Sclerosteosis in a Spanish male: first report in a person of Mediterranean origin


Abstract
We report the first observation of sclerosteosis in Spain. To the best of our knowledge, this is the first case of sclerosteosis in a person of Mediterranean origin with no known Dutch ancestors. He has the characteristic phenotype of the disease with right facial nerve palsy and syndactyly and the typical radiological features, including generalised bone sclerosis and cortical widening of the tubular bones.

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Sclerosteosis (MIM 269500) is a rare autosomal recessive skeletal dysplasia, included in the group with increased bone density osteochondrodysplasias and the subgroup with endosteal hyperostoses. The term sclerosteosis was coined by Hansen in 1967 and since then about 60 cases have been reported, the majority in persons from the Afrikaner community of South Africa. The few reports from outside South Africa have come from the United States, Japan, Switzerland, and Brazil.

Here we describe the first case of sclerosteosis in Spain, a Mediterranean country, in a male with reportedly no Dutch ancestry.

Case report
The patient, a 12 year 10 month old male, was referred to us for evaluation of a right facial nerve palsy of five months' duration. He is the first and only child of unrelated Spanish parents. The pregnancy and delivery were uncomplicated. At birth he was noticed to have oligodactyly and syndactyly of the hands and bilateral 3-4 syndactyly of the toes. The family history was unremarkable and all family members were natives of Spain with no known Dutch ancestors.

Clinical examination showed a height of 167 cm (>97%) and a head circumference of 59 cm (>98%). He had facial asymmetry owing to right facial palsy (fig 1), mild frontal bossing, midfacial hypoplasia, dental malocclusion, and marked prognathism. He had oligodactyly with 3-4 syndactyly in both hands (fig 2), the distal portions of some of the fingers were short and radially deviated, and they had hypoplastic nails. He also had mild bilateral 3-4 syndactyly of the toes. No other physical anomalies were found at that time. Intelligence seemed appropriate for his age.

Radiographical findings in the skull included a widened and uniformly sclerotic calvarium with a very dense base (fig 3). Other findings were prominent vascular marks, expanded sella turcica, enlarged frontal sinuses, enlarged sclerotic mandible, widened and dense clavicles and ribs, and sclerotic vertebral end plates and pedicles. The sclerotic pelvic bones were not expanded. All long bones were involved and showed cortical hyperostosis with moderate alteration of their external contours. In the hands bilateral 3-4 bony syndactyly with shortening and radial deviation of several phalanges were noted (fig 4). No skeletal anomalies were observed in the toes. CT scan of the head confirmed thickening and sclerosis of the skull diploë, with no evidence of cerebral anomalies or signs of increased intracranial pressure.
Functional studies, including audiograms, showed intact cranial nerve pairs, except for the seventh pair. Electromyogram and conduction velocity studies of the facial nerve showed basal grouped discharges with a normal reinervation pattern. Plasma levels of growth hormone (hGH), LHRR, and testosterone were within normal limits, as well as plasma and urine phosphate and calcium.

The patient at the age of 17 is 188 cm tall (>97%) and his head circumference is 60 cm (>98%). Besides his mild right facial palsy he is otherwise asymptomatic.

Discussion
Sclerosteosis is a rare autosomal recessive disorder and the majority of the cases have been described in the Afrikaner community of South Africa. Affected persons show signs of overgrowth and sclerosis of the bones, especially involving the skull and the facial bones. The anomalies may be present in early childhood and they are progressive. Usually, cranial nerve dysfunction is an early symptom and it is frequently accompanied by tall stature, macrocephaly, prognathism, and distortion of the face. Unlike van Buchem disease, considered an allelic variant of sclerosteosis, persons with sclerosteosis have syndactyly, generally involving the second and third fingers and associated with nail anomalies.10

To date, sclerosteosis has been reported in about 60 patients from the Afrikaner (Dutch ancestry) community of South Africa, with a minimum prevalence estimated at about 1/60 000 and a gene frequency of 0·004.14 Interestingly, only a few cases have been reported in other countries, including two kindreds in New York and Maryland in the United States,14 a 21 year old woman in Japan, a young woman from Switzerland, and two relatives from a highly consanguineous Brazilian family of Dutch ancestry.6 We report here the first case of sclerosteosis in Spain, a Mediterranean country, in a male who reportedly had no Dutch ancestors. It seems likely that cases of sclerosteosis in persons with no evidence of Dutch ancestry, such as the case of Sugiuara and Yasuhara7 and our case, could be explained by the random appearance of spontaneous mutations of the sclerosteosis gene in the world population. In their paper, Sugiuara and Yasuhara7 provide no data about the affected woman’s ancestry. It will be very important, once the molecular defect is found, to determine if the mutation is the same in the different populations.

Clinically, our patient has most of the major manifestations of the disease: tall stature, cranial nerve dysfunction (facial palsy), dental malocclusion, prognathism, syndactyly, and generalised hyperostosis and sclerosis of the skeleton, including the skull. At the age of 17, he still has no hearing impairment and no signs of increased intracranial pressure.

In the past, many patients with sclerosteosis were erroneously diagnosed as having Albers-Schönberg disease (osteopetrosis). However, there is now no reason for confusion after excellent clinical descriptions1 and its confirmation as a separate entity by the International Working Group on Constitutional Diseases of Bone.3

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