDC or the Kniest and Stickler dysplasias. This family also presented with an unusual combination of phenotypic traits. The most consistent were vitreoretinopathy (all are at risk of retinal detachment) and the developmental abnormalities that occurred in the fingers. Other aspects of the skeletal phenotype were variable; in some subjects severe premature osteoarthritis necessitated hip replacement, the youngest being at 22 years. However, there were no spinal dysplastic features. The intriguing developmental changes that occur to the fingers make this family highly unusual and are most likely the result of the novel nature of the mutation in the C-propeptide.

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Authors' affiliations

A J Richards, Department of Pathology, University of Cambridge, Cambridge, UK

J Morgan, Department of Ophthalmology, University Hospital of Wales, Cardiff, UK

P W P Bearcroft, Department of Radiology, Addenbrooke’s NHS Trust, Cambridge, UK

E Pickering, H Hughes, Department of Clinical Genetics, Ysbyty Glan Clwyd, N Wales, UK

M J Owen, N Williams, Department of Neuropsychiatric Genetics, University of Wales College of Medicine, Cardiff, UK

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