Prenatally Evolving Ectopia Cordis with Successful Surgical Treatment

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Case Report

The 28-year-old healthy patient (G1, P0) had no history of congenital anomalies. Screening ultrasound performed at 6 and 10 weeks of pregnancy were described as ‘normal’. NT evaluation and nasal bone at gestational week 12 were normal. Subsequent prenatal ultrasonography at gestational week 18 showed omphalocele and ectopic heart located partially outside the thorax. Fetal echocardiography revealed ventricular septal defect and enlargement of the right atrium and ventricle, insufficiency of the tricuspid valve and hypoplastic root of the aorta (fig. 1). The pregnant women was offered cytogenetic evaluation and termination of pregnancy; however, she decided to continue her pregnancy and delivered in a tertiary obstetrical and cardiac center. Fetal echocardiography was performed at gestational weeks 19, 24, 33 and 38 (fig. 2). The cardiovascular profile score was always 10 and did not change during the fetal monitoring. Heart anatomy assessment was similar as in the first echocardiography; however, the position of the fetal heart and size of omphalocele changed: there was an improvement in the fetal atria and ‘niche’ in the fetal chest and a decrease in size of fetal omphalocele (fig. 3). At gestational week 39, an elective cesarean section was performed: a boy, weighting 3,400 g, was delivered and immediately transferred from the delivery ward to the cardiosurgery theater. Physical and echocardiographic examination pointed out ectopic heart with double outlet tract from the right ventricle, ventricular septal defect and small epigastric omphalocele (fig. 4). The cardiac surgeon performed a single-stage closure of the abdominal defect, displacement of the ectopic heart and banding of the pulmonary artery. The operation succeeded without any complications (fig. 5).
After 53 days the child was discharged from the hospital. At 6 months of age he was doing fine and was scheduled for final cardiosurgery repair at 10–12 months of age.

Discussion

Ectopia cordis (EC) is a rare malformation, occurring in 5.5–7.9 per million live births due to failure of maturation of the midline mesodermal components of the chest and abdomen. It can be defined as a complete or partial displacement of the heart outside the thoracic cavity. EC can be classified into four types according to the cardiac location: thoracic (60%), abdominal (30%), thoracoabdominal (7%) and cervical (3%) [1]. Ultrasound is the primary diagnostic modality for the diagnosis of fetal anomalies, allowing the detection of complex lesions such as EC at an early stage of gestation. EC comprises 0.1% of congenital heart diseases. Common cardiac anomalies associated with this malformation are: ventricular septal
defect (100%), atrial septal defect (53%), and tetralogy of Fallot (20%) [2]. Extracardiac anomalies associated with EC described in the literature are amniotic band syndrome, diaphragmatic hernia, body stalk syndrome, omphalocele, cleft lip and palate, and skeletal malformations such as kyphosis. EC with sternal cleft, omphalocele and the ventral diaphragmatic hernia raises the possibility of pentalogy of Cantrell (table 1). The defect of the abdominal wall can range from simple diastasis to huge omphalocele with bowel, liver, and heart in the defect covered by a translucent membrane [3].

When several anomalies are present, the fetal prognosis is usually dismal despite normal fetal karyotype and we would consider EC as a lethal anomaly. Despite advances in neonatal cardiac surgery, complete thoracic EC remains a surgical challenge with only few long-term survivors. The possibility and efficacy of surgery in a surviving neonate depends on the degree of EC and co-existing congenital heart defects and/or absence of extracardiac malformations. The aims of surgical treatment are to provide a soft tissue cover of the heart, to reduce the heart into the thoracic cavity, palliation or repair of intracardiac anomalies and reconstruction of the chest wall. The major peri- and postoperative problem is avoidance of high abdominal and intrathoracic pressures occurring because of repositioning of the heart in a small thorax [4].

During fetal life the EC may change with time, as presented in our case: there was significant regression of the huge omphalocele at mid-gestation compared with the end of pregnancy; a similar positive evaluation was described by Zidere and Allan [5].

The presented case is important in two ways: firstly, it showed the possibility of the fetal heart changing position during pregnancy, and secondly that immediate postnatal surgery might be planned based on prenatal findings. Detailed fetal echocardiography has helped in counseling and choosing the optimal term, place and method of delivery and the optimal postnatal care in this unique case, originally presented to the parents as a lethal malformation.

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<td>Beckwith-Wiedemann syndrome</td>
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References