CASE PRESENTATION

An Uncommon Congenital Abnormality Discovered Using Multimodality Cardiac Imaging in an Elder Hospitalized For Decompensated Heart Failure

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ABSTRACT
We report the case of a 77-year-old patient admitted for worsening dyspnea, palpitations and dizziness. The electrocardiogram (ECG) showed atrial fibrillation (AF) and left bundle branch block (LBBB). Clinical examination showed hypoxemia (SaO₂ 87% room air), jugular venous distension, pitting edema, arrhythmic heart sounds, a splitting of the second heart sound and systolic murmur (III-IV/VI) in the tricuspid area. Transthoracic echocardiography (TTE) showed right heart dilation, moderate tricuspid regurgitation, biventricular hypertrophy, and a dilated coronary sinus (CS). Agitated saline injection in the left basilar vein raised a suspicion of persistent left superior vena cava (PLSVC) to coronary sinus fistula. Transesophageal echocardiography (TOE) showed a superior sinus venous defect, with bidirectional shunt, and persistence of LSVC. CT angiography confirmed and detailed the malformation features. A low suspicion for congenital defects in older adults presents a significant diagnostic challenge. Multimodality cardiac imaging is critical in determining the etiology of right heart dilation, and knowledge of the anatomy and physiology of various shunt lesions is essential for clinicians.

Keywords: heart failure, coronary sinus, persistent left superior vena cava, sinus venous defect, multimodality imaging.

REZUMAT
Prezentăm cazul unei paciente în vârstă de 77 de ani, care se internează pentru agravarea dispneei, palpității și vertii. Electrocardiograma (ECG) a surprins fibrilație atrială și bloc de ramură stângă. Examenul clinic a decelat hipoxemie (SaO₂=87% în a.a.), turgescență jugulară, edeme gambiere, zgomote cardiace aritmice, dedublarea zgomotului 2, suflu sistolic gradul III-IV/VI în focarul tricuspidian. Ecocardiografia transtoracică (ETT) a vizualizat dilatarea cordului drept, insuficiență tricuspidiană moderată, hipertrofie biventriculară și sinus coronar (SC) dilatat. Testul cu soluție salină barbotată injectată la nivelul venei bazilice stângi ridică suspicunea de persistență de venă cavă superioară stângă (PVCS). Ecocardiografia transesofigiană (ETO) a decelat defect septal atrial tip sinus venos superior cu șunt bidirecțional și PVCS. Examinarea angio-CT a confirmat și detaliat malformația. Defectele congenitale desoperite la vârstnici sunt cauze rare de insuficiență cardiacă și trebuie caracterizate prin metode imagistice multimodale. Aceste tehnici sunt indispensabile în depistarea etiologiei dilatații cordului drept și cunoașterea anatomei și fiziologiei leziunilor cu șunt este esențială pentru clinicieni. 

Cuvinte cheie: insuficiență cardiacă, sinus coronar, persistență venă cavă superioară stângă, defect tip sinus venos, imagistică multimodată.

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BACKGROUND
A low suspicion for congenital defects in older adults admitted for decompen­sated heart failure (HF) pre­sents a significant diagnostic challenge. We present the incidental discovery, in an elderly patient with a history of coronary disease (atypical angina) and left bundle branch block (LBBB), admitted for acute dyspnea in the setting of paroxysmal atrial fibrillation (AF), of a dilated right ventricle which was proven to be caused by the superior sinus venous defect (SVD), as well as a persistent left superior vena cava (PLSVC) with drainage into coronary sinus (CS). PLSVC, the most common venous anomaly in thorax, without an associated adult congenital heart disease (ACHD) is usually asymptomatic and discovered incidentally. Among the ACHD frequently associated with PLSVC (based on odds ratio) is the atrial septal defect (ASD). ASD can be diagnosed incidentally in asymptomatic adults examined for other comorbidities. However, a right heart dilatation discovered in the setting of unrepaired ASD can be symptomatic as right HF, atrial arrhythmias and pulmonary hypertension. ACHD are frequently asso­ciated with HF, which represents the main cause of mortality in these patients.

CASE REPORT
A 77-year-old woman is admitted to the Emergency Department (ED) for progressive exertional dyspnea, with a day of dyspnea at rest, palpitations and dizzi­ness. A SARS CoV2 infection was ruled out after RT-PCR and chest X ray (CXR) in the ED. The patient’s medical history was relevant for NYHA class II heart failure (HF) diagnosed in an ambulatory setting six months earlier, coronary disease (atypical angina), left bundle branch block (LBBB), documented on ECG two years before the actual admission, and chronic he­patitis C. She was treated with spironolactone 50 mg o.d., furosemide 40 mg o.d., bisoprolol 5 mg o.d. and a heptoprotective medication for the last six months. Clinical examination upon admission revealed peri­pheral cyanosis, jugular venous distension, irregularly irregular heart sounds with a split S2, systolic murmur (III-IV/VI) in the tricuspid area, painful hepatomegaly, and hypoxemia (SaO2 87% room air), that improved to 95% with 4L of oxygen through a nasal canula. ECG performed upon admission in the ED showed atrial fibrillation (AF) and LBBB (Figure 1A). CXR show­ed cardiomegaly, and bilateral lung hypervascularity (Figure 1B). Bloodwork showed pancytopenia (RBC 3.31 mil/mm3, WBC=3.040/mm3, PLT 46.000/mm3), hepatic cytolysis and cholestasis, an eGFR of 42 mL/min/1.73m2, and NT-proBNP 2189 pg/mL. The tran­s­thoracic echocardiography (TTE) revealed a dilated coronary sinus (CS) with a maximal diameter of 33 mm (Figure 2 A,B), enlarged right heart chambers, a dilated left atrium (LA), an interatrial septum bulging towards the LA, and a subtle enlargement of the main and right pulmonary artery (PA). Other noticeable findings were a right ventricular (RV) hypertrophy (7 mm), with normal systolic function (TAPSE 21 mm), an interventricular septum (IVS) of 13 mm flattened in dyps­ole (Video 1), a preserved systolic function of the LV (51%), and pulmonary hypertension (estimated PA systolic pressure 48 mmHg, PA acceleration time 87 msec). Also, a moderate tricuspid regurgitation, mild mitral and aortic regurgitation, and a dilated inferior vena cava 30 mm which collapses 50% with inspira­tion, a calcification of the mitral annulus, aortic lea­flets and ascending aorta were noticed. Agitated saline contrast injection from the left basilic vein (Video 2) showed initial opacification of the CS and then of the right atrium (RA), with contrast rapidly moving to LA (Figure 2 C), which raised the suspicion of a persistent left superior vena cava (PLSVC) to CS as well as a right to left shunt. A TOE showed a sinus venous defect (SVD), with a bidirectional shunt (Figure 2D), PLSVC, and dilation of the left superior pulmonary vein. CT angiography confirmed the PLSVC (Figure 3 A,C,D) with drainage into CS, with a bridging vein of 8 mm caliber (Figure 3 B) and a normal pulmonary venous return. CT angiography also showed a dilated main PA (40 mm), and its branches (left PA 28 mm and ri­ght PA 31 mm), revealing coronary and aortic calcified atheromatosis, and ruled out a pulmonary embolism or other arteriovenous malformations. During hos­pitalIZATION, the invasive measurement of pulmonary hypertension and vascular resistance was not possible due to technical difficulties.

The patients’ symptoms improved with low dose angiotensin receptor blockers, increased dose of di­uretics, alpha-beta-blockers and oral anticoagulation. She however continued to have episodes of desat­urations as well as exertional dyspnea. Patient was not compliant to follow-up appointment.

The particularity of this case lies in the incidental finding of a rare cardiac congenital malformation in an elderly patient hospitalized for acute decompensati­
on of HF, probably favored by recent onset AF. To our knowledge, this might be the eldest patient with association of SVD and PL SVC complicated with Eisenmenger syndrome (ES) reported in literature.

DISCUSSION

The incident discovery of congenital heart disease in an older patient with known ischemic heart disease, admitted for acute on chronic decompensation of HF, is rare. In patients with ACHD, RV failure is a common phenomenon with an incidence of up to 70% in selected patient populations. The most common causes of right heart dilation are congenital tricuspid valve disease and shunt lesions. Superior SVD amounts to up to 5% of all atrial septal defects (ASD). PL SVC has a prevalence of 0.2-3% in the healthy general population, but up to 11% in people with other cardiac congenital malformations. These malformations are occasionally discovered in adults with right heart dilation. Persistence of left anterior cardinal vein joining the CS which is grossly dilated in calibre and draining into RA. D. Thoracic CT angiography coronal section, mediastinal view: LS SVC between Ao and LPA, the RS SVC in normal position (green arrows).

PL SVC plays a considerable role in the induction and maintenance of AF in nearly half of the patients. If PL SVC is detected as the trigger or driver of AF, it can be an additional target during electrophysiologic study and ablation. In our case, the onset of AF cannot be attributed only to the presence of LS SVC but also to the RA and right heart failure.

Our patient had classic physical examination findings suggesting an ASD. However, ECG displayed a LBBB instead of a classical incomplete right bundle branch block, perhaps due to her ischemic heart disease. Right heart dilation, without another etiology, in the setting of unrepaired ASD is considered a risk for progression toward symptomatic right heart failure, atrial arrhythmias, and potential development of pulmonary arterial hypertension. This was also the case with our patient.

Changes in atrial size and pressures in patients with ASD have been shown to induce and maintain atrial tachyarrhythmias by various mechanisms. An increase in RV pressure by acute volume overload significantly prolongs RA effective refractory period, interventricular conduction time and increases the propensity of the atria to fibrillate by rapid atrial pacing. Pulmonary hypertension develops in up to 10% of patients with unrepaired ASD, and ES is a rare complication. In our case, the patient developed moderate pulmonary hypertension and ES.

The 2020 ESC guidelines on the management of ACHD states that in patients with shunt lesions and pulmonary hypertension, the closure of shunt lesion remains a matter of debate. A decision for surgery should be preceded by a meticulous evaluation, including invasive measurement of the pulmonary vascular resistance (PVR). In patients with ES, pulmonary hypertension and desaturation with exercise the closure of the lesion should not be performed.

Considering the late diagnosis of this malformation and the ES, and the lack of invasive measurement of PVR, surgical repair is not a solution for this patient. Long term optimal therapy is that of a chronic HF and AF. Also, use of selective pulmonary vasodilators and diuretics can be effective for symptom management in the setting of pulmonary hypertension with significant RV volume loading and also beneficial in ES.

CONCLUSION

We wish to highlight that multiple cardiac congenital anomalies can be diagnosed in elderly patients, even with a prior history of acquired heart disease as a cause for or contributor to HF. When a dilated CS is seen, PL SVC should be sought, as they may contribute to risk of atrial arrhythmias. A careful assessment of the entire interatrial septum using multimodality cardiac imaging is warranted for patients with right heart enlargement, in particular if a PL SVC is present, to evaluate other potentially associated malformations, and to determine adequate therapy.