A Rare Case of Aplasia Cutis Congenita

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Abstract- Aplasia cutis congenita is a rare anomaly presenting with the absence of skin. No definite etiology is available. The most common site is the scalp. We present an instance with ACC occurring symmetrically in both sides of the body from chest to flank.

Keywords: Aplasia cutis congenital; Giant; Nonscalp

Introduction

Aplasia cutis congenita (ACC) is an uncommon anomaly of absence of skin. It may be localized or widespread. It most commonly presents as a solitary lesion of the scalp and may be as large as 70 percent of that site.

Although usually benign, they may be associated with other physical abnormalities and syndromes. Frieden classified them into 9 groups based on the number and the presence or absence of other anomalies (1). Nearly 86 percent belong to the first group with a solitary lesion. Involved areas are well-circumscribed, not inflamed, and vary in size from 0.5 to 10 cm or larger. At birth lesions may appear as scars or ulcers (1). On the scalp they may appear as parchment-like scars with alopecia.

The cause is not certain as it is the result of more than one disease process. Genetic factors, teratogens, compromised vasculature to the skin, and trauma are all implicated (1,2). There is no racial or sexual predilection. It is present from birth.

Maximum tensile force during the development of scalp hair whorl is implicated for the scalp lesion. Early rupture of amniotic membrane forming amniotic bands may also be responsible (1).

According to our knowledge, there is not any report of symmetric body involvement around the world. We present an instance with ACC occurring symmetrically in both side of the body from chest to flank.

Case Report

Clinical synopsis

A 3-day-old girl of nonconsanguinous parents presented at Ali-ebn-abitaleb Hospital (Zahedan, Iran) for a skin lesion that was present on both sides of her body birth. The lesions were quadrangular ulcers extending from the chest anteriorly to the flank posteriorly. Examination revealed to 15 x 5 cm ulcers within which there was a total absence of skin (Figures 1, 2).

There were no other organ abnormalities on clinical examination. Radiological examination and ultrasonography of the abdomen revealed no abnormalities. Routine investigations were within normal limits.
Aplasia cutis congenita

Discussion

Aplasia cutis congenita (ACC) is an uncommon anomaly of absence of skin. Frieden (1) created a classification system for aplasia cutis congenita consisting of 9 groups based on the number and location of the lesions and the presence or absence of associated malformations:

Group 1-This is scalp aplasia cutis congenita without multiple anomalies (3). Nearly 86% of all solitary lesions occur on the scalp. A collar of hair is often seen around the defect. It can be autosomal dominant (4) or sporadic (5).

Group 2-This is scalp involvement with limb anomalies.(6, 7, 8, 9) Adams-Oliver syndrome(10-14) is a distinct subtype in which distal limb reduction abnormalities are found in association with solitary midline scalp defects. More than 15 such cases have been reported, usually with an autosomal dominant inheritance pattern and variable genetic expression. The scalp lesions tend to be large. The most common limb malformation is hypoplastic or absent distal phalanges. Other anomalies may include cutis marmorata telangiectatica congenita, hemangiomas, cranial arteriovenous malformation, skin tags, supernumerary nipples, and woolly hair.

Group 3-This is scalp aplasia cutis congenita with epidermal and sebaceous (organoid) nevi (15,16), which also involve the scalp, usually adjacent to the cutis aplasia. Some patients have also had ophthalmic and neurologic findings typical of epidermal nevus syndrome, including seizures, mental retardation, corneal opacities, and eyelids colobomas. Inheritance is sporadic.

Group 4-This is aplasia cutis congenita often with a hair collar overlying deeper embryologic malformations.(17-20) Examples include meningoencephalocele, porencephaly, leptomeningeal angiomatosis, cranial stenosis, spinal dysraphism, gastrochisis, and omphalocele. The inheritance pattern in this group varies with the associated underlying condition.

Group 5-This is aplasia cutis congenita associated with fetus papyraceous or placental infarct (21-25). Extensive truncal and limb aplasia cutis congenita in a linear or stellate configuration is associated with the presence of fetus papyraceous. Fetus papyraceous is found at the time of delivery and results from the death of a twin fetus early in the second trimester. The surviving fetus is affected with aplasia cutis congenita and usually is otherwise normal.

Group 6-This is aplasia cutis congenita associated with simplex, junctional, or dystrophic types of epidermolysis bullosa (EB) (26-28). Many reports describe aplasia cutis congenita, usually occurring on the lower extremities, in patients eventually diagnosed with EB. Initially described as Bart syndrome, this type of presentation represents a variant of dystrophic EB. A subgroup includes the association of pyloric or duodenal atresia, ureteral stenosis, renal abnormalities, craniofacial abnormalities, nail dystrophy, and aplasia cutis congenita.

Group 7-This is aplasia cutis congenita localized to the extremities without EB (29-31). At least 2 families have been reported in which multiple members have had extensive aplasia cutis congenita on the pretibial lower extremities and the dorsal aspects of the hands and feet.

Group 8-This is aplasia cutis congenita due to teratogens. A few cases of aplasia cutis congenita have been linked to intrauterine infection with herpes simplex virus or varicella-zoster virus or to exposure to methimazole (32-35) in the treatment of maternal thyrotoxicosis during pregnancy. Imperforate anus has been associated with methimazole or carbimazole exposure during gestation.

Group 9-This is aplasia cutis congenita associated with malformation syndromes (36,37). Aplasia cutis congenita has been reported as a characteristic in many syndromes and more will be reported. Various syndromes (38-41) and dysplasias include trisomy 13 (Patau syndrome) with large membranous scalp defects, 4p- (Wolf-Hirschhorn) syndrome with midline scalp defects, Setleis syndrome with bitemporal aplasia cutis congenita and abnormal eyelashes, Johanson-Blizzard syndrome with stellate scalp defects, focal dermal hypoplasia (Goltz syndrome), amniotic band disruption
complex, oculocerebrocutaneous (Dellemann) syndrome, scalp-ear-nipple syndrome (Finlay-Mark syndrome), Kabuki syndrome (42), and 46XY gonadal dysgenesis. Reticulolinear aplasia cutis congenita on the face and the neck is a distinctive cutaneous manifestation in several syndromes linked to Xp22.

As mentioned above, there is no report of symmetric body involvement around the world. We suggest adding another group to Frieden’s classification system to include this rare form of ACC.

References


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