The clinical features of Ehlers-Danlos syndrome type VIIB resulting from a base substitution at the splice acceptor site of intron 5 of the COL1A2 gene


Abstract
The features of a 32 year old woman with Ehlers-Danlos syndrome type VIIB and affected members of her family, resulting from a mutation in one COL1A2 allele, were studied. Her dermal type I collagen contained α2(I) chains and mutant pN-α2(I) chains in which the amino-terminal propeptide remained attached to the α2(I) chain. She was heterozygous for an AG→AC mutation at the splice acceptor site of intron 5 of the COL1A2 gene. The mutation activated a cryptic AG splice acceptor site corresponding to positions +14 and +15 of exon 6 of the COL1A2 gene. In contrast to previous reports only five, rather than all 18, amino acids encoded by exon 6 were deleted in the proband. The deleted peptide removed the amino-proteinase cleavage site, but not the nearby lysine cross linking site, of the amino-telopeptide of the α2(I) chain.

She was born with bilateral hip dislocations, knee subluxations, and generalised joint hypermobility. Bilateral inguinal herniae and an umbilical hernia were present at birth. Facial features included a depressed nasal bridge with prominent paranasal folds. The skin was soft, moderately hyperelastic, and sagged over the face. Skin fragility and easy bruising were apparent from childhood. Skin wounds healed slowly and with broad, paper thin scars. Throughout her life, she had multiple fractures of the small bones of her hands and feet following moderate trauma.

Electron microscopy of the proband's dermis as well as deep fascia and hip joint capsule from her affected brother showed that collagen fibrils in transverse section were nearly circular but with irregular margins. Light microscopy of bone from her affected brother and son showed normal Haversian systems and lamellae bone. All of these tissues contained approximately equal amounts of the normal and mutant α2(I) chains.

The findings of this study confirm that loss of the amino-proteinase cleavage site of the pro α2(I) collagen chains, owing to anomalous splicing of exon 6 sequences in the conversion of pre-mRNA to mRNA, produces the clinical features of Ehlers-Danlos syndrome type VIIB. The history of frequent fractures found in this family is atypical and indicates an overlap with osteogenesis imperfecta.

Type I collagen is the major fibrillar collagen of dermis, ligament, tendon, and bone. It is synthesised as a larger precursor, procollagen, consisting of two pro α1(I) and one pro α2(I) chains. During, or shortly after, secretion the amino(N)- and carboxy(C)-terminal propeptide extensions of the pro α chains are removed by N- and C-proteinases, respectively, to yield the mature chains of type I collagen. Genetic defects of the N-proteinase cleavage sites or of the N-proteinase enzyme produce the type VII variant of the Ehlers-Danlos syndrome (EDS). The main phenotypic features of this variant of EDS are congenital joint dislocations and severe generalised joint hypermobility. EDS type VIIIA results from mutations that remove the N-proteinase cleavage site of the pro α1(I) chains. EDS type VIIIB results from similar defects involving the pro α2(I) chains. EDS type VIIIC results from a deficiency of N-proteinase.

The two reported cases of the VIIA variant were the result of a G→A transition at the −1 position of the splice donor site of intron 6 of the pro α1(I) gene (COL1A1). This mutation results in alternative splicing with products lacking exon 6 encoded sequences, which includes the N-proteinase cleavage site, or including exon 6 sequences and a substitution of methionine-159 by isoleucine. The clinical features of these two cases were similar.

Ten cases of the type VIIIB variant have been described; three were the result of G→A transition at the −1 position of intron 6, as in the type VIIA cases, except that they involved the COL1A2 gene. Six cases had point mutations of the GT dimerucleotide splice donor site of intron 6 which resulted in the complete loss of exon 6 encoded sequences from the α2(I) mRNA and chain. The deleted sequence included the N-proteinase cleavage site and the cross linking lysine site in the N-telopeptide of the α2(I) chain. In the case reported here, there was partial loss of exon 6 encoded sequences owing to a base substitution at the splice acceptor site of intron 5 of the COL1A2 gene which activated a cryptic splice site at positions +14 and +15 of exon 6. The N-proteinase cleavage site was lost but the
The clinical features of Ehlers-Danlos syndrome type VII B

nearby cross linking lysine site in the N-terminal peptide was retained.

Dermatosparaxis (EDS type VII C) was first recognised as a recessively inherited disorder of cartilage and was subsequently described in dogs, cats, and sheep.15-20 Two human cases of type VII C, owing to N-proteinase deficiency with retention of the N-propeptides of pro α1(I) and pro α2(I) chains, resemble clinically and histologically the animal equivalent, dermatosparaxis.21, 22

We report the phenotypic features of EDS type VII B resulting from the partial loss of exon 6 encoded sequences from the COL1A2 gene. The findings suggest that the retention of the N-propeptide is the main determinant of the phenotype.

Case report

CLINICAL FEATURES

The proband was the eighth child of unrelated parents (fig 1). She was born at 37 weeks’ gestation by normal vaginal delivery and weighed 2500 g. She had bilateral hip dislocations which were treated by spinline and later by serial plaster of Paris casts. Both knees were hyperextended with the tibiae subluxed anteriorly. She also had bilateral inguinal herniae and an umbilical hernia at birth; they were managed without surgery.

Her height was between the 25th and the 50th centiles throughout growth. Walking was delayed until she was 2.5 years old. In early childhood, her generalised ligamentous laxity produced frequent, painful patellar and shoulder dislocations. She was usually able to reduce them herself. By the age of 7 years she was independently mobile. She had learnt to reduce the frequency of joint dislocations and subluxations by avoiding running and other activities. She had mild postural pes plano-abducto-valgus deformities of the feet. Her finger joints, particularly the interphalangeal joints, were hypermobile with swan neck deformities. During puberty she developed a mild, non-progressive thoracolumbar scoliosis measuring 25°.

She had frequent fractures throughout childhood, particularly of the small bones of her hands and the distal radius and ulna. Approximately 20 to 30 fractures occurred after moderate trauma. Her susceptibility to fractures also prevented her from participating in sports.

Her skin was lax, velvety, and easily stretched. She had paper thin scars, particularly in the pretibial region. Her skin healed slowly after even minor trauma. Throughout her life she bruised easily and for prolonged periods. After the birth of each of her children, she developed post partum haemorrhage requiring dilatation and curettage of the uterus.

Her sight and hearing were normal. She had recurrent chest infections as a child and asthma for which she received intermittent antibiotics and bronchodilators.

At the age of 29 years she developed, over a period of 48 hours, respiratory symptoms and progressive weakness in her arms and legs and retention of urine. A diagnosis of transverse myelitis, probably unrelated to her EDS, was made. Her condition improved gradually over the next two years.

She was reviewed at the age of 32 years. Her facial appearance showed mild depression of the nasal bridge with prominent paranasal folds (fig 2). She continued to have stretchable, fragile skin (fig 3). She was confined to a wheelchair because of weakness of her arms and legs that resulted from a spinal cord lesion at the C5 level. In the presence of these neurological problems her generalised joint hypermobility became less troublesome. Shoulder movements were full, but there was marked laxity of the shoulders with excessive anteroposterior translation and a groove beneath the acromian indicating inferior subluxation. The superior radioulnar joints were subluxed. Her finger joints were lax (fig 4) and she was able to

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Figure 1. The family pedigree.

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Figure 2. Facies of the proband showing a depressed nasal bridge with slightly prominent paranasal folds.
Figure 3 Appearance of the skin on the dorsum of the proband’s hands showing hyperelasticity.

Figure 4 Appearance of the proband’s fingers showing marked laxity of the interphalangeal joints which could be manipulated into marked hyperextension.

Figure 5 Appearance of the proband’s hands showing swan neck deformities of the fingers.

Figure 6 Appearance of the proband’s feet showing plano-ab ducto-valgus deformities.

Figure 7 Anteroposterior radiograph of the pelvis of the proband’s affected brother showing a dislocation of the hip with gross deformity of the proximal femur and narrowing of the joint space. A proximal femoral osteotomy had been performed in childhood.

voluntarily produce swan neck deformities of her fingers (fig 5). Her hip movements were restricted in flexion and rotation while her knees flexed fully but hyperextended 20°. The patellofemoral joints were excessively lax and she was apprehensive about lateral movements of her patellae. She had plano-abducto-valgus deformities of the feet (fig 6).

FAMILY HISTORY

The family pedigree showed affected members in three generations consistent with autosomal dominant inheritance of the syndrome (fig 1). The proband’s affected father was also born with bilateral hip dislocations. He had marked joint hypermobility with intermittent patellar, interphalangeal, wrist, and shoulder dislocations and subluxations throughout his life. He bruised easily and produced poor, thin scars after trauma.

The proband had two affected sibs with similar clinical phenotypes. The female sib also sustained multiple fractures in childhood, mainly of the small bones of the hand, but also of the distal radius and olecranon.

The affected male sib was reviewed at 35 years of age when he underwent a total hip joint replacement. He was born with bilateral hip dislocations which required open surgical reductions in the first few years. One hip became osteoarthritic (fig 7) and was replaced at the age of 35 years. He was also disabled by the marked swan neck deformities of his hands and underwent reconstructive surgery in an attempt to reduce the deformities (fig 8). He was of normal height. He had easy bruising, skin hyperelasticity, and multiple fractures of the metacarpals, the distal radius, and distal ulna. He also sustained a fracture of the patella and olecranon. The frequency of fractures reduced markedly after his teenage years. His nasal bridge was mildly depressed and he had mild paranasal folds (fig 9).

The proband had three children of whom only the first was affected by the syndrome. He
The clinical features of Ehlers-Danlos syndrome type VIIB

The clinical features of Ehlers-Danlos syndrome type VIIB were characterized by a combination of skin, joint, and vascular abnormalities. The proband, a boy aged 36 weeks, was born by vaginal delivery at 36 weeks’ gestation after premature rupture of membranes. He had bilateral hip and knee dislocations and generalized joint hypermobility. His hip dislocations were treated with plaster of Paris spica casts. An umbilical hernia was present at birth. He died suddenly at 2.5 months of age. Necropsy showed hugely dilated large intestine proximal to a kink of the sigmoid colon at the level of the pelvic brim. This was the only abnormality found to account for his death.

PATHOLOGICAL FEATURES
Specimens of skin from the proband, skin, deep fascia, and joint capsule from her affected brother, and similar tissue from a control were fixed in 2.5% glutaraldehyde, postfixed in osmium tetroxide, and embedded in Spurr’s resin. Thin sections were stained with uranyl acetate, lead acetate, and tannic acid.

Electron microscopy showed that the proband’s dermal collagen fibrils were round with only mild surface irregularities in transverse section (fig 10A,C), much less than the irregularities observed in a case of Ehlers-Danlos syndrome type VIIA (fig 10D). The affected and control specimens from the joint capsule and deep fascia showed some areas with a single population of fibrils and other areas with two populations of fibrils (fig 11A,B). The transverse sections did not show significant irregularity of the margins of the fibrils in either area.

In the proband’s son who died at 2.5 months of age, the sigmoid colon was kinked with proximal dilatation of the colon. Light microscopic sections from the thoracic and abdominal viscera and a vertebra did not show any anomaly of the connective tissues. Re-processing of the paraffin embedded pericardium for transmission electron microscopy showed that collagen fibrils, in transverse section, had mainly smooth rounded outlines (fig 10B).

Light microscopy of bone from the femoral head of the proband’s brother showed normal lamellar bone and well formed Haversian systems.

BIOCHEMICAL FEATURES
From the proband’s affected brother, bone and fascial collagens were prepared using previously described methods and shown to contain approximately equal amounts of normal α2(I) and mutant pN α2(I) chains (fig 12). Previous studies showed that the DNA extracted from formalin fixed and paraffin embedded samples of spleen and decalcified bone from the proband’s affected son contained the same mutation.

Discussion
The proband and her affected relatives had the clinical features of Ehlers-Danlos syndrome type VIIB with stretchable, fragile skin and gross joint hypermobility with multiple dislocations that were obvious at birth. These features as well as the electron microscopic finding of mainly rounded collagen fibrils were similar to those reported in other cases of EDS type VIIB. A major difference, however, was the high frequency of fractures in affected members of the current family. Transverse myelitis in the proband has not previously been reported in EDS. Its aetiology was not determined but it was probably unrelated to EDS as the onset and progress of the illness were typical of transverse myelitis.

The findings of approximately equal amounts of normal and mutant α2(I) chains in the dermis, deep fascia, joint capsule, and bone suggest that all type I collagen containing tissues had a similar composition. This proposal is supported by similar observations in
Figure 10  Transmission electron micrographs of collagen fibrils in the dermis of the proband (A), pericardium of the affected son (B), and dermis of the affected brother (C). The fibrils from affected persons with EDS type VIIIB are nearly circular in cross section. They are more irregular than the control fibrils but less irregular than the fibrils of a case of EDS VIIA.29

Figure 11  Transmission electron micrographs of collagen fibrils in the hip joint capsule from the proband’s brother (A) and a control (B). A double population of fibrils of near circular outline is shown in both micrographs.

Figure 12  Pepsin digested collagens from the bone and deep fascia of the proband’s affected brother were resolved by polyacrylamide gel electrophoresis. Proteins were stained with Coomassie blue. Lane 1 is bone and lane 2 deep fascia. The identities of the type I collagen chains are shown and the mutant chain is designated pN α2(I). The results are consistent with the tissue containing equal amounts of abnormal pN α2(I) and normal α2(I) chains.
The recipient of mutations involving type EDS fragility overlap an examination of the skin and joint abnormalities observed in EDS type VII. The abnormal collagen composition of the bone also provides an explanation for the bone fragility that was a feature of affected members of the present family. However, the bone, in keeping with the dermis, fascia, and joint capsule, did not show any light microscopic changes in the extracellular matrix. Similarly, no abnormalities in trabecular architecture or density of the bone were evident from standard radiographs.

The bone fragility observed in the present family may reflect the uniqueness of the mutation which resulted in the loss of the N-proteinase cleavage site and retention of the nearby lysine cross linking site in the N-terminal of the α2(I) chains. Alternatively, bone fragility may have been overlooked in other reported cases of EDS type VII. In support of the latter proposal, a further review of a previously reported case of EDS type VIIA showed that the child had sustained numerous fractures with minor injuries.

The similar clinical and histological features of the reported cases and the current case of EDS VIIB suggest that the loss of the N-proteinase cleavage site with retention of the N-propeptide is of greater importance in the pathogenesis of the disorder than the loss of the cross linking lysine residue in the N-terminal of the α2(I) chain. This proposal is supported by the features of EDS type VIIIC (dermatosparaxis) in which a deficiency of N-proteinase activity results in the retention of the N-propeptides without loss of the cross linking lysine sites in the N-telopeptide. Affected persons showed severe laxity and fragility of the skin, inguinal herniae, blue sclerae, micrognathia, and growth retardation. The collagen fibrobls show hieroglyphic forms in cross section which probably result from the adverse effect of the persistent pN α2(I) and pN α1(I) chains on collagen fibrillogenesis.

The current features of the current family indicate an overlap between EDS type VII and osteogenesis imperfecta. In the latter disorder, bone fragility is the main clinical feature but ligament laxity is also frequent and may be severe, so producing mixed osteogenesis imperfecta and EDS phenotypes. In such cases, mutations involving the triple helical domain of type I collagen misalign the N-proteinase cleavage sites and retard the processing of pN collagen to collagen.