EMERGENCY CESAREAN SECTION IN CASE OF ANEURYSM OF THE MUSCULAR INTRAVENTRICULAR SEPTUM WITH NEONATAL FOLLOW-UP - A CASE REPORT AND REVIEW OF THE LITERATURE

Abstract
We present the case of aneurysm of the muscular intraventricular septum with accompanying cardiomegaly and abnormal venous flow patterns requiring emergency cesarean section and specialized neonatal treatment. At the age of 8 months our patient has no clinical symptoms with a 6 mm scar in intraventricular muscular septum.

Key words: cesarean section, fetal echo, aneurysm of the septum, cardiomegaly

INTRODUCTION
In the literature regarding fetal cardiology there are several case reports about prenatal detection of aneurysms or diverticulum of the fetal heart at different levels.

Usually they do not influence fetal growth, mainly their nature is benign with the tendency to resolve.

Here we report a special case with dramatic presentation in the third trimester, which thanks to the good cooperation between perinatologists, had a successful outcome.

CASE REPORT
A 20 year old prima gravida suffering from bilateral hydronephrosis observed since 20 weeks of gestation, that underwent two DJ shunt implantations, repeatedly treated with antibiotics, was referred and transported by medical transport from Opole to our prenatal cardiology center in Lodz at 35 weeks of gestation due to suspected fetal congenital heart malformation.

The previous US examinations at 8, 16, 20, 24, 28, 30, 31, 33 weeks of gestation were described as normal.

The abdominal US examination of the gravida showed a strongly dilated pelvicalyceal system of the right kidney, duplication of the left kidney pelvicalyceal system with moderate dilatation. The gravida was hospitalized several times, decompressions of the hydronephrosis were conducted twice without effect and at 33 weeks of gestation the gravida refused reimplantation of the DJ shunt and nephrostomy. Oral antibiotics were administered on an outpatient basis.

Due to cervical insufficiency at 35 weeks the gravida was admitted for hospitalization in the obstetrical department and upon fetal ultrasound examination significant cardiomegaly, abnormal axis of the heart, and ventricular septal defect were suspected. The gravida was transported from her city to our center (215km) for an urgent consultation.

The ultrasound exam in our unit revealed a single male fetus in cephalic longitudinal position with LGA biometry (34.5hbd by LMP, 38.1 hbd by ultrasound) and Heart area/ Chest area ratio of 0.5. The four chamber view was grossly abnormal showing two asymmetrical atria and two asymmetrical ventricles with no contraction of RV. The foramen ovale was 4 mm, systemic and pulmonary venous return were normal. Abnormal flow through the ductus venosus and pulmonary veins were observed with reversal

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flow $v=37\text{cm/s}$ suggesting postnatal hemodynamic and respiratory distress. The CVPS was 6 points (minus 2 for heart size, minus 1 for RV function, minus 1 for abnormal DV flow). The initial diagnosis of acute ischemia of the right ventricle and septum was made in differentiation with severe myocarditis. The gravida was referred to the emergency department with an urgent recommendation of delivery by cesarean section, which was performed on the same day.

Male neonate, weighing 3000g received 5/6/8/8 points in the Apgar scale, required assisted mechanical ventilation for 2 days and further nCPAP support for 3 days.

Echocardiography hour after birth confirmed a significant degree of right ventricular dilatation with hypokinesis and abnormal structure and motion of the lower muscular part of the interventricular septum: bulging aneurysmal structure into the right ventricle.

Screening for infection was conducted, and broad-spectrum antibiotics modified by antibiogram were administered, as well as dopamine and Corotope.

Laboratory studies showed an increase in Troponin T 128.5 (pg/ml), (maximum 597.5pg/ml on 24th day of
life), NTproBNP=30038 (pg/ml), CK-MB (93 IU/L). During the hospitalization gradual improvement in clinical status was observed along with the normalization of test results. Upon discharge: CK-MB 57IU/L, Troponin T 228.6(pg/ml), NTproBNP 1491 (pg/ml). Echocardiography showed improvement in contractility within the septum and the gradual reduction in the size of the right ventricle improving its function.

ECG also showed variation from the initial dominant features of large pre-excitation to the dominant rhythm with normal PQ and the image of the right bundle branch block. Holter examination (24-hour ECG) showed 2% of single extrasystoles of atrial origin. The baby boy in good general condition, with no signs of heart insufficiency was discharged home at 3 weeks of postnatal life with a body weight of 3500g, with recommendation of continuation of pharmacological treatment with Enareal and Acesan. At the age of 8 months baby boy is asymptomatic and on echo has a 6 mm scar at the level of lower part of intermuscular septum (Fig. 5,6)

**DISCUSSION**

Congenital malformations of the muscular intraventricular septum are rare, however described in prenatal cardiology [1,2,3,4,5,6]. Left and right ventricular outpouchings are also described [7,8].

Aneurysms of the ventricular septum are usually considered to be acquired, a result of infarction, cardiomyopathy, anomalies of the coronary arteries, Kawasaki disease, sarcoidosis or as complications from cardiac surgery [4,6]. More common are aneurysms of the membranous section of the intraventricular septum and are often associated with ventricular septal defects, with
<table>
<thead>
<tr>
<th>Name of author</th>
<th>Year of publication</th>
<th>Prenatal diagnosis</th>
<th>Postnatal diagnosis</th>
<th>Vaginal delivery or CS</th>
<th>Follow-up</th>
</tr>
</thead>
<tbody>
<tr>
<td>Alday 14</td>
<td>1976</td>
<td>No</td>
<td>20th day Aneurysm</td>
<td>Unknown</td>
<td>Cardiac cath, 2nd year open heart aneurysmectomy</td>
</tr>
<tr>
<td>Chen 3</td>
<td>1991</td>
<td>No</td>
<td>29 year old man and his 4 year old son</td>
<td>Both symptom free</td>
<td></td>
</tr>
<tr>
<td>Mohan &amp; Kumar 15</td>
<td>1992</td>
<td>No</td>
<td>18 year old (male)</td>
<td>Asymptomatic</td>
<td></td>
</tr>
<tr>
<td>Prabhakar 5</td>
<td>1993</td>
<td>No</td>
<td>1 Year old</td>
<td>Ventricular septal aneurysmectomy performed successfully</td>
<td></td>
</tr>
<tr>
<td>Eriksson 2</td>
<td>1996 3 cases</td>
<td>No</td>
<td>One family</td>
<td>Asymptomatic</td>
<td></td>
</tr>
<tr>
<td>Fujiwara 4</td>
<td>2001 2 cases</td>
<td>26 wks- VSD</td>
<td>Aneurysm</td>
<td>Vaginal delivery</td>
<td>Asymptomatic</td>
</tr>
<tr>
<td>Case 2</td>
<td>28 wks</td>
<td>Aneurysm</td>
<td>Vaginal delivery</td>
<td>Asymptomatic</td>
<td></td>
</tr>
<tr>
<td>Donofrio 10</td>
<td>2002</td>
<td>26 wks</td>
<td>Confirmed</td>
<td>Vaginal delivery at term</td>
<td>Cardiac catheterization on 5th day, digoxin</td>
</tr>
<tr>
<td>Baspinar 16</td>
<td>2005</td>
<td>No</td>
<td>5 months - aneurysm</td>
<td>Vaginal delivery at term</td>
<td>Pathological ST changes + WPW + minimal RVOT stenosis + RV hypertrophy, No clinical symptoms</td>
</tr>
<tr>
<td>Wong 17</td>
<td>2007</td>
<td>No</td>
<td>5 days- aneurysm</td>
<td>Vaginal delivery at term</td>
<td>L &amp; RBBB + abnormality of the short arm of chromosome 20, (46, XX, add (20)(p13) progressing well from a cardiac perspective but has poor muscle tone and developmental delay</td>
</tr>
<tr>
<td>Nicolae 18</td>
<td>2007</td>
<td>No</td>
<td>Yes</td>
<td>unknown</td>
<td>Surgical repair</td>
</tr>
<tr>
<td>Nguyen A 19</td>
<td>2008</td>
<td>28t wks</td>
<td>Large PDA, moderate biventricular dysfunction, Severe thinning of basa to mild ventricular septum resulting in RVOTO &amp; critical tricuspid stenosis with absent pulmonary valve</td>
<td>unknown</td>
<td>Died</td>
</tr>
<tr>
<td>Nguyen B</td>
<td>2008</td>
<td>23rd wks</td>
<td>Severe TV stenosis, hypertrophied RV free wall, multiple RV sinusoids</td>
<td>unknown</td>
<td>BTS, BDG, modified Fontan, resolution of septal aneurysm at 2yr of age, alive at 5 yr</td>
</tr>
</tbody>
</table>

Table 1: Echocardiographic findings and neonatal outcome in analysed population (part 1)
<table>
<thead>
<tr>
<th>Nguyen C</th>
<th>Thin ventricular septum aneurysm bulging into LVOT without obstruction and pulmonary atresia with IVS</th>
<th>Hypoplastic TV stenosis, hypertrophied RV free wall, multiple RV sinusoids</th>
<th>unknown</th>
<th>BTS, BDG, modified Fontan, alive at 3yr, bulging aneurysm without LVOTO</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nguyen D 2008</td>
<td>Large ventricular septal aneurysm bulging from the body of RV into the LV causing mild LVOTO and pulmonary atresia with nearly IVS</td>
<td>Severe TV stenosis, additional small VSD, single LCA, multiple RV sinusoids</td>
<td>unknown</td>
<td>BTS, septal plication of aneurysm, BDG, modified Fontan, alive at 5yr, no LV obstruction</td>
</tr>
<tr>
<td>Carr 20 2008</td>
<td>19 wks</td>
<td>Confirmed aneurysm</td>
<td>unknown</td>
<td>Dilated cardiomyopathy requiring heart transplant</td>
</tr>
<tr>
<td>Lalit 21 2015</td>
<td>Male 48 years</td>
<td>Incidental findings</td>
<td>---</td>
<td>---</td>
</tr>
<tr>
<td>Our case 2015/2016</td>
<td>8, 16, 20, 24, 28, 30, 31, 33 US no abnormalities 35 wks RV akinesis</td>
<td>Aneurysm</td>
<td>URGENT Cesarean Section</td>
<td>Alive &amp; Well, no surgery, Echogenic “scar” in lower muscular part of the IVS</td>
</tr>
</tbody>
</table>

Table 1: Echocardiographic findings and neonatal outcome in analysed population (part 2)

reports by echocardiography reaching the year 1969.\(^9\)

To our knowledge we report the first case of fetal aneurysm of the muscular interventricular septum, requiring immediate intervention and delivery by caesarian section due to acute cardiomegaly and virtually asystole of right ventricle.

The first prenatal diagnosis of an aneurysm of the muscular IVS was that presented by Fujiwara et al.\(^4\) (Table 1) in two brothers diagnosed at 26 and 28 weeks of gestation. In both cases the neonatal condition upon delivery was good and neither brother presented with cardiac or respiratory failure, as opposed to our patient.

Upon literature review it is theorized that this type of aneurysm may be a result of an embryologic defect of the myocardium.\(^1,2,3,4,5\) The first case report suggesting the familial occurrence of muscular aneurysms of IVS was made by Chen et al.\(^3\) in the year 1991: in this case both father and son were diagnosed. Such a theory is supported by another study by Eriksson et al.\(^2\) and Donofrio 2002. Eriksson\(^2\) described the familial occurrence of such aneurysms in 2 brothers and their father. These reports seem to support the concept that muscular aneurysms are congenital embryological defects of the myocardium, although the exact inheritance mechanism is yet to be determined. As in the cited cases our patient was also male. According to adult cardiovascular assessment, the grandfather’s heart was anatomically normal, father’s medical history is not available at the current moment.

It seems that fortunately our patient was referred to our center at the last moment, which resulted in immediate CS due to acute cardiomegaly, grossly impaired RV function and Doppler flow patterns (6 points CVPS)\(^11,12\) allowing for intensive, targeted neonatal therapy in the tertiary medical center for the baby (advantage of “in utero” transport of 215km instead of by neonatal ambulance).

In the literature upon prenatal screening the initial diagnosis that was made was that of a ventricular septal defect.\(^4\) In our case the referring obstetrician also made this false diagnosis. However what was striking in the presented case was fetal heart cardiomegaly with large disproportion of the four chambers in favor of the right side of the heart. Previous cases described good clinical state upon delivery.\(^4\) Our patient after birth received a poor Apgar score, initially required nCPAP FiO2 0.35 and was ultimately intubated because of increasing respiratory failure mainly due to cardiomegaly, and not only because of the aneurysm of the septum. As in the cases presented by Eriksson et al.,\(^2\) our patient also presented with abnormal ECG, which evolved in postnatal life. It is too early to state further prognosis and future impact. The other postnatal finding in the second month of life was of course the abnormal presentation of muscular intraventricular septum-
a scar after prenatal ischemic event?

Taking into consideration this case and our previous experience [13], and also the data from literature (table1) which shows that the postnatal course of fetal muscular septum aneurysm could be very different from cases that are asymptomatic for a long time to those resulting in early death, requiring cardiac surgery or heart transplantation,²,³,4,5,10,14,15,16,17,18,19,20 this is further proof that with the current knowledge and ultrasound machines widely available, third trimester fetal echocardiography is justified. The gravida was examined 8 times before the final diagnosis was made, 8 ultrasound examinations were described as normal, the 9th revealed cardiomegaly which was the main cause for targeted echocardiographic examination. Maybe it is worth considering again introducing mandatory third trimester echocardiographic examinations? It is important to remember that this III trimester examination is not only for the late diagnosis of congenital malformations but mainly to assess the hemodynamic status of the fetus and detect acute cardiovascular changes that will require immediate obstetrical intervention and targeted neonatal treatment. Thus “late echocardiography” may allow, like in this case, for transport in utero and optimal management of neonate upon delivery.
Emergency cesarean section in case of aneurysm of the muscular intraventricular septum with neonatal follow-up - a case report and review of the literature

References


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Division of work:
J. Plużańska: preparing the manuscript , review of the literature, submitting the manuscript

Co-authors: work with the manuscript
Maria Respondek-Liberska: idea of the article, diagnosis of the case, documentation of the case, correction of the manuscript

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