Case Report

Prenatal Diagnosis of Cardiac Diverticulum with Pericardial Effusion in the First Trimester of Pregnancy with Resolution after Early Pericardiocentesis

Raquel Garcia Rodriguez, Azahara Rodriguez Guedes, Raquel Garcia Delgado, Lourdes Roldan Gutierrez, Margarita Medina Castellano, and Jose Angel Garcia Hernandez

Prenatal Diagnosis and Fetal Therapy Unit, Department of Obstetrics and Gynecology, University Maternity Hospital of Canaries, Las Palmas, 35004 Gran Canaria, Spain

Correspondence should be addressed to Raquel Garcia Rodriguez; raquelgarciaod@gmail.com

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Cardiac diverticulum is a rare anomaly, which may present in association with pericardial effusion. Only a few cases diagnosed during fetal life have been published and only in 12 cases pericardiocentesis was made with good postnatal outcomes in 83% of the cases. In the first trimester of pregnancy only 6 cases were reported. We described the largest series of cases published. We describe a case of cardiac diverticulum complicated with pericardial effusion, which was diagnosed at 14 weeks of pregnancy and resolved by intrauterine pericardiocentesis at 17 weeks of pregnancy. We made a systematic review of the literature with the cases reported of cardiac diverticulum, management, and outcomes.

1. Introduction

Cardiac diverticulum is a rare anomaly, which may present in association with pericardial effusion and may produce complications such as pulmonary hypoplasia or hydrops fetalis. Only a few cases diagnosed during fetal life have been published and 10 cases during the first trimester of pregnancy.

We describe a case of cardiac diverticulum complicated with pericardial effusion, which was diagnosed at 14 weeks of pregnancy and resolved by intrauterine pericardiocentesis at 17 weeks of pregnancy.

We review the ultrasound findings, management, and outcomes in 34 cases published in the literature, 21 of which were associated with pericardial effusion, and in 12 cases a pericardiocentesis was performed. In this group of pericardiocentesis, 10 cases have good postnatal outcomes.

In light of the good neonatal outcome of this procedure, we propose that early pericardiocentesis should be considered as a therapeutic option in these cases that do not resolve spontaneously.

2. Case Report

A 22-year-old sub-Saharan woman, gravida 2 para 1, was referred to the Unit of Prenatal Diagnosis and Fetal Therapy at 12 + 4 weeks of gestation for a fetal hydrothorax. The ultrasonographic study revealed the presence of severe pericardial effusion. The first trimester screening showed 2.9 mm nuchal translucency, reversed ductus venosus a-wave, absent nasal bone, and absence of tricuspid regurgitation. All other anatomical and echocardiographic findings were normal. Biochemical markers were β-hCG 0.14 MoM and PAPP-A 0.72 MoM. The screening results indicated high risk of chromosomopathy. Therefore, a chorionic villus sampling biopsy study was performed, which revealed a normal 46 XY karyotype. On week 14 of pregnancy, pericardial effusion was still observable. Furthermore, a two-millimeter anechoic image was found on the heart apex, with a narrow base and expanding from the ventricular wall into the pericardium. Blood flow could be observed through this structure with color Doppler
sonography. On this basis, the diagnosis of a cardiac diverticulum was made (Figure 1). The rest of the echocardiography was normal.

Serial ultrasound studies were conducted until week 17 of pregnancy (Figure 2). Given that the pericardial effusion persisted, pericardiocentesis with a 20-gauge needle (like amniocentesis) was performed and a clear serum-like fluid was removed. After pericardiocentesis, both lungs expanded and only a small amount of residual fluid remained. Results of the infections study of the amniotic fluid (Toxoplasma, cytomegalovirus, rubella, herpes virus, and parvovirus B19) were negative. A week after pericardiocentesis, no new pericardial effusion was detected. The clinical picture remained stable until week 25, when cardiomegaly and thickening of the ventricular walls were observed, thus leading to a diagnosis of hypertrophic myocardopathy. Ultrasound studies showed normal ventricular function without signs of cardiac failure. On week 40, a boy was born in eutocic delivery, with 3150-gram body weight, 9/10 Apgar score, and 7.29 umbilical artery pH. The newborn was admitted to hospital for cardiological study. Subsequent serial examinations ruled out hypertrophic myocardopathy, although they showed residual pericardial effusion, an apical cardiac diverticulum, persistent small ductus venosus, and permeable oval foramen. At present, the child is four years old, remains asymptomatic, and is on a prophylactic treatment with acetylsalicylic acid. He is being monitored in annual follow-up visits to the Service of Pediatric Cardiology.
Table I: Differential diagnosis of the different types of diverticulum and aneurism.

| Etiopathogeny           | Isolated apical diverticulum | Nonapical diverticulum | Aneurism |
|-------------------------|-----------------------------|------------------------|----------|
| Implantation base on   | Narrow base                 | Narrow base            | Wide base|
| ventriculum             |                             |                        |          |
| Size                    | Small                       | Small                  | Large    |
| Development during     | Constant size throughout   | Constant size throughout pregnancy | Enlarging with gestational age |
| pregnancy               | pregnancy                   |                        |          |
| Histology               | Myocardium in walls; usually presenting the three layers (myocardium, pericardium, and endocardium) | Myocardium in walls, usually presenting the three layers (myocardium, pericardium, and endocardium) | Myocardium disruption, usually presenting thin myocardium and fibrous tissue |
| Kinesis                 | Normal                      | Normal                 | Akinetic, hypokinetic, and dyskinetic |
| Contractility           | Contractility in synchrony with heart's rhythm | Contractility in synchrony with heart's rhythm | Contractility paradoxical with heart's rhythm |
| Complications           | Depending on the associated anomalies | Usually not occurring | Arrhythmia, thromboembolism, heart failure, and rupture |
| Prognosis               | Good                        | Good                   | Bad      |

3. Discussion

A cardiac diverticulum is a protrusion located on the ventricular wall, in close communication with it. This rare entity has been scarcely reported in the literature. Prenatal diagnosis may be difficult when it appears isolated [1]. Since such formations are closely communicated with the heart ventriculum, bidirectional blood flow may be observed through them with color or pulsed Doppler sonography [1].

Two types of diverticula are known: apical and nonapical [2, 3]; their characteristics are shown in Table I. Apical diverticulum of the left ventriculum may be of three different types: apical isolated diverticula, which are not associated with other malformations; multiple diverticula, which are located on the diaphragmatic or anterior surface of the ventricle; and large apical diverticula, which are associated with midline thoracoabdominal malformations or with the Pentalogy of Cantrell [4]. Although their etiology is not known, they seem to be caused by local weakening of the ventricular wall, which may in turn be due to possible embryogenesis defects [5], secondary to infections, or caused by local ischemia resulting from coronary anomalies, such as stenosis, hypoplasia, intimal proliferation, and thrombosis [6–8]. Differential diagnosis should include aneurisms (Table I) [6, 9–12], myocardioathy, Ebstein’s anomaly, and auriculoventricular regurgitation [9].

We made a systematic review of the literature with PubMed and Embase database search in English, French, or German which was performed without any restriction of publication date or journal, using the following key words: fetal cardiac diverticulum. The last search was updated in December 2012.

For each included case we focused on the following criteria: time of diagnosis, pericardial effusion association, management, intrauterine evolution, associated complications, and time of delivery fetal outcome (intrauterine death/ stillbirth, death after delivery, and survival). We excluded the cases of cardiac aneurysm. After reviewing 27 articles, 34 cases published between 1990 and 2012 met the above criteria in which cases of fetal cardiac diverticulum were presented (Table 2) with our case included.

It was most frequently diagnosed during the second trimester of pregnancy (51%). During the first trimester, 29% of cases were diagnosed; three of them presented increased nuchal translucency. This entity affects male fetuses more than female ones (3:1); its most frequent location is on the right ventriculum (70%) and the most frequently involved area is the heart apex (57%). Ultrasonographic findings associated with diverticula include pericardial effusion, cardiomegaly, septal defects and arrhythmia with fetal death before delivery, and hydrops [6, 13, 14]. Pericardial effusion is the most frequently associated finding (63%) and should be considered an indirect sign of the presence of cardiac diverticula. Although the etiology of effusion is not known, it has been proposed to result from the diverticulum rubbing the pericardial walls or from heart failure. Thus, the observation of pericardial effusion makes it necessary to examine the cardiac function [1, 6, 15].

A problem associated with pericardial effusion is that the resulting compression may produce heart failure and pulmonary hypoplasia. The management of such cases (Table 3) usually varies from performing pericardiocentesis to adopting an expectant approach. In a published series of 22 cases, pericardiocentesis was performed in 12 (56%) fetuses of 14 to 25 weeks’ gestational age, with an outcome of 10 cases that progressed favorably (83%) and 2 intrauterine deaths (17%) [16, 17]. In the cases of fetal death, the pericardial fluid was blood. In the same series, 7 cases were managed with an expectant approach (32%). Five of them showed spontaneous
Table 2: Description of the cases of cardiac diverticulum reported in the literature.

| Author                     | GA di | Size     | Sex  | Location                          | Karyotype | Associated anomalies | Intervention | Prenatal progression | Neonatal                  | Follow-up                           |
|----------------------------|-------|----------|------|-----------------------------------|-----------|----------------------|--------------|-----------------------|-------------------------------|-------------------------------------|
| 1  Kitchiner et al. (1990) [13] | 33    | —        | Female | Apex VI                           | —         | cardiomegaly         | No           | Stable                | Vaginal delivery 40 w; cardiomegaly, tachypnea, heart murmur; muscular IVC, and mild mitral regurgitation | Asymptomatic at 3.5 months of life |
| 2  Hornberger et al. (1994) [9] | 31    | —        | —     | Lateral wall below tricuspid valve (RV) | —         | —                    | No           | Stable                | Vaginal delivery at term       | Asymptomatic at 12 months of life |
| 3  Carles et al. (1995) [24]  | 13    | —        | Male  | Apex LV                           | —         | pericardial effusion | TOP 14 w    | —                     | —                            | —                                   |
| 4  Cesko et al. (1998) [25]  | 17    | —        | Male  | Apex RV                           | 46XY      | pericardial effusion | TOP 22 w    | Stable                | —                            | —                                   |
| 5  Cavallé-Garrido et al. (1997) [6] | 20    | Large    | Female | Lateral wall below mitral valve (LV) | Trisomy 18 | ventricular septal defect, hydrops | No          | Fetal death 26 w     | —                            | —                                   |
| 6  Cavallé-Garrido et al. (1997) [6] | 19    | Small    | Female | Apex RV                           | —         | No                   | No          | Stable spontaneous resolution at 34 w | Asymptomatic                  | Asymptomatic at 22 months of life |
| 7  Cavallé-Garrido et al. (1997) [6] | 19    | Small    | —     | Apex RV                           | —         | pericardial effusion | PC 20 w     | Stable                | Asymptomatic                  | Asymptomatic at 12 months of life |
| 8  Cavallé-Garrido et al. (1997) [6] | 36    | Small    | Male  | Lateral wall below tricuspid valve (RV) | —         | pericardial effusion | —           | —                    | Asymptomatic at 18 months of life | —                                   |
| 9  Johnson et al. (1996) [16] | 19    | 3 mm     | Female | Apex RV                           | 46XX      | pericardial effusion | PC 20 w     | No relapse after PC, no growth | Eutocic delivery 41 w; weight 3700 grams; asymptomatic | Asymptomatic at 16 months of life |
| 10 Brachlow et al. (2006) [23] | 32    | —        | —     | Apex LV                           | —         | cardiomegaly         | No          | Stable                | —                            | Asymptomatic at 6 months of life |
| 11 Bernasconi et al. (2004) [26] | 22    | 10 × 5 mm | Male  | LV lateral wall below mitral valve* | 46XY      | pericardial effusion | PC 22 w     | —                    | Fetal death 26 w, probably due to diverticulum rupture | —                                   |
| 12 McAuliffe et al. (2004) [27] | 13    | 4 × 6 mm | Male  | Apex RV                           | 46XY      | First trimester NT 4.2 mm pericardial effusion | PC 16 w     | Resolution of the effusion; CD stable | Eutocic delivery 38 w; weight 3070 grams; asymptomatic | Asymptomatic at 10 months of life |
| Author                       | GA di | Size       | Sex | Location   | Karyotype | Associated anomalies                          | Intervention                                    | Prenatal progression | Neonatal                                      | Follow-up                  |
|------------------------------|-------|------------|-----|------------|-----------|-----------------------------------------------|-----------------------------------------------|----------------------|----------------------------------------------|----------------------------|
| 13 McAuliffe et al. (2004) [27] | 13    | 4 × 3 mm   | Male| Apex RV    | 46XY      | First trimester NT 2 mm Pericardial effusion   | PC 14 w Resolution of the effusion; CD stable | Eutocic delivery | 38 w; weight 3150 grams; asymptomatic    | Asymptomatic at 8 months of life |
| 14 Prefumo et al. (2005) [1]  | 14    | 5 × 5      | Male| Apex RV    | 46XY      | First trimester NT 3.7 mm Pericardial effusion, ascites, and skin edema | PC 16 w Resolution of the effusion and hydrops; CD stable | Vaginal full-term eutocic delivery; asymptomatic | Asymptomatic at 22 months of life |
| 15 Prefumo et al. (2005) [1]  | 12    | 1 mm       | —   | Apex RV    | —         | First trimester NT 1.2 mm Pericardial effusion | No Spontaneous resolution of PE with 21 w; CD stable | Full-term eutocic delivery; asymptomatic | Asymptomatic at 17 months of life |
| 16 Gardiner et al. (2005) [19] | 14    | 2-3 mm     | —   | Apex RV    | Normal    | Pericardial effusion PC 14 w | Resolution of the effusion and hydrops; CD collapsed | Asymptomatic at birth | —                                    |
| 17 Gardiner et al. (2005) [19] | 14    | 2-3 mm     | —   | Apex RV    | Normal    | Pericardial effusion TOP | — | —                                    |
| 18 Del Rio et al. (2005) [18] | 13    | 5 × 5      | Female| Apex RV    | 46XX      | Pericardial effusion, septal defect AV** | No | Spontaneous resolution at 28 w | Eutocic delivery | 40 w; weight 3400 grams, asymptomatic at birth | Correction of septal defect at 3 months of life, resection of diverticulum; asymptomatic at 8 months of life |
| 19 Wax et al. (2007) [14]     | 20    | 6 × 9 mm   | Male| Junction base RV-infundibulum | —         | | No | No | Stable | Full-term eutocic delivery; weight 3689 grams; asymptomatic; small permeable FO | Asymptomatic at 18 months of life |
| 20 Koshiishi et al. (2007) [17] | 24    | 7 × 10 mm  | —   | Lateral wall below tricuspid valve (RV) | —         | Mild pericardial effusion; MC pregnancy with laser intervention for TTTS at week 20 where donor fetus died | No | Stable | Prenatal fetal death at 29 w | —                                    |
| 21 Pradhan et al. (2007) [28] | 28    | —         | —   | Apex LV    | —         | Fetal arrhythmia Hydrops fetalis Medical treatment (digoxin) | — | Vaginal delivery | 40 w | Asymptomatic at 12 months of life | —                                    |
| Author                  | GA di | Size   | Sex  | Location | Karyotype    | Associated anomalies                  | Intervention | Prenatal progression          | Neonatal | Follow-up                       |
|------------------------|-------|--------|------|----------|--------------|---------------------------------------|--------------|-------------------------------|----------|---------------------------------|
| Barberato et al.       | 16    | 5 × 5.7 mm | —    | Apex LV  | —            | Mild pericardial effusion              | PC 20 w      | Discrete enlargement of PE with normal heart function | —        | Prenatal fetal death 37 w     |
| Barberato et al.       | 30    | 12 × 13 mm | —    | Mitral subvalvular | —            | LV dilatation and reduced systolic function | No          | Stable                        | —        | Asymptomatic at 6 months of life |
| Davidson et al.        | 20    | —      | —    | Apex RV  | —            | Pericardial effusion                  | No          | Spontaneous resolution         | —        | Surgical treatment             |
| Williams et al.        | 21    | 5 × 5.5 mm | Male | RV       | —            | Pericardial effusion                  | PC 24 w      | Mild tricuspid regurgitation at 31; CD stable | Full-term delivery | Asymptomatic at a year of life |
| Perlitz et al.         | 22    | 7 × 4 mm | Male | RV lateral wall | —            | No                                    | No          | Stable, CD growth up to 9 × 9 mm | Eutocic delivery week 40; weight 4010 grams; asymptomatic at birth | Asymptomatic at a year of life |
| Menahem                | 19    | —      | —    | Apex LV  | —            | Pericardial effusion                  | No controls performed | Full-term live birth | —        | Asymptomatic at 10 months of life |
| Carrard et al.         | 13    | 2.6 × 2.9 mm | Male | RV lateral wall | 46XY       | First trimester NT 2.2 mm Pericardial effusion | Resolution after PC; CD collapsed at 26 w | Eutocic delivery 40 w, 2780 grams | Asymptomatic at 11 months of life |
| Abi-Nader et al.       | 22    | 3-4 mm | Male | RV       | 46XY        | Pericardial effusion                  | No          | Resolution at 32-33 w           | PROM 34 w; Intubation due to prematurity; caesarean section; weight 2460 gr; 2 muscle IVCs | Asymptomatic at 14 months of life |
| Abi-Nader et al.       | 21    | 11 × 15 mm | Male | RV lateral wall below tricuspid valve | —            | Isolated                              | —           | Eutocic delivery; weight 2780 gr; asymptomatic at birth | Caesarean section 38 + 5 w; weight 3270 grams; mild reduction of systolic function and premature ventricular contractions at birth | Asymptomatic at 16 months of life |
| Abi-Nader et al.       | 25    | 26 × 16 mm (37 s) | Male | RV       | —            | Arrhythmia and reduced systolic function | Induced delivery | —                             | Caesarean section 38 + 5 w; weight 3270 grams; mild reduction of systolic function and premature ventricular contractions at birth | Asymptomatic at 3 years of life, on prophylactic treatment with acetyl salicylic acid |

Table 2: Continued.
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| Author | GA di | Size | Sex | Location | Karyotype | Associated anomalies | Intervention | Prenatal progression | Neonatal | Follow-up |
|--------|-------|------|-----|----------|-----------|----------------------|--------------|----------------------|----------|-----------|
| 32     | 17    | —    | —   | Apex LV  | Normal    | Mesocardia, perimembranous IVC | No           | Stable               | Full-term live birth | Asymptomatic at 2 years of life |
| 33     | 21    | 1.6 × 0.4 mm | — | Apex LV  | Normal    | Defect on thoracoabdominal midline | TOP          | —                    | —        | —         |
| 34     | 31    | 12 mm (postnatal) | — | RV lateral wall below tricuspid valve | —         | —                    | No           | Ventricular septal defect | Full-term live birth; asymptomatic at birth; symptoms at 45 days of life: closure of septal defect at 3 months of life | Asymptomatic at 10 months of life |
| 35     | 14    | 2 mm | Male | Apex RV | 46XY      | Pericardial effusion | PE resolution after treatment; CD stable; moderate cardiomegaly; normal heart function | Full-term live birth; spontaneous eutocic delivery 40 + 1 w; weight 3150 grams | Asymptomatic at 4 years of life |

GA di: gestational age at diagnosis; RV: right ventricle; LV: left ventricle; w: weeks of pregnancy; TOP: termination of pregnancy; PC: pericardiocentesis; CD: cardiac diverticulum; IVC: interventricular communication PE: pericardial effusion; PROM: premature rupture of membranes; NT: nuchal translucency.  
*Diagnosis was made during the pathological examination after death. **Diagnosis of the ventricular septal defect was made after birth.
Table 3: Management and outcomes of the cases with cardiac diverticulum and pericardial effusion.

| Reference     | GA (w) | PE | Di | Loc.                  | Size (mm) | Intervention | PE findings | Prenatal progression | Postnatal progression |
|---------------|--------|----|----|-----------------------|-----------|--------------|-------------|----------------------|-----------------------|
| Carles et al. [24] | 13     | —  | —  | Apex LV               | —         | TOP 14 w     | —           | —                    | —                     |
| Cesko et al. [25] | 17     | AP | —  | Apex RV               | 3 mm      | TOP 22 w     | —           | —                    | —                     |
| McAuliffe et al. [27] | 14     | 14 | —  | Apex RV               | 2-3 mm    | TOP          | —           | —                    | —                     |
| Cavallé-Garrido et al. [6] | 19     | —  | —  | RV                    | 3 mm      | No           | —           | Spontaneous resolution at 34 w | Asymptomatic at 22 months |
| Cavallé-Garrido et al. [6] | 20     | —  | —  | LV lateral wall below mitral valve | Large    | No           | —           | Prenatal fetal death at 26 w, trisomy 18 | —                     |
| Prefumo et al. [1] | 12     | 12 | —  | Apex LV               | 1 mm      | No           | —           | Spontaneous resolution, effusion disappeared at 14 weeks; CD did not grow | Asymptomatic at birth; effusion or diverticulum not visible; asymptomatic at 17 months' follow-up |
| McAuliffe et al. [27] | 13     | 13 | —  | Apex RV               | 5 × 5 mm  | No           | —           | Spontaneous resolution; CD did not grow | IVC and IAC (postnatal); asymptomatic up to 3 months of age; surgical treatment; asymptomatic at 8 months of age |
| Pradhan et al. [28] | 20     | 20 | —  | Apex RV               | —         | No           | —           | Spontaneous resolution CD did not grow | Surgical treatment at birth |
| McAuliffe et al. [27] | 21     | 24 | —  | RV lateral wall       | 7 × 10 mm | No           | —           | Fetal death on week 29 | —                     |
| Perlitz et al. [30] | 19     | 19 | —  | Apex LV               | —         | No           | —           | No control performed | Full-term live birth; asymptomatic at 10 months of age; heart murmur; no treatment |
| Cavallé-Garrido et al. [6] | 19     | —  | —  | Apex RV               | —         | PC 20 w      | —           | No PE relapse, CD did not grow | Full-term live birth; asymptomatic at 12 months of age |
| Carles et al. [24] | 19     | 19 | —  | Apex RV               | 3 mm      | PC 20 w      | 7 cm$^3$ yellow fluid, 20 gr/L proteins (transudate), acellular | No PE relapse, CD did not grow | Full-term live birth; asymptomatic at 16 months of age; no treatment |
| Cesko et al. [25] | 22     | AP | —  | Pared lateral LV      | 10 × 5 mm | PC 25 w      | 25 mL old blood fluid | Intrauterine fetal death at 26 weeks (CD rupture) | —                     |
| Reference          | GA PE | GA Di | Loc. | Size (mm) | Puncture | PE findings                              | Prenatal progression                                      | Postnatal progression         |
|--------------------|-------|-------|------|-----------|----------|------------------------------------------|-----------------------------------------------------------|-----------------------------|
| 14 Brachlow et al. [23] | 13    | 13    | Apex RV | 4 × 6 mm | PC 16 w  | 3 mL sero-hematic fluid, 18 g/L proteins (transudate), lymphocytes, and mesothelial cells | No PE relapse or enlarging; CD was not visible on week 37 | Full-term live birth; asymptomatic at 10 months of age; no treatment |
| 15 Brachlow et al. [23] | 13    | 13    | Apex RV | 4 × 3 mm | PC 14 w  | 0.8 mL sero-hematic fluid, 15 g/L proteins (transudate) | No PE relapse; CD did not grow | Full-term live birth; asymptomatic at 8 months of age; no treatment |
| 16 Prefumo et al. [1]  | 14    | 14    | Apex RV | 5 × 5 mm | PC 16 w  | 5 mL clear fluid                          | No PE relapse; CD did not grow; mild cardiomegaly       | Full-term live birth; asymptomatic at 22 months of age; no treatment |
| 17 Gradiner et al. [19] | 14    | 14    | Apex RV | 2-3 mm   | PC 14 w  | 2 mL yellow fluid                         | No PE relapse; CD did not grow                            | Full-term live birth; asymptomatic; no treatment |
| 18 Carrard et al. [32] | 13    | 15    | Apex RV | 2.6 × 2.9 | PC 17 w  | 4 mL clear fluid, 21 g/L proteins (transudate) | No PE relapse; diverticulum was not visible from week 26 on | Full-term live birth; asymptomatic at 11 months of age; no treatment |
| 19 Williams et al. [3] | 21    | 21    | Apex RV | 5 × 4.5  | PC 24 w  | Yellow fluid 10 mL, 15.4 g/L proteins (transudate), lymphocytes | Complete resolution one week after PC CD did not grow | Full-term live birth; asymptomatic at one year of age; no treatment |
| 20 Barberato et al. [29] | 16    | 16    | —     | —        | PC 20 w  | Blood-stained fluid                       | Moderate growth of PE size as compared with postpuncture effusion; expectant approach Intrauterine fetal death on week 37 | —                           |
| 21 Abi-Nader et al. [2] | 12    | 22    | Apex RV | —        | PC 18 w  | —                                        | Relapse one week later and subsequent spontaneous resolution on weeks 32-33 | —                           |
| 22 Our case           | 12    | 14    | Apex RV | 2 mm     | PC 17 w  | Clear yellow fluid, acellular, transudate | No PE relapse; CD did not grow; moderate cardiomegaly     | Full-term live birth; asymptomatic at birth; treatment with ASA; asymptomatic at 4 years of age |

GA PE: gestational age at pericardial effusion; GA di: gestational age at diverticulum diagnosis; RV: right ventricle, LV: left ventricle; w: weeks of pregnancy; PC: pericardiocentesis; CD: cardiac diverticulum; IVC: interventricular communication; PE: pericardial effusion.
resolution (71%) and 2 resulted in intrauterine death (29%): one of them, which occurred on week 26, was associated with trisomy 18 and the other, which occurred on week 29, was associated with treated twin-to-twin transfusion syndrome and death of one of the twins after treatment [6,18].

Our review is the largest series published in the literature with 34 cases of fetal cardiac diverticula. If we exclude the cases of termination of pregnancy, chromosomal abnormalities, and pregnancy complications (twin-to-twin transfusion syndrome), the cardiac diverticulum is a benign condition with good postnatal outcomes, with a mortality of 0.6%. In the cases with pericardial effusion, a pericardiocentesis, performed from 16 weeks of gestation with a 20-gauge needle, seems to be a safe procedure with good outcomes in 83% of cases. The evolution of the pericardial effusion and the diverticula (spontaneous resolution, stability, or progression) is not known because there are few cases published, but the risk of cardiac insufficiency, hydrops fetalis, or pulmonary hypoplasia could be avoided with this procedure.

In our case we made an early diagnosis and treatment with pericardiocentesis with resolution of the pericardial effusion with an excellent outcome. Only 10 cases (29%) were reported during the 11–14 weeks’ scan and only 6 cases (17%) with pericardial effusion associated during this period.

Cardiac diverticula are rarely associated with chromosomal abnormalities especially when other malformations are not present [5]. Cardiac abnormalities usually associated with diverticula include ventricular or auricular septal defects, permeable oval foramen, tricuspid atresia, tetralogy of Fallot, persistent left superior vena cava, pulmonary artery hypoplasia, and coarctation of the aorta [5]; cardiomegaly is the most frequently associated complication in the third trimester, like in our case [6,19]. Usually associated extracardiac malformations include midline thoracoabdominal defects (omphalocele, ectopia cordis, Pentalogy of Cantrell, etc.) [2, 5, 8, 20–22].

The prognosis of this entity is generally good, although the outcome largely depends on the size and location of associated anomalies. Cases of rupture, both pre- and postnatal, arrhythmia, fetal death, heart failure, and coronary insufficiency have been described [9,16,18,21,23]. In these patients, serial control examinations are necessary to detect possible complications. In general, postnatal progression is good and surgery is not necessary in asymptomatic cases [19].

In conclusion, a cardiac diverticulum is a rare entity. The most frequently associated complication is pericardial effusion, which may result in cardiac decompensation, hydrops fetalis, or pulmonary hypoplasia. Although spontaneous resolution of effusion has been reported, pericardiocentesis is a safe and effective technique, which may be used to reduce the risk of secondary disorders in selected cases. Nowadays there are few cases published and the evidence is poor to establish the appropriated management.

## Conflict of Interests

The authors declare no conflict of interests.

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