Brachymorphism-onychodysplasia-dysphalangism syndrome

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Abstract
Three unrelated children are reported with intrauterine proportionate growth retardation and facial dysmorphism (broad nose, flat malar area, large mouth, pointed chin), microcephaly, hypo/aplasia of the terminal fifth digits, and (sub)normal intelligence. Radiological findings include hypo/aplasia or fusion of the distal phalanges of the fifth finger and toe, brachymesophalangism V, and nail dysplasia or aplasia. One child had cystic adenomatoid disease of the lung. The pattern of anomalies presented by these children closely resembles a syndrome incompletely delineated in 1971 by Senior in six children, which has often been considered to be a mild form of Coffin-Siris syndrome. We suggest that this is an independent entity (BOD syndrome). The aetiology is still unknown. Differential diagnosis and nosological difficulties are discussed.

(J Med Genet 1993;30:158-61)

In 1971, Senior described six 'short children with tiny toenails'. Pre- and postnatal short stature, hypoplastic fifth digits with abnormal phalanges and tiny toenails, facial dysmorphism, and, in some, mild intellectual impairment were observed. Except for a single report, no other cases have been recorded. We describe here three unrelated children fitting the 'Senior syndrome'.

Case reports
Case 1
This girl was born at 41 weeks of a pregnancy marked by intermittent bleeding. There was no exposure to alcohol or known teratogens. Birth weight was 2100 g, birth length 46 cm, and head circumference 31 cm (all below the 10th centile). Postnatal growth remained below the 3rd centile. A cholesteatoma of the right ear was surgically removed. Mental development was not delayed. She attended normal school and had difficulties but never repeated grades. When we saw her at the age of 14 years she was 144 cm tall (–2.5 SD) and weighed 45 kg. Head circumference was 47 cm (< –4 SD). Puberty began at 12 years but menstruation did not occur. She had a brachycephalic skull, narrow forehead, large nose with a broad base, flattened malar area, long philtrum, large mouth with somewhat protruding lips, and pointed chin (fig 1A,B). She had bilateral aplasia of the distal phalanges of the fifth digits (fig 1C), which had no nail. The fifth toes were very hypoplastic and lacked nails (fig 1D). The nails of the second, third, and fourth toes were small, dystrophic, and soft. Radiologically, the acral anomalies were restricted to the fifth digits and included minute terminal phalanges with probable symphalangism 2-3 in the right hand (fig 2A), aplasia of P3 on the left, aplasia of P2 and P3 bilaterally in the fifth toes, and a single, short distal phalanx in toes 4 and 5 (fig 2B). Other radiological anomalies were partial fusion of L5 and the sacrum and large sella turcica (fig 2C). Brain CT scan was normal. There was no pituitary dysfunction and no visceral malformations. G banded karyotype was normal.

The family history is unremarkable. Her parents were tall and had normal head circumferences. They were aged 30 and 29 at the birth of the proband. She had two unaffected sibs.

Case 2
This boy was born at 39 weeks weighed 2400 g (<10th centile) and had a birth length of 45 cm (10th centile). The pregnancy was uneventful and there was no exposure to drugs or teratogens. A cystic adenomatoid malformation of the lung was surgically removed at 1 month. Subsequent development was normal and not delayed. When evaluated at 2 years 5 months, he was 84 cm tall (–2 SD), weighed 9750 g, and had a head circumference of 47.5 cm (< –2 SD). He had a flat, oval facies (fig 3A); with a high forehead, horizontal palpebral fissures, epicanthus, broad nasal root, flat malar area, upturned nose (fig 3B), long, smooth philtrum, high arched palate, and small, somewhat receding, pointed chin. The hands were stubby, with hypoplastic fifth nails, clinodactyly V, and bilateral transverse palmar creases. The left fifth nail was absent and the left fourth and right fifth nails were very hypoplastic. Radiographic investigations showed short distal phalanges in the fifth fingers (fig 4A) and absence of ossification of the median and distal phalanges of the fifth toes (fig 4B). On the left, the third phalanx of the fifth finger and toe was absent and only one ossification centre was visible in the left second and fourth toes. No other anomalies of the bones were recorded, except for two parietal lacunae (fig 4C). No visceral malformation was present. The karyotype was normal.

The family history was unremarkable. The father was 180 cm and the mother 163 cm tall. They were not consanguineous. An older child was in good health.
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CASE 3
This girl was born at 34 weeks of gestation. The pregnancy was normal and there was no exposure to known teratogens. Birth weight was 1730 g (10th to 25th centile) and birth length was 41 cm (10th centile) with a head circumference of 31.3 cm (mean). An ASD ostium secundum type was noted at birth. At 14 months, her mental development was normal. She was 66 cm tall (−3.5 SD), weighed 7200 g, and had a head circumference of 43 cm (−2.9 SD). She had a rounded facies with a pointed chin, upward slanting palpebral fissures, broad nasal root, epicanthus, large nose, and a long philtrum (fig 5). The distal phalanx of the left fifth finger was absent. The left third and fourth and the right fifth fingernails were hypoplastic, soft, concave, and thickened. The fifth toenails were hypoplastic bilaterally, as...
were the left third and fourth toenails. X rays showed bilateral absence of the third phalanges in the fifth fingers (fig 6A) and absence of ossification of the two distal phalanges of the fifth toes (fig 6B). Family history was unremarkable. The mother, aged 34, was short (160 cm), as were her own mother and sisters. The father, aged 25, was 168 cm tall. They had no other children.

Discussion
Senior,1 in 1971, reported six unrelated children with growth retardation (below the 3rd centile), peculiar facies, and acral anomalies. Five of them presented with absent (1/5) or fused (4/5) middle and distal phalanges and short fifth fingers, with brachyphalangism more pronounced in the middle ones. The sixth child was less convincing, as he had only tiny toenails. No information was given regarding prenatal exposure to drugs or microcephaly. Mace and Gotlin2 suggested the same diagnosis for a case with brachymetatarsia IV. The hand of another child with absent third phalanx was depicted in Poznanski’s textbook.3

The three children reported here show a concordant pattern of anomalies of the extremities and general growth failure, which fits the description of Senior.1 Brachydactyly is of the B type.4 The face has a similar triangular or oval shape with pointed chin and apparently large mouth (in patients 1 and 2). Absolute or relative microcephaly, an important feature in our cases, was not mentioned by Senior.1 Cystic malformation of the lung is a sporadic developmental anomaly, which could be fortuitous in case 2. Children with the syndrome are of low normal intelligence or even slightly mentally impaired, as seen in patient 1. Patients 2 and 3 are too young to determine moderate mental disturbance. The aetiology of Senior syndrome is not known and all cases were sporadic. The nine cases with available family data had 25 unaffected sibs. Consanguinity between parents has not been reported.

The delineation, and even the existence, of Senior syndrome remain controversial, particularly as the original cases were relatively poorly described. Senior himself stated that “the listed features of these patients are neither particularly distinctive, nor individually rare”, and possible prenatal exposure to alcohol or phenytoin was not discussed. The syndrome is presented as a separate entry in two textbooks of radiology.5,6 It is discussed with Coffin-Siris syndrome in Gorlin’s monograph7 (but the authors acknowledged that Senior syndrome was likely to be different). Senior’s paper is quoted under the Coffin-Siris heading in the McKusick catalog7 and in LDDB. It is not included in POSSUM.

The digital anomalies correspond to type B brachydactyly (apical dystrophy),4 which can include symphalangism and nail hypoplasia,
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and usually has a dominant mode of inheritance. However, it differs from the ‘common’ apical dystrophy by an unusual gradient of severity from the preaxial to the postaxial rays. Growth retardation and low intelligence are not associated with this type of brachydactyly. Around 50 syndromes may be extracted from LDDB or POSSUM, using a combination of ungual dysplasia/hypoplasia/aplasia, hypoplasia/aplasia of phalanges or terminal hypoplasia of fingers. In relation to Senior syndrome, most of them are easily rejected, because of their associated features (for example, Fryns syndrome and several ectodermal dysplasias). Ungual hypoplasia/dysplasia occurs in a form of brachydactyly classified as A4 by Poznanski or A5 by McKusick (autosomal dominant absence of middle phalanges with duplication of the terminal phalanx of the thumb). Apical dystrophy has been observed in association with coloboma in Sorsby syndrome. Schott reported a dominantly inherited syndrome of hypo/aplasia of the terminal phalanges, sparing the thumb, associated with nail dysplasia. The feet were not affected. Goldshkag-Cooks et al. reported nail dystrophy and terminal hypoplasia of the toes and fifth fingers in a two generation family. Some affected patients had triphalangeal thumbs. Ballard syndrome consists of camptodactyly, finger-like thumbs, distal phalangeal hypoplasia, nail dysplasia, and brachymetacarpia. None of these syndromes are associated with short stature, microcephaly, or mental subnormality. Our patients do not appear similar enough to be considered as having one of these syndromes. Fetal hydantoin syndrome may show terminal hypoplasia and growth retardation, but the facial features are different and no exposure to anticonvulsivant drugs was noted (at least in our cases). Dup(9p) syndrome shows the same acral anomalies but severe mental impairment is a constant feature and chromosomes were normal in our patients 1 and 2.

Distal digital hypoplasia of the fifth ray is a hallmark of Coffin-Siris syndrome. This recessively inherited disease is characterised by microcephaly, growth retardation, constant, and usually severe, mental retardation with hypotonia, dorsal hirsutism, alopecia, patellar hypoplasia, and coarse facial features. Although considering Senior syndrome as the mildest form of this syndrome cannot be rejected, it is difficult to combine in a single entity our mildly affected patients and the ‘common’ cases of Coffin-Siris syndrome. At least for clinical and prognostic reasons, the splitting of both entities seems reasonable.

At the moment, this syndrome seems best characterised as a recurrent pattern of anomalies of unknown genesis, with some similarities to Coffin-Siris syndrome, but distinguishable by a milder phenotype and much less severe mental handicap. Aetiological heterogeneity among the cases reported by Senior cannot be ruled out, or that some of them may in fact belong to one of the previously discussed syndromes. Nevertheless, we suggest that our cases (and at least some of those described by Senior) represent a genuine and clinically distinguishable entity. We would recommend the use of ‘Brachymorphism-onychodysplasia-dysphalangism (BOD) syndrome’ to refer to it, rather than the eponymous ‘Senior syndrome’, to prevent confusion with the Senior-Loken syndrome.


Figure 6 Patient 3 at birth. (A) Right hand, (B) feet.