Case Report

Intracardiac Mass Due to Fibrosing Mediastinitis: The First Reported Case

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ABSTRACT

We report a case of a woman, aged 53 years, presenting with a right atrial mass due to idiopathic fibrosing mediastinitis with periaortic involvement. This challenging diagnosis was confirmed by different imaging modalities and histopathologic analysis. The diagnosis of cardiac tumors is often difficult. To our knowledge, this is the first reported case of an intracavitary cardiac mass due to fibrosing mediastinitis. This rare disorder, which is characterized by invasive proliferation of fibrous tissue within the mediastinum, should be included in the differential diagnosis of intracardiac tumors.

The diagnosis of cardiac tumors is often challenging because of their rarity and diverse etiology. To our knowledge fibrosing mediastinitis (FM) has not yet been reported as a cause of an intracavitary cardiac mass. We report a case of a right atrial mass and periaortic involvement due to this unexpected condition.

Case Report

A woman, aged 53 years, with no relevant past medical history, except for mild hypertension, went to the emergency department because of retrosternal pain radiating to the back, which had started 2 weeks before. The electrocardiogram, chest radiograph, and blood analysis were normal, and she was discharged. She remained asymptomatic, but a transthoracic echocardiogram revealed an unexpected round, echodense, and heterogeneous 20 × 14-mm right atrial mass, without significant mobility, distinct from the valvular plane, in addition to a thickened aorta. The chest-abdomen-pelvis computed tomography scan showed a partially calcified circumferential thickening of the ascending aorta extending to the right atrioventricular groove, where it continued with the atrial mass, in addition to mediastinal lymphadenopathy mainly in the right paratracheal group. The cardiac magnetic resonance imaging further characterized the periaortic and atrial masses, which were in continuity (Fig. 1; Video 1, view video online). The hemogram, coagulation, biochemistry, autoimmune screening, and microbiological examinations were normal. Because of the possibility of malignancy, a percutaneous atrial mass biopsy via jugular approach, guided by a transesophageal echocardiogram, was performed (Video 2, view video online). The histopathology revealed marked fibrosis, but malignancy could not be excluded. A video-mediastinoscopy attempting to biopsy mediastinal lymphadenopathy was unsuccessful because of extensive mediastinal fibrosis. Finally, a median sternotomy was performed, and a nonvascularized white solid mass involving the ascending aorta was identified. An extensive sampling of the periaortic tissue was performed, and the histologic and immunohistochemical analysis confirmed the diagnosis of FM (Fig. 2). The periaortic tissue was identical to the right atrial mass tissue, confirming that the atrial mass, which was in continuity with the periaortic mass, was also due to FM. Both samples

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revealed no bacteria or fungi (Gram and periodic acid-Schiff staining, tissue culture) or increased number of IgG4-positive cells. The Histoplasma serology and the tuberculin skin test were normal. Systemic corticosteroids were started, considering it was an idiopathic presentation. Nine months later, the patient is still on corticosteroids and asymptomatic. The periaortic and atrial masses are smaller (Supplemental Fig. S1; Video 3, view video online).

Discussion

A cardiac mass detected on transthoracic echocardiography can be further characterized by other imaging techniques, but histologic examination is usually necessary to establish a definitive diagnosis, as it was in this case. The dimension of the atrial mass sample obtained percutaneously was insufficient to exclude malignancy, as previously reported. Thus, an extensive sampling via sternotomy was performed, confirming this challenging diagnosis.

FM is a rare benign disorder characterized by invasive proliferation of fibrous tissue within the mediastinum. It commonly involves the superior vena cava, pulmonary vessels, tracheobronchial tree, or esophagus, and the clinical presentation is usually related to obstruction or compression of these structures. Less-frequently-involved structures include aorta and aortic branch vessels, pericardium, coronary arteries, lymph nodes, nerves, neck, thyroid, pleura, and spinal cord, and FM may be associated with retroperitoneal fibrosis.

FM has an unpredictable but often benign course. The main treatment options are systemic antifungal agents or corticosteroids, surgical resection, and local therapy. The response to corticosteroid therapy is highly variable; while many patients do not improve, there are cases of effective treatment, particularly in idiopathic cases or cases with an active inflammatory process. Surgical and local interventions are usually reserved for symptomatic patients for relief of compression. It was decided not to excise the atrial mass, because the patient was asymptomatic and the excision could be associated with increased morbidity and mortality, particularly in this case, where excision would be technically difficult without extreme wall replacement. In addition, the
atrial mass was not likely to embolize or cause obstruction, given its nonfragile appearance, high adherence, limited mobility, and the small dimensions. Moreover, the atrial and periaortic masses diminished with corticosteroids. The role of anticoagulation is not yet established.

To our knowledge, FM has not yet been reported as a cause of intracardiac tumours. This condition should be included in the differential diagnosis of intracardiac masses.

Disclosures

The authors have no conflicts of interest to disclose.

References


Supplementary Material

To access the supplementary material accompanying this article, visit the online version of the Canadian Journal of Cardiology at www.onlinejc.ca and at http://dx.doi.org/10.1016/j.cjca.2012.11.020.