Single Atrium and Miscarriages

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Introduction

Single atrium is a rare heart condition comprising under 1% of all congenital heart disease, in which the inter atrial septum is missing. These congenital heart defects usually are surgically corrected after birth.

Case Presentation

We present a successful surgical repair of a single atrium (SA) in a 27-year-old woman, who has a complete missing of the atrial septum, without coexisting valvular pathology. The SA diameter was 9.97 × 6.18 cm and the Systemic blood flow/pulmonary blood flow (Qp/Qs) was 4.1 due to the single atrium. Surgical correction consisted of creating a new atrial septum, using a double-velor patch. At first- and second-year follow up, the patient was in a very good condition and without any symptoms present. We consider that the diagnoses of single atrium, especially in child bearing women, should be done in appropriate time, to avoid miscarriages, preterm births and other complications associated.

Conclusion

There are many case reports that show good surgical repair of single atrium and without late complications. In our case, we have obtained a very good surgical repair, without AV bundle damage and no residual shunt demonstrated on intraoperative transesophageal echocardiography.

Keywords

► single atrium
► common atrium
► miscarriages
► preterm births
► heart defect
► pregnancy

Abstract

Background A single atrium is a very rare heart condition, in which the inter atrial septum is missing. These congenital heart defects usually are surgically corrected after birth.

Case Presentation We present a successful surgical repair of a single atrium (SA) in a 27-year-old woman, who has a complete missing of the atrial septum, without coexisting valvular pathology. The SA diameter was 9.97 × 6.18 cm and the Systemic blood flow/pulmonary blood flow (Qp/Qs) was 4.1 due to the single atrium. Surgical correction consisted of creating a new atrial septum, using a double-velor patch. At first- and second-year follow up, the patient was in a very good condition and without any symptoms present. We consider that the diagnoses of single atrium, especially in child bearing women, should be done in appropriate time, to avoid miscarriages, preterm births and other complications associated.

Conclusion There are many case reports that show good surgical repair of single atrium and without late complications. In our case, we have obtained a very good surgical repair, without AV bundle damage and no residual shunt demonstrated on intraoperative transesophageal echocardiography.
was not diagnosed with heterotaxy syndrome, Ellis-van Crevald or trisomy 21.

In our clinic, she presented with mild cyanosis of her fingers and lips. She told us that her feet and hands have been cold since childhood and on exercise she had difficulty in breathing.

Saturations on room air 91%, on arterial gasses partial pressure of oxygen (pO₂) 69 mm Hg, SaO₂ 89%.

Preoperative echocardiography and TEE described a single atrium, diameter 9.97 cm with 6.18 cm. Left ventricular ejection fraction (LVEF) 60%. Mitral and tricuspid valve insufficiency grades I to II, same level offset, and pulmonary arterial pressure (PAP) 40 to 45 mm Hg. Dilated right ventricle (33 mm), and tricuspid annular plane systolic excursion (TAPSE) 22 mm. Systemic blood flow/pulmonary blood flow (Qp/Qs) is 4.1 (►Figs. 1B, 2A1, 2A2).

Preoperative Imaging. (A) Chest X-ray: demonstrating cardiomegaly, right atrial and right ventricular enlargement and a prominent pulmonary trunk (B) Echocardiography: four chamber view, single atrium with a diameter 9.9 cm × 6.1 cm. IVS, inteventricular septum; LV, left ventricle; MV, mitral valve; RV, right ventricle; SA, single atrium; TV, tricuspid valve.

Fig. 1 Preoperative Imaging. (A) Chest X-ray: demonstrating cardiomegaly, right atrial and right ventricular enlargement and a prominent pulmonary trunk (B) Echocardiography: four chamber view, single atrium with a diameter 9.9 cm × 6.1 cm. IVS, inteventricular septum; LV, left ventricle; MV, mitral valve; RV, right ventricle; SA, single atrium; TV, tricuspid valve.

Fig. 2 Preoperative Imaging. (A) TEE: A1, tricuspid Valve with a short septal leaflet and mild insufficiency; A2, mitral valve with mild insufficiency and light myxomatous leaflets. (B) CT angiography: B1 is the transversal view; single atrium with normal drainage of pulomonary veins to the left side. B2 is the frontal view; huge single atrium with contrast from superior vena cava drained to right side of the atrium and prominent pulmonary artery trunk. A, aorta; CT, computed tomography; LPV, left pulmonal veins; LV, left ventricle; RPV, right pulmonal veins; RV, right ventricle; SA, single atrium; TP, truncus pulmonalis; VCS, vena cava superior.
TEE showed a left shift of the interventricular septum, aortic valve with three normal leaflets, mitral valve with myxomatous leaflets, and mild insufficiency, mild tricuspid valve insufficiency.

Computed tomography (CT) angiography showed a single atrium with normal drainage of pulmonary veins to the left side of atrium and with the superior and inferior vena cava drained to the right side of atrium. Truncus pulmonalis was prominent (►Fig. 2B1, 2B2, ►Video 2)

Video 1


Surgical Procedure

Surgical procedure for this study was standard midline sternotomy and aortic and bicaval cannulation. After opening of the right atrium, we noticed complete absence of the interatrial septum (►Fig. 3A, B). Superior and inferior pulmonary veins from the right and left lung drained into what supposed to be left atrium. Mitral and tricuspid valves had normal morphology. The coronary sinus opened normally into right atrium. After careful inspection of the entire atrium, we started to create an interatrial septum, with double pericardial patch. As described by Jing et al (second technique2), we started sutures (4–0 Prolen) from the middle of the ventricular septal crest downward to the borderline between the tricuspid septal valve annulus to the left inner side of the coronary sinus to the remaining border of the atrial septum. The patch was sutured with running sutures and extra fixating mattress suture without pledgets were taken every 2 to 3 cm.

Results

Intraoperative and postoperative echocardiography showed that there was a mild mitral and tricuspid valve insufficiency. There was no interatrial defect remained. Right atrium with normal size, however, mildly enlarged left atrium.

Postoperatively, there was no atrial fibrillation or AV block diagnosed; however, saturations dropped to a range of 84 to 85%. Mitral valve had mild insufficiency, v.contracta 0.2 cm on echocardiography postoperative and in follow up after 2 years (►Fig. 3C, ►Video 1). The pulmonary systolic arterial pressure (PSAP) postoperatively was 29 mm Hg and the right

Fig. 3  Intraoperative imaging and TTE at follow-up after 2 years. (A) TV and MV with the patch in the middle and the suture line where the patch was sutured at the posterior border of CS. (B) Right atrium, view of tricuspid valve and artificial septum with patch. (C) Echocardiography, follow up after 2 years. Mild insufficiency of mitral valve. Vena contracta 0.2. CS, coronary sinus; LA, left atrium; LV, left ventricle; MV, mitral valve; RA, right atrium; RPV, right pulmonal veins; RV, right ventricle; TTE, transthorakal echokardiografy; TV, tricuspid valve; VCI, vena cava inferior.
ventricular systolic pressure (RVSP) was 23 mm Hg. Anticoagulation with cumarin (international normalized ratio [INR]: 2.5–3.0) was given for a period of 6 months. Follow-up was performed after first and second year, the patient was in very good condition result.

In the initial recovery period, we noticed a reduction of saturation in room air (SaO2), from 93% preoperatively to 85% during the first 30 days postoperatively. The SaO2 values stabilized within 1 year from surgery. At present, her SaO2 values are maintained at 97%, with partial pressure of oxygen (pO2) 66 mm Hg, partial pressure of dioxidecarbon (pCO2) 28.4 mm Hg and on follow-up echocardiography PASP reduced to 29 mm Hg from 45 mm Hg, without medical therapy. She keeps very active and for the first time she does not complain of tiredness and there is no cyanosis present.

Discussion and Conclusion

There are few case reported involving common atrium also known as single atrium.1

Rastelli and associates3 reported few cases were the absence of the atrial septum was accompanied by presence of the cleft into the anterior leaflet of the mitral valve and the absence of interventricular communication. On the other hand, Levy and associates4 reported a case where the complete absence of the atrial septum was not accompanied by other defects as described by Rastelli et al, suggesting that this condition can exist alone as a specific entity. Our case supports the recommendation by Levy and associates; the term "single atrium" is used in this report.

The term miscarriage is defined as pregnancy that ended spontaneously with the loss of a non-registerable fetus under < 24 weeks of gestation. Preterm birth is defined as a birth before 37 weeks of gestation.5 In March of Dimes, the miscarriage is defined as early pregnancy loss before 20 weeks of pregnancy.

The clinical picture of our patient, cyanosis, rhythm disturbances, weight loss, and reduced exercise tolerance started years back. This clinical presentation did not differ from patients diagnosed with a large atrial defect at the fossa ovalis.4

In single atrium, the blood (arterial and venous) is mixed, which explains mild cyanosis, shortness of breath, increased fatigue,6 and reduced exercise tolerance early in life.7

Surgical correction should be done early in life to prevent later complications. Our patient unfortunately was diagnosed very late and only after going through three preterm births.5 Surgical repair consists in creating two atria, using a patch, which could be a Dacron or an autologous pericardial patch. For reconstruction of the atrium, an artificial patch, as we could not obtain a fixing solution for native pericardium. The suture method was as described by Jing et al (second technique).1

We cannot explain the drop in oxygen saturations in the initial postoperative period. There was no postoperative shunt present and the blood was not mixed in the atra. Maybe, this was as a result of postoperative pain, impeding patient to take deep breaths and perform respiratory exercises resulting in mild atelectasis or maybe that was triggered by pulmonary hypertensive attack. In these situations, use of sildenafil would be recommended. We are open to discussion about this problem and maybe some other authors have encountered this in their cases.

Prematurity is a health priority in many countries. Howson et al8 described as a second leading cause of death in children under 5 years. The Organization March of Dimes reports that premature birth is a first cause of death of babies in the United States.5

World Health Organization (WHO) in the Global Action Report “Born Too Soon”9 want to support all families who have been touched by preterm births and said that over three quarters of premature babies could be saved.

Congenital Heart Disease Guidelines from 2018 recommended a pregnancy heart team: include pediatric cardiologist, gynecologist–obstetrician, anesthetist, and cardiothoracic surgeon should discuss and follow prepregnancy patients who can have complications during pregnancy and offer adequate treatment.

Our patient presented with a heart defect which was undiagnosed for years and costs our patient three preterm births and lately divorce. This history makes us push forward the need for guidelines to be followed and implemented in every institution which treat pregnant women. Patients like ours will not only bear the consequences of losing three pregnancies, but losing her family too, which will have psychological implications.

We can conclude that presence of single atrium in child bearing women, increases the risk of early births and endangering a woman’s life. After successful surgical repair, long-term survival is very good and quality of life improves enormously.

Consent for Publication
Written informed consent for publication of his clinical details and clinical images was obtained from the patient.

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Conflict of Interest
None.

References