PRENATAL ULTRASONOGRAPHIC DIAGNOSIS AND CLINICAL MANAGEMENT OF ACARDIAC TWIN

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ABSTRACT

Acardiac twinning is a rare complication of monozygotic twin gestation occurring in 1/35,000 pregnancies, in 1/100 of all monozygotic twins and 1/30 monozygotic triplets. Twin condition is characterized by partial or complete lack of development of the heart in one of the twins, and requires that the normal twin (pump twin) provides circulation for itself as well as the acardiac sibling (recipient twin) by means of reverse circulation through large artery-to-artery and vein-to-vein anastomoses. The acardiac twin is grossly abnormal and the outcome is invariably fatal. Perinatal mortality for the normal twin is about 50-70%, as a result of congestive heart failure, polyhydramnios and preterm delivery.

In this report, we present a case of acardiac twin prenatally diagnosed by ultrasound. The principal sonographic features for prenatal diagnosis and the clinical management are briefly discussed.

Key words: acardiac acephalic twin, clinical management, prenatal ultrasonographic diagnosis

INTRODUCTION

The development of the acardiac anomaly is a rare complication of monozygotic multiple pregnancies. It was first described in the 16th century (Benedetti, 1533) and occurs in nearly 1/100 of monozygotic twin pregnancies an in one out of 35,000 deliveries and the condition has also been reported to occur in 1/30 monozygotic triplets(1,2).

It has been hypothesized that this form results from
the fused placentation of monochorionic twins, in which vascular anastomoses arise between the arterial circulation of the hemodynamically advantaged twin (pump twin) to the other one (recipient twin) by means retrograde flow. The recipient twin may display severe and lethal anomalies, including acardia and acephalus. The pump twin is structurally normal but, its expanded cardiac failure and without treatment dies in 50 to 75 percent of cases (3,4).

CASE REPORT

A 27 year-old woman, gravida 4, para 3; was referred to our institution at 23 weeks of gestation because of a twin pregnancy in which one fetus was suspected to be dead. Real-time ultrasound examination demonstrated monochorionic monoamniotic twin pregnancy with a single posterior placenta; twin A was in breech presentation with fetal measurements consistent with 23 weeks. No abnormalities were noted in this twin. Twin B, had no cardiac activity, had a large amorphus mass containing multiple cystic structures predominantly in the upper portion of the body and generalized edema. There was no evidence of a fetal head structure and of a central cardiac structure. Amniotic fluid volume was increased. (Figure 1) The diagnosis of a fetus acardius acephalus was made. The patient was admitted because of legal termination. Labour was induced by prostoglandin E1 analogs and 12 hours later twins were delivered. The normal twin weighed 600 gr and was a female. APGAR scores at one and five minutes were 1 and 0, respectively.

The acardiac twin weighed 680 gr (Figure 2). Upper extremites were absent, the spine and pelvic bones are abnormal; lower limbs were apparent (Figure 3). Eyes were identified, in addition rudimentary ears, mouth and nose were present (Figure 4) and ambiguous genitalia (Figure 5) was observed. Skeleton deformation on X-ray is presented in Figure 6. Autopsy couldn’t be done because the family were not give the permission for the autopsy.
Many different terms have been used to describe the acardiac twin; including monster, parasitic twin, holoacardiac fetus, amorphous twin and some other colorful descriptive names\(^\text{(5,6)}\). It is very hard to tell a woman that she is carrying a monster or a parasite. Since all these fetuses have no or only a rudimentary cardiac structure and since the etiology is possibly cardiac maldevelopment, the term ‘acardiac twin’ is appropriate to describe this anomaly.

The acardiac twin is a unique complication of monozygotic twin gestation in which the normal twin (pump twin) provides circulation for itself as well as the acardiac sibling by means of reverse circulation through large artery-to-artery and vein-to-vein anastomoses\(^\text{(7)}\). Although the etiology of this condition remains unknown, the most widely accepted hypothesis is the vascular reversal perfusion theory. According to this, large vascular anastomoses develop during early embryogenesis leading to competition between the two circulations. When arterial pressure in one twin exceeds that of the other, reversed circulation in one twin develops, with secondary disruption and reduction of morphogenesis resulting in the cardiac anomaly. Since the perfused twin has no direct vascular connection with the placenta, the blood enters directly through a single umbilical artery and exits through the umbilical vein. Thus pattern of circulation, known as the Twin Reversed Arterial Perfusion (TRAP) sequence, has been recently verified with doppler sonography by several authors. Since the lower part of the body receives better oxygenated venous blood through the hypogastric artery the most severe abnormalities are always present in the upper part of the body\(^\text{(8-11)}\).

The acardiac twin has been classified according to the anomalies found in the four groups: The clinician must be aware that classification may be difficult in certain cases.

1. Acardius acephalus: the most frequent form. Cranial portion is absent, upper extremities may not be present and there are rudimentary intrathoracic and intrabdominal organs.
2. Acardius anceps: There is partially developed head and brain tissue, imperfect face, the body and extremities are usually present.
4. Acardius acormus: The head is present but the body is absent with an umbilical cord attached to the head or the head directly attached to the placenta.

The categorization into one of the four types does not affect the management and is mainly of academic interest\(^\text{(2,9,12,13)}\).

Obstetric ultrasonography has made prenatal diagnosis of the acardiac twin possible. Sonographic findings includes absence of cardiac pulsation, amorphus shape of the cephalic pole, poor definition of trunk and
extremities, diffuse subcutaneous edema and abnormal
cystic areas in the upper part of the body. Differential
diagnoses include large sacrococcygeal teratomas,
omphaloceles and intrauterine fetal demise(14). However,
prenatal diagnosis of this condition can sometimes be
extremely difficult; frequently the acardiac fetus is mistaken
for a single fetal demise in a twin pregnancy, and serial
sonography is required to establish the correct diagnosis
by documentation of growth in the suspected ‘dead
twin’. Sometimes a heart beat may be seen, which may
be a rudimentary heart or the result of the normal twin’s
cardiac function(14-16).

Many obstetric and neonatal complications are
associated with the presence of an acardiac twin.
Obstetric complications are: polyhydramnios, preterm
labour, cord accidents, dystocia, hydrops fetalis, rupture
of the uterus, increased rate of operative deliveries and
intrauterine fetal demise. Neonatal complications are:
prematurity, twin-to-twin transfusion syndrome. As a
result, the normal fetus may develop congestive heart
failure and eventually hydrops fetalis(4,17-19).

Once the diagnosis of acardiac twin is made,
management must be focused on the normal twin, who
had a perinatal mortality rate of 50% or more(20,21).
Recently Moore et al. have proposed that the weight
ratio of acardius to the normal twin may predict
congestive heart failure and polyhydramnios in the
normal (pump) twin(20). With a ratio below 70% it is
possible to select a conservative management. If polyhydramnios develops, maternal administration of
indomethacin and/or serial amniocentesis have been
proposed to recuce amniotic fluid load. If congestive
cardiac heart failure develops in normal twin, maternal
administration of digoxin can be used succesfulled.
On the other hand, aggressive management options
which could improve the survival rate of the normal
(pump) twin are available. Selective termination of
one fetus is a treatment option in well-selected cases
of complicated monochorionic twin pregnancy. Case
selection appropriate timing of the procedure, and
chose of the optimal technique to be used remain
uncertain. When the umbilical cord or major fetal
evessels are subjected to obliteration, this should be
effected and permanent to arrest fetofetal transfusion
and to preclude postmortem interfetal hemorrhage.
The different reported techniques balance invasiveness
and coplexity versus efficacy, and no method clearly
superior. Embolization with different thrombogenic

substance has been described, but intrauterine death of
both twins has often been observed. Probably caused by
incomplete obliteration of all umbilical cord vessels or
migration of the substance(21,22). Conventional fetocide
techniques with intracardiac injection of potassium chloride
are not an option in monochorionic twin pregnancies
because the substance could embolize to the nonaffected
twin through the virtually ever-present placental vascular
anastomosis. More appropriate techniques aim at arresting
the umbilical cord flow completely and permanently. If
the occlusion is incomplete or becomes patent over time,
persistent fetofetal transfusion or acute agonal interfetal
hemorrhage may occur, which is a common event in case
of intrauterine fetal death in monochorionic twins.

Embolorization, neodymium/yttrium-aluminum-
garnet(Nd:YAG) laser, monopolar thermodacoagulation,
bipolar thermodacoagulation and fetoscopic ligation are
acceptably invasive procedures and have been suggested
for this condition; however none of them are universally
successful(21-25).

Nd:YAG is an infrared colored and 1,064-1,381 nm
wavelengthed laser beam. The energy of this laser,
provides tissue coagulation efficiently, is able to
penetrate human tissue to depths of 4mm or more.
Nd:YAG laser coagulation of the cord can be achieved
relatively quickly and easily by using a double-lumen
needle to accomodate a fetoscope and a laser fiber.
However, it has been shown to have a high failure rate
above 20 to 22 weeks of gestational age. Later in
gestation, the cord may become too hydropic, or the
vessels may be too large to allow successful
coagulation(26,27).

Monopolar thermodacoagulation of the cord has also
been attempted once, but it did not arrest umbilical
cord flow completely. Whereas, bipolar coagulation is
a new alternative technique to occlude the umbilical
cord by using available bipolar forceps. This technique
simultaneously obliterates both umbilical arteries and
the vein, causing immediate cessation of flow, and if
effective, it prevents postmortem interfetal hemorrhage
through a vessel remaining patent. Also, the procedure
can be done through one port. Moreover the technique
relies on existing instrumentation and has been shown
to be reproducible in different hands. Fetal medicine
specialists are familiar with performing invasive
procedures under real-time ultrasononographic guidance.
Ultrasonographic guidance also permits the procedures
inconditions in which fetoscopy is difficult such as
particles or blood. Bipolar coagulation is a well-known, easy and readily available energy modality and needed generators are present in any modern operating room. Unlike the monopolar coagulation technique, it also has the theoretical advantage that the electrical current does not travel through the umbilical cord, the placenta, or the body of the other twin because bipolar current passes only between the two blades of the instrument. Compared with the cord ligation, the duration of the procedure is markedly reduced (28-31). Fetoscopic cord ligation achieves complete and permanent arterial and venous occlusion, with survival rates of about 70%. The procedure is relatively complex and lengthy, and even in experienced hands it fails in about 10% of cases. Moreover postoperative rupture of the membranes occurs in over 30% of cases and is therefore a major drawback. This is much higher than the 10% preterm prelabor rupture of the membranes rate observed after other fetoscopic interventions, such as laser coagulation of the chorionic plate vessels for twin-to-twin transfusion syndrome (32). It has been speculated that a higher risk of preterm prelabor rupture of the membranes after cord ligation is related to the complexity of the procedure, the higher number of ports and longer operating times (33). For all of these reasons, a simpler but equally effective technique for the cord obliteration would be welcomed.

Alcohol injection is a noninvasive technique which has been used for management of acardiac twin pregnancies. Sepulveda et al. were the first to report ablation of a 23-week acardiac twin by alcohol injection into the intraabdominal portion of the umbilical artery under direct ultrasonographic guidance (34). The relative lack of invasiveness with this percutaneous technique requiring no special instrumentation was compelling. The risks of this technique are minimal because it is analogous to other end-organ or appendage ablations used by interventional radiology. The possibility of aliquots of alcohol migrating upstream from a properly positioned needle into critical areas of the pump twin’s circulation is remote (35,36).

Other technique which is selective delivery of the acardiac twin by hysterotomy and continuation of the pregnancy for the normal twin has been reported (29). Chromosomal analysis of the acardiac twin has usually been normal. However several cases of abnormal karyotypes have been reported on (37). In our case, we found a normal (46 XX) karyotype. In summary, the acardiac anomaly is a high-risk condition with exceedingly high perinatal mortality for the normal twin. No general consensus exists on the management of a pregnancy complicated by an acardiac twin. Many of these pregnancies are delivered prematurely because of polyhydramnios, preterm labour and premature rupture of membranes. A conservative approach seems reasonable. Serial amniocentesis may be needed for symptomatic polyhydramnios. Preterm labour precautions are necessary, as with any other kind of twin pregnancy. Fetal therapy may be indicated for complications secondary to the twin to twin transfusion syndrome. This report emphasizes the role of ultrasonography for early intratuterine diagnosis of acardiac accephalic twin and the management procedure of this syndrome; Early diagnosis may enhance rational management scheme for improving the prognosis of the normal (pump) twin.

REFERENCES


