Case Report

Cantu syndrome in an Egyptian child

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A R T I C L E   I N F O

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A B S T R A C T

We report a 3 month old female, third in order of birth of non consanguineous Egyptian parents with the typical features of Cantú syndrome including coarse features, low frontal hairline, hairy forehead, broad flat nasal bridge, anteverted nares, long philtrum, small low set ears, high arched palate, excess hair on the cheeks, short neck and excess hair over extremities and back. The patient had patent ductus arteriosus ligation, and mild pulmonary hypertension. Our patient has an affected mother which is consistent with autosomal dominant inheritance.

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1. Introduction

Cantu syndrome (CS) is a rare congenital systemic disease characterized by congenital hypertrichosis, a distinctive facial appearance, osteochondrodysplasia and cardiomegaly [1]. Affected individuals show great variability in clinical manifestations and severity. Cantú syndrome was first recognized in Mexico in 1982. It is an extremely rare genetic disease and only about 50 cases have been reported worldwide [2].

We report a case with the typical features of Cantú syndrome after taking consent of the parents.

2. Case report

A 3 month old female, third in order of birth of non consanguineous Egyptian parents. The patient was delivered at full term by cesarean section. Her birth weight was 3.850 kg. No problems were noted by the mother during pregnancy. The patient was referred to the Genetics Clinic, Pediatric Hospital, Ain Shams University complaining of abnormal features.

At birth, the patient had jaundice which necessitated admission to neonatal intensive care unit (NICU). Her birth weight was 3.850 kg. No problems were noted by the mother during pregnancy. The patient was referred to the Genetics Clinic, Pediatric Hospital, Ain Shams University complaining of abnormal features.

At birth, the patient had jaundice which necessitated admission to neonatal intensive care unit (NICU). Congenital heart disease was discovered by echocardiography during admission. At the age of 1.5 months, she had operation to correct patent ductus arteriosus (PDA). The patient had repeated chest infections and was admitted to hospital at the age of 2 months with pneumonia.

The mother had similar coarse features. Family history showed a previous sib death at the age of 40 days with congenital heart disease and the same features.

On examination, her weight was 6 kg (75th percentile), her height was 60 cm (50th percentile), her span was 60 cm and her skull circumference was 41 cm (75th percentile).

The patient had coarse features, low frontal hairline, hairy forehead, broad flat nasal bridge, anteverted nares, long philtrum, small low set ears, high arched palate, excess hair on the cheeks and short neck (Figs. 1 and 2).

She also had excess hair over extremities and back (Fig. 3). The patient had left posterolateral thoracotomy incision scar for PDA ligation (Fig. 4).

Cardiac examination was apparently normal. Abdominal examination revealed umbilical hernia. Neurologic examination and genital examination were apparently normal.

Abdomino-pelvic ultrasonography was normal. ECHO cardiography revealed PDA ligation, patent foramen ovale (PFO), and mild pulmonary hypertension. Skeletal survey was normal.

3. Discussion

Cantu syndrome is a rare condition. About 50 individuals have been reported previously [2]. Cantú syndrome is characterized by congenital hypertrichosis; distinctive coarse facial features: enlarged heart; additional cardiovascular abnormalities that may include patent ductus arteriosus, pericardial effusion, aortic aneurysm and skeletal abnormalities [2].

We report a 3 month old female with the typical facial features of Cantú syndrome including coarse features, low frontal hairline, hairy forehead, broad nasal bridge, anteverted nares, long phil-
All newborns with Cantú syndrome have hypertrichosis with thick scalp hair and excessive hair growth on the forehead that grows down onto the cheeks in front of the ears, back, and extremities as detected in our patient. The hypertrichosis usually persists over time [2].

The facial changes of Cantú syndrome are consistent and evolve over time (broad nasal bridge, epicanthal folds and full mouth). With age the face lengthens, the forehead becoming tall, and the chin prominent [3]. The mother of our patient had similar features.

Many newborns have macrosomia. Ultimate adult height is usually within the normal range; however, short stature has been seen in few individuals [2].

Macrocephaly, often present at birth, typically persists throughout life as detected in our patient. Some individuals who do not have macrocephaly at birth have developed progressive macrocephaly in childhood [2].

Characteristic skeletal abnormalities include thickening of the calvaria, broad ribs, platyspondyly, ovoid vertebral bodies, scoliosis, narrow thorax and shoulders, pectus carinatum, hypoplastic ischium and pubic bones, Erlenmeyer-flask-like long bones with metaphyseal flaring, narrow obturator foramen, and coxa vara. Generalized osteopenia, delayed bone age, and craniosynostosis have also been described [4]. Our patient had normal skeletal survey.

Many infants with Cantú syndrome are born with heart defects. Onset and severity of each heart problem is quite variable among affected individuals [5]. Cardiac manifestations such as patent ductus arteriosus, ventricular hypertrophy, pulmonary hypertension, and pericardial effusions are present in approximately 80% of cases [4]. Our patient had PDA ligation, PFO, mild pulmonary hypertension. Vascular abnormalities reported in this syndrome and which were not detected in our patient include tortuous retinal vessels and multiple tortuous pulmonary arteriovenous communications [3]. Abnormal vasculature in the brain
has also been seen in individuals with a pathogenic variant in ABCC9 [3]. Generalized edema, which may be present at birth, spontaneously resolves. Subsequently, edema involving the lower extremities and occasionally the arms and hands may develop over time, usually in adolescence or early adulthood [6].

Less frequent features include umbilical hernia, pyloric stenosis, gastroesophageal reflux, increased frequency of infections, tortuous arteries in the circle of Willis and internal carotids, multiple tortuous venous collaterals and lack of flow in the inferior sagittal sinus, cerebral atrophy and thin corpus callosum. Our patient had umbilical hernia and repeated chest infections.

Although the majority of affected individuals have normal intelligence, mild learning disabilities and/or developmental delays have been also observed, including delay in acquisition of early motor milestones and delay in speech development [7]. Our patient had motor milestones delay however; early motor developmental delay was gradually improved during follow-up periods. Behavioral problems can include anxiety, mood swings, obsessive-compulsive disorder, tics and rarely autism spectrum features [7].

The diagnosis of Cantú syndrome is established in a proband with characteristic clinical features and is confirmed by detection of a heterozygous pathogenic variant in ABCC9 or KCNJ8 [8]. The majority of genotyped CS patients are heterozygous for mutations in the genes encoding ABCC9 [9] in most cases, and KCNJ8 [10] in others. ABCC9 and KCNJ8 encode the channel regulator, SUR2, and the channel pore-forming subunits, Kir6.1 respectively [11].

Cantú syndrome was initially thought to be an autosomal recessive condition with affected sibs in the family of Cantú et al. and an affected child born to consanguineous parents [1]. However Robertson et al. performed a segregation analysis based on all reported families and concluded that the data made autosomal recessive inheritance unlikely. Lazalde et al. [12] suggested autosomal dominant inheritance is more likely than autosomal recessive inheritance and the affected sibs reported by [1] Cantu et al. could be explained by parental gonadal mosaicism. Grange et al. [7] reported a mother and 2 daughters with Cantu syndrome,
consistent with autosomal dominant inheritance. Our patient has an affected mother which is consistent with autosomal dominant inheritance.

Yearly echocardiogram and electrocardiogram are recommended in patients with Cantú syndrome to monitor cardiac size and function, as well as for evidence of pericardial effusion. These studies should be started in infancy, or as soon as the diagnosis is made, and continued throughout life. Consider brain MRI with magnetic resonance angiogram (MRA) and magnetic resonance venography (MRV) to evaluate persistent headaches or other neurological symptoms, given the increased risk for cerebral vascular abnormalities [2].

References


Fig. 4. Left posterolateral thoracotomy incision scar for PDA ligation (arrows) and hypertrichosis.