Sotos' Syndrome (Cerebral Gigantism) with Peripheral Dysostosis

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Evans, P. R. (1971). Archives of Disease in Childhood, 46, 199. Sotos' syndrome (cerebral gigantism) with peripheral dysostosis. A girl had short stumpy hands and feet due to osseous dystrophy; in other respects she showed all the main and many of the minor features of Sotos' syndrome during infancy. Then she was on the 97th centile for height, but she grew to be a woman on the 10th centile. Menarche was at 12.5 years; growth in height stopped before 14.5 though epiphysial fusion had not occurred; educational difficulties were still being fought at 18 years but she was well adjusted and happy.

This case is presented because (1) the combination of cerebral gigantism and gross manupedal dysostosis is unusual, perhaps unique; and (2) growth has been followed from infancy into adult life.

Case Report

The patient was a girl born in 1952, only child of non-consanguineous parents; her father was an epileptic house-painter taking phenobarbitone. Pregnancy and delivery were normal, birthweight was 2270 g; she was given oxygen for one week and kept in hospital for one month. She was breast-fed for 8 months, got her first tooth at 6 months, sat and stood normally, walked at 18 months, spoke her first word at 2 years; sphincter control was normal.

She was brought to hospital at 3 years because of clumsiness of hands and tendency to run on toes instead of walking, to trip and to fall forward. She was a tall, long-legged, heavy girl with rosy cheeks: upper to lower segment ratio was 1.16 (average for age 1.35, for height 1.24); span was 96.5 cm (average 90.5 cm for age, 99 cm for height); head circumference was 53 cm (average for age 51 cm), chest circumference was 58 cm. Hands and feet were broad (Fig. 1a). She had 17 teeth, 3 more had been removed for caries. She had a depressed nasal bridge and a staring expression (Fig. 2a). Tonsils were infected. BP 120/80 mm Hg. Speech was indistinct, perhaps in part due to a hearing loss of 10–20 decibels at 250–4000 c.p.s. The following results were normal: nuclear sexing of leucocytes; glucose tolerance test; subcutaneous insulin sensitivity test; blood plasma levels of bicarbonate, Na, Cl, K, Ca, P, alkaline phosphatase, free and ester cholesterol; urinary secretion in 24 hours of 17 ketosteroids and 17 hydroxycorticoids; urea clearance; radiographs of skull,
At 5 years, tonsils and adenoids were removed. At 6 years, IQ was estimated as 89. At 10 years, she transferred to a school for educationally subnormal children as she had poor auditory discrimination and was clumsy. At 12 years, the pituitary fossa was examined radiographically because of the possibility of acromegaly, but it was normal; the vault of the skull was thick. There was valgus deformity of the feet, varus of the ankles, and dental caries was evident. At 12\(\frac{1}{2}\) years she first menstruated. At 13 years, a mixed parotid tumour was excised. EEG showed increased delta activity and some symmetrical following under photostimulation, with a resting record of 4–8 c.p.s., and there was high voltage activity with underlying low voltage activity without focal features. At 14 years, shoes were 22 cm long and 11 cm wide; shoulder movements were stiff so that she could not undo her dress at the back. When last seen at 17 years she was a healthy, happy girl, short in stature but not abnormally so, large of hip and thigh compared with chest and shoulder (Fig. 2b).

She had compensated well for her earlier clumsiness and was doing fine work as a Quick-unpick assembler (Quick-unpick unpicks hems). She could not pick small objects up from a flat surface but when asked to take a pin from a hard table, she neatly swept it to the edge where her stubby fingers were able to close over it as it turned the corner. She was waiting for a plastic operation to construct a bridge in her nose as her

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Fig. 2.—(a) At 3 years. (b) At 17 years. The face (deleted) has matured, corresponding to her age.

spine, and chest. Metatarsals, metacarpals, and phalanges were short and wide (Fig. 3a), with cupped ends and no calcified epiphyses, yet the carpus was like that of a child twice her age.

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Fig. 3.—(a) X-ray of hand at 3 years. (b) X-ray at 18 years. Ulnar and radial epiphyses have not fused, while metacarpal and phalangeal epiphyses have not even appeared.
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Discussion

'Cerebral gigantism' is a term introduced by Sotos et al. (1964). 'Sotos' syndrome consists of 'excessively rapid growth dating from infancy, acromegalic features and a non-progressive neurological disorder manifested by clumsiness and a dull intelligence' (Abraham and Snodgrass, 1969). The definition fits this case except that the hands (Fig. 1) and feet were short, i.e. there was no acromegaly; her facial appearance (Fig. 2a) resembles that seen in the cases of Abraham and Snodgrass (1969).

The dysostosis occurring in this case (Fig. 3 and 4) is more difficult to name, but Dr. A. R. Chrispin has pointed out that it is similar to the 'peripheral dysostosis' described by Garces et al. (1969). The patients described by Sotos et al. (1964) showed advanced bone age, large hands and feet, and slight tufting of terminal phalanges in 2 patients out of 5, but no gross abnormality.

These authors described very rapid growth in the first 4 years of life, followed by growth parallel to the normal curve. Gigantism is perhaps too big a word, but they certainly are large children, usually around the 97th centile in height. In the patient described here height was first recorded at 3 years, it remained at the 97th centile until the age of 10 years, but increased little after that (Fig. 4a). Adult height had been reached by 14½ years, and was on the 10th centile, a paradoxical swing from the giant infant to the little woman. Weight followed a comparable course (Fig. 4b) allowing for her being a fat child.

Abraham and Snodgrass (1969) note features reported in accounts of 43 cases of Sotos' syndrome; of those occurring in a third or more the following were present in this case: birth at term, neonatal respiratory difficulty, delayed walking, delayed speech, IQ below 90, macrocrania, hypertelorism (probably), antimongoloid slant of eyes, facial plethora, prognathism, clumsiness, abnormal EEG, normal pituitary fossa. Missing features were: high birthweight, jaundice, feeding problems,
aggressiveness, kyphosis or scoliosis, large hands and feet, convulsions, drooling, and high 17-ketosteroids.

REFERENCES


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