Late presentation of Ellis van Creveld Syndrome with Common atrium anomaly: A case report in a Nepalese Adult

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ABSTRACT

Ellis van Creveld syndrome (EVC) is a rare genetic disorder having autosomal recessive inheritance characterized by chondrodys trophy, polydactyly, ectodermal dysplasia, and various cardiac anomalies. Acromelic shortening of upper and lower limbs, genu valgum, deformed teeth, short ribs and narrow thorax and other systemic anomalies complete the picture of this syndrome. The patients with the syndrome rarely survive into adulthood. Here, we report a 30 year old male with EVC presenting for the first time in middle age with Common atrium anomaly.

INTRODUCTION

Ellis van Creveld syndrome (EVC) is a rare genetic disorder with disproportionate dwarfism characterized by postaxial polydactyly, several skeletal, oral mucosal and dental anomalies, nail dysplasia and presence of congenital cardiac defects. The syndrome, also known as chondroectodermal dysplasia, is an autosomal recessive disorder with mutations of the EVC-1 and EVC-2 genes located on chromosome 4P16. The first full description of the syndrome was given by Richard Ellis and Simon Van Creveld in 1940. A large number of cases have been reported in the Amish Community of Lancaster, Pennsylvania, USA, by McKuisk in 1964. Here, we report a case of EVC who presented for the first time at 30 years of age.

Case Report:

30 year old male presented with progressive shortness of breath since adolescence (Class- II- III NYHA) for which he was evaluated at a primary care centre. There was no history of consanguinity in parents. There was no detail documents available, however he was evaluated in a primary health centre and was prescribed with diuretics without detailed cardiac evaluation and remained on the same treatment for several years. This admission, he presented with history of progressive breathlessness for 15 days with associated orthopnea and paroxysmal nocturnal dyspnea for the same duration. He also developed progressive abdominal swelling and swelling of both limbs for 10 days duration along with increased bluish discoloration of his face and hands and feet.

On examination, he had a short stature of 130 cm and weight of 36 kg. His examination revealed six digits in each hand (X-rays as shown in Fig. 1 and 2). The distal and middle segments of hands and feet were short. Clinodactyly of fifth fingers were present along with dystrophic and hypoplastic nails (Fig 2). There was widened space between hallux and the rest of the toes as well as genu valgum being present (Fig 3). His oromaxillary examination revealed deficient mucocutaneous fold with ankyloglossia and dental malocclusion defects (Fig 4). The patient did his schooling till 10 th standard but had normal psychomotor and cognitive development. He was cyanosed (oxygen saturation ~80%) at room air. The jugular
venous pulse (JVP) was elevated with prominent ‘a’ wave. The patient had long and narrow appearing thorax with a precordial bulge. There was cardiomegaly, with apex beat at left 6th intercostal space, 4cm lateral to the mid-clavicular line. Grade III/III left parasternal heave was present. The first heart sound was loud; the second sound was widely split and fixed. The pulmonary component was loud. There was a grade 3/6 pansystolic murmur with inspiratory accentuation audible at left lower sternal border. Chest X-ray revealed cardiomegaly with right atrial enlargement while his ECG revealed normal P wave axis and right axis deviation, p’ pulmonale with RVH. His echocardiographic examination revealed situs solitus, enlarged right ventricle and a common atrial chamber without any interatrial septum (Fig. 5). The right and left components of the common AV valves were at the same place with evidence of regurgitation through both the components of the valve. There was evidence of pulmonary arterial hypertension (PAH) in the form of right ventriculo-atrial gradient of 89 mm Hg. A diagnosis of common atrium anomaly with severe PAH was made.

Fig : 1  Polydactyly

Fig 2 : Polydactyly with nail dystrophy

Fig 3 : Limb deformity with genu valgum, talipus equinovarus

Fig 4 : Oromaxillary defect with dental malocclusion, ankyloglossia

Fig 5 : Common atrium defect with TR jet and PAH
Discussion:

Ellis van Creveld syndrome (EVC) otherwise known as chondroectodermal dysplasia has autosomal recessive inheritance. It is a syndrome found in Amish population of Pennsylvania in USA, affecting male and female equally. Familial history is significant with parental consanguinity or with affected siblings or family members. Polydactyly is a constant finding in the hands, and is usually bilateral, postaxial and on the ulnar side. Polydactyly of the feet is present in only 10% of the patients. Recently, the syndrome has been included in a new class of human genetic disorders called ‘Ciliopathies’, where the underlying defect may be dysfunctional molecular mechanism in the primary cilia of cells. The phenotype of the syndrome is variable and can affect multiple organs.

Other features include genu valga, curvature of the humerus, talipes equinovarus pectus carinatum with long narrow chest. Congenital heart malformations are described in a 50~60% of patients. The anomalies include defects of the mitral and tricuspid valves, patent ductus arteriosus, ventricular septal defect, atrial septal defect and hypoplastic left heart syndrome which are the principal causes of decreased life expectancy in these patients. The disease has characteristic oral manifestations that help early diagnosis at birth or during early childhood. The most common among them include fusion of the upper lip to the gingival margin resulting in the absence of mucobuccal fold, broad maxillary labial frenum described as partial harelip, multiple small accessory frenula, ankyloglossia, malocclusion etc.

In the present index case, he had short stature, genu valgum, cyanosis, postaxial polydactyly of hands, hypoplastic and deformed nails and toes, and multiple frenula of lower lip were present. The patient had congenital heart disease in form of common atrium with severe mitral and tricuspid regurgitation and severe pulmonary hypertension. The syndrome is characterized by high mortality in infancy and early life because of severe restriction imposed by short ribs and narrow thorax. Presence of congenital heart diseases like common atrium, AV canal defect and VSD also contribute to early mortality.

Rudnik-Schoneborn et al described about 2 patients with EVC syndrome who were 18 and 30 years old respectively. Verbeek et al summarized growth data of 101 patients with EVC syndrome, including 10 patients above 20 years of age. We experienced this first case report from Nepal documented in literature of an adult with Ellis van Creveld syndrome presenting as complex congenital heart disease with Common atrium anomaly. One third of these patients die at the early age or at infancy from cardiorespiratory problems and those who survive require multidisciplinary approach for treatment i.e. orthopedic correction of genu valgum, amputation of extra digits, surgical repair of cardiac malformations and dental intervention for high caries risk individuals.

Conclusion:

A case has been reported here with classic features of Ellis van Creveld syndrome who presented late in life with little complications developed until presentation to the hospital. It is a rare congenital disorder, with a high mortality in early life, 1/3 of these patients die in infancy from cardiac and respiratory problems and those who survive require multidisciplinary approach for treatment.

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