Ellis Van-Creveld Syndrome in an Ethnic Group in Nepal: Case Report With Review of Literature

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Abstract

Ellis van-Creveld syndrome is a rare autosomal recessive disorder characterized by acromesomelic dwarfism, postaxial polydactyly, oro-dental abnormalities and congenital heart disease. We are reporting three cases in two families of the same ethnic group from Western Nepal. Genetic testing should be done in this population in order to identify the mutation in the gene causing the syndrome.

Key words: Ellis van-Creveld syndrome, ethnic groups, Nepal

Introduction

Ellis van-Creveld syndrome is a rare autosomal recessive disorder comprising of abnormalities of the ectodermal, mesodermal and endodermal derivatives. Characteristic features are disproportionate short stature, postaxial polydactyly, hypoplastic nails, hypodontia, conical teeth, harelip, multiple frenula attaching the alveolar margin to the inner side of lips and heart defects¹. We report three cases in two families of the same ethnicity from Western Region of Nepal. This will be the second report of the case from a Nepalese family; the first being from a Gurkha family reported from Hong Kong².

Case 1: A 17 month old female, an only child of unaffected parents, with normal developmental history was brought to the Pediatric department for respiratory symptoms. She was born out of non-consanguinous union at term with a birth weight of 2.5 kg. Her present weight was 7 kg (< 3rd percentile for age); length 74 cm (between 3rd and 10th percentile); weight/length < 3rd percentile; head circumference 44 cm (3rd to 10th percentile). General physical examination revealed flat nasal bridge, high-arched palate, bilateral hexadactyly of hands (Figure 1), small phalanges, short limbs, absence of medial and lateral incisors, conical peg-shaped teeth, fusion of the middle part of the upper lip to the maxillary gingival margin creating a V-shaped upper lip, small dysplastic nails and long narrow thorax. Radiological investigations showed acromesomelia with flaring of metaphysis of lower ends of long bones, absence of terminal phalanges of hands, absence of carpal bones, bilateral postaxial polydactyly in hands with polymetacarpalia, flattened ilium with left tibial exostosis, delayed appearance of distal fibular ossification centres, normal skull and spine. Echocardiogram and abdominal ultrasound examination were normal.
**Case 2:** A newborn baby was admitted to the neonatal intensive care unit at 12 hours of life for severe respiratory distress. The baby was born to a 34 year old mother out of non-consanguinous marriage at term gestation by vaginal delivery with clear amniotic fluid. Family history showed an elder sister (Case 3) having short stature with dysmorphisms. There was a history of similar dysmorphisms in the paternal uncle's male child who had died at six months of age, apparently due to a cardiac defect and heart failure. Weight was 3.75 kg (75th to 90th percentile), length-50 cm (50th to 75th percentile) with US: LS ratio1.63:1 and head circumference 34 cm (50th percentile). General examination revealed bilateral post-axial hexadactyly and clinodactyly in the hands with syndactyly of second, third and fourth toes bilaterally with hypoplastic nails in all limbs. Hands and fingers were short. There was a hare-lip with upper lip fused to the upper alveolar margin by multiple frenula. The thorax was long and narrow. Respiratory system examination showed bilateral crackles in the chest; cardiovascular system showed tachycardia and a pansystolic murmur in the left lower sternal border, grade 4/6 clinically consistent with ventricular septal defect. Abdominal examination showed liver palpable 7 cm and spleen 3 cm under the respective costal margins. X-rays showed small thoracic cage with horizontally oriented ribs, thickened, irregular costochondral junctions, handle-bar shaped clavicles (Figure 2), shortened humerus and femurs bilaterally, dumb-bell shaped femur with flaring of metaphysis of lower ends of femur, squared iliac wings, flattened acetabulum with three downward projecting spikes (Figure 3). The baby had hypocalcemia and mild derangements in liver function tests. The baby was kept on continuous positive airway pressure and died 48 hours after admission before any other investigations could be performed.

**Case 3:** An eight years female child, elder sibling of Case 2; having a history of recurrent chest infections till the age of three years, with average intelligence and height of 106 cm (< 3rd centile; expected 124 cm) and US:LS ratio 1.46:1; bilateral post-axial hexadactyly in the hands, clinodactyly, short and stout hands, hypoplastic nails, high-arched palate, absence of upper and lower incisors, conical teeth, fusion of upper lip to alveolar margin and lower lip to lower alveolar margin by multiple prominent frenula (Figure 4). Cardiovascular system, skin and hair were normal. Skeletal survey showed highly placed handle-bar clavicles, acromesomelia of upper and lower extremities, dumb-bell shaped deformity of humerus and femur, mild bowing of humerus with exaggeration of epiphyseal depression by pointed metaphysis of humerus (Figure 5), flattened ilium, decreased angulation of femoral neck, broadening of upper tibial metaphysis, medially placed tibial epiphysis with hypoplasia in its' lateral aspect with notching in the middle (Figure 6), genu vulgam, congenital vertical talus (Figure 6), fusion of capitate and hamate in the hands (Figure 7), postaxial polydactyly (hands), polymetacarpalia, short broad metacarpals with decreased density of bones, normal skull, spine and chest. Other investigations like abdominal sonogram, echocardiogram, liver and renal function tests were normal.
Discussion

In 1940, Richard Ellis and Simon van Creveld described a syndrome comprising of chondrodysplasia, bilateral postaxial polydactyly of hands, small and dysplastic extremities and congenital heart defects. They termed the syndrome “Chondroectodermal dysplasia”, now known as Ellis van Creveld syndrome (OMIM 225500). In 1933, McIntosh had described a child with similar features but without heart defect. His case was included in the cases reported by Ellis and van Creveld. Since then, several cases have been described, with and without heart malformations. This is a rare condition; there have been only ~ 150 cases reported so far. Of these, ectodermal dysplasia is seen in ~93% cases. In 1964, McKusick et al reported 52 cases in the Amish community of Lancaster, Pa. The incidence in the Amish is 5 in 1000 while in the non-Amish population, it is estimated to be 1 in 60,000. Today, the syndrome has been found to affect all races especially where consanguinity is practiced. A history
of consanguinity is found in 30% cases. Genetic studies have identified mutations in two genes, EVC and LBN (EVC2) located head-to-head on chromosome 4p16.2.4 which produce similar clinical manifestations with variable expression6,7.

EvC syndrome presents with a characteristic tetrad:
1. Disproportionate dwarfism. Short stature may be present at birth or becomes more apparent with subsequent growth. Surviving adults have an ultimate stature of 110-150 cms. The severity of limb shortening increases from the proximal to the distal portions
2. Bilateral postaxial polydactyly (100% cases) of the hands, occasionally toes (10%)
3. Ectodermal dysplasia with dystrophic, small nails, thin sparse hair, hypodontia and abnormally formed teeth
4. Congenital cardiac malformations in 50% to 60% of cases, the most common being a single atrium and a ventricular septal defect. Congenital heart defect is considered to be the least constant of the tetrad3. Half of these children die during infancy because of cardiovascular-related complications. Several authors have reported cases where cardiac defect is absent leading to normal life spans8-11. In our case, two of three cases did not have an associated cardiac defect.

The most consistent oral manifestations are: fusion of the upper lip to the maxillary gingival margin, causing a slight V-notch in the middle (hare-lip). The anterior portion of the lower alveolar margin is serrated and multiple small labial frenula are present. Teeth tend to be small and conical, molars have abnormal cusps or accessory grooves and enamel may be hypoplastic. Other abnormalities such as genu valgum, genitourinary, hepatic, renal and CNS abnormalities are occasionally seen. The skin and sweat glands are usually normal. Most patients have average intelligence.

The chief differential diagnosis includes Jeune syndrome, McKusick-Kaufman syndrome and Weyer’s syndrome. Distinguishing features are significant respiratory distress, less severe short stature, renal and hepatic involvement, absence of ectodermal dysplasias and heart defect in Jeune syndrome12. Our second case (Case 2) had several features suggestive of Jeune syndrome; in addition he also had a cardiac defect and ectodermal defects which went in favour of EvC syndrome. McKusick-Kaufman syndrome can be differentiated by the presence of hydrometrocolpos. Weyer’s syndrome, in addition to facial abnormalities has many features similar to that of EvC syndrome but of a milder nature and is inherited in autosomal dominant manner. In some cases, there may be overlapping features of all these syndromes. The syndrome can be diagnosed in-utero from 18th week of gestation by ultrasound13,14. Increased fetal nuchal translucency at 13th week has also been associated with EvC15. Definitive diagnosis is molecular genetic testing for mutation of EVC and EVC2 gene4. Radiological findings suggestive of EvC syndrome are: narrow thoracic cage with short, horizontally aligned ribs, progressive distalward shortening of the long bones with slight curvature of humerus, dumb-bell appearance of metaphysis of lower ends of long bones, postaxial polydactyly of hands, polymetacarpalia, fusion of the hamate and capitate in older children, cone-shaped epiphysis of proximal phalanges, short broad middle phalanges with hypoplastic distal phalanges; premature ossification of femoral epiphysis at birth, short iliac wings with flattened acetabuli and hook-like downward projecting spikes on its’ medial and lateral aspects3,15. There can be hypoplasia of lateral proximal tibial ossification centres and exostosis at medial aspect of proximal tibia3,7. The metaphyseal border of the tibia appears angulated in the form of a double-sloped roof, with the outer slope being longer than the inner- this finding is considered to be pathognomonic of EvC syndrome3. All three of our cases had the radiological features suggestive of EvC syndrome, with the third case having the pathognomonic feature of tibial plateau deformity and carpal bones fusion with polymetacarpalia. In addition, she also had congenital vertical talus which has not been reported in other cases of EvC syndrome.

Management of a child with EvC syndrome is multidisciplinary; care should be provided by the neonatologist, odontologist, cardiologist/ cardiothoracic surgeon, orthopaedic surgeon for correction of bony deformities etc. The longevity usually depends on the presence of cardiac lesion. Growth hormone therapy for short stature has to be considered although it was found to be ineffective by some authors5. Genetic counselling is important as the recurrence risk in the next pregnancy is 25%.

Conclusion
EvC syndrome is a rare condition. Here we reported three cases in children from the same community and ethnicity. Genetic testing is necessary in these ethnic groups in order to identify the pocket of population in Nepal having the mutated gene.
Ellis Van Creveld Syndrome

References


