A novel form of syndromic cutis laxa with facial dysmorphism, cleft palate, and mental retardation

D Genevieve, C Baumann, C Huber, L Faivre, D Sanlaville, C Bodemer, S Hadj-Rabia, A Assoumou, A Verloes, F Raqbi, A Munnich, V Cormier-Daire

C utis laxa (CL) is a rare congenital disorder characterised by loose and redundant skin. Three groups of CL have been recognised, according to their mode of inheritance.

The first group, accepted as an autosomal dominant trait, is relatively benign, with late onset skin manifestations and subnormal life span. Other manifestations occasionally include pulmonary artery stenosis, emphysema, bronchiectasis, hernia, and genital prolapse. This subtype is ascribed to mutations in the elastin gene. The second group (also called Ehlers-Danlos type IX syndrome or mild Menkes syndrome) undergoes an X-linked recessive inheritance and is ascribed to ATP7A deficiency. The third group, inherited as an autosomal recessive (AR) trait, includes type I and type II CL, wrinkly skin syndrome, and De Bary syndrome. Type I AR CL is characterised by early infantile pulmonary emphysema, hernias, multiple diverticulae, and a poor prognosis. Recently, filbin 5 mutations have been reported in Turkish patients with CL AR type I and supravalvular aortic stenosis. Type II AR CL is associated with growth and developmental delay, joint laxity, peculiar face with frontal bossing, large fontanelle, and skeletal dysplasia including congenital dislocations of the hips. A deficiency in lysyl oxidase has been suggested in CL AR type II, which is closely related to wrinkly skin syndrome. Finally, De Bary syndrome is associated with mental retardation, short stature, and corneal clouding. Here, we report on a novel type of AR CL with facial dysmorphism, hygroma in early pregnancy, cleft hard palate, ventricular septal defect, and moderate mental retardation, which is distinct from previously reported AR CL.

CASE REPORT

Patient 1, a girl, was the fourth child of first cousin healthy Tunisian parents (fig 1). The pregnancy was uneventful. Moderate hypotonia and cutis laxa were noted in the neonatal period (birth length 44 cm, –2.5 SD; weight 2520 g, –2 SD; OFC 34 cm, mean). Clinical examination showed cleft hard palate, increased joint laxity, facial dysmorphism including square and flat face, high and broad forehead, very short nose, blue sclerae, sagging cheeks, and small mouth and chin. Palms and soles were normal. The patient could sit unaided aged 1 year and walk at 2.5 years. At 7 years age she had mildy delayed growth development (–1 SD), and mental retardation requiring special schooling. Cutis laxa improved with age and facial dysmorphism became more pronounced, with a round flat face, very short nose, blue sclerae, and small mouth (fig 1). Drooping cheeks disappeared with age and joint laxity was noted. Thoracic x ray revealed slight pulmonary emphysema.

Patient 2, a boy, was the fifth child from the same sibship (fig 2). Prenatal hygroma and a moderate hydramnios were noted at 19 weeks of gestation. He was small for his age (length 46 cm, –2 SD; weight 3000 g, –1 SD; OFC 35 cm, mean). Severe hypotonia, increased joint laxity, broad hands and feet, wide anterior fontanelle, and cutis laxa were noted in the neonatal period. Left hip dislocation and ulnar deviation of hands were also noted. His facial dysmorphism was similar to that of his sister, including square and flat face, high and broad forehead, very short nose, sagging cheeks, small chin, and normal palate. Excess of skin was noted on the neck, occiput, and limbs (fig 2). Examination of palms and soles was unremarkable. He died 25 days postpartum of severe emphysema and multiple pneumothorax. The family refused postmortem examination.

Patient 3, a girl, was the seventh child of the sibship (fig 3). Hygroma and hydramnios were detected at 14 weeks of gestation. She was born at term with severe hypotonia, increased joint laxity, complete Pierre–Robin sequence, and cutis laxa. Her length was 49 cm, mean; weight 2990 g, –1 SD; and OFC 35 cm, +1 SD). Facial features were similar to those of her affected sibs, namely square and flat face, high and broad forehead, very short nose with anteverted nostrils, small mouth and chin, cleft hard palate, hypertelorism, blue sclerae, and sagging cheeks (fig 3). She had an excess of skin on her neck, forehead, and abdomen, but palms and soles were normal. Tracheotomy was performed for respiratory distress. At 10 months of age she could not sit unaided, and

Key points

- Cutis laxa (CL) is a clinically and genetically heterogeneous condition characterised by loose and redundant skin with lack of elasticity.
- CL was found to be associated with facial dysmorphism, cleft palate, and moderate mental retardation in three siblings, born to first cousin healthy parents. Facial dysmorphism included flat and square face, very short nose, sagging cheeks, blue sclerae, hypertelorism, and small chin.
- Additional features included hygroma during pregnancy (2/3), cleft hard palate (2/3), and ventricular septal defect (2/3). Skin histology showed a lack of elastic fibres (1/3). X ray and ophthalmological examinations were normal.
- The clinical features observed in our patients are compared with those described in other CL syndromes in the literature. Cleft palate, facial dysmorphism, joint laxity, mental retardation, and emphysema have not been previously reported in association with CL.
- It is concluded, therefore, that this distinctive association represents a novel syndromic form of CL.

she had severe hypotonia and developmental delay (growth and length = -2 SD, OFC = +2 SD). She died at 10 months of age from progressive emphysema complicated by a pulmonary infection.

METHODS
With the written consent of the parents genomic DNA was purified from peripheral blood leucocytes according to standard techniques,¹³ and microsatellite markers of the Genethon map (http://www.genethon.fr/) were PCR amplified¹⁴ and analysed on 6% polyacrylamide denaturing gel. To test linkage to the elastin gene,² microsatellite DNA markers of chromosome 7q11.2 were used (D7S2472, D7S1870, D7S2470, D7S669, ELN, D7S524, D7S630, and D7S657). Microsatellite markers were also used to test linkage to the lysyl oxidase gene at chromosome 5q23.3-q31.2 (D5S433, D5S2027, D5S2055, D5S471, LOX, D5S2057, and D5S2115). Linkage to fibulin 5 was tested using an intragenic microsatellite DNA marker (FBLN 5' GGT ACA ATG AGC TGA GAT CGT G, FBLN 3' ACT CAA CTC TCG GCT TGG TTA G).

RESULTS
Laboratory investigations
Cerebral MRI was normal in patient 1. In patient 3, cerebral CT showed cerebral atrophy, and thoracic CT showed diffuse pulmonary emphysema. Heart ultrasound showed a ventricular septal defect in 2/3 patients. Skeletal x rays showed advanced bone age (2/3) and a butterfly shaped hemivertebra in patient 2. Histological study of skin biopsy performed for patient 2 showed paucity of pre-elastic fibres and lack of mature elastic fibres (fig 4). Kidney ultrasound, ophthalmological examination, and high resolution blood chromosomes (including 22q11.2 in situ hybridisation), amino acid, and organic acid chromatography were normal in all three cases. Copper metabolism was also normal in case 3.

Figure 2   Facial features in patient 2 at day 25 (A and B). Note square and flat face, high and broad forehead, short nose, small chin, small mouth, and excess of skin on arms, neck, and occiput.

Figure 3   Facial features in patient 3 at day 4 (A) and 2 months of age (B and C). Note facial dysmorphism including square and flat face, high and broad forehead, hypertelorism, blue sclerae, sagging cheeks, small nose with anteverted nares, excess of skin, and small mouth with small chin.
Electronic letter

Figure 4  (A) Skin biopsy of patient 2 at 23 days old. (Catechin-fuchsin staining, magnification × 52). The pre-elastic fibres (oxytalan + elaunin) in the superficial dermis are rarefied, and the elastic fibres are almost completely absent in the centre part of the dermis. (B) Skin biopsy of a control subject at 3 months of age (Catechin-fuchsin staining, magnification × 64). Pre-elastic fibres (oxytalan + elaunin) are well defined in the superficial dermis and well structured with typical arborisation. Dermal elastic fibres are numerous and well positioned.

Molecular studies
In view of recent evidence for the involvement of elastin, lysyl oxidase, and fibulin 5 in AR CL, we first tested these candidate genes by linkage analysis in affected children using flanking or intragenic microsatellites. Haplotype construction and segregation analyses excluded linkage to the elastin gene (fig 5), fibulin 5, lysyl oxidase, or COL2A1 genes (data not shown).

DISCUSSION
This study reports on three sibs with a novel form of syndromic cutis laxa with facial dysmorphism, cleft hard palate, and mental retardation. The observation of CL in three sibs born to first cousin parents supports an AR mode of inheritance in this family. Molecular studies excluded the three genes known to be involved in CL, namely the elastin, lysyl oxidase, and fibulin 5 genes. Cleft hard palate and facial dysmorphism (square and flat face, short nose, small mouth and chin, and blue sclerae) have not been described before in any other AR CL syndrome. Table 1 compares the clinical findings in our patients with those of other AR CL syndromes which do not, in our opinion, account for the features observed in our family. Type I cutis laxa is characterised by emphysema, multiple pneumothorax, diverticulae, and hernias; emphysema and multiple pneumothorax were indeed observed in 2/3 of our cases, but their other features, namely cleft palate, mental retardation, and facial dysmorphism, have not been reported in type I CL and involvement of fibulin 5 gene could be excluded.

De Barsy syndrome, combining mental retardation and increased joint laxity, could also be considered, but the absence of corneal clouding rules out this diagnosis. In fact, the cases in the present report share several clinical features with AR CL type II ascribed to lysyl oxidase deficiency. However, the characteristic pre and postnatal growth retardation and loose skin on palms and soles were not present, and involvement of lysyl oxidase gene could be excluded in our family. The normality of palms and soles in our patients excludes wrinkly skin syndrome, which seems closely related (if not similar) to AR CL type II. Similarly, the association of Pierre–Robin sequence, with flat face, small chin, small nose, and prominent eyes, prompted us to test and eventually exclude COL2A1.

In conclusion, we suggest that the association of CL, cleft hard palate, ventricular septal defect, and striking facial dysmorphism represents a novel form of autosomal recessive cutis laxa distinct from types I and II AR CL. Additional observations are needed to decide whether this association indeed represents a distinct and novel entity.

### Table 1 Features observed in our cases in comparison with other autosomal recessive cutis laxa syndromes

<table>
<thead>
<tr>
<th>Clinical features</th>
<th>Present cases</th>
<th>Case 1</th>
<th>Case 2</th>
<th>Case 3</th>
<th>CL I</th>
<th>CL II</th>
<th>DB</th>
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<tbody>
<tr>
<td>Hygroma/ hydramnios</td>
<td>?</td>
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<td>(+14)</td>
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<tr>
<td>Neonatal hypotonia</td>
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<td>Cuts laxa</td>
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<td>Increased joint laxity</td>
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<td>Wide anterior fontanelle</td>
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<tr>
<td>Mental retardation</td>
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<td>NA</td>
<td>+</td>
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<td>+</td>
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<tr>
<td>Square and flat face</td>
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<td>Very short nose</td>
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<td>Blue sclerae</td>
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<td>Sagging cheeks</td>
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<td>PR/cleft hard palate/small chin</td>
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<td>Diverticulae and hernias</td>
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<td>+</td>
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<td>Prenatal growth deficiency</td>
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<td>+</td>
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<td>Postnatal growth deficiency</td>
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<td>Wrinkly skin on palms and soles</td>
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<td>Corneal clouding</td>
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<td>Advanced bone age</td>
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<td>Emphysema</td>
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<td>Ventricular septal defect</td>
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CL, cutis laxa; DB, de Barsy syndrome; NA, not applicable; PR, Pierre–Robin sequence.
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