Pulmonary Arterial Hypertension Associated with Adult Congenital Heart Disease, when Inoperable becomes Operable: A Case Report

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Abstract

Pulmonary arterial hypertension associated with congenital heart disease (PAH-CHD) is one of the most common forms of pulmonary arterial hypertension (PAH). Unlike the other forms, PAH-CHD can be treated surgically closing the extra- or intracardiac shunt if the shunt is left-to-right and the pulmonary vascular resistance (PVR) is low enough (<2,3-operable, 2,3-4,6 WU-borderline operable). If the PAH postoperatively persists the prognosis is even worse than for non-operated patients. Due to the previously mentioned fact the criteria for operability in such cases are often discussed. We report a case of a 54-year-old man who suffers from coronary artery disease (SYNTAX score-23,5) and PAH-CHD. As the patient required coronary artery bypass grafting (CABG) surgery, the atrial septal defect (ASD) was closed (surgically with a patch followed by transcatheter closure with an occluder) as well despite the fact that according to the current guidelines his pulmonary vascular resistance (4,7 WU) was considered too high for the defect to be operable. The patient was given PAH-specific treatment preoperatively in order to lower his PVR. This article describes the case and discusses the disease, its history, the new classification, diagnostics and treatment options.

Keywords: Pulmonary hypertension; Atrial septal defect; Surgical repair; Sildenafil; Congenital heart disease

Background

Pulmonary arterial hypertension (PAH) is a specific subgroup of pulmonary hypertension (PH). It is characterized by an increased pulmonary pressure and pulmonary vascular resistance which can lead to right ventricular failure [1-4]. PAH is a common complication of congenital heart disease (CHD), which occurs when an intra- or extracardiac shunt exists (Table 1) [5,6]. Such cardiac defects cause PAH by increasing pressure and volume overload in the right side of the heart, as intimal proliferation and medial hypertrophy increase the resistance of the pulmonary vessels [4]. The UK National PH Audit reported in 2012 that the prevalence of PAH-CHD (30.2%) was similar to idiopathic PAH (33.6%) and connective tissue disease-related PAH (28.3%) of all PAH patients [7]. Atrial septal defect (ASD) is one of the most common congenital heart defects [8]. If PAH is left untreated it can be life-threatening [4]. Therefore we would like to report a case of a 54 year-old male who was diagnosed with PAH due to an ASD. A high PVR (4.7 WU) was detected. And despite the fact that the current guidelines prohibit such patients (PVR>4.6 WU) to undergo surgery, the patient was operated on. The patient's overall status significantly improved.

Case Presentation

A 54 year-old male who had been experiencing palpitations due to atrial flutter was admitted to the cardiology ward in June 2008. A transesophageal rhythm restoration was performed. Transesophageal echocardiography (TEE) revealed right atrial and ventricular dilatation, pericardial fluid (1.7 cm) and an ASD (2 cm) with a bidirectional shunt, the right ventricular systolic pressure (RVSP) was 85-90 mmHg therefore pulmonary hypertension was suspected. Laboratory findings showed a high normal erythrocyte level (5.80x1012/L) and a slightly elevated Hb level (178 g/l). High levels of total cholesterol (7.1 mmol/l), triglycerides (2.8 mmol/l) and low density lipoprotein (4.77 mmol/l) were detected. High density lipoprotein level was normal (1.06 mmol/l). His ECG showed sinus rhythm, right axis deviation, severe hypertrophy of the right ventricle and signs of right atrial and ventricular overload (Figure 1). His spirometry test results showed a normal forced vital capacity (FVC) of 80.1% and a mildly lower forced expiratory volume in 1 second (FEV1) of 67.8%. Anomalous pulmonary venous return was excluded on the computed tomography (CT) scan but dilatation of the right atrium with hypertrophy, dilatation of both pulmonary arteries and pericardial fluid (0.8-1.3 cm) was seen. Coronary angiography was performed and an occlusion of the proximal third of the left anterior descending artery (LAD), a stenosis (>90%) of the second obtuse marginal branch (OM2) of the left circumflex artery (LCX), and a stenosis (>75%) in the mid third of the LCX was detected. Right heart catheterization revealed an atrial septal defect with a left-

Table 1: Anatomical classification of congenital systemic-to-pulmonary shunts associated with PAH [6]. ASD: Atrial Septal Defect; VSD: Ventricular Septal Defect; AVSD: Atrioventricular Septal Defect.

| Simple pre-tricuspid shunts | Simple post-tricuspid shunts | Combined shunts | Complex CHD |
|-----------------------------|-----------------------------|-----------------|-------------|
| ASD (Ostium secundum) | VSD | Combination of shunts | AVSD (partial or complete) |
| ASD (Sinus venosus) | Patien ductus arteriosus | Truncus arteriosus |
| Total or partial unobstructed anomalous pulmonary venous return | Single ventricle physiology with unobstructed pulmonary blood flow |
| | Transposition of the great arteries with VSD (without pulmonary stenosis) and/or patent ductus arteriosus |

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to-right shunt. The mean pulmonary arterial pressure (mPAP) was 67 mmHg, PVR-4.7 WU, PVRI-2.27 WU/m², Qp-9.8l/min, Qs-3.8l/min, SVR-22.6 WU, SVRI-10.7 WU/m². The patient’s oxygen saturation assessed in the femoral artery was 95% and 94% in the left pulmonary vein. The patient was recommended to be operated on (ASD closure and coronary artery bypass grafting). PH specific therapy in particular sildenafil 20 TID was prescribed a month prior to surgery and was continued until the defect was closed completely.

In November 2008 percutaneous transcatheter coronary angioplasty (PTCA) was performed in the LCX and the OM2. The patient underwent surgery in February 2009. A longitudinal sternotomy with aortocoronary shunting (LITA-LAD) and closure of the atrial septal defect with a patch (4 × 4 cm) leaving a residual 1cm in diameter shunt was done. The patient was in the intensive care unit for a day and the rest of the postoperative period went well. After the surgery the patient’s right ventricular systolic pressure (RVSP) estimated by echocardiography decreased to 34 mmHg.

A follow-up echocardiography was done in June and September 2009. A residual left-to-right shunt of 6 mm in diameter and dilatation of the left heart was seen. Tricuspid regurgitation II that was seen in June was no longer detectable in September.

In 2010 the patient underwent percutaneous residual shunt closure with an occluder. On echocardiography in 2012 no residual shunt was seen, there was mild dilatation of the right ventricle with the systolic function being intact, both atria mildly dilatated, RVSP-40-45 mmHg, tricuspid regurgitation I-II (Table 2).

The patient was seen on a follow-up visit in November 2015. Right now the patient doesn’t take PH-specific medication, he takes acetylsalicic acid and atorvastatin 40 mg QD. The patient hasn’t experienced any of the complications associated with ASD and pulmonary arterial hypertension, has no dyspnea, cyanosis or edema.

### Table 2: Consequent results of the patient’s echocardiogram.

| Sep 2008 (TEE) | Jun 2009 | Sep 2009 | Sep 2012 |
|---------------|----------|----------|----------|
| RA            | ¶         | 53x59mm  | ¶         | 30 cm²   |
| RV(mm)        | ¶         | 46       | 47       | 50       |
| LA (mm)       | -         | 55       | 55       | 58       |
| RVSP (mmHg)   | 85-90     | 75-80    | -        | 40-45    |
| TR (I-IV)     | II        | II       | 0        | I-II     |

6 minute walk test results were excellent 519 m. His saturation was 93% before the test and 97% after the test; he rated his dyspnea as a 2 according to the BORG scale.

### Discussion

Pulmonary arterial hypertension is a progressive disease that is characterized by an increased mean pulmonary arterial pressure of at least 25 mmHg at rest and an elevation of the pulmonary vascular resistance (PVR) of more than 3 WU with a normal pulmonary artery wedge pressure of ≤ 15 mmHg [1,9,10].

Currently it is thought that the pathophysiological cornerstones of pulmonary hypertension are vascular remodelling caused by imbalance between cell proliferation and apoptosis, thrombosis, and the prevalence of vasoconstriction [9].

In 1973 the WHO organized an international conference on PH. Right now according to the WHO classification there are 5 groups of PH [9]. Since 1998 when the working symposia of the WHO was held in Evian, pulmonary arterial hypertension associated with congenital heart disease (PH-CHD) has been classified under the WHO group I, pulmonary arterial hypertension (PAH), along with idiopathic PH,
familial PH, toxin induced PH, persistent neonatal PH and PH due to autoimmune, infectious or hepatic and hematologic causes [11].

One of the most common congenital heart defects is an atrial septal defect (ASD) [8]. It makes 10% of all CHD and 22-40% of CHD in adults. The most common is ostium secundum defect (60-70%). ASD occurs twice as often in women as in men [12]. It has been reported that out of all patients with pulmonary arterial hypertension 5-10% have CHD [1]. Yet the prevalence of PAH in patients with CHD can vary as the location and size of the defect matter [5]. PAH can be prevented if the cardiac defect is successfully closed early but operability can only be considered for patients with a prevalent left-to-right shunting [1,5]. Due to the rapid progress in paediatric cardiology and surgery the number of patients with PAH-CHD in western countries has noticeably decreased and the number of patients with CHD who have survived into adulthood has increased [5].

A large ASD is associated with clinical presentation during childhood but mostly the patients are asymptomatic and the disease tends to manifest as the patient gets older [8,12]. But even when the patient is asymptomatic, such severe complications as right ventricular failure, atrial arrhythmias, paradoxical embolization, cerebral abscess formation, and pulmonary arterial hypertension that can become irreversible and lead to right-to-left shunting which is called Eisenmenger syndrome can occur [13]. 90% of the untreated patients develop such symptoms as dyspnea, fatigue, palpitation and persistent arrhythmia by the age of 40 [12]. In case of left-to-right shunting via ASD, symptoms appear when right ventricular volume overload and excessive circulation in the pulmonary system has developed [8].

The physician should always suspect the diagnosis of ASD when the right heart is enlarged without another explanation [14]. Echocardiography can detect the size and location of the ASD, as well as evaluate the shunt and the degree of PAH [12]. Signs of PAH on a chest x-ray include increased lung vascularity, pulmonary vascular congestion, and rib notching. Computed tomography (CT) or magnetic resonance (MRI) can improve the description of the defect. Cardiac catheterization remains the gold standard in detecting the necessary pressures, cardiac output, and PVR and most importantly cardiac catheterization offers the opportunity to correct the defect or to determine whether the patient is suitable for surgery [14].

In the past 10 years PAH treatment options have improved noticeably [15]. Currently there are many oral, intravenous and inhaled drugs. The main pharmacologic groups are calcium channel blockers (CCBs), prostanooids, endothelin-1 receptor antagonists and phosphodiesterase-5 inhibitors [9]. Patients with PAH-CHD should avoid using CCBs as there is no available data that support their benefit. In patients with Eisenmenger's syndrome CCBs are contraindicated due to the systemic hypertensive effect which can intensify the right-to-left shunt, thus increasing the risk of syncope or even sudden death. According to the BREATHE-5 trial bosentan improves exercise capacity, haemodynamics and functional class in patients with Eisenmenger's syndrome. Sildenafil has been proven to increase exercise tolerability, dyspnea, functional class, the quality of life and haemodynamics in patients with PAH-CHD as well as patients with Eisenmenger's syndrome. Recently there have been many studies investigating whether combined PAH-specific therapy could be beneficial but so far the results are inconclusive. Yet the available data show that PAH-specific treatment definitely improves the outcome [5].

An ASD is closed with a surgical suture or a patch, or via right heart catheterization [13]. It is indicated to close an ASD if an enlargement of the right atrium or the right ventricle is present even in the absence of symptoms. For patients <25 years of age the prognosis is excellent after ASD closure. Unfortunately the prognosis for patients of age >40 years is poorer than in the general population as this group is predisposed for postoperative arrhythmias [14]. Still it is reported that regardless of age most symptomatic adults experience clinical improvement [8]. According to the 2015 ESC/ERS Guidelines for the diagnosis and treatment of pulmonary hypertension CHD closure is contraindicated when the PVR is >4.6, PVRI >8. And in order to achieve the necessary criteria for a surgical defect correction PAH-specific therapy is not recommended [1]. Despite the fact that the patients with a high mPAP and PVR have notable changes of the pulmonary vasculature, there have been several case reports of ASD closure in patients who were borderline operable or inoperable thus the interest in the “treat-to-close” approach has increased [5,9,16]. The aim of this approach is to use PAH-specific therapy in order to reduce the PVR thus improving operability.

Conclusion

This case report emphasizes the importance of the case-by-case approach for evaluation of the PAH-CHD patient's suitability for operation as surgical correction is the most effective way to limit the progression of PAH. PAH-specific treatment should always be considered not only to improve the patient's quality of life and decrease the mPAP but also to decrease the PVR thereby potentially readdressing the operability criteria.

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