Unknown syndrome: peculiar face, severe hypodontia of permanent teeth, and precocious choroid calcifications

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Abstract
We describe a mother and her twin daughters affected with severe hypodontia of the permanent teeth, precocious calcification of the choroid plexus, and minor digital anomalies. The presence of inner telecanthus, broad and flattened nasal bridge, mild ocular proptosis, small nose with antverted nostrils, and slight microretrognathia gives them an unusual appearance. All three affected persons are of normal intelligence.


Keywords: oligodontia; persistent deciduous teeth; choroid plexus calcifications

Both hypodontia and anodontia are well known disorders. Complete anodontia is very rare; according to Bartsocas,\(^1\) Pyrrhus (318-272 BC), King of Epirus, may have been affected. It is inherited as an autosomal recessive trait.\(^2\)

Witkop\(^1\) suggested that the trait of small, peg shaped, or missing maxillary lateral incisors not associated with a syndrome is best explained as an autosomal dominant trait modified by a polygenic component, and that agenesia of succedaneous teeth is a recessive trait that results from the homozygous state of this gene.\(^1\)

Dolamore\(^2\) described a case of persistent deciduous canines with absence of permanent successors. Gorlin (personal communication) pointed out that about one third of the general population lack one or more of the third molars (wisdom teeth), the teeth most often missing, followed by the premolars. Anodontia and hypodontia are often the expression of complex situations involving defective formation of structures of ectodermal origin.

We describe a family who have, to our knowledge, a completely new condition.

Case report
We report a family which includes a 39 year old affected mother and her 20 year old twin daughters. At the time of conception their father was 24 years old. They are of southern Italian origin, their mother, the third of five children (three males and two females), coming from Sicily and their father, the second of three brothers, from Abruzzo; the parents are not related. The family history on both sides was negative for miscarriages, malformations, and genetic or metabolic disease. The girls were born at 31 weeks of gestation and were registered as monochorionic identical twins. Their growth was appropriate for age (90th centile), they were negative for transplacental infections (TORCH), and their psychomotor development was absolutely normal. The blood group of all three patients was O Rh(D)+. They came to our attention at the age of 12 years through genetic counselling. Their early medical history, like that of their mother, was unremarkable; growth parameters were still around the 90th centile and their intelligence was well above average.

The twins’ faces showed very similar characteristics to that of their mother (figs 1, 2, and 3), although more accentuated, and included inner telecanthus (ICD >97th centile, IPD >95th centile, OCD >75th centile), downward slanting palpebral fissures, mild ocular proptosis, flattened, wide nasal bridge, a small nose with forward slanting nostrils, slight microretrognathia, and severe oligodontia of the permanent dentition, the upper central incisors (11, 21) and first molars being present (16, 26, 36, 46) with persistence of the remaining deciduous teeth. The resulting dental formula was:

| 16 | 55 | 54 | 53 | 52 | 11 | 21 | 62 | 63 | 64 | 65 |
| 46 | | | | | | | 71 | 72 | 73 | 74 | 75 | 36 |

Furthermore, questionable loss of the interdental vertical dimension, full lips, pectus carinatum, clinodactyly of the fifth finger, and hypoplastic distal phalanges of the second toe were observed. The mother had a right single transverse palmar crease.

In all three family members, cranial radiographs showed two bilateral and symmetrical,
rounded, calcified masses (fig 4, top row) apparently localised at the level of the basal ganglia, hypoplastic greater wings of the sphenoid, narrowed, cribiform plate of the ethmoid bone, fused clinoid processes, hypoplastic nasal septum, and agenesis of all permanent teeth, except for the upper middle incisors and first upper and lower molars. These findings were all confirmed three months later on CT scan, except for the large calcifications, which were situated at the level of the choroid glomus (fig 4, bottom row) but not accompanied by signs of intracranial hypertension. The metacarpophalangeal profile (MPP) showed some shortening of all the second phalanges, more accentuated in the fifth finger. X rays also confirmed the shortening of the distal phalanx of the second toe. The remaining skeletal, ophthalmological, and cardiac examinations were normal.

Dermatoglyphic analysis showed: (mother) right: Wd, Wd, Wd, W, W; palmar a-b ridge count (a-b)=40; atd=40°; total finger ridge count (TFRC)=148; mainline formula=11.9.7.5'.13''-t, left: Wd, Lr, Wd, Wd, Lr; a-b=43; atd=40°; TFRC=130; mainline formula=9.7.5'.13'-t. (1st girl) right: Lu, Lr, Lu, W, Wd; a-b=44; atd=47°; TFRC=97; mainline formula=11. 9.7.5'.13'-t, left: W, Lr, Lr, W, W; a-b=37; atd=43°; TFRC=107; mainline formula=9.9.5'.13'-t. (2nd girl) right: Wd, Lr, A, Wd, Wd; a-b=40; atd=44°; TFRC=97; mainline formula=11. 9. 7. 5'. 13'-t, left: Wd, Lr, Lr, W, Wd; a-b=43; atd=44°; TFRC=93; mainline formula=11. 9. 7. 5'. 13'-t.

Direct counting of the patients' sweat pores in different areas of the palm and fingers was normal. The karyotype was normal. In spite of the absence of any neurological or bone density abnormality, the intracerebral calcifications led us to perform several laboratory tests, including calcium and parathormone level, which all proved to be normal.

We followed up the family for over eight years. During this period nothing of particular clinical interest was observed. The problem of the dental anomalies was solved by extracting the deciduous teeth and applying a permanent prosthesis in the mother and by applying a
conservative prosthesis to the deciduous teeth in the daughters. The growth parameters have remained on the 90th centile. School results confirmed the high intelligence level of the twins.

Discussion

Although the twins had been referred to at birth as monochorionic and identical, we also performed the following: dermatoglyphic analysis, blood groups, the Goldsmith twin similarity questionnaire, and the ponderal index (Pondex). The conclusions drawn from all the results were consistent with monozygosity.

The patients reported here showed not only dental abnormalities (the permanent teeth consisting merely of the upper central incisors and first molars with persistence of the remaining deciduous teeth), but also ocular proptosis resulting from malformations of the sphenoid and ethmoid and hypoplasia of the nasal septum. These anomalies, together with the inner telecanthus and full lips, give them a quaint, flat profile. The bone anomalies of the limbs are not typical, consisting of clinodactyly of the fifth finger, hypoplasia of the terminal phalanx of the second toe, and shortening of the second phalanx of the fingers at the MPP.

The total finger ridge count in the mother was 283, in the first twin 204, and in the second 204, all high values compared to the average 143.17 of a female population in southern Italy.

As shown in table 1, the expression of the gene defect is identical in all three subjects. The bilateral and symmetrical calcifications of the choroid glomus were very interesting. The latter were already evident when the girls were 12 years old and, judging by their size, which was more or less the same in the daughters and mother, were presumably present long before the age of 10 years; they were bilateral and symmetrical.

Similar calcifications may also be found in metabolic diseases, such as pseudohypoparathyroidism. Although the cyclic AMP after infusion of exogenous parathormone in serum was not measured because of the unavailability of this hormone, the suspected hypoparathyroidism or pseudohypoparathyroidism was excluded by the absence of biological or clinical signs, including the skeletal anomalies.

In the absence of any clinical pathological signs, the calcifications of the choroid glomus in all three patients may be considered physiological. Calcifications with similar characteristics in the first decade of life have either not been found at all or very rarely, in 0.5%. It is well known that although these formations can occur as early as the first three years of life, they are very uncommon in patients under the age of 10.

It is an established fact that familial calcifications of the choroidal plexus are present in some genetic diseases. However, a thorough review, using computerised dysmophologic databases (both London and POSSUM), did not show any association of choroid glomus calcifications with hypodontia or facial anomalies or both that was similar to the features in our family.

To our knowledge, oligodontia with persistence of the deciduous teeth and intracerebral calcifications has so far only been reported once, in two daughters of a consanguineous marriage showing the typical onion skin phenomenon, pathognomonic of lipoid proteinosis. The condition presented here is clearly different.

Thus, in spite of follow-up lasting over eight years and in the absence of other similar cases, only a familial factor or possibly a new malformation syndrome, clearly of autosomal dominant inheritance, can be proposed.

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