Intravascular Leiomyomatosis of the Lower Vein Cava Extended to the Right Atrium

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Summary

Intravenous leiomyomatosis is a rare condition that presents extension uterine gonadal veins entering the inferior vena cava to the right heart chambers, patients are usually asymptomatic and cause significant symptoms for pulmonary embolism or valve lesions by intracardiac obstruction. With high mortality in these cases.

Case: 38-year-old who presents with abnormal uterine bleeding, within their preoperative studies; transthoracic echocardiogram evidenced by mass in the right atrium, mobile that is inserted into the right ventricle; possible origin of atrial wall. It takes abdominal CT angiography which reports mass extending from the left iliac vein and travels through the inferior vena cava to the right atrium. It is carried in a first cardiac surgery with no evidence sternotomy thereof in any of the heart chambers. It is carried back a second surgical time when hysterectomy is performed, and exploration of the left common iliac vein is performed; showing soft dough, white, mobile color. Simultaneously transesophageal echocardiogram where the mass is reaching the right atrium and retracts into the inferior vena cava is performed. It is possible to remove intravascular mass of approximately 18 cm in length. Echocardiography is performed with no evidence of residual mass. Pathology report which is positive for intravascular leiomyomatosis. Adequate postoperative course.

Conclusion: This is an important case because of its rarity and operation set of various specialties and complexity of the surgical management with adequate postoperative course.

Keywords: Intravascular Leiomyomatosis; Right Atrium

Introduction

Intravascular leiomyomatosis is a rare condition that consists of proliferation of the smooth muscle which extends from the gonadal veins and can create large masses [1,2] with invasion from the mimes with a path from them; passing through the inferior cava vein and with compromise of the cardiac cavities [3], this can occur in 45% of the reported cases, it may be associated with trombosis of the adjacent tissue [4,6].
Additionally, the symptomatology presented by the patients is not clear; this type of mass is usually evidenced incidentally [4, 5]. At the moment in which the patients present symptoms they are related to the complications which are severe given by massive pulmonary embolisms; Tricuspid valve obstruction and dysfunction [5, 7].

The multidisciplinary diagnosis as well as the surgical management which and in the cases reported is made in two stages; having as a first intervention the resection or cardiac examination and after this the joint performance of hysterectomy and definitive intravascular resection by abdominal route as the one performed in our case.

We present the following case report of intravascular leiomyomatosis.

**Clinical Case**

This is a 38-year-old female patient who consults due to a clinical picture of abnormal uterine hemorrhage. She is evaluated by the gynecology department who programs for hysterectomy; within pre-surgical studies and taking into account the multiple comorbidities of the patient. Echocardiogram after thoracic is performed, showing mass of regular edges in the right atrium of approximately 8 cm x 6 cm, mobile; (Figure 2), of possible neoplastic origin adhered to the atrial wall. After this, abdominal angiotac is taken, where there is evidence of mass that occupies intravascular space from the right common iliac vein extending through the inferior vena cava and reaching the right atrium (Figure 1).

![Figure 1: Abdominal CT scan showing mass in the Inferior cava vein to the right atrium.](image_url)
In the first stage, cardiac surgery is performed by performing sternotomy and exploration of the right atrium without evidence of such mass, or signs of intracardiac thrombosis. It is evaluated with a new echocardiogram finding persistence of the atrial mass without another significant sign. It is carried out to a second surgical time by abdominal route in a multidisciplinary way by oncological gynecology for hysterectomy and subsequent exploration in a controlled way of the left common iliac vein; simultaneously, it is evaluated with transesophageal echocardiography to monitor possible emboli and documentation of intravascular mass travel. Complete extraction of the dough with a length of approximately 18 cm in a cylindrical shape is achieved. (Figure 3). The complete extraction and non-occupation of the cardiac cavities are evaluated with echocardiography. Another fragment is extracted from one of the gonadal veins of the same characteristics.

Patient with adequate postoperative evolution and is in controls.

Final pathology report which shows and confirms intravascular leiomyomatosis (Figure 4).
**Discussion**

Intravascular leiomyomatosis is an entity derived from uterine myomatosis which grows with invasion into the vessels, it is a proliferation of smooth muscle cells. This tissue proliferation is benign. The first cases were described in 1975 by Parmley [1, 3, 6]. Because it is a rare entity, it is not known what its etiology is. It should be taken into account that the complications given by massive pulmonary embolization and cardiac lesions caused by mass mobilization; they are severe and cause the death of patients [1, 5, 7].

The clinic of the patients is nonspecific given by dyspnea, syncope, arrhythmias, dysmenorrhea in most cases and as ours are found incidentally [4, 5, 8].

The treatment is surgical, the complete extraction of the intravascular mass must be done due to the high embolic irrigation [3]. In our case, a first surgical time was performed as described with intracardiac examination to extract it; entering through the right atrium. A second time by abdominal route to perform hysterectomy and controlled venous exploration, complete extraction of the mass and verification by cardiology with echocardiography and to be certain of complete resection and non-pulmonary embolization of the same [2, 3].

The management of this pathology should be done interdisciplinarity given the complexity. In our case, an intervention was carried out by the services of oncological gynecology, cardiology and cardiovascular surgery which led to the intravascular and cardiac extraction of the mass [3, 5].
**Conclusion**

It is a complex pathology with a low incidence, multidisciplinary management, with a nonspecific clinic and mostly incidental diagnosis. In this way it shows an important challenge for the diagnosis and definitive management. High mortality due to pulmonary and cardiac complications must be taken into account.

**References**
