Permanent transvenous pacemaker implantation in a patient with Cor triatriatum dextrum

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Abstract

Cor triatriatum dextrum is an extremely rare congenital heart abnormality in which the right atrium is separated into two chambers by a persistent fibrous membrane. A transvenous approach to place a dual-chamber pacemaker in such patients is technically challenging. We report the first case of a transvenous permanent pacemaker placement in a patient with cor triatriatum dextrum. An 87-year-old woman was diagnosed with paroxysmal atrial fibrillation. She was accidentally found to have cor triatriatum dextrum during the transesophageal echocardiography (TEE) prior to cardioversion. Later during her hospital stay, it was indicated to place a permanent pacemaker due to high grade atrioventricular block. After thorough reviewing TEE imagings, a transvenous catheter-based approach was decided feasible. Patient successfully received a dual chamber pacemaker through left subclavian venous approach. Furthermore in our case, using specially designed pacemaker leads and cautious intra-procedural maneuvering under fluoroscopic guidance ensured procedural success. In summary, a thorough pre-operative evaluation with transesophageal echocardiogram is critical for the planning and eventual success of the transvenous placement of right-sided leads.

Key words: Congenital heart defect; Complete heart block; Inter-atrial membrane; Dual-chamber pacemaker

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Core tip: Cor triatriatum dextrum is an extremely rare congenital heart abnormality in which the right atrium is separated by a persistent fibrous membrane. This membrane poses a technical challenge for dual-chamber pacemaker placement through the transvenous approach. Here we report the first transvenous pacemaker placement in a patient with cor triatriatum dextrum. A thorough pre-operative evaluation by transesophageal echocardiogram was critical for the planning of transvenous catheter-based right-sided leads placement. Using specially designed pacemaker leads and cautious...
CASE REPORT

An 87-year-old woman presented to the hospital with complaints of dyspnea on exertion. She has no significant medical history other than prior cigarette smoking. Upon her arrival to the emergency room she was found to be in atrial fibrillation with rapid ventricular response. She was treated with heart rate control medications and anticoagulation. A transesophageal echocardiogram (TEE) was performed prior to cardioversion to normal sinus rhythm given poor rate control. This revealed the presence of a well-defined transverse membrane dividing the right atrium (Figure 1) consistent with the diagnosis of cor triatriatum dextrum. Color Doppler evaluation indicated separation of blood flow in the two divisions of the right atrium. However the division of the right atrium by this membrane was not complete. Color Doppler confirmed partial obstruction in the superior sinus venosus valve. It is uncommon to encounter such patients requiring placement of a permanent pacemaker. We report the placement of a dual-chamber pacemaker through the transvenous approach in a patient with cor triatriatum dextrum. To our knowledge, there are no similar reports in the published literature.

DISCUSSION

The prevalence of adult congenital heart disease has increased in the past 10 years and its management has proposed new challenges to current cardiology practice. We presented a case of successful pacemaker lead placement in a patient with non-obstructive cor triatriatum dextrum. Cor triatriatum dextrum is an extremely rare congenital heart abnormality in which the right atrium is separated into two chambers by the persistence of the right sinus venosus valve. The superior chamber receives the venous blood from both vena cava and the inferior chamber is in contact with the tricuspid valve and the right atrial appendage. The size of the communicating orifice between the superior and inferior atrial chambers determines the natural course of cor triatriatum dextrum. If the communicating orifice is small, the patient shows symptoms of congestive heart failure during infancy or childhood and usually requires surgical intervention for survival. If the connection is large and non-obstructive, patient may remain asymptomatic for many years, as in our case. The clinical presentation therefore is somewhat variable. Patients with cor triatriatum dextrum may present with recurrent supraventricular tachycardia, right-side heart failure, or cyanosis in the presence of ASD with right-to-left shunt. In our case, patient presented initially with atrial fibrillation with rapid ventricular response. The disorder can be treated surgically in symptomatic patients by

intra-procedural maneuvering under fluoroscopic guidance ensured procedural success.

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removing the membrane dividing the atrium.

During normal embroygenesis, the right atrium is formed by two different portions joining together: the right horn of the sinus venosus that forms the smooth posterior portion, and the original embryologic right atrium that forms the trabeculated anterior portion. The connection between these two portions is called sinoatrial orifice. The sinoatrial orifice is sided by two valvular folds that are called the right and left venous valves. During the development of right atrium, the right valve of the right horn of the sinus venosus forms a membranous valve that divides the right atrium in two parts. This valve directs oxygenated venous return from the inferior vena cava across the foramen ovale to the left side of the heart. This membranous valve normally regresses by the 12th week of gestation. Incomplete regression of the superior portion of right venous valve forms membranes attached to the crista terminalis, while remnant of the inferior portion results in the Eustachian valve of the inferior vena cava, web-like remnant as Chiari network or the Thebesian valve of the coronary sinus. Failure of regression of this membrane causing persistent partition between the venous (smooth) and trabeculated portions of the right atrium leads to the formation of cor triatriatum dextrum.

Cor triatriatum dextrum has been associated with other congenital abnormalities, including ASD, patent foramen ovale, ventricular septal defect, hypoplastic right ventricle, hypoplastic tricuspid valve, and pulmonary atresia. In our case, the presence of a small ASD posed the risk of crossing into the left atrium during manipulation of the lead across the atrial membrane. We also found an enlarged coronary sinus, measured diameter 15.8 mm (normal range 6.6 ± 1.5 mm). This is likely related to the elevated right atrial pressure. A rare but important congenital vascular anomaly associated with an enlarged coronary sinus is a persistent
left superior vena cava draining into the coronary sinus. To investigate this possibility, an agitated saline injection via the left arm was performed during TEE and showed no saline contrast in the coronary sinus, indicating the absence of a persistent left superior vena cava.

Considering the rarity of cor triatriatum dextrum, a patient with such a congenital abnormality who requires a permanent pacemaker is unique. To our knowledge, this is the first reported case of a permanent pacemaker placement through a transvenous approach in a patient with cor triatriatum dextrum. Although it is technically challenging, transvenous catheter-based approach is feasible if the membrane in the right atrium is non-obstructive and caution exercised during the procedure. A thorough pre-operative evaluation by transesophageal echocardiogram was critical for the planning of the transvenous catheter-based right-sided leads placement. Using specially designed pacemaker leads and cautious intra-procedural maneuvering under fluoroscopic guidance ensured procedural success.

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