Cor triatriatum dexter (CTD) is a rare congenital heart disease resulting from persistence of the right valve of the sinus venosus. The persistent valve forms a membrane that divides the right atrium into a proximal and a distal chamber. This disorder exhibits varying clinical manifestations depending on the degree of partitioning or septation of the right atrium. In asymptomatic patients, the disease may be discovered during surgical procedures, diagnostic testing such as echocardiography, or hemodynamic monitoring. Severe septation abnormalities may cause right-sided heart failure and elevated central venous pressures due to obstruction of the tricuspid valve, right ventricular outflow tract, or inferior vena cava. Here we report a case of CTD in a patient presenting with symptoms of dyspnea and fatigue, followed by a short discussion of the embryology and clinical implications of this congenital disease, as well as current advances in management.

**Keywords:** Cor triatriatum dexter; Right-sided heart failure; Congenital heart disease

**Introduction**

Cor triatriatum dexter (CTD) is an extremely rare congenital heart disease that results from the continued presence of the right valve of the sinus venosus. This valve forms a membrane that divides the right atrium into a proximal and a distal chamber [1]. This disorder has varying clinical manifestations, from asymptomatic disease to severe right-sided heart failure and elevated central venous pressures.

A 70-year-old Asian male presented to the hospital with worsening dyspnea on exertion and severe fatigue. He also reported an “issue with his heart” and a murmur which had been present since childhood. The patient noted absence of chest pain, orthopnea, or palpitations. He had a past medical history consistent with essential hypertension and cataracts but denied any history of diabetes, dyslipidemia or smoking. Family history of cardiovascular-related disease was non-existent.

On examination, his vitals were stable. The patient weighed 45 kg with a body mass index (BMI) of 17.2 kg/m². Cardiovascular exam was significant for a 4/6 systolic murmur with a palpable thrill best heard at the left lower sternal border and a right ventricular heave.

Transthoracic echocardiography was significant for a left ventricular ejection fraction (LVEF) of 65%, as well as moderate to severe mitral and tricuspid regurgitation (Figs. 1, 2). Transthoracic echocardiography revealed a normal left ventricular size and systolic function with an LVEF of 55-65%. It also showed evidence of a septum within the right atrium diagnosed as CTD. The mitral valve had moderate to severe regurgitation and the tricuspid valve had severe regurgitation. There was an intra-atrial septal aneurysm with a small patent foramen ovale and a right-to-left shunt (Figs. 3, 4).

Left heart catheterization was significant for a vertically rotated heart. He had several redundant large caliber vessels in the left circumflex and right coronary artery, in addition to a large caliber wrap-around left anterior descending coronary artery with no stenosis noted in these arteries, just luminal irregularities.

The patient was discharged to be followed up in the cardiology clinic for continued evaluation of his valvular disease and cor triatriatum. Together, they appeared to be the main contributors to his fatigue and shortness of breath. In contrast, his coronary arteries were large in caliber, with no disease that could be attributing to his symptoms. The patient would require monitoring for his mitral and tricuspid regurgitation with the possibility of surgical/percutaneous valve repair if deemed necessary in the future. He would also require monitoring for right ventricular failure secondary to CTD with the possibility of surgical resection/percutaneous disruption of the membrane if needed.

**Discussion**

CTD results from partitioning of the right atrium to form a tria-
trial heart. Unlike cor triatriatum sinister (left atrial septation), which is more common, CTD is an extremely rare congenital anomaly. It has an incidence of approximately 0.1% of congenital heart malformations [2].

During embryogenesis, the right horn of the sinus venosus gradually incorporates into the right atrium to form the smooth posterior portion of the right atrium, whereas the original embryologic right atrium forms the trabeculated anterior portion. The right valve of the right horn of the sinus venosus divides the right atrium into two. This right valve forms a sheet that serves to direct the oxygenated venous return from the inferior vena cava across the foramen ovale to the left side of the heart during the fetal life [1]. Normally, the valve regresses at approximately 12 weeks gestation and leaves behind the crista terminalis proximally, and the eustachian valve of the inferior vena cava and the thebesian valve of the coronary sinus distally. Complete persistence of the right sinus valve results in a separation between the smooth and trabeculated portions of the right atrium, resulting in

Figure 1. Transthoracic echocardiography in an apical four-chamber view revealing a left ventricular ejection fraction of 65% and a possible septation abnormality within the right atrium.

Figure 2. Transthoracic echocardiography in an apical four-chamber view with color Doppler revealing moderate to severe tricuspid regurgitation.
CTD. The proximal portion receives the venous blood from both vena cavae and the distal portion is in contact with the tricuspid valve and the right atrial appendage [3].

CTD can present in a myriad of forms depending on the degree of partitioning or septation/fenestration of the right atrium. When the septation is mild, the condition is often asymptomatic, and is found incidentally during surgery to correct other cardiac abnormalities or during echocardiography. More severe septation can cause right-sided heart failure and elevated central venous pressures due to obstruction of the tricuspid valve, the right ventricular outflow tract, or the inferior vena cava. CTD can occur as an isolated cardiac anomaly or

Figure 3. Transesophageal echocardiography at the level of the aortic valve with the scope rotated to 54° showing evidence of a septum within the right atrium diagnosed as cor triatriatum dexter.

Figure 4. Transesophageal echocardiography at the level of the aortic valve with the scope rotated to 29° during a bubble study, revealing an intra-atrial septal aneurysm with a small patent foramen ovale and a right-to-left shunt as evidenced by the bubbles within the left atrium.
associated with other malformations of right heart structures, including pulmonary artery stenosis or atresia, tricuspid valve abnormalities, atrial septal defects, and/or Ebstein anomaly. Cyanosis, however, is a very rare presentation of this condition. Even though our patient had an accompanying patent foramen ovale, this was small in size and would not have resulted in significant right to left shunting or cyanosis.

Detection of CTD is often based on echocardiography. However, magnetic resonance imaging (MRI) was previously found to have a higher detection rate (95%) when compared with echocardiography and cardiac angiography in the evaluation of pulmonary venous anomalies, which included cases of cor triatriatum (69% for angiography and 38% for echocardiography) [4-7]. Further studies are needed to evaluate detection rates between transthoracic and transesophageal echocardiography.

Right heart catheterization may reveal elevated pressure in the proximal right atrial chamber. It is also capable of detecting a gradient across the accessory membrane. One proposed complication with right heart catheterization is the entrapment of the guidewire or catheter within the fenestrations. Therefore, care must be paid with procedures within the right atrium of such patients. Other complications include misplacement, or leftward bowing of atrial devices for management of concomitant atrial septal defects. In addition, proper deployment and device stability might be prevented, overlap of the valve remnant, and under- or oversizing of devices could occur. This is due to the CTD membrane obstructing the view and preventing adequate definition of the margins of an associated atrial septal defect [8, 9].

Asymptomatic patients are generally not treated unless they are undergoing cardiac surgery for other reasons. In symptomatic patients, control of fluid retention and rate-control of atrial arrhythmias (e.g. atrial fibrillation) may be required. Inotropic agents may be needed in patients with severe right sided heart failure. In the past, the mainstay of treatment for symptomatic patients was surgical resection of the dividing membrane [7]. Recently, however, percutaneous catheter disruption of the membrane has been suggested as a preferred alternative to open heart surgery [10, 11].

Conclusion

CTD is extremely rare; however, this diagnosis should be considered in patients presenting with symptoms of right-sided heart failure. Due to the association it has with right-sided congenital heart abnormalities, it should also be considered prior to interventions within the right atrium. CTD can be easily corrected percutaneously or by surgical excision of the membrane.

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Conflict of Interest

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Informed Consent

Informed consent was obtained.

Author Contributions

GH: drafting article, writing manuscript; JS: writing manuscript, obtaining consent; SI: obtaining appropriate images and interpretation of images; PH: correspondence, general supervision, manuscript editing and manuscript review.

Data Availability

The authors declare that data supporting the findings of this study are available within the article.

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