Pacemaker Lead Associated Superior Vena Cava Syndrome Treated By Veno-Atrial Bypass

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Abstract

Pacemaker lead associated venous thrombosis is common. However, due to collateral venous drainage those who are affected rarely become symptomatic. Superior vena cava (SVC) syndrome is a constellation of signs and symptoms that occurs as a result of obstruction to flow in the superior vena cava. In this case report we present a patient who developed this complication following multiple transvenous pacemaker wires. After failed percutaneous venoplasty he was treated successfully with a right internal jugular vein to right atrial appendage bypass.

Keywords

SVC syndrome, Pacemaker leads, Lead associated thrombosis

Case Report

A 57-year-old male patient who was on a permanent pacemaker due to complete heart block presented with slowly progressive swelling of the face and neck. He had severe congestion of the head and neck area when bending down. His first pacing device was implanted in 1992. Due to malfunction, the device was revised twice, in 2005 and 2012. At each revision the existing pacing wires were not extracted.

Physical examination revealed engorged neck veins. Upper limbs were not oedematous. There were no prominent venous collaterals over the anterior chest wall. His basic blood investigations were normal and the recent 2D echocardiogram showed good biventricular function. Chest X-ray confirmed the presence of multiple pacemaker wires (Figure 1).

According to the clinical findings SVC obstruction was suspected. A CT venogram was done which showed dilated bilateral internal jugular, superior intercostal and azygos veins. The SVC was patent with no definite level of obstruction. However, as the clinical suspicion was high the patient was started on warfarin and oral frusemide. Even though medical therapy for continued more than 6 months, the symptoms progressed to a point where he had to sleep seated due to unbearable congestive symptoms on lying flat.

Figure 1: CXR with multiple pacemaker wires.
common. It has an incidence of 30-64% and the affected majority are asymptomatic [1,2]. However pacemaker leads causing full blown superior vena cava (SVC) syndrome is rare. It occurs in < 0.1% [3].

The first recorded case of SVC syndrome was by William Hunter in 1757 [4]. The syndrome encompasses a collection of signs and symptoms due to obstruction of blood flow through the SVC [5]. Intrathoracic malignancy is the leading cause for this syndrome and accounts for > 90% of all cases. Historically thoracic aortic aneurysms and mediastinal fibrosis following infection were the commonest benign etiologies. At present iatrogenic SVC syndrome has assumed a more prominent role due to increasing use of intravascular devices such as central venous lines, cardiac pacemakers and implantable cardiac defibrillators [4].

Endothelial injury due to mechanical trauma induced by pacemaker wires triggers the thrombotic process within the vein. Fibrin deposits on the leads and an inflammatory reaction within the vein wall itself contributes to the thrombosis. The end result is total occlusion or fibrotic stenosis of the SVC [6]. The disease process is fairly slow so it allows adequate time for collaterals to develop. Patients become symptomatic when adequate collateralization does not occur [4].

Risk factors for SVC thrombosis in a patient who has

Discussion

Pacemaker lead associated venous thrombosis is
undergone transvenous pacing are thrombophilic conditions, lead infection, use of hormonal contraceptives, use of a temporary pacing wire before implantation of the permanent device and the presence of multiple active or non-functional pacemaker leads [7]. Our patient had multiple leads within the SVC due to repeated device revisions.

Symptomatic patients with SVC syndrome present with swelling of the face, neck and upper limbs, shortness of breath and orthopnea. Headache, dysphagia, dizziness are other infrequently reported symptoms. The symptoms are the result of impaired venous drainage of head and neck region and upper limbs due to the obstruction in the SVC [4]. Symptoms typically become worse when bending over and lying down. This was noted in our patient as well.

When the diagnosis of SVC syndrome is clinically suspected it should be confirmed by imaging to delineate the location, cause and the extent of the blockage. Chest X-ray is not specific and may show a mediastinal mass in those with obstruction due to tumor or aneurysm [4]. CT venography is the first choice of imaging at present and is mandatory to exclude or characterize extrinsic compression of the SVC. Contrast venography is considered the gold standard of imaging but it is invasive. When there is a discordance between clinical findings and CT images as in our case, contrast venography has a place [7]. Magnetic resonance venography is also an imaging option provided the pacemaker device is MRI compatible.

Symptomatic patients with pacemaker lead associated SVC syndrome require treatment and the options are medical management, percutaneous venoplasty or surgery [8]. Medical management includes anticoagulation to prevent clot propagation, diuretics for relief of peripheral oedema, steroids for cerebral/laryngeal oedema and head end elevation of the bed [4]. Those with persistent symptoms despite conservative therapy will need invasive treatment.

The place for routine extraction of non-functional transvenous pacemaker leads in the absence of infection is controversial. The procedure has an associated mortality of 0.4-0.6% and a morbidity of 1-2% [9]. Lead associated SVC syndrome is considered an indication for extraction of non-functional, superfluous pacemaker leads [9,10]. Removal of the leads per se will not restore the venous patency, so it is unlikely to result in adequate symptom relief [2]. However the channel created in the process of extraction may provide a valuable pathway for a new lead implantation in those with a nonfunctional device and limited access [2,9]. When the gap between the lead implantation and the attempted extraction is prolonged, the lead tends to be well adherent to the vein wall by scar tissue, and pulling the lead out risks damage to the vein wall [2,11]. Considering the fact that some of the leads in our patient was implanted > 2 decades back and as the current device was functioning well, we decided it was unwise to attempt removal.

Percutaneous intervention with balloon venoplasty with or without stenting is the preferred first line intervention for SVC syndrome at present [3,5]. According to Sfyroeras, et al. 80% of those who were treated percutaneously required a stent [5]. Thrombolysis can be used as an adjunct before venoplasty in selected cases. Percutaneous interventions in the presence of pacemaker leads can be technically challenging. There is a risk of lead damage and device malfunction which may be life threatening. Retrieval of leads if they get infected becomes difficult when a stent is deployed over them. Covering the procedure with temporary pacing wires, extracting and replacing the leads after the venoplasty and stenting are done are steps that can be taken to circumvent these problems. Patients are usually put on long term anticoagulation therapy following percutaneous treatment [7].

Surgery in the form of bypass grafting has a secondary role at present. It is considered when percutaneous venoplasty fails or when there are complex obstructive lesions not amenable for venoplasty [5]. Native vein grafts, vein allografts, aortic allografts, pericardial tubes and prosthetic grafts have been used as conduits. The long term patencies of native vein grafts are superior to prosthetic grafts. For technical success of bypass grafting size of the conduit should match the inflow vein and the conduit should be of optimum length to avoid redundancy [4]. Patients will need long term anticoagulation after open surgery in order to maintain patency in a low flow bypass.

Short and medium term patency rates after open and after endovascular treatment for SVC syndrome are comparable, but long term patency is better after open surgery. Avoiding a sternotomy is the major advantage of percutaneous intervention over open surgery [5].

Conclusions

Pacemaker lead associated SVC syndrome is rare. Those with debilitating symptoms despite conservative management will need intervention. Percutaneous balloon venoplasty with or without stenting is the first line approach and surgery assumes a secondary role. Patients will need long term anticoagulation therapy after either form of intervention.

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Conflicts of Interest

None.

Authors Contribution

Thilina Gunawardena designed the manuscript. Thi-
Ilina Gunawardena, Balasubramanium Saseekaran and Gayan Bandara wrote the manuscript. Rezni Cassim and Mandika Wijeyaratne supervised the project.

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