Two Novel Large ANKH Deletion Mutations in Sporadic Cases With Craniometaphyseal Dysplasia

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Two novel large ANKH deletion mutations in sporadic cases with craniometaphyseal dysplasia

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To the Editor:

Craniometaphyseal dysplasia (CMD; OMIM #123000) is a rare genetic disorder characterized by hyperostosis of craniofacial bones and metaphyseal flaring of long bones. Progressive bone thickening causes foraminal stenosis, which leads to the compression of cranial nerves and resultant facial palsy, hearing loss, and blindness. CMD is inherited as an autosomal dominant trait or occurs sporadically with apparent de novo mutations (1, 2). Here, we present two novel large deletions in exons 7 and 10 of ANKH from two unrelated patients without the family history of CMD manifesting severe forms of the disorder.

Female patient 1 (Fig. 1a) had acute-onset left facial nerve palsy and subacute upper airway obstruction at 10 weeks of age. At 8 months, a temporal bone computed tomography (CT) revealed significant sclerosis of the skull base, which led to her diagnosis. At 9 months, she was diagnosed with hearing loss bilaterally necessitating hearing aids. Ophthalmologic evaluation at 10 months noted poor visual fixation and tracking in her left eye, left exotropia, and primary optic atrophy. She developed nystagmus at 12 months and subsequently had two episodes of orbital cellulitis. Progressive overgrowth of her inferior and middle turbinates prompted surgical resection to relieve obstruction, which was complicated by the marked density of the bone. By 17 months, she had the eruption of six normal teeth with hypertrophic appearing gums. With length consistently about the 20th percentile (~1 SD), her head circumference measured 1.62, 2.54, 3.07, and 3.87 SDs above the mean for age at 4, 7, 10, and 17 months without hydrocephalus or narrowing of the foramen magnum. Alkaline phosphatase levels were persistently elevated, ranging from 427 to 1944 U/l (normal: 150–400). Other evaluations included normal total calcium levels, ionized calcium, which was normal at 1 week but elevated to 5.53 mg/dl (normal: 4.48–5.28) at 8 months, and phosphorus, which was 2.7 mg/dl (normal 4.5–6.7) at 2 months but normal at 8 months.
Patient 2 (Fig. 1b) is a 5-year-old female who was evaluated for thickened calvarium and temporal bone with bony sclerosis of the internal auditory canals, cochleas, and semicircular canals, bilateral obliteration of mastoid air cells, and mixed bilateral hearing loss. Progressive hearing loss was diagnosed at 2 years of age. At 5 years, height was 115.3 cm (95th centile) and head circumference was 54.5 cm (3.4 SD above the mean). She has macrocephaly, tall broad forehead, hypertelorism, telecanthus, epiblepharon, paranasal bossing, excess vertical face height, prominent maxillary alveolar ridge, and borderline low-set posteriorly rotated ears. Dental examination showed normal primary dentition with normal occlusion. Neurologic examination showed mild left facial palsy. Alkaline phosphatase was minimally elevated at 324 U/l (normal for age 93–309). Serum calcium was normal, 9.4 mg/dl (normal 8.9–10.4), as was phosphorus, 4.3 mg/dl (normal 3.0–6.0).

Sequence analysis of ANKH in patient 1 detected an extensive in-frame deletion of 18 nucleotides in exon 7 (c853-870del) affecting amino acids 285 to 290 (pV285-Y290del) (GenBank #BC014526) (Fig. 1c,d). Patient 2 had an extensive 12-base pair in-frame mutation in exon 10 of ANKH (c1178-1189del) affecting amino acids 393 to 396 (pT393-K396del) (Fig. 1e,f). Both deletions map to putative cytosolic regions of ANK. These mutations were not identified in the parents of the patients.

These novel 18 and 12 base pair in-frame deletions are the largest ANKH mutations causing CMD identified to date, in addition to the recently identified complex mutation of two missense point mutations and a 12-bp deletion (3). All other previously identified mutations were inframe deletions of a single amino acid, an insertion of one amino acid due to a splicing defect, or amino acid substitutions due to point mutations (1, 2, 4). Most of the mutations in ANKH that we have identified in more than 40 families and sporadic cases previously reported single amino acid deletions p.Phe377del and p.Ser375del. In two cases, we found a previously identified alanine insertion (p.Pro380insAla) and in two other cases we detected point mutations (p.Cys331Arg, p.Trp292Arg). We noted variable expressivity in families and in sporadic cases with the same mutation.

CMD mutations in ANKH affect exons 7–10, while mutations for chondrocalcinosis (CCAL2) involve exons 1, 2, and 12 (5, 6). In contrast to CMD, chondrocalcinosis is characterized by arthritic changes in joints due to the accumulation of calcium pyrophosphate dehydrate crystals, which form only in the presence of high pyrophosphate (PPi) levels. Interestingly, a novel mutation in exon 6 of ANKH (Leu244Ser) has recently been associated with the first human progressive ankylosis phenotype with mental retardation, hearing loss, ankylosis, periarticular ligament ossification, enthesopathy, and dentinogenesis imperfecta (7). To date, the structural and functional impacts of the distinct mutations in ANKH remain speculative.

In summary, we report two novel in-frame deletions in patients with non-familial CMD affecting multiple amino acids in ANKH. Although both patients described here are severely affected, the size of the deletion or class of mutation does not appear to correlate with the severity of the disease.

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References


Fig. 1.
(a) Cranio-metaphyseal dysplasia (CMD) patient 1 presents with frontal bossing, macrocephaly, and a left facial palsy. Her distal femur and proximal tibia show typical metaphyseal flaring. (b) CMD patient 2 presents with characteristic facies, hyperostotic cranial bones, and flaring of long bone metaphyses. (c) Electropherogram showing a heterozygous deletion in exon 7 of the ANKH genomic DNA from CMD patient 1. (d) Sequence comparison to wild type sequence shows an 18 bp/6 aa in-frame deletion. (e) Electropherogram showing a deletion in exon 10 of the ANKH genomic DNA from CMD patient 2. (f) Sequence comparison to wild type sequence shows a 12 bp/4 aa in-frame deletion.
deletion. (g) Mutations for CMD located in exons 7, 8, 9, and 10 of ANKH. New mutations indicated in red.