A case of Adams-Oliver Syndrome

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Abstract
A late preterm, 2.3kg, Male child was born to P2L2 mother by Normal Vaginal Delivery. Baby had a large scalp defect, acrania with hypoplastic and absent digits. Acrania portion of the skull was covered by a thick membrane. X-ray skull showed absence of skull vault, X-ray of extremities showed hypoplastic and absent digits. Neurosonogram was normal. Echocardiography done showed moderate PDA with left to right shunt. CT brain was normal. Child was diagnosed as a case of Adams– Oliver syndrome. It is an autosomal dominant disorder comprises aplasia cutis congenita with terminal transverse limb defects.

Key words: Aplasia cutis congenita, Hypoplastic and absent digits

Introduction
Adams and Oliver described eight members of a family with this disorder in 1945. More than 100 affected individuals have been reported[1]. Adams-Oliver syndrome is an autosomal dominant disorder with aplasia cutis congenita and terminal transverse limb defects[2]. We report a case of Adams Oliver syndrome with typical skin and limb defects without any major internal organ anomalies.

Case Report
A late preterm, 2.3kg, Male child was born to P2L2 mother by Normal Vaginal delivery to non-consanguineously married parents. Baby cried immediately after birth. Apgar was 5 and 8 at 1 min and 5 min respectively. No adverse perinatal events occurred during birth. Elder sibling was alive and healthy, had no scalp or limb defects. Both the parents had no scalp or limb defects. Baby’s vitals were within normal limits. Head circumference is 33cm. Baby had a large scalp defect, acrania with hypoplastic and absent digits. Acrania portion of the skull was covered by a thick membrane.

Systemic examination findings were within normal limits. Baseline investigations were within normal limits. On radiological examination skull showed absence of skull vault, and extremities showed hypoplastic and absent digits. Neurosonogram showed absence of any intracranial anomalies. CT brain taken showed normal intracranial structures. Baby had moderate PDA with left to right shunt in Echocardiography.

Discussion
In 1945, Forrest Adams and C. Peter Oliver from Minneapolis first reported this condition. This condition is transmitted by Autosomal dominant mode of transmission, with presence of vertex cranial defects resembling aplasia cutis congenita and terminal limb malformations. More than 100 affected individuals have been reported[3].
Figure 1: Acrania portion of scalp

Figure 2: Cutis marmorata

Figure 3: Hypoplastic digits in hand

Figure 4: Hypoplastic digits in feet

Figure 5: X-ray hand showing absent phalanges

Figure 6 and 7: CT brain showing normal intracranial structures
Mild growth deficiency (3rd to 10th percentile) is seen. Aplasia cutis congenita is seen over posterior parietal region, with or without an underlying bony defect. These skin defects of the scalp can typically be found as solitary or multiple, round-oval hairless scars.

Terminal transverse limb defects, including those involving lower legs, feet, hands, fingers, toes, or distal phalanges, short fingers and small toe nails. Cardiac defects are seen in 20% of affected individuals, including ASD, VSD, Coarctation of Aorta, obstructive lesions of the left heart, hypoplastic left and right ventricles, DORV, and DOLV. Cutis marmorata telangiectasia seen in 20% of cases.

An autosomal dominant inheritance pattern with marked variability in expression and lack of penetrance is seen in majority of cases. Gain-of-function mutations of ARHGAP31, a Cdc42/Rac1 GTPase regulator are responsible for the defects.

Autosomal recessive inheritance has been suggested in a few families. Recessive mutations in the dedicator of cytokinesis 6 (DOCK6) gene have been identified in some cases. Far more likely to have a severe phenotype with neurologic abnormalities and intellectual disability[1].

The pathophysiologic mechanism of these defects remains unclear, but several mechanisms have been proposed, including predisposition to amniotic rupture sequence, other forms of extrinsic trauma or compression, and vascular compromise. An intrinsic predisposition to interference with normal tissue development seems a likely etiology.

The association of cutis marmorata and the dilated and or tortuous scalp veins may be additional indicators pointing to an underlying predisposition to vascular compromise in “watershed areas such as the cranial vertex and limbs[2].

The male-to-female ratio of the affected cases is about 2:3 (in earlier reports of sporadic cases = m:f = 5:13; in earlier reports of familial cases = 16:26; in the present cases = 7:3; all together = 28:42).

Genetic counseling regarding the inheritance of this syndrome should be given to all parents. In genetic counseling autosomal dominant inheritance with great variability in expression of the syndrome of congenital scalp defect and distal reduction defects of the limbs should be stressed.

However, genetic counseling of families with a sporadic manifestation of the syndrome may be difficult. An unaffected parent may represent a nonpenetrant individual. Thus, the unaffected parents of an affected child always have some risk of having another affected child. Ultrasonic examination might be indicated in all potential affected pregnancies.

Larger scalp defects with underlying defects of bone, where the superior sagittal sinus or dura are exposed there is an increased risk of hemorrhage or meningitis. Early surgical intervention with grafting is indicated. Cases in which the sagittal sinus or dura is not exposed, healing without need for grafting almost always occurs.

Several syndromes with congenital skin and limb defects have to be differentiated from the syndrome of scalp defect with polydactyly. Distal deficiencies occur in the aglossia-adactyly anomaly (Hanhart syndrome), Poland complex, and as a part of the limb defects in focal dermal hypoplasia.

A characteristic pattern of congenital scalp defects can be seen in the syndrome of scalp defects and postaxial polydactyly, the syndrome of scalp defects and split-hand defect, trisomy 13, and Johanson-Blizzard syndrome.

The skin defects of the amniotic band sequence are rarely found as localized defects on the scalp. In epidermolysis bullosa dystrophica type Bart the defects are typically on the lower legs, and in focal dermal hypoplasia irregular atrophic areas are observed[4].
Conclusion
Adams Oliver syndrome is a rare disorder that can be associated with lethal major organ anomalies. Prognosis is good in majority of cases, especially those without any major organ anomalies. We report a case of AOS, most likely to be sporadic. The patient presented with isolated aplasia cutis congenital and terminal transverse limb defects.

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References


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