A Case of Progressive Ductal Constriction in a Fetus

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The ductus arteriosus is a normal and essential structure in fetal circulation. Since the introduction of fetal echocardiography, there have been reports of ductal constriction, many of which were related to maternal use of some medications. However, there have been some reports of idiopathic ductal constriction, which usually present in later gestation. Recently we experienced a case, which initially showed an S-shaped ductus with mild narrowing at 23 weeks and 27 weeks gestation and developed severe ductal constriction at 33 weeks. Soon after birth, ductus was searched for but no ductus was found in 2-D and color images. The neonate required mechanical ventilation with supplemental oxygen for 3 days. All echocardiographic abnormalities were normalized in 7 months. We report progressive ductal constriction in an S-shaped ductus and emphasize the importance of continuous follow up extending to the third trimester and even immediately after birth. (Korean Circ J 2013;43:774-781)

KEY WORDS: Ductus arteriosus; Constriction, pathologic; Ultrasonography, prenatal; Echocardiography; Prenatal diagnosis.

Introduction

The ductus arteriosus (DA) is a normal and essential structure in fetal circulation, and carries most of the right ventricular output to the descending aorta. In fetal life, ductal patency is actively maintained by factors such as prostaglandin E₂ produced in the placenta, endothelial nitric oxide (NO) synthase and low oxygen tension of the blood.

Since the introduction of fetal echocardiography, there have been reports of ductal constriction, many of which are related to maternal use of some medications. However, there have been some reports of idiopathic ductal constriction, which usually present in later gestation. Recently we experienced a case, which initially showed an S-shaped ductus with mild narrowing at 23 weeks and 27 weeks gestation and developed severe ductal constriction at 33 weeks. Soon after birth, ductus was searched for but no ductus was found in 2-D and color images. The neonate required mechanical ventilation with supplemental oxygen for 3 days. All echocardiographic abnormalities were normalized in 7 months. We report progressive ductal constriction in an S-shaped ductus and emphasize the importance of continuous follow up extending to the third trimester and even immediately after birth. (Korean Circ J 2013;43:774-781)

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Case

A 22-year-old woman, gravid 2, para 1-0-0-1, was referred to our institution for fetal echocardiography at 23 weeks of gestation because of abnormally looking great arterial arches. The pregnancy had not been complicated. The mother had not had any diseases or taken any drugs or herbal medications. There was no family history of congenital heart disease. Obstetric evaluation showed normal fetal growth, normal amount of amniotic fluid and normal placenta. The fetal echocardiography revealed the normal four chamber view, where two atria and two ventricles were normal in size and thickness (Fig. 1A). The two atrioventricular valves moved normally. Two
great arteries arose normally from the appropriate ventricles. The pulmonary and aortic valves looked normal and the size of the main pulmonary artery and aortic root was within normal limits. The aortic arch was normal in size and location, but the ductal arch was not visualized in any views. The main pulmonary artery bifurcated into two branch pulmonary arteries but was not continuous with the DA (Fig. 1B). However color Doppler examination showed a tortuous ‘S’ shaped ductus cuddled inside the aortic arch (Fig. 1C and D). The DA joined the descending aorta with an obtuse angle and showed a flow signal from the main pulmonary artery to the descending aorta (Fig. 1E and F). Color aliasing occurred along the course of the ductus but pulsed Doppler measurement of velocity was not attempted because of the poor angle of insonation. Flow velocity and waveforms of the two great arteries were within normal limits.

Fig. 1. The fetal echocardiography was done at 23 weeks of gestation. A: the four chamber view shows normal ventricular size and wall thickness. B: aortic arch was normal in size and location, but the ductal arch which connected with the main pulmonary artery (MPA) was not visualized. C: the main pulmonary artery bifurcated into the left pulmonary artery (LPA) and the right pulmonary artery (RPA) but it had no visible continuity with the ductus arteriosus (DA). D: only the color Doppler examination showed the tortuous S shaped DA (arrow). E: the ductus joined the descending aorta at an obtuse angle. F: color aliasing occurred through the DA (arrow).
A diagnosis of S-shaped DA with mild narrowing was made and follow up examination was arranged. A repeat examination was performed at 27 weeks of gestation and showed nearly the same findings as before. The RV was normal in size and thickness and no TR was noted. Because the DA was S-shaped, it was very difficult to assess size along the whole range of the DA. However, the color Doppler examination showed tortuous turbulent flow in the ductus and suggested mild obstruction. The sagittal section of the aortic arch showed a ductal junction through which low velocity forward flow from the DA to the descending aorta was noted. Obstetric evaluation showed that fetal growth was appropriate and the amniotic fluid index was adequate. No signs of hydrops were noted.

The third examination was performed at 33 weeks of gestation. The right atrium was big and the RV was hypertrophied and globular in shape. The intracavitary volume of the RV looked slightly smaller than normal because of the thick ventricular wall and hypertrophy of the trabeculations. The right ventricular endocardial lining and the chordae of the tricuspid valve were very bright (Fig. 2A). Color Doppler examination showed moderate TR and its velocity was 4.5 m/sec (Fig. 2B and C). Every attempt failed to demonstrate any flow through the ductus. In spite of the right ventricular overload, fetal growth was within normal range and there was no evidence of fetal hydrops. After a discussion, we decided to deliver the baby. A female infant weighing 2625 g was born at 34 weeks gestation by induced delivery. Apgar scores were 7 at 1 minute and 8 at 5 minutes. Symptoms and signs immediately after birth were tachypnea, chest retraction, grunting sound and low oxygen saturation. An echocardiography performed at 20 minutes after birth showed a small hypertrophied RV and increased echogenicity of the moderator band and chordae of tricuspid valve (Fig. 3A). There was a moderate TR with a velocity of 3.7 m/sec (Fig. 3B and C). The short axis view showed a hypertrophied RV (Fig. 3D). The ductus was searched for but no ductus was found in 2-D and color images (Fig. 3E). Shunt flow through the foramen ovale was mainly right to left (Fig. 3F). She required respiratory support with a mechanical ventilator for the first 3 days. She was successfully weaned from the ventilator on her fourth day and discharged at 12 days old. Follow-up examination at the outpatient clinic showed that she did not have any cardiac symptoms or signs. Echocardiography taken at 2 months of age showed much improvement. The RV was nearly normal in size and thickness. Increased echogenicity of the RV was also nearly normalized. There remained TR with a velocity of 3.0 m/sec. At 7 months of age, all echocardiographic findings returned to normal with resolution of TR.

Discussion

Since the introduction of fetal echocardiography, various abnormalities of the DA have been described. Among ductal diseases in the fetus, ductal constriction or premature closure is one of the most important diseases because it is associated with significant morbidity or mortality. Traditionally, ductal constriction is defined as a systolic velocity higher than 1.4 m/s and a diastolic velocity higher than 0.3 m/s in conjunction with a pulsatility index smaller than 1.9. This criterion is based on clinical observations in which normal fetuses were compared to a ductal constriction group and a terbutaline group. In their study, the number of normal fetuses was only 41 and Doppler waveforms were obtained by the continuous Doppler technique. Mielke and Benda measured ductal velocity by pulsed Doppler technique in normal fetuses and showed that the 95 percentile of systolic velocity was well above 1.4 m/s and a diastolic velocity higher than 0.3 m/s in conjunction with a pulsatility index smaller than 1.9. This criterion is based on clinical observations in which normal fetuses were compared to a ductal constriction group and a terbutaline group. In their study, the number of normal fetuses was only 41 and Doppler waveforms were obtained by the continuous Doppler technique. Mielke and Benda measured ductal velocity by pulsed Doppler technique in normal fetuses and showed that the 95 percentile of systolic velocity was well above 1.4 m/s in later gestation. Ductal constriction literally means that the ductal luminal diameter is smaller than normal. However, ductal constriction is usually defined by Doppler velocity due to difficulties of measuring the luminal diameter. Measurement of the dimensions is more dif-
Doppler velocity is not without fault. Not to mention the technical difficulties of obtaining a Doppler spectrum from fetal small vessels, Doppler velocity is also flow dependent. The pulsatility index might be useful to differentiate ductal constriction from increased flow. However, in cases with decreased flow, ductal velocity could be normal in the presence of severe constriction.

In order to assess ductal constriction in a more comprehensible way, the estimated pressure of the RV, flow through the right side of heart and the response of the RV to pressure overload should be added. Hemodynamically significant ductal constriction makes the systemic and pulmonary circulation independent and right ventricular pressure becomes higher than left ventricular pressure when right sided flow is maintained. However, right ventricular pressure may not be high if flow is decreased. Right ventricular pressure and flow through the main pulmonary artery can be measured by the Doppler technique. Diameter and flow of the main pulmonary artery can be compared to those of the aorta. The fetal RV seems to respond to pressure overload in two ways, one is a strain pattern and the other is a failure pattern. In RV strain, the RV is hypertrophied and the endocardium becomes echogenic. The right ventricular cavity is normal or slightly decreased. This case corresponds to this category. Till the 2nd trimester, we could not find any evidence of RV dysfunction. However, in the 3rd trimester and the immediate postnatal period, we deduced RV diastolic dysfunction from the echocardiographic images. In contrast, the RV failure pattern shows a dilated and poorly contracting ventricle. Some of the failure group might have increased echogenicity. Premature closure of the ductus is diagnosed when there is no luminal patency and no flow through the ductus. Premature closure should be differentiated from absence of the ductus. Ductal absence is thought to be an early developmental anomaly and is associated with certain cardiac anomalies such as absent pulmonary valve syndrome and truncus arteriosus.

Fetal ductal constriction or premature closure is mostly associated with the use of cyclooxygenase-inhibiting drugs. Recently, other drugs and special foods have also been associated with ductal constriction. However, the idiopathic cases of fetal ductal constriction are few. Our case was diagnosed to have idiopathic severe ductal constriction or premature closure at 33 weeks and it was associated with suprasystemic RV pressure and RV strain. However,
| Authors                     | Number of patients | GA at diagnosis | Ductal velocity systole/diastole (Pulsatility index) | RV response | TR Degree/velocity (m/s) | Fetal symptoms                  | Course                      | Postnatal outcome |
|---------------------------|--------------------|-----------------|------------------------------------------------------|-------------|--------------------------|---------------------------------|-------------------------------|-------------------|
| Mielke et al.             | 1                  | 32              | 1.49/0.26 (2.62)                                     | Failure     | Severe/3.2               | No hydrops or fetal distress    | Progression of TR at 35 weeks | Alive, well       |
| Luchese et al.            | 12                 |                 |                                                      |             |                          |                                 |                               |                   |
|                           |                    | 33              | 1.8/1.51                                             | Dilated and hypertrophic RV | No           | Oligohydramnios/LUGR       | PH at birth                  |                   |
|                           |                    | 35              | 2.19/0.72 (0.88)                                     | Failure     | Moderate                 | Gemellary/hydrocephali          | Severe IT at 37 weeks          | PH at birth       |
|                           |                    | 37              | 2.43/0.69 (1.38)                                     | Dilated RV  | Minimum                  |                                 | Normal at birth               |                   |
|                           |                    | 34              | 2/1.77                                               | Mildly dilated RV | Minimum      |                                 | Improved at 35 weeks          |                   |
|                           |                    | 31              | 1.87 (1.83)                                          | Normal      | No                       |                                 | Normal at 37 weeks            |                   |
|                           |                    | 34              | 2.69/0.68 (1.73)                                     | Dilated RV  | No                       |                                 | Normal at 37 weeks            |                   |
|                           |                    | 32              | 2.12/0.61 (1.38)                                     | Dilated and hypercontractile RV | No         |                                 | Normal at 36 weeks            |                   |
|                           |                    | 28              | 2.14/0.66 (1.15)                                     | Dilated RV  | Moderate                 | Severe hyrops                   | Severe hydrops                 | Neonatal death    |
|                           |                    | 38              | 2.64/0.96 (1.47)                                     | Normal      | No                       |                                 | Good neonate evolvement        |                   |
| Trevett and Cotton        | 1                  | 33              | 2.5/0.8 (1.1)                                         | Normal      | Moderate                 |                                 | Normal at 34 weeks            |                   |
|                           |                    | 33              | 2.36/0.5 (1.54)                                      | Dilated and hypertrophied RV | No        | Fetal arrhythmia           |                                | Good neonate evolvement    |                   |
| Soslow et al.             | 1                  | 33              | 1.66/0.45 (1.55)                                     | Normal      | No                       |                                 | Normal at 34 weeks            |                   |
| Shima et al.              | 1                  | 31              | Continuous flow                                       | Moderately dilated and hypertrophied RV | ND/4.1     | Small pericardial effusion | ↑ TR velocity at 32 weeks      | Alive, well       |
| Enzensberger et al.       | 3                  | 32              | 305/150 (0.7)                                         | Hypercontractile RV | TR/        | Small pericardial effusion | Immediate delivery            | Alive, well       |
|                           |                    | 33              | 219/110 (0.7)                                         | Dilated RV, right heart volume load | ND/4.5     | No hydrops, normal venous flow pattern | No change till 37 weeks       | Alive, well       |
|                           |                    | 36              | 360                                                   | Right heart hypertrophy | Positive | No hydrops                  | Immediate delivery            | Alive, well       |
| Current case              | 1                  | 33              | Not measured                                          | Strain      | Moderate/4.5             | No hydrops                      | No progression at 34 weeks    | Alive, well       |

ND: no data available, RV: right ventricle, TR: tricuspid regurgitation, GA: gestational age
Table 2. Cases with idiopathic premature closure of the ductus arteriosus

| Authors          | Number of patients | GA at Dx | RV response | TR Degree/velocity (m/s) | Flow | Fetal symptoms | Course | Postnatal outcome |
|------------------|--------------------|----------|-------------|--------------------------|------|----------------|--------|-------------------|
| Chao et al.      | 1                  | 31       | Normal      | Mild                     | Decreased right side flow | Nonstress test: nonreactive, no hydrops | Immediate delivery | Alive, well       |
| Hofstadler et al.| 4                  | 37       | Failure     | ND/5.25                  | No hydrops                  | Labor induction          | Alive, well       |
|                  | 37 Failure         | ND/3.3   | No hydrops, abnormal UV pulsation | Labor induction | Alive, well       |
|                  | 38 Failure         | ND/3.25  | Aseites, abnormal UV pulsation | Immediate CS delivery | Alive, well       |
|                  | 33.5 Failure       | ND/2.9   | No hydrops  | Progression at 34.5 weeks | Alive, well       |
| Leal et al.      | 3 (5)              |          | Mild to moderate | Absent flow in the DA | Polyhydramnios, no fetal hydrops, abnormal IV and ductus venosus flow | No further progression during the next few days | Alive and well       |
| Mielke et al.    | 1                  | 38       | RV made hypertrophy, dilated, poor contraction | No | TV, MPA flow ↓ | Polyhydramnios, no fetal hydrops, abnormal IV and ductus venosus flow | No further progression during the next few days | Alive and well       |
| Luchese et al.   | 1 (20)             | 36       | Dilated and hypertrophic RV | Severe | Occluded DA | Good neonate evolvement |        |
| Nygaard et al.   | 1 (3)              | 41       | Failure with strain | + | Absent flow in the DA | Polyhydramnios, no fetal hydrops, abnormal IV and ductus venosus flow | No further progression during the next few days | Alive and well       |
| Gewillig et al.  | 5 (8)              | 37       | RV dysfunction | Moderate to severe | No | Spontaneous delivery at 38 weeks | Alive |        |
|                  | 27                 | RVH      | Moderate to severe | Hydrops | Premature induced delivery at 30 weeks | Alive with cardiomyopathy |        |
|                  | 32                 | Dilated and hypertrophic RV | Moderate to severe | Progressive TR, premature induced delivery at 34 weeks | Alive |        |
|                  | 20                 | Dilated and hypertrophic RV | Moderate to severe | Cystic lung | Spontaneous delivery at 39 weeks | Died due to respiratory failure? |        |
|                  | 28                 | RV hypertrophy | Moderate to severe | No | Induced delivery at 37 weeks | Homograft at pulmonary valve position at 30 months |        |
| Wei et al.       | 1                  | 38       | RV hypertrophy, RV failure | Severe/5.1 | Abnormal ductus venosus flow | No | Induced delivery | Alive and well       |
| Ishida et al.    | 1 (3)              | 32       | Failure     | Mild TR/3.1             | Hydrops                  | Immediate delivery       | Alive and well       |

UV: umbilical vein, MC: inferior vena cava, CS: cesarean section, MPA: main pulmonary artery, RVH: right ventricular hypertrophy, GA: gestational age, TR: tricuspid regurgitation, ND: no data available, RV: right ventricle
previous two examinations at 23 and 27 weeks gestation showed only an S-shaped ductus with possible mild narrowing but without any right ventricular abnormalities. Our case is very similar to a case reported by Trevett and Cotton.6 That case also showed an S-shaped ductus at 27 weeks without any cardiac abnormalities but severe ductal constriction with RV strain developed at 33 weeks. These two cases certainly prove that DA may develop progressive dense calcification in the 3rd trimester.

In order to review idiopathic ductal constriction or premature closure, a literature search was performed in PubMed and PubMed Central. In some reports, both idiopathic cases and drug-related constriction and premature closure were included.6 All idiopathic cases were selected and were divided into a constriction group or a premature closure group and tabulated in Table 1 and 2.

There have been 20 cases of idiopathic ductal constriction including our case (Table 1). Luchese et al.6 reported 13 cases of idiopathic ductal constriction, and used traditional diagnostic criteria for inclusion. Their cases seemed to include ductal constriction of variable degree, from mild to severe, and from asymptomatic to symptomatic. However all others are reports of symptomatic cases. The diagnosis of symptomatic ductal constriction was usually made between 28 and 38 weeks, and mostly after 31 weeks. Significant TR was a commonly associated finding. The RV response was either failure or a strain pattern. Among 30 cases of idiopathic premature closure of the DA (Table 2),12-20 two cases in Gewilling’s report20 seemed to have ductal absence rather than premature closure. The diagnosis of premature closure is usually made between 28-41 weeks. Fetal symptoms, degree of TR, and the right ventricular response in cases with premature closure seem to be similar to those with severe symptomatic ductal constriction. However, the velocity of TR was more variable and ranged from 2.9 to 5.25 m/sec, probably depending on the flow through the right heart and contractility of the RV, and degree of TR.

With regard to management, fetuses with right ventricular failure and other signs of fetal distress such as hydrops are usually managed by immediate delivery if the fetuses are not too premature. However, if the fetuses are too premature, a decision on delivery should balance fetal risk and post-natal risk. The fetus without fetal distress can be followed under careful observation. Pulmonary hypertension and right ventricular failure are major concerns after delivery, but the post-natal prognosis seems to be good. Severely affected babies require supplemental oxygen or mechanical ventilation. Pulmonary hypertension can be managed with NO. Right heart abnormalities seem to resolve within a couple of weeks, but right ventricular wall thickness and diastolic dysfunction take several months to recover.

In summary, we report a case in which a fetal S-shaped ductus with mild narrowing at 23 and 27 weeks gestation showed progressive ductal constriction or premature closure at 33 weeks, and emphasize the importance of follow up well into the 3rd trimester. It is also suggested that the diagnosis of ductal constriction should be based on not only the ductal diameter, flow velocity, and pulsatility index, but also on right ventricular pressure, flow through right side of the heart and the right ventricular response to pressure overload.

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