Twin Pregnancy with One Twin Acardius-Amorphus

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ABSTRACT
Acephalus Acardia is extremely rare complication of multiple pregnancies and very uncommon congenital malformation disclosed in fetuses and newborns due to twin reverse arterial perfusion syndrome. We report a case of twin pregnancy with one normal fetus and other Acardius Amorphus.

Key Words: Acephalus Acardia, Twin reverse arterial perfusion syndrome, Monozygotic twins.

INTRODUCTION
Multiple pregnancy accounts for 1.5% of all pregnancies, with approximate perinatal morbidity and mortality of 10%.1 Multiple pregnancy is complicated by congenital malformations twice as often as with singletons. Acephalus Acardia is extremely rare complication of multiple pregnancies and very uncommon congenital malformation disclosed in fetuses and newborns due to twin reverse arterial perfusion syndrome. In this condition the donor twin provides circulation for itself and the recipient acardiac twin. The acardiac twin is grossly abnormal with severe reduction anomalies of the upper part of the body. Acardiac “recipient” fetuses may have different morphological forms and often their birth weight is higher than that of the “donor” fetuses. The latter represent morphologic features of circulatory overload due to the necessity of supplying blood to the acardius amorphus fetus. They are usually born preterm with circulatory insufficiency, intrauterine hypoxia and with developmental anomalies. Survival of one of the “donor” fetus implies the necessity of early diagnosis to detect pregnancy pathology, possible intrauterine intervention and monitoring of the healthy twin.

CASE REPORT
A 25-year-old gravida 1 was referred to us with suspected twin pregnancy at 34 weeks of gestation. Examination revealed a term-sized gravid uterus with multiple fetal parts and an increased amount of amniotic fluid. Fetal heart sounds could not be localized. Investigations included complete blood count, urinalysis, syphilis screening, and blood glucose measurement. Ultrasonography revealed a monozygotic twin pregnancy with 1 acardiac fetus and 1 normal fetus with polyhydramnios (amniotic fluid index -24). The acardiac twin had absent upper limbs with absent head while the other twin had no visible malformation or features of congestive cardiac failure. Estimated fetal weight was 2.5Kg for the pump twin and 2.7Kg for the recipient twin. The mother was plan for caesarean section in view of both breech presentation, followed by delivery of 2 male fetuses weighing 2.4 and 2.6kg (twin A and twin B, respectively), with a normal single placenta. Twin B was acardius-amorphous (Figure 1), the head and neck were absent, and the trunk and upper extremities were not developed. The other twin was normal and had no features of hydrops (Figure 2). There was no obvious placental anomaly. The post-operative period was uneventful and the patient was discharged on 6th postoperative day.

DISCUSSION
Acardiac twinning is a rare congenital anomaly characterized by formation of a malformed fetus with an absent or rudimentary (but nonfunctional) heart. A variety of acardiac twins have been described based on the degree of cephalic and truncal maldevelopment. The acardiac-acephalus fetus has no cephalic development, while an acardius-anceps fetus has some cranial structures and/or neural tissue development. The acardius-amorphus fetus has cephalic structures with limited or no truncal development. The fourth type, the acardius-amorphus fetus, has the most severe malformation and lacks all cephalic and truncal differentiation.2

Acardiac anomaly usually occurs in monozygotic twins, although there are a few reports of dizygotic twins with a fused placenta. It is more common in female twins, and because the disorder is monozygotic, the twins are usually of same gender. The etiopathogenesis of this anomaly is abnormal placental vascular communication between the twins, leading to imbalance of interfetal circulation. Reversed blood flow in the umbilical artery of the acardiac twin causes atrophy of the heart and other organs.3 This flow pattern has been termed “twin reversed-arterial-
perfusion” (TRAP) sequence, in which the lower body of the fetus receives blood with more oxygen saturation and nutrients than the upper body, leading to maldevelopment of the head, neck, and upper extremities. The twin with cardiac activity may be hydropic or malformed in 9% of cases. This twin is often at high risk for congestive cardiac failure due to increased cardiovascular demand, resulting in preterm delivery. Perinatal mortality for the pump twin has been estimated to be up to 55%, while it is usually fatal for the acardiac twin.

Termination of pregnancy should be considered if acardiac twinning is diagnosed early in the pregnancy. However, early recognition and active management of the TRAP syndrome with maternal digoxin and indomethacin therapy can salvage the pump twin. This protocol has been reported to prevent congestive cardiac failure in the pump twin and polyhydramnios in the mother. Following are the strategies for managing the TRAP syndrome based on the published literature.

After exclusion of chromosomal aberrations and malformations of the pump twin, close sonographic follow-up may be initiated. Cord occlusion has been attempted by embolization, cord ligature, laser coagulation, bipolar diathermy, and monopolar diathermy, while intrafetal ablation has been performed with alcohol, monopolar diathermy, interstitial laser, and radiofrequency. With radiofrequency ablation, greater than 90% survival can be achieved in monochorionic diamniotic pregnancies complicated by TRAP sequence. An exact preoperative evaluation of the vascular flow including Doppler sonography is, however, mandatory. By virtually disconnecting the twins, transfusion of blood between them can be stopped. In a recent analysis, intrafetal ablation was associated with a longer duration of treatment delivery interval, greater gestational age at delivery, lower technical failure rate, lower rate of preterm deliveries, reduced rupture of membranes before 32 weeks, and a higher rate of clinical success than cord occlusion techniques.

CONCLUSION
Termination of pregnancy should be considered if acardiac twinning is diagnosed early in the pregnancy. Conservative treatment is best suited for salvation of the pump twin when the acardiac twin is less than one fourth the weight of the pump twin and there are no signs of impending heart failure. Invasive intervention is justified when the acardiac twin exceeds 70% of that of the pump twin; however, the prognosis appears to be poor. Intrafetal ablation is the treatment of choice because it is simpler, safer, and more effective than cord occlusion techniques.

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REFERENCES

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