TWIN REVERSED ARTERIAL PERFUSION SEQUENCE (ACARDIAC TWIN): A RARE CASE REPORT

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ABSTRACT

Acardiac twin is a rare congenital anomaly and is exclusively associated with monochorionic twin pregnancy. It is due to abnormal placental vascular communication between the two fetuses in the form of arterioarterial and venovenous communication (reversed perfusion). Acardiac twin receives blood from the pump twin. Twin Reversed Arterial Perfusion (TRAP) is a rare complication of monochorionic twins. TRAP sequence is known as acardius or chorioangiopagus parasiticus. It occurs in 1% of monochorionic twin pregnancies and in 1 in 35,000 pregnancies. We report a rare case of acardiac twin, the incidence among 1 in 80229 of total deliveries and 1 in 1922 of twin pregnancies (From January 2010 to September 2015) in our institution. Hence, we present this rare interesting case of acardiac twin (TRAP SEQUENCE) diagnosed and delivered at our institution.

KEYWORDS

Acardiac twin, TRAP Sequence.


INTRODUCTION

Acardiac twin or TRAP sequence is a rare complication of monochorionic twin pregnancies. Multiple pregnancy accounts for 1.5% of all pregnancies, acardiac twin incidence is 1 per 35,000 birth among 1% of monozygotic twins. It results from abnormal placental vascular anastomosis, blood from the normal pump twin reaches the perfused twin through arterioarterial anastomosis, flows through acardiac twin in reverse course, then returns to pump twin through venovenous anastomosis (TRAP). There is a high risk of cardiac failure and death of normal twin. Etiology is also due to primary defect in cardiac embryogenesis.

CASE REPORT

A 23-year-old female gravida1 abortion,1 booked case in a primary health centre was diagnosed to have acardiac twin at 16 weeks of gestation. A series of USG was done (Tabulated below). She came to our institution for antenatal check up at 25 weeks of gestation. Radiofrequency ablation was done at 28 weeks of gestation as the serial ultrasound showed there was an increased volume of the perfused twin. Following ultrasound showed there was a decrease in the volume of the acardiac twin. Patient admitted with bleeding per vagina at 34 weeks in our institution. Emergency LSCS was performed. First the surviving fetus was delivered in cephalic presentation, male baby weighing 1.760 grams, APGAR-7/10 followed by extraction of the acardiac twin weighing 300 grams. The acardiac twin had a malformed head, both upper and lower limbs were present. The male baby was shifted to neonatal care unit for close monitoring. Mother and the baby was transferred to the ward and discharged in satisfactory condition.

Table 1

<table>
<thead>
<tr>
<th>Date of USG done</th>
<th>Fetus A</th>
<th>Fetus B</th>
</tr>
</thead>
<tbody>
<tr>
<td>27/5/15 MCDA</td>
<td>GA-16W+5D, 14.2gms, single umbilical artery, common placenta, normal liquor, normal-doppler</td>
<td>Acardiac twin/TRAP, Volume of 28cc</td>
</tr>
<tr>
<td>9/6/15 MCDA</td>
<td>GA-18 Weeks+4 Days, 19.4gms</td>
<td>Volume- 44cc</td>
</tr>
<tr>
<td>22/6/15 MCDA</td>
<td>GA-20 Weeks+3 Days, 28.9gms</td>
<td>Volume-99cc</td>
</tr>
<tr>
<td>13/7/15 MCDA</td>
<td>GA-23 Weeks+3 Days, 44.3gms</td>
<td>Volume-170cc</td>
</tr>
<tr>
<td>27/7/15 MCDA</td>
<td>GA-25 Weeks+3 Days, 66.5gms</td>
<td>Volume-370cc</td>
</tr>
<tr>
<td>24/8/15 RFA</td>
<td>GA-28 Weeks</td>
<td></td>
</tr>
<tr>
<td>9/9/15 GA-31 Weeks+5 Days, 138.2gms, cephalic presentation</td>
<td>Volume-103cc</td>
<td></td>
</tr>
<tr>
<td>24/9/15 GA-33 Weeks+6 Days, 175.6gms, hydramnios</td>
<td>Volume-129cc</td>
<td></td>
</tr>
</tbody>
</table>

MCDA-Monochorionic diamniotic twin, RFA- Radiofrequency ablation.

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Blood from the normal pump twin reaches the perfused twin through arterio-arterial anastomosis flows through acardiac twin in reverse course, then returns to pump twin through veno-venous anastomosis (TRAP). Another theory states that there may be a primary defect in cardiac embryogenesis.\textsuperscript{3}

Acardiac twin is classified based on degree of cephalic and truncal maldevelopment. 1. Acardiac-acephalus has no cephalic development. 2. Acardius-anceps has rudimentary cranial structures. 3. Acardius-amorphous has severe malformation and lacks all cephalic and truncal differentiation. 4. Acardius-acromus has cephalic structures, all other structures are absent.\textsuperscript{4}

Diagnosis of acardiac twin should be made as early by ultrasound and Doppler. It helps to show the reversed blood flow through the umbilical artery to the affected fetus and to assess the weight ratio of the twins.\textsuperscript{5}

Moore et al. study concluded that perinatal mortality of 50%-55% is due to polyhydramnios leading to premature delivery secondary to congestive cardiac failure. If the weight ratio of acardiac twin to the weight ratio of donor fetus was >70%, the incidence of preterm delivery was 90%, hydramnios was 40%, congestive cardiac failure was 30% and if the ratio was <70% the incidence was 75%, 30%, 10% respectively.\textsuperscript{6}

CONCLUSION
Diagnosis of acardiac twin can be made in the first trimester itself by USG and Doppler. Early diagnosis of acardiac twin helps in prevention of preterm labour and diagnosing of cardiac failure in the pump twin.

REFERENCES