Scimitar Syndrome: Infra-Diaphragmatic Form a Novel Surgical Approach

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Background

Scimitar syndrome is a complex malformation in which the main feature is a partial anomalous venous drainage of the right side toward the inferior vena cava or the portal system. There are anomalies associated in bronchopulmonary segmentation and vascular lung connections [1]. In 1836 in specimens of autopsy Cooper in London published the first description and later Chassinat in Paris [2, 3]. In 1949 Dotter et al through angiocardiography shown the first clinical diagnosis where noted the particular appearance of the anomalous vein on x-ray of thorax [4]. Later Neill et al in 1960 called the scimitar sign for describing this anomaly [5]. Drake and Lynch in 1950 performed the first surgical management of this disease [6]. Later the first physiological correction of anomalous venous drainage was published in 1956 by Kiklin et al [7]. Finally Neill CA et al. published the familial occurrence of Scimitar syndrome.

The objective of this report is to inform of a case successfully repaired using an exterior tube. We analyzed the current guidelines for diagnosis and management, particularly in the adult patient.

Clinical Case

Female 38 years of age, who presented a pulmonary murmur, it was detected at 13 years of age, without follow-up; in 2002 began to be coopered in study by suspicion of pulmonary hypertension, her principal symptom was dyspnea with Class II of NYHA. In 2015 reset study protocol for congenital heart disease and moderate pulmonary arterial hypertension. Laboratory: Hb 13 g/dL, Hto 32%, Leucos 6,000 mm3, platelets 203,000 mm3, Creatinine 0.59 mg/dL | BUN 12 µg/dL | Glucemia 88 mg/dL | INR 1.1, NT – PRO BNP: 235. Rx-Thorax and EKG are shown in Figure 1. The echocardiography report shown an anomalous pulmonary venous drainage and suggested a Scimitar syndrome Table 1. Surgical Findings: Cardiomegaly grade II at the expense of right cavities, aorto-pulmonary ratio 1:3, extra cardiac anomalous venous drainage with venous duct in “H” which empties into the inferior vena cava, correlation with scimitar syndrome, right atrium dilated without atrial septal defect, without presence of any pulmonary drainage Figure 2, 3. A Dacron tube was used as extra cardiac conduit to make surgical repair Figure 4. After unclamping patient presented ventricular fibrillation which required three defibrillation shock of 10, 20 and 20 Joules reaching a normal rhythm; Cardiopulmonary Bypass time: 181 minutes; bleeding: 320 ml; recovered: 259 ml; blood products: one globular package and two fresh frozen plasma. After a week she was discharged without any complication. An angio-tomography performed later was able to shown the final result Figure 5.

Table 1: Echocardiogram parameters

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
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<tbody>
<tr>
<td>LA</td>
<td>33x31x44mm</td>
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<tr>
<td>RA</td>
<td>41x48x46mm</td>
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<tr>
<td>LVEF</td>
<td>64%</td>
</tr>
<tr>
<td>Septum</td>
<td>10 mm</td>
</tr>
<tr>
<td>Posterior Wall</td>
<td>10 mm</td>
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<tr>
<td>TAPSE</td>
<td>18</td>
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<tr>
<td>Systolic Pulmonary Pressure</td>
<td>75 mmHg</td>
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Discussion

Developing early pulmonary venous sinus communicates through small channels with systemic veins of the embryo. This primitive drainage disappears when the common pulmonary vein of the left atrium is connected with such senous of this vein Agenesia determines the persistence of the first channels which causes different types of anomalous connection [8, 9].

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Figure 1: A: RX Chest: Describes the typical image of a scimitar in the right side margin
B: EKG Sinus rhythm, AQRS +60°, Growth of right cavities and systolic overloading, growth of right atrium

Figure 2: A: Angiotomography; B & C: Third dimension Angiotomography, where the passage of the drainage through the diaphragm is shown, arriving at the suprahepatic vein, which is an event that takes place in very rare form; D: Cardiac catheterization. No cyanogen Congenital Heart Disease with anomalous venous drainage of the Pulmonary veins; Right upper and lower Pulmonary veins draining toward a collector (rear, right), which drains into the union of the inferior cava vein with right atrium; Left Pulmonary veins well-connected; LV normal with LVEF 60%, LVDEP 10 mmHg; Mitral valve; Aortic components | Left Aortic Arch | Origin and normal distribution of coronary arteries; Dilated Pulmonary arch, Severe pulmonary hypertension (90 mm Hg) which descending to 70 mmHg after oxygen supplement. Arrow : right pulmonary branch (scimitar)
Figure 3: PA: Pulmonary Artery; AO: Aorta; SCV: Superior cava vein; RA: right atrium; ICV: Inferior Cava Vein; C: Scimitar, RV: Right Ventricle, LR: Round Ligament.
A & B: Anomalous vein throwing diaphragm (arrow Scimitar)

Figure 4: SCV: superior cava vein, ICV: Inferior cava vein. IAS: inter-atrial Septum
A: Normal interatrial septum.
B & C: right atrium-Dacron graft anastomosis
D: Formation of Dacron hemi tube ceiling
Scimitar syndrome pulmonary venous sinus presents double connection, on the left with the left atrium and the right side with the supra-hepatic segment of the inferior vena cava, derived from the right vitelline vein of the embryo; This favors the separation of pulmonary venous sinus in two portions, left joins to the left atrium and right is continued with the curved manifold Figure 6.

The connection of the right pulmonary veins to level infra-diaphragmatic either to the vena porta or inferior vena cava types that deviate from the scimitar syndrome. Partial anomalous pulmonary venous connection to the right atrium without curved manifold and right pulmonary hypoplasia should be excluded from the scimitar syndrome.

Scimitar syndrome is classified in three groups according to the age of the patients as it was suggested in a multicenter study of 122 patients with ages between 1 and 58 years old. The Group I correspond to adult patients without pulmonary hypertension (PH) and with interatrial septal communication; group II, It has complex congenital abnormalities affecting the symptomatology and the natural history of this syndrome; Group III the child is characterized by severe and poor PH prognosis [10-13]. The most common symptoms present in all patients were breathing and progressive dyspnea [11, 12, 14]. In some cases the radiographic sign of the scimitar is not observed as a consequence of the cardiac dextrorotation or because of the venous collector is not curved or wide rather than straight, slim or multiple. This sign is presented by 70% of patients. It is due to hypoplasia of the right lung and it is absent in adults [10, 15].

The echocardiographic diagnosis had a good correlation with cardiac catheterization. Transthoracic echocardiography has more value in children than adults with this syndrome; and, in adults the diagnosis must be complemented with Transesophageal Echocardiogram to assess structures as the atrial septum position, the size, number, morphology and localization of the defects and the connection of the Pulmonary veins that cannot
be well valued by Trans thoracic technique. Trans esophageal echocardiography is also useful for detecting obstructions in tubes placed for the re-implantation as happened in one of the cases studied was corroborated by resonance magnetica [10, 13, 16].

It is very important to determine the levels of the supra connection for a better surgical management. A surgical alternative could be the re-implantation of the manifold to the rear wall of the left atrium without extracorporeal circulation, as Brown and collaborators carried out it [12, 14, 17-20].

**Conclusions**

On the basis of our case-study we concluded that scimitar syndrome is rare, whose diagnosis is performed in the majority of cases by echocardiography. With the development of techniques not only in some cases invasive catheterization diagnosis is complementary [18, 19]. A correlation anatomo-echocardiographic is useful because it shows the correspondence between the anatomical feature and its image diagnostic which confers a degree of precision to the diagnostic echocardiography, which is essential for the surgical decision, however computed, with three-dimensional reconstruction tomography images, allowing perfectly provides the surgery planning.

Surgical treatment of scimitar syndrome during the pediatric age, allows direct anastomosis of the right inferior pulmonary vein to the left atrium, referring cases without cardiopulmonary bypass [20]. During the adult stage the flexibility of the tissue makes it more difficult this situation it is necessary to interpose a Dacron graft as it was met in this case.

Another important particularity of this case was the small left atrium, so it was necessary to perform the anastomosis of the dacron tube into the right atrium, and subsequently open the atrial septum, and place a Dacron tube in dome-shaped, allowing re-direct the flow to the left atrium.

**References**