Ehlers-Danlos syndrome has varied molecular mechanisms

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Ehlers-Danlos syndrome (EDS) is a group of variable clinical entities which share a propensity to skin fragility, joint laxity, and ligamentous fragility or shortening. Tschernekobow, Ehlers, and Danlos independently described unusual bruising, excessive cutaneous extensibility, and molluscoid pseudotumours. The eponym of Ehlers-Danlos syndrome (EDS) was first suggested by Pomeau-Delille and Soulé. Spectacular early examples were retrospectively commented upon by Beighton. Morris described a notable British patient who Beighton re-examined more than 60 years later. The proband had classical EDS. Beighton not only showed impressive pictures of the so-called “Elastic Lady” but also included the original lithograph of a Spaniard with unilateral cutaneous hyperextensibility investigated in Leyden almost 300 years before, who was very probably a somatic mosaic. In Ehlers’s time there was considerable difficulty in distinguishing clearly between cutis hyperelastica (overextensible skin), dermatorrhesis (easy splitting of skin), and both localised and generalised dermatochalasis (pendulous or redundant skin). To this day the similarities of such physical signs still cause confusion. In practice the designations hyperelastic, pendulous, and lax skin often overlap and can even occur at different times in the same patient. For example, EDS skin is hyperelastic early in life but later becomes lax or drooping, especially in old age. The systemic implications, complications, and associations of EDS were largely unrecognised for many years although the association of congenital hip dislocations was described early. In 1960 Mories first described a four-fold increase in fetal prematurity. He also noticed that the dermis was collagen depleted and elastin rich. Thirdly, he described catastrophic lethal arterial bleeding in a 15 year old adolescent boy with a traumatic arterial tear which was surgically unreparable because of the extreme venous and arterial fragility. Very probably the patients with prematurity had EDS I/II while the vascular fragility was caused by acrogeric EDS IV. Subsequent clinical, molecular, and genetic progress has been rapid. For example, Barabas first proposed three distinct subsets. These included classical forms with prematurity, a milder type with venous varicosity, and a third, potentially lethal, form with minor cutaneous and joint changes but extreme arterial fragility (which we now call EDS IV). Soon afterwards, Beighton also clearly described the frequent prematurity. He added an X linked form, while dividing Barabas’s first group into “gravis” and “mitis” variants, which we now call EDS I and II respectively, to make five subtypes. He also delineated the wide clinical heterogeneity with various rheumatological, orthopaedic, surgical, and cutaneous complications and was also fully familiar with such surgical complications as arterial aneurysms, venous varicosities, arteriovenous fistulas, and inguinal, umbilical, and hiatus hernias. He also documented reflux hydronephrosis, bladder neck obstruction, colonic diverticulae with perforation, and pleuroperitoneal rupture (leading to pneumo- or haemothorax). These complications are now recognised as particularly common in EDS types IV, VI, and VII. In 1972, McKusick added two further subtypes (types VI and VII). We now know that the former is caused by lysyl underhydroxylation, while the latter results from the misprocessing of procollagen (to collagen). In 1988 Beighton et al published an International Nosology of Connective Tissue Disease where nine subcategories of EDS were defined. Subsequently, he correlated the subtypes with the various biochemical and molecular abnormalities (table 1) and also published the equivalent Molecular Nosology.

Keywords: Ehlers-Danlos syndrome; molecular mechanisms

Genetics of EDS (table 1) Autosomal dominant or autosomal recessive inheritance occurs in EDS I, II, III, IV, VI, and VII. Compound heterozygosity (see later) is also very likely. X linked forms have been described but are highly unusual. Autosomal dominant inheritance was first mentioned by Kopp. Wiener described 12 affected subjects in three generations. Stuart and Coe and Silver both described the vertical transmission particularly of EDS type I. Johnson and Falls published a very large five generation American family of British-Canadian extraction with typical EDS I and with two separate
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homoygotes. They also speculated upon autosomal recessive inheritance while both Weber and Aitken22 and Ronchese21 separately described sporadic examples of EDS I with affected first cousin parents.

Autosomal recessive inheritance occurs in EDS VI although compound allelic heterozygosity may be more common than homozygosity except in consanguineous or inbred populations. Beighton1 first suggested autosomal recessive inheritance in an affected brother and sister whose parents and children were normal. Convincing autosomal recessive inheritance also occurs in procollagen peptidase deficiency. In humans this causes EDS VIIc and in animals dermatosparaxis. While the animal skin is excessively fragile, in humans there is extreme joint laxity and blepharochalasis/cutis laxa (CL). The original EDS VII patient of Lichtenstein et al2 had congenital hip dislocations and short stature. Skin from EDS VIIc patients, cattle, sheep, and cats show severely disorganised and misassembled hieroglyphic collagen fibrils when examined by transmission electron microscopy. Homozygosity or double heterozygosity has also been postulated in EDS IV.12 23-27 The latter patient had a normal collagen III profile. The family of Beasley and Cohen28 with joint laxity, lop ears, and an unusual face may also be an example of autosomal recessive inheritance but was unclassifiable clinically. Apparent face synomy of EDS X also occurs in EDS I. Some families with clinical phenotype resembling EDS II. One family contained six affected males and three possible carrier females in four generations. Patients with EDS X may have joint laxity, hypermobile fingers, and ecchymosis, and families with EDS X are now well understood. Most disrupt collagen fibril assembly by altering certain crucial molecular components, the anatomical distribution of which dictates the subsequent molecular pathology. Thus the molecular composition of regions such as skin, vasculature (arteries, veins, and capillaries), pleuroperitoneum, intestinal walls, ligaments, tendons, cartilage, and eyes (the cornea and vitreous) may be severely weakened and disorganised.

Many types of EDS have specific mutations, while in others, such as EDS III, VIII, and X, the abnormality is either unknown or inconsistent. Generally, there are two classes of mutations, one in which structural genes are faulty and the other in which their processing enzymes are disturbed.27 Examples of the former are mutations of the COL1A1, 1A2, 3A1, and 5A1 genes and of the latter lysyl oxidase dependent form of EDS has not been confirmed.

Molecular abnormalities causing EDS (table 1)
In keeping with recent rapid advances, the molecular mechanisms of many EDS subtypes are now well understood. Most disrupt collagen fibril assembly by altering certain crucial molecular components, the anatomical distribution of which dictates the subsequent molecular pathology. Thus the molecular composition of regions such as skin, vasculature (arteries, veins, and capillaries), pleuroperitoneum, intestinal walls, ligaments, tendons, cartilage, and eyes (the cornea and vitreous) may be severely weakened and disorganised.

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Table 1  Ehlers-Danlos syndrome (cutis hyperelastica) after Beighton et al22

<table>
<thead>
<tr>
<th>Type</th>
<th>Synonym</th>
<th>McKusick No</th>
<th>Special features</th>
<th>Histology electron microscopy</th>
<th>Basic defect</th>
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<td>EDS I</td>
<td>Gravis type</td>
<td>AD 130000</td>
<td>Widespread scarring and bruising, especially forehead, chin, and shin. Molluscoid pseudotumours</td>
<td>Cauliflower fibrils</td>
<td>COL5A1 linked in some families. Mutations include exon skips and a translocation and a cysteine substitution</td>
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<td>EDS II</td>
<td>Mitis type</td>
<td>AD 130010</td>
<td>Similar but less severe</td>
<td>Cauliflower fibrils</td>
<td>Unknown</td>
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<td>EDS III</td>
<td>Hypermobile type</td>
<td>AD 130020</td>
<td>No cutaneous scars</td>
<td>Collagen depletion, variation of fibre size</td>
<td>COL3A1 mutations. Numerous point mutations and exon skips, rarely deletions</td>
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<td>EDS IV</td>
<td>Vascular type</td>
<td>AD 130050</td>
<td>Nearly always type III collagen deficient</td>
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<tr>
<td></td>
<td>1VA acrogeric</td>
<td>AR 22535</td>
<td>Risk of arterial rupture highest in acrogeric subtypes</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>1VB acrogeric</td>
<td></td>
<td>Very rare. Not lysyl oxidase deficient. Reminisces EDS I, II, and III</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>IVC ecchymotic</td>
<td>AD 130050</td>
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<tr>
<td>EDS V</td>
<td>X linked type</td>
<td>XL 305200</td>
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<td>EDS VI</td>
<td>Ocular-scoliotic</td>
<td>AR 225400</td>
<td>Muscular hypotonia, often muscular dystrophy suspected (slow motor milestones)</td>
<td>Non-specific</td>
<td>Lysyl hydroxylase point mutations or exon skips (homozygosity and double heterozygosity)</td>
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<td></td>
<td>VIA decreased lysyl</td>
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<td>hydroxylase levels</td>
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<td>VB normal levels</td>
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<tr>
<td></td>
<td>Arthrochalasis</td>
<td>AD 130060</td>
<td>Persistent premature scoliosis. Arterial rupture in 30%</td>
<td>Fibrils vary from angular in A and B to hieroglyphic in C</td>
<td>Types A &amp; B specific exon 6 skips or deletions of COL1A1, 1A2 either pNa1(1) or pNa2(1) retained. Type C procollagen peptidase deficiency. Both pNa1(1) and pNa2(1) extensions retained</td>
</tr>
<tr>
<td></td>
<td>multiplex congerita</td>
<td></td>
<td></td>
<td></td>
<td>Some are COL3A1 others not</td>
</tr>
<tr>
<td>EDS VII</td>
<td>Periodontitis type</td>
<td>AD 130080</td>
<td>Allelic variation with variable expression</td>
<td>Non-specific</td>
<td>Association with FN may be coincidental</td>
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<td>EDS X</td>
<td>Fibronectin abnormality</td>
<td>AR 225310</td>
<td>Fibronectin deficient</td>
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Typical clinical features of EDS type I and II are similar but milder. Shown are the characteristic facial scars (A), extreme cutaneous extensibility (B), and palmar and pretilial scars (C). Broadened feet and hands are also a feature in some families, as shown in this affected father and child (D).

Clinical features of EDS types I and II (fig 1)

The five cardinal clinical elements of EDS include hyperextensible, doughy skin, atrophic scars, joint hypermobility, connective tissue fragility, and bruising (table 1). The clinical phenotypes of EDS I and II overlap substantially and are probably allelic autosomal dominant mutations with variable penetrance. Compound heterozygosity is also possible. EDS type I differs from type II only in the degree of skin fragility and ligamentous laxity. The skin splits over prominences such as the forehead, chin, elbows, knees, and shins. Broadened hands and feet and a wide body build (mesomorphism) are also common. Other notable clinical signs include cutaneous varicosities, myopia, and late onset osteoarthritis (OA). Since collagen α1(V) chains occur in cartilage and vitreous humour, the OA is probably a direct complication rather than primarily mechanical. In EDS II the clinical phenotype is milder and skin fragility much less impressive; epicanthic folds, fibrous nodules, and mesomorphism are infrequent or even absent.

Collagen type V and the role of COL5A1/COL5A2 genes

There is now strong evidence to implicate type V collagen α1 chains in the aetiology of EDS I and II. Clinical features include excessive skin fragility, ligamentous laxity, and relatively benign prognosis. The collagen is disorganised and forms so-called cauliflower fibrils.

Like collagens I, II, III, and XI, collagen V has an uninterrupted triple helix with N and C terminal extensions. There are two or possibly three forms of collagen V proteins, at least two of which, α1(V) and α2(V), are coded by separate non-allelic genes. Firstly, on clinical grounds, there are very severely disorganised collagen fibrils (cauliflower) (fig 2D). Secondly, abnormalities of both collagen type V α1 chains and the COL5A1 gene have been described in a sporadic EDS I/II patient with increased skin fragility and joint laxity with corneal flattening. Furthermore, transgenic homozygous mutant mice with an in frame exonic deletion in Col5a2 also had fragile skin, dermal thinning, and corneal disorganisation as well as severe skeletal and ligamentous deformities. An equivalent rabbit model shows similar pathology. Several independent groups have described linkage of human COL5A1 markers in families with EDS I or II which are therefore allelic. Other COL5A1 mutations have also recently been reported. In one, a translocation interrupts the COL5A1 gene at intron 24 and in the second an exon 65 skip was found in a three generation family. A third had a mutation of a highly conserved C terminal cysteine. There are therefore at least three published COL5A1 mutations with either EDS I or II phenotypes. In other similar families, linkage to COL5A1 has been excluded (Burrows et al, unpublished data). Somehow, collagen type V controls type I collagen fibril packing. This credibility explains the gross fibrillar disorganisation which typifies EDS I and II and implies that the disorganisation is a direct effect of the mutant type V collagen. Allegedly, collagen type XI analogously regulates the thickness of collagen type II fibrils and collagen V and XI coassociate with or even substitute for one another in vitreous humour. Corneal shape is also collagen V dependent. Mutations of either collagen type I gene (COL11A1 or COL11A2) can cause variants of the Stickler syndrome (SS) in humans and mice. Since COL11A1 but not 11A2 is expressed in the vitreous, only mutations of the former cause vitreous pathology. Collagen α2(V) chains coassociate with α2(XI) chains, which may both coassociate with collagen type II. It is therefore not surprising that corneal or vitreous abnormalities and high myopia cosegregate with premature OA in some EDS families. Furthermore, COL5A2 mutations are a potential cause of arterial fragility, suggesting the possibility that α2(V) chains and collagen type III interact analogously to α1(V) chains and type I collagen chains. Lastly, several
different collagen mutations cause disordered fibrillogenesis in vivo. COL5A1 homozygotes are either exceptionally rare or genetic lethals. Good animal models for COL5A1 mutation will be essential for the future detailed analysis of the chemical composition and 3D organisation of tissues such as articular cartilage, ligament, cornea, vitreous humour, arteries, veins, and capillaries.

Clinical features of EDS type IV (vascular Ehlers-Danlos syndrome) (fig 3)

Vascular/arterial EDS differs from other varities by virtue of severe vascular fragility, both venous and arterial. It was originally described by Barabas and Sack. In echymotic EDS there are often post-haemorrhagic haemosiderin deposits over the knees, shins, and elbows, which may also complicate EDS I/II and VIII. Acrogeria is specific to EDS IV and is characterised by generally thinned skin; the extremities (hands, feet, and face) are prematurely aged (fig 3A, B). Other suggestive features include prominent capillaries and keloidal or elastic scars (often with elastosis perforans serpiginosa). Large eyes, lobeless ears, and a Madonna-like face are also typical. Although short stature is usual, occasional slim and tall people overlap with MFS. Other clinical features include acro-osteolysis, diffuse alopecia of the scalp, and certain orthopaedic complications such as congenital talipes or hip dislocations and tendon contractures, particularly of the Achilles or extensor tendons of the feet and toes. Pleuropertoneal or colonic rupture are more non-specific (and may also complicate other EDS or MFS phenotypes).

Vascular pathology includes aneurysms of small to medium sized arteries such as renal, splenic, axillary, brachial, femoral, popliteal, and internal carotid vessels. The internal carotid system may be compromised by dilatation, arteriovenous malformations, aneurysms, or dissections caused by arterial thinning and fragility from collagen III deficiency. Occasionally the general phenotype is EDS III/BHS. For example, we observed a Gly->Ser 637 substitution in a large autosomal dominant pedigree with generalised joint laxity and premature osteoarthritis.

In EDS IV, skin histology shows striking dermal thinning (from one-third to two-thirds of normal), collagen depletion, and elastin proliferation. The mechanism of the latter is unclear (and may be primarily that secondary to collagen depletion). Transmission electron microscopy typically shows collagen fibril disorganisation with irregularity and a bimodal size distribution (fig 2E). Somehow collagen III/IV ratios influence collagen fibril diameter and interactions and, if mutated or diminished, seriously impair long term arterial strength and stability.

Protein chemistry

There is very strong clinical, histological, biochemical, and molecular evidence that faulty collagen III causes dermal atrophy and vascular and gastrointestinal fragility. Firstly, collagen III predominates in skin blood vessels and ligaments. It is a unique interstitial collagen, analogous to types I, II, V, and XI. It has a specific role in the strength and stability of blood vessels where it exerts its maximum clinical effect. The protein is a homotrimer \( \alpha 1(III) \), with two intrachain cysteine cross links at the C terminus. Collagen III deficiency, in the main, causes vascular EDS but may also cause other less specific phenotypes such as EDS III.

Two patterns are usual. The first (group 1) exhibits poorly secreted and overhydroxylated collagen III proteins caused by either helical glycine substitutions or exon skips. The stoichiometry dictates that only one-eighth of the triple helices are normal. The remaining seven-eighths contain either one, two, or three faulty \( \alpha 1(III) \) chains which are consequently retained intracellularly or poorly secreted and degraded. Not surprisingly, patients with such severe biochemical deficiencies have severely abnormal clinical phenotypes. The second abnormal pattern includes null alleles (group 2). Here reduced collagen type III secretion...
collagen III, although poorly secreted, is not overmodified. In other words the residual collagen III molecules are usually normal. Thus, the clinical phenotype is milder, more difficult to diagnose, and overlaps with EDS III/BHS or even merges into normality. Similarly, the arterial phenotype varies; occasionally there is premature aortic rupture, internal carotid or cerebral arterial dilatation, or arterial rupture. In contrast, the more spectacular carotid-cavernous-sinus aneurysms segregate with group I mutations. The group 2 (null allele) phenotype of arterial dilatation and cutaneous striae also overlaps clinically with atypical forms of MFS, from which it may be clinically indistinguishable.

Molecular pathology of COL3A1 mutants (fig 4)
Since 1991, there have been rapid advances in phenotype/genotype correlations. Generally, 3' mutations are acrogeric whereas the phenotypes of 5' or middle helical exon skips are harder to recognise. Unfortunately, even the latter are at risk of arterial rupture which is nevertheless much more frequent in mutations at the 3' end. Arterial rupture is difficult, if not impossible, to predict, occurring as early as the second decade (rare) to early or late middle age (very common). Like analogous COL1A1, 1A2, or 2A1 mutations, they are virtually private to affected families. They include autosomal dominant glycine substitutions or exon skips. So far, only three large in frame deletions ranging from 0.5 to 5 kb have been reported. A smaller 27 bp intronic deletion was caused by spliced mispairing. All disrupt the collagen triple helix by a dominant negative mechanism. Here, each trimer is either wild type, or contains three, two, or one abnormal chains in homotrimers such as collagen II and III or in heterotrimers such as collagen I. Thus, COL3A1 mutations cause severe disturbance with seven-eighths of the protein molecules abnormal compared to heterotrimers such as type I collagen where between half and three-quarters are faulty, depending whether the mutation affects COL1A1 or COL1A2. Both homozygosity and heterozygosity for various COL3A1 mutations or compound heterozygosity with other collagen or extracellular matrix molecules are also theoretically possible, although to date there have been no published examples. Similar considerations also apply to COL1A1 and COL1A2 (OI mutants) in which only occasional double mutations have been identified. Perhaps such double hits would be catastrophic in widespread tissue components such as collagen I or III. In contrast, double mutations of minority components like collagen VII are relatively common. Thus, double glycine substitutions cause severe recessive dystrophic epidermolysis bullosa while in contrast single glycines are either clinically silent or can cause minor autosomal dominant epidermolysis bullosa.

Reliable genetic counselling and prenatal diagnosis are available for most vascular EDS patients. This requires a combination of histology, electron microscopy, and type III collagen
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A

COL3A1 point mutations

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B

COL3A1 splicing and deletions

![Diagram of COL3A1 splicing and deletions]

Figure 4 Map of recently published COL3A1 mutations. (A) Glycine substitutions, arranged by residue number and (B) exon skips or large deletions (studies of Kujanemii et al. and Pope et al.).

protein or COL3A1 gene analysis. Prenatal diagnosis by amniocentesis, chorionic villus biopsy, or termination of pregnancy is potentially hazardous in affected females owing to the inherent fragility of cervical, uterine, or abdominal blood vessels. On the other hand, opinions vary as to the safety or dangers of pregnancy in this subset. There is little doubt that COL3A1 mutations do cause such defects but how frequently remains to be determined.

Collagen type I mutations (fig 5)

Collagen type I is a heterotrimer (α(I)1α(I)2α(I)) of two different α chains. The genes COL1A1 and COL1A2 are located on chromosomes 17 and 7 respectively. Both are interstitial collagens with uninterrupted (GlyXY) triple helices and globular, removable N and C termini; other fibrillar collagens include types II, III, V, and XI, all with cross banded fibrils. Mutations of COL1A1 and COL1A2 produce distinctive but overlapping clinical phenotypes which are position, type, and domain related. Mutations of either N terminal extensions each cause EDS VII (with distinctive clinical features) while, in contrast, helical and C terminal mutations cause OI. As expected, OI and EDS VII overlap clinically and biochemically. In EDS VII, cutaneous fragility and ligamentous laxity predominate, while in OI the bones are abnormally fragile. OI families with loose ligaments and fragile, delicate blood vessels and skin are uncommon, while otherwise typical EDS families sometimes have bone fragility/osteoporosis. The clinical phenotype is strictly dictated by mutational position and EDS VII is an especially focused example. Its mutations all cause skipping of exon 6 of either the COL1A1 or COL1A2 gene. By contrast, mutations which cause OI are more diffusely distributed between exons 7 and 52. In general the phenotype worsens with higher exon numbers (3' end or C terminal locations). EDS VII is caused by two distinct but related mechanisms. Either there is a structural abnormality of the peptidase cleavage site or the cleaving enzyme is faulty.

EDS type VII (fig 6E)

There are two distinctive clinical phenotypes, one overlapping with EDS I/II and the other with congenital cutis laxa. The former is mild and autosomal dominant, while the latter is severe and autosomal recessive. EDS VII A and B result from structural mutations of type I collagen, while EDS VIIIC is caused by the enzyme deficiency. Clinical signs include short stature and excessive premature ligamentous laxity (such as congenital dislocation of the hips). There may also be excessive cutaneous fragility reminiscent of EDS I/II, or mild cutis laxa (CL). Spinal and ligamentous deformities such as kyphoscoliosis are relatively infrequent. In contrast to EDS I/II, in EDS VII A and B collagen fibril morphology is only marginally abnormal and the fibres are angular in transverse section rather than forming cauliflower (fig 2A, B). On the other hand, EDS VIIIC shows grossly distorted, hieroglyphic fibrils.
Deletion of the peptidase cleavage site

Since collagen type I is a heterotrimer, loss of either the pNt1 or pNt2 cleavage sites can cause EDS VII. When pNt1 sequences are uncleaved three-quarters of collagen type I heterotrimers possess either one or two persistent pNt1 chains. With pNt2 defects, half the type I collagen retains abnormal pNt2 sequences while the rest become normal α2 chains. In both circumstances collagen fibril assembly is compromised and normal head to tail packing of individual triple helical molecules is severely disrupted. The persistent pN sequences seriously distort gap regions. Such misassembled fibrils are visible by transmission electron microscopy, which shows slightly irregular to angulated structures in cross section. When thicker than 10 nm, both initial nucleation and subsequent fibril growth are seriously distorted by the presence of persistent pNt1 sequences which are longer than the aggregating α chains. Exon skips also disrupt type I collagen α chains since both the pepnis and peptidase cleavage sites are deleted. Protein electrophoresis of pepnisised α chains therefore shows elongated molecules with abnormally persistent pN sequences in addition to the correctly cleaved normal α chains. Such abnormal pN proteins can also be purified from NaCl extracted tissues. Procollagen “snapshots” also show inefficient processing patterns. Following the original descriptions of Lichtenstein et al. of faulty conversion of procollagen to collagen in EDS VII, persistent pNt1 and pNt2 components were separately demonstrated by Steinmann et al. and Cole et al. Subsequently, various missplicing mutations of exon 6 sequences caused by faulty splice acceptor or donor sequences have been discovered. Most are near to the obligate G of the 5’ intron acceptor GT sequence in intron 6 (fig 5).

Mutations of procollagen peptidase (PP)

PP deficiency of cows was first described by Lapierre et al. The animals had excessively fragile skin (dermatosparaxis), which was a genetic lethal. Transmission EM of the dermis showed numerous bizarrely shaped collagen fibrils which formed flanged rods instead of normal cylinders. In transverse section these appeared as hieroglyphs while in longitudinal section they were severely disaggregated and twisted. Biochemical analysis showed abnormally elongated collagen pNt1 and α2 chains which were extractable with salt or acid solutions, implying abnormal cross linking. At the time, procollagen was thought to have single N terminal extensions and no C terminal additions. In contrast to EDS type VIIA and B (see above), every component of type I triple helix is abnormal, that is, contains both pNt1(1) and pNt2(1) chains. Consequently, no normal collagen triple helices are formed: instead fibrils assembled form [pNt1(I)], pNt2(1) triple helices. Consequently, fibrillar packing is very seriously disrupted (fig 2C). The equivalent human phenotype was not identified until recently. Various other animal models such as sheep and Himalayan cats were described previously, but included very convincing clinical and structural data, and the expected hieroglyphic fibrils and deranged biochemistry. The human clinical phenotype unexpectedly showed premature cutis laxa (CL) with blepharochalasis. So far three examples have been published, two with exceptionally clear biochemical data and the third with good illustrations of the clinical phenotype. Both the French and American descriptions included disturbed procollagen processing, pNt1(1) or α2(1) molecules, hieroglyphic fibrils, and deficient enzyme levels. The phenotype of Smith et al. included premature delivery, soft, fragile, and easily bruised skin, and patchy sagging skin, while in the patient of Nusgens et al. skin fragility, bruising, and generalised osteoporosis were notable. Reardon et al. in contrast, described severe premature generalised CL with blepharochalasis, redundant skin, and fragility. Every patient had typical hieroglyphic fibrils. Transient CL also occurs with the milder EDS VII type A and B phenotypes in which skin fragility varies. Even though less disrupted than dermatosparaxis, the fibrils of EDS VII A and B can be manipulated to produce hieroglyphics under suitable experimental conditions. The faulty angular fibrils characteristic of exon 6 deletions (which cause EDS VII A and B) are one step in the progression from normal cylinders to hieroglyphs. Theoretically both more localised and extensive mutations within and around exon 6 of the N propeptide coding region could produce a wider spectrum of fibrillar angularity and clinical features caused by faulty splice acceptor or donor sequences have been discovered. Most are near to the obligate
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Other enzyme abnormalities causing EDS

Other crucial post-translational modifications of fibrillar collagen triple helices include enzymatic hydroxylation of certain lysines and excision of the C propeptides in addition to trimming by the specific N propeptidases. Lysyl oxidase then promotes cross links by decondensing hydroxylsines to aldehydes. Mutated enzymes cause very specific forms of EDS. Thus, lysyl hydroxylase (LH) deficiency causes EDS VIA\textsuperscript{95,96}, while procollagen peptidase deficiency causes EDS VIIC\textsuperscript{95-97} (see also above). The role of lysyl oxidase is more controversial\textsuperscript{105,106} and so far the diseases caused by C terminal propeptide mutations are unknown. Since the C propeptide is essential for chain association, homozygous enzyme mutations might be genetically lethal. Bone morphogenic protein 1 (BMP1) and the C propeptidase are now known to be identical.\textsuperscript{107,108}

Ehlers-Danlos syndrome type VI (lysyl hydroxylase deficiency) (fig 6C)

The disorder was first recognised as unique by McKusick,\textsuperscript{12} although Beighton et al\textsuperscript{13} had observed autosomal recessive inheritance in a family with severe spinal deformities, ligamentous laxity, ocular fragility, and retinal detachment. There is extreme ligamentous laxity with severe ocular fragility, retinal detachment, and scleromalacia. The severe ligamentous problems present in early infancy usually with motor delay caused by severe muscular hypotonia. Sometimes muscular dystrophy is (wrongly) suspected but nearly all affected children eventually walk normally. The difficulties are caused by excessive ligamentous laxity which young muscles cannot adequately control. In adult life or middle age, there may be aortic dilatation or arterial rupture.

Krane et al\textsuperscript{95} and Pinnell et al\textsuperscript{94} then discovered underhydroxylation of lysine and deficient collagen lysyl hydroxylase in two affected sisters with severe infantile hypotonia, premature scoliosis, and ocular fragility. Other significant features included soft, hyperextensible, easily scarred skin and premature rupture of the membranes. There are two similar clinical phenotypes, type VIA with enzyme deficiency and type VIB with normal enzyme. There may also be a third, predominantly ocular, variant.

Biochemistry and molecular pathology (table 2)

Krane et al\textsuperscript{95} and Pinnell et al\textsuperscript{94} measured enzyme activity using tritium labelled chick procollagen as substrate. Nearly 20 years later, both the chick and human genes were cloned and sequenced.\textsuperscript{101-106} LH or procollagen lysyl 2 oxoglutarate 5 dioxygenase requires copper, Fe\textsuperscript{2+}, and O\textsubscript{2} as cofactors. EDS VI is caused by either deficient or faulty enzymes with variable Km or V max.\textsuperscript{101} The mutations are often complex and include Alu-Alu compound recombinations and homozygous or double heterozygous exon skips.\textsuperscript{96,102} Homozygous deletions tend to cluster in inbred families. The
hydroxylsine content of defective tissues varies between 5% and 50% being lowest in skin, bone, and tendon. Collagen I is more underhydroxylated than collagen III, while collagens II, IV, and V are normally hydroxylated. This implies more than one hydroxylase with variable specificity for other collagen types, so that other clinical phenotypes are also likely. Underhydroxylation of collagen α1(1) and α2(1) chains causes overmigration and impairs condensation cross linking of adjacent hydroxylsines which can be measured in urine.\textsuperscript{63,105} Urine sampling and type I collagen α chain electrophoresis are simple and useful screens for enzyme deficiency.

### The C terminal propeptide (BMP1) deficiency

It is now apparent that the C terminal propeptidase (CPP) is identical to bone morphogenetic protein 1 (BMP1). The latter is homologous to the \textit{Drosophila} pattern formation genes TLD and TRI with analogues in insects and sea urchins, such as the BP 10 and SpAN genes which are TGFβ activators.\textsuperscript{107} Homozygous enzyme deficiencies would either be genetic lethals or severely crippling in mechanical terms. Persistence of pC extensions would severely impair collagen triple formation just as persistent pN sequences interfere with molecular packing. Errors of the pC extensions would seriously disrupt both helical winding and chain association; quite possibly collagen triple helices would not form at all. The effects of C propeptidase deficiency should be worse than N propeptidase deficiency which causes EDS VIIA-C. Holmes \textit{et al}\textsuperscript{102} studied the stepwise removal of pC or pN propeptides before collagen fibrillogenesis. Retention of either extension by faulty processing inhibits fibril formation as do helical mutations of the fully processed molecule.

Heterozygotes for CPP deficiency might also have impaired fibrillogenesis. Homozygosity could easily cause severe clinical phenotypes, such as lethal OI, and certain severe chondrodysplasias or EDS variants either as allelic homozygotes or as double heterozygotes in combination with other ECM propeptide mutations such as pC I, II, III, V, or VI abnormalities respectively.

### EDS variants of unknown cause

This applies to EDS types III, VIII, and X in which the causes are non-specific (EDS III), variable (EDS VIII), or represented by a single one-off example (EDS VIII).

#### EDS type III (fig 6 A, B)

Unlike EDS types I and II there is no significant cutaneous scarring despite obvious joint hypermobility and doughy skin. This clinical subtype merges with the so-called benign hypomobile syndrome (BHS) but which has normal skin texture. This milder clinical phenotype can also segregate in typical EDS I/II families probably owing to incomplete penetrance. Similar clinical features may also accompany COL3A1 mutations,\textsuperscript{15,10} such as MFS and SS. Heterozygotes for autosomal recessive disorders such as PXE and lysyl hydroxylase deficiency (EDS VI) have very similar clinical phenotypes.\textsuperscript{109} There can also be overlap (clinical and biochemical) with vascular EDS (see above).

### EDS type VIII

First described by Stewart \textit{et al}\textsuperscript{110} this phenotype is clinically, biochemically, and allelically heterogeneous. Distinguishing clinical features include chronically inflamed, heavily pigmented, discrete, pretilial plaques and premature periodontal recession. There is collagen degradation as judged by gum resorption and the cutaneous inflammation. In EDS VIII no consistent biochemical or structural changes are detectable.\textsuperscript{110-114} Certain mutant collagens may be abnormally proteinase susceptible. Alternatively, faulty collagenase inhibitors such as TIMP might be implicated, although all of our EDS VIII patients have normal TIMP profiles. We have observed both vertical and horizontal transmission of early adult gum recession, sometimes accompanied by collagen III deficiency. Similar features may accompany EDS I, II, and IV. Others have also noted this phenotype.\textsuperscript{111,114} The distinctive clinical phenotype\textsuperscript{106} is therefore biochemically and genetically heterogeneous. Histology of the skin lesions shows a granulomatous collagen degeneration resembling necrobiosis lipoidica.\textsuperscript{115}

### Ehlers-Danlos syndrome with fibropectin deficiency (EDS X)

Only one family with this phenotype has been described.\textsuperscript{116} This included four affected sibs in one generation but with normal parents, which McKusick later classified as autosomal recessive. The phenotype includes thin, fragile, easily scarred skin, joint hypermobility, and excessive bruising and therefore resembles EDS II/III. Platelet aggregation was abnormal but correctable with normal fibronectin. In other EDS families a variety of clotting abnormalities have been described which are too inconsistent to be directly related.\textsuperscript{117,118}

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