Right heart dilatation in a fetus with an abnormal foramen ovale valve: an indicator of interatrial communication restriction

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Abstract

Aims: Foramen ovale (FO) valve with a shape or motion abnormality is frequently detected during routine obstetric ultrasonic examinations. However, the hemodynamics mechanism of this entity remains unclear. The purpose of the study is to determine the relevance of interatrial communication restriction and resultant morphological modifications. Materials and methods: We reviewed the echocardiographic records of fetuses with isolated abnormal FO valve evaluated between January of 2010 and December of 2016. The size (DFO) of the FO orifice, opening angle (α) of the FO valve, and dimensions of cardiac chambers, FO channel outlet (DOUT) and inferior vena cava (DIVC) were measured. We evaluated their (DFO, DOUT, α) relationships to the diameters of RA and DIVC. Five hundred and seventy normal fetuses were selected to establish the normal range of the DOUT/DIVC ratio so as to provide a criterion for restriction. Results: An abnormal FO valve was identified in 89 fetuses without congenital heart disease, with restriction noted in 62 fetuses (45 fetuses with RA dilatation, 12 fetuses with RA and RV dilatation, and 5 fetuses with no RA dilatation). There were no significant correlations between RA/LA and DFO/DIVC, RA/LA and α. RA/LA was negatively correlated with DOUT/DIVC (R²=0.97, p<0.01). Conclusions: For a fetus with an abnormal FO valve, right heart dilatation could be considered as an indicator of interatrial communication restriction, which could be assessed by evaluating the FO channel outlet. The degree of right atrium dilatation indicates the severity of the restriction. Keywords: fetal echocardiography; right atrial dilatation; restriction; foramen ovale valve; hemodynamics

Introduction

The atrial septum is an important intra-cardiac structure formed by the septum secundum of the right atrial side and the septum primum of the left atrial side [1]. The foramen ovale (FO) is formed on the septum secundum and must remain open during intrauterine life to maintain normal fetal circulation. The entrance of FO into the left atrium (LA) is covered by the FO valve, which is a vesti-
ten et al showed that the outlet portion of FO channel represents the restricting structure during normal intrauterine development [10].

In the current study, we aimed to determine if an abnormal FO valve has relevance to the interatrial communication restriction. We also analyzed the hemodynamics of those fetuses with abnormal FO valve for evaluating the factors responsible for the severity of interatrial communication restriction.

Materials and methods

Study Population

We retrospectively reviewed the imaging data obtained from the fetal echocardiography center in our hospital from January of 2010 to December of 2016. The subjects comprised a normal group and a case group. During the study period, the echocardiographic records of normal fetuses diagnosed in our hospital were reviewed. Those without high-quality images of the sagittal views for measuring the diameter of the FO channel outlet were excluded. Fetuses of the gravida with smoking, diabetes, hypertension, or any general chronic disease were also excluded. All fetuses were singletons. The case group was selected from cases in which the FO valve had a shape or motion abnormality. The FO valve was diagnosed as a shape abnormality when it appeared to be either aneurysmal or hypermobile as previously described [11-13]; or motion abnormality, when it appeared to be thick and flat without a “flapping” motion normally seen in the uterus [14,15]. The cases diagnosed with congenital heart diseases or with no clear images were excluded. Outcome data was obtained from postnatal echocardiograms.

All fetuses were examined using the Voluson 730, E8, and E10, GE Healthcare, Kretztechnik, Zipf, Austria ultrasound systems. Routine obstetric sonography was initially performed to detect extra-cardiac malformations. Then, a detailed cardiac examination was performed by an experienced fetal echocardiographer (Z.Y.). The visceral and cardiac positional relationship was determined as previously described [16]. Four transverse views including the four-chamber view (4CV), the left and right outflow tract views, and the three-vessel-trachea view were scanned. In addition, three sagittal views including the bivacal view, the aortic arch view, and the ductal arch view were also obtained. High-definition flow imaging (HDFI) was performed when necessary. All imaging data was saved as video clips for later analysis.

Study design and fetal cardiac image interpretation

For the normal group, the distance (D_{N-OUT}) between the final point of the FO valve and the atrial septum and the dimension (D_{IVC}) of the inferior vena cava (IVC) were measured in the sagittal views. Data of D_{N-OUT}/D_{IVC} was plotted using a linear regression with 95% distribution range for individual observations. Diagnosis of interatrial communication restriction was made on the basis of the criterion that the ratio is less than the lower limit of 95% reference range.

According to our criterion, the case group was separated into two groups including a restriction group and a non-restriction group. Figure 1a shows the measurement of the diameters of cardiac chambers. As established by Tan J, et al [6], right atrium (RA) dilatation between 26 and 32 gestational weeks (GW) was defined as an estimated RA/LA >1.1 and defined as >1.2 at 32 GW and above. Right ventricle (RV) dilatation was defined as an estimated RV/LV >1.3 (normal ratio=1.18) [17]. Fetuses in the restriction group were assigned to three subgroups according to alterations of the cardiac chambers: isolated RA dilatation (Group A), RV and RA dilatation (Group B), and no RA or RV dilatation (Group C). The size (D_{FO}) of the FO orifice was measured in the 4CV. Besides, we also measured an angle (α) in the sagittal views, which could represent the shape or extent of the distension of FO valve in some degree. A sketch map shows this method of measuring α (fig 1b). The maximal value of the FO channel outlet (D_{OUT}) and D_{IVC} were obtained in the sagittal views. The relationships of RA/LA (and RV/LV)
to $D_{FO}/D_{IVC}$, $RA/LA$ (and $RV/LV$) to $a$, and $RA/LA$ (and $RV/LV$) to $D_{OUT}/D_{IVC}$ were analyzed. HDFI technique was used to detect the full course of the IVC draining into the LA through the outlet.

In addition, the mean $a$, the mean $D_{FO}/D_{IVC}$ ratio and the mean $D_{OUT}/D_{IVC}$ ratio were calculated and compared among groups (Groups A, B, C, and non-restriction group). Data of $D_{OUT}/D_{IVC}$ for fetuses with RA dilatation (Groups A and Group B) was also represented graphically. The data was plotted against GW and compared with that of normal group.

Statistical Analysis
Data was expressed as mean±standard deviation (SD). Data was analyzed using Prism (GraphPad, La Jolla, CA). A simple linear regression analysis was used to evaluate relationships between the variables of interest. Student’s t-test was used to compare mean values. The chi-squared test was used to assess statistical relationships between categorical data. Statistical relationships were considered significant at $p<0.05$.

Results
Abnormal FO valve characteristics
There were two predominant types of abnormal FO valve demonstrated in our study. A type of motion abnormality, which was thick and did not have the typical swinging motion during the cardiac cycle, was documented in 23 (25.8%) fetuses. Two video clips illustrate this in greater detail (video 1 and video 2, on the journal site). The FO channel outlet was the orifice formed by the single fixed FO valve and the septum secundum, which could be observed in the 4CV and the sagittal views (fig 2a,b). Fig 2c shows only a small amount of flow entering the LA through a narrow outlet in a sagittal view by HDFI. An additional video clip illustrates this in greater detail (video 3, on the journal site). The structure of the outlet is best illustrated by the sketch map (fig 2d). The remaining 66 (74.2%) fetuses demonstrated the typical presentations of atrial septal aneurysm, which was formed by two hypermobile and thin valves and appeared to balloon into the LA (fig 3a,b). An additional video clip illustrates this in greater detail (video 4, on the journal site). The FO channel outlet was formed by the final portions of two FO valves, which adhered to the septum secundum. An additional video clip illustrates this in greater detail (video 5, on the journal site). A sketch map illustrates the structure of this type of outlet (fig 3c).

Distribution of all groups and outcome of statistical analysis
In figure 4 the subjects, distribution of all groups, and statistical analysis described in the study design are
shown. During the study period, 1637 fetuses underwent cardiac evaluation. No structural heart defect was found in 960 fetuses and 570 fetuses were selected for the normal group. Age in GW ranged from 21 to 39 (median, 30.0). Maternal age ranged from 20 to 33 years old (median, 28.0). Abnormal FO valves were found in 119 fetuses resulting in a prevalence of 12.4% among this highly selected cohort; 89 fetuses were selected into case group.

In Table I are detailed the number the fetuses selected in the five groups and the data of $\alpha$, $D_{FO}/DIVC$, and $D_{OUT}/DIVC$ obtained in these groups. There were no remarkable differences in mean $\alpha$ and mean $D_{FO}/DIVC$ ratio among Groups A, B, C and non-restriction group. However, the mean $D_{OUT}/DIVC$ ratio in Group B was lower than that observed in Group A and Group C ($p<0.01$, respectively). Also, the mean $D_{OUT}/DIVC$ ratio in Groups A, B, C was lower than that of non-restriction group ($p<0.01$, respectively).

For fetuses with RA dilatation (Groups A and Group B), there were no significant correlations between RA/LA and $D_{FO}/DIVC$ ($R^2=0.03$, $p=0.18$) or RA/LA and $\alpha$ ($R^2=0.01$, $p=0.57$). However, RA/LA was negatively correlated with $D_{OUT}/DIVC$ ($R^2=0.97$, $p<0.01$). Notably, for fetuses with RV dilatation (Group B), the RV/LV ratio had no significant correlation with $D_{FO}/DIVC$ ($R^2=0.003$), $\alpha$ ($R^2=0.02$), or $D_{OUT}/DIVC$ ($R^2<0.01$). This analysis was limited by the number of samples.

Linear regressions between $D_{OUT}/DIVC$ ($D_{OUT}/DIVC$) and GW were performed. The lower limit of 95% reference range remained at 0.5-0.6 during second pregnancy. All cases with RA dilatation (Group A and Group B) were under the lower limit of 95% reference range.

Follow-up and prognosis

For fetuses in the case group, fetal echocardiogram was performed on a regular basis at least every week for potential existence of restriction and later evolutive signs. For the non-restriction group, GW at first diagnosis ranged from 26 to 39 weeks (median, 32.7). There was no RA dilatation or other signs of restriction during the follow-up period. For the restriction group, GW at first diagnosis ranged from 26 to 36 weeks (median, 30.4). Five cases that were lost to follow-up were excluded. There were 40 fetuses with stable hemodynamic state and 17 cases with evolutive signs of restriction (Table II).

In our study, the prognosis of all fetuses with abnormal FO valve was good. Fifty-two of 57 fetuses with RA dilatation showed a normal RA within the first 2 weeks of life, as determined by the postnatal echocardiogram.

<table>
<thead>
<tr>
<th>Group</th>
<th>n</th>
<th>$\alpha$ (°)</th>
<th>$D_{FO}$/DIVC</th>
<th>$D_{OUT(N-OUT)}/DIVC$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Restriction group</td>
<td>45</td>
<td>56.56±16.40</td>
<td>1.12±0.15</td>
<td>0.45±0.08†</td>
</tr>
<tr>
<td></td>
<td>12</td>
<td>59.92±16.28</td>
<td>1.01±0.14</td>
<td>0.32±0.14†</td>
</tr>
<tr>
<td></td>
<td>5</td>
<td>52.40±3.85</td>
<td>1.11±0.15</td>
<td>0.51±0.06†</td>
</tr>
<tr>
<td>Non-restriction group</td>
<td>27</td>
<td>55.48±9.21</td>
<td>1.02±0.11</td>
<td>1.02±0.19†</td>
</tr>
<tr>
<td>Normal group</td>
<td>570</td>
<td>NA</td>
<td>NA</td>
<td>0.95±0.24</td>
</tr>
<tr>
<td>p value</td>
<td></td>
<td>NA</td>
<td>NA</td>
<td>S†</td>
</tr>
</tbody>
</table>

Group A, fetuses with isolated RA dilatation; Group B, fetuses with RA and RV dilatation; Group C, fetuses with no RA dilatation; $\alpha$, an angle indicating the opening range of the FO valve; $DOUT$, the maximal value of the FO channel outlet in case group; FO, foramen ovale; LA, left atrium; RA, right atrium. 

Fig 4. Subjects, group distribution and statistical analysis in our study. $\alpha$, an angle indicating the degree of distention of the FO valve; $DFO$, the size of the FO orifice; DIVC, the dimension of the inferior vena cava; DN-OUT, the distance between the final point of the FO valve and the atrial septum in normal group; DOUT, the maximal value of the FO channel outlet in case group; FO, foramen ovale; LA, left atrium; RA, right atrium.
Another 5 neonates showed an enlarged RA with varying degrees of TR. Table III show these results in detail.

**Discussions**

FO valve with shape or motion abnormality is frequently detected during routine obstetric ultrasonic examinations as it is easy to be visualized in the 4CV. However, there remains some confusion whether abnormal FO valve belongs to normal variations or may affect fetal cardiac hemodynamic. The results of the current study indicated that interatrial communication restriction may exist in parts of these fetuses with resultant right heart dilatation. We also demonstrated that a restriction is not present when the cardiac chamber is within the normal range during the late pregnancies. Accurate evaluation of the FO channel outlet is the key factor to decide whether a restriction exists when abnormal FO valve are present. In fact, restrictive FO has long been noticed by fetal echocardiographers. Previous reports mainly focused on the dimension of the FO orifice [7-9]. However, these diagnosis standards of restrictive FO are obviously not satisfactory as they could not show the full hemodynamics of interatrial communication. The measurement of blood velocity across FO was a development. However, accuracy might be affected as Doppler shift is angle-dependent. Recently, Patten et al [10] proposed the conception of the FO channel and indicated that the outlet portion of this channel could make the assessment of the interatrial communication function more complete. On this basis, Kiserud et al [18] established the normal range of the area ratio between the FO channel outlet and IVC. In our study, we measured the diameter of the FO channel outlet. A relatively large number of normal fetuses were included to establish the reference range. The results showed a high consistency with that of Kiserud et al. As
the diameter of both the FO channel outlet and IVC could be easily measured when compared with measuring the area, the obstetric sonographers and fetal echocardiographers are apt to use our newly proposed standard for evaluating fetal interatrial communication.

As is well known, the physiologic interatrial shunt enables the oxygenated blood entering the right heart to be ejected to the left heart, which is important to ensure a stable hemodynamics state between the left heart and the right heart [19]. Intrauterine restriction of interatrial communication might cause the increased blood flow into the right heart and decreased flow into the left heart with resultant morphologic abnormalities, such as right heart dilatation [20]. Theoretically, the chambers may not dilate when only mild restriction is present. However, RA will dilate, and then with the presence of the enlargement of RV, when the restriction is moderate or severe. The restriction could only be diagnosed when the FO channel outlet was accurate-ly evaluated. We believe that the sagittal views are best suited for evaluating the FO channel outlet, particularly since it is based on the actual flow direction in the atrial area as evidenced by prior studies [18,21,22]. Besides, according to our experience, it is impossible to identify the FO channel outlet in the 4CV in most cases.

However, it is a challenge for an obstetric sonographer to scan qualified sagittal views and to evaluate the FO channel during routine examinations. Instead, the alterations of cardiac chambers could be easily identified. According to our results, the majority fetuses with abnormal FO valve and interatrial communication restriction showed dilated RA, or combined with RV dilatation. On the contrary, fetuses with abnormal FO valve while with no restriction showed no alterations of cardiac chambers. This suggests that RA/LA could be used to predict the existence of restriction, when abnormal FO valve presents.

It was worth noting that the RA volume overload, RA pressure overload, or ventricular disproportion may develop as a progression of earlier restriction. In the current study, 5 fetuses enrolled in the second trimester evidenced symmetrical left and right chambers. However, RA dilatation appeared in late pregnancy. We speculate that the normal cardiac dimension at the initial diagnosis resembles the compensatory ability of the right heart during the early stage of restriction.

In fact, a close follow-up is necessary when evidence shows interatrial communication restriction, whether RA dilatation exists or not. The examiner should be concerned with the appearance of RA dilatation, or progressive right heart dilatation with fetal hydrops, and tricuspid valve insufficiency [4,5,12]. In addition, the prognosis of interatrial communication restriction also depends at the point of time when RA dilatation appears and the severity of restriction [4]. The earlier and the more severe RA dilatation appears, the easier progressive restriction may present. However, obstetricians should not be too pessimistic about the prognosis of interatrial communication restriction. Hagen et al. reported that, as long as the fetus remained hemodynamically stable, the prognosis of an interatrial communication restriction was good and the pregnancy should be allowed to continue until term with close monitoring [1]. In the current study, all fetuses with restriction had a good result. Various degrees of RA dilatation, or combined with RV enlargement presented, resembled right heart volume overload. During the follow-up period, most cases maintained a stable hemodynamic state. Signs of progressive right heart volume overload presented in only 17 fetuses, 7 of which delivered at full term with close monitoring, while the rest delivered prematurely with adequate gestational age and lung maturity. For intrauterine interatrial communication restriction, obstetricians and sonographers should identify the abnormality as early as possible. We believe that a close monitoring ensures proper strategies for the management of these fetuses, and in turn, guarantees a good prognosis.

The main limitation of this study is that it was a mono-center respective review. Furthermore, fetuses without high-quality sagittal views were excluded from the study which may cause potential selective bias.

Conclusions

We proposed an easy method with reference range herein to identify intrauterine restriction of interatrial communication. For fetuses with abnormal shape or motion of FO valve, the dilatation of the right heart could be considered as an indicator of interatrial communication restriction. The larger RA/LA ratio indicates a more severe restriction. Evaluation of the FO channel outlet in the sagittal views guarantees an accurate assessment of the restriction.
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Conflict of interest: none

References

Additional Files

Video 1. 2D imaging showing the 4CV of a fetus with a single fixed FO valve at 33 weeks gestation. A narrow gap (indicated by an icon) formed by a single fixed FO valve and the septum secundum is visualized. FO, foramen ovale; LV, left ventricle; RA, right atrium; RV, right ventricle.

Video 2. 2D imaging showing a sagittal view of the same fetus as in Video 1. A clear outlet (indicated by an icon) formed by a single fixed FO valve and the septum secundum is visualized. The FO valve did not have the typical swinging motion. FO, foramen ovale; IVC, inferior vena cava; RA, right atrium.

Video 3. Simultaneous imaging of gray-scale and high-definition flow imaging showing a sagittal view of the same fetus as in Video Files 1 & 2. By high-definition flow imaging, only a small amount of flow (indicated by an arrow) could be identified entering into the LA through the narrow outlet formed by a single fixed FO valve and the septum secundum. FO, foramen ovale; LA, left atrium; RA, right atrium.

Video 4. 2D imaging showing the 4CV of a fetus with an atrial septal aneurysm at 30 weeks gestation. Two FO valves
with an indistinct gap (indicated by an arrow) at their final portions are visualized. AO, aorta; FO, foramen ovale; LV, left ventricle; RA, right atrium; RV, right ventricle.

**Video 5.** 2D imaging showing a sagittal view of the same fetus as in Video File 4. A clear outlet (indicated by an icon) formed by the ends of two FO valves is visualized. AO, aorta; FO, foramen ovale; IVC, inferior vena cava; RA, right atrium.