Giant Lipoma in Superior Vena Cava: A Case Report and Literature Review

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Abstract

Background: Intravascular lipomas were a rare occurrence, especially in major vessels. This tumor is composed of adipocytes in a fibrous capsule that had a slow growth rate and usually shows no symptoms. There were only eight reports in the literature regarding intravascular lipoma located in the superior vena cava.

Case presentation: A 54-year-old man had episodes of supraventricular tachycardia and atrial flutter for over a year. Radiological findings preoperative showed a giant mass that arose from superior vena cava to right atrium and biopsy catheter showed that there were no signs of malignancy. The patient then underwent surgery through a median sternotomy and the mass was extirpated on the highest part of the stalk that could be reached. The patient was stable and remains to show no symptoms or evidence of residual mass or stalk in 2 years follow-up.

Conclusion: The surgical approach in excising lipoma in SVC should be considered wisely with the support of adequate preoperative diagnostic. Extensive manipulation that could increase surgical technique difficulty or postoperative morbidity and mortality is not necessary since lipoma is a very slow-growing tumor.

Background

Primary benign tumors that originate from the intravascular wall are considered a rare occurrence. Among all of them, lipomas were found to be extremely rare, especially the ones that occur in superior vena cava (SVC).1 Currently there were only 30 pieces of literature found on PubMed by using search terms “intravascular lipoma”, “superior vena cava lipoma” and “intravenous lipoma”. There were only eight cases found regarding intravascular lipoma located in the SVC.2

Intravascular lipomas were composed of adipocytes in a fibrous capsule that had a slow growth rate. It usually shows no symptoms that it commonly diagnosed after an incidental finding.3 Even though only large-sized of this tumor could cause obstructive symptoms, many had believed that it better be surgically removed due to its probability of causing turbulent blood flow and subsequently thrombotic complication in the venous portal system.4

Case Presentation

A 54-year-old man had episodes of supraventricular tachycardia and atrial flutter for over a year. The patient was an ex-smoker with a history of hypertension, dislipidemia, and a family history of sudden cardiac death. The physical examination results were unremarkable. Transthoracic echocardiography (TTE) examination shows a large mass at right atrium (RA) with the size 4,7 x 3,6 cm, occupy more than half of RA chambers. The left ventricle is normal in size and function, with an ejection fraction of 73%. Other findings in TTE were normal. A computed tomography scan (CT) showed an elongated lesion with low density that arose from SVC to RA (Fig. 1a). Magnetic resonance imaging (MRI) examination
confirmed a big capsulated mass that arose from SVC to RA with the size of 12x4x4 cm (Fig. 1b). The mass was confirmed as a fat-rich content and diagnosed as lipoma. Given the size and patient's age, malignancy could not be excluded. Therefore the patient underwent biopsy by catheterization and the results showed that there were no signs of malignancy. The patient was prepared for extirpation and underwent catheterization. It was found that the left main artery had 20–30% stenosis on the distal, left anterior descending artery had 40–50% stenosis on the middle, left circumflex artery had 60–70% stenosis on the distal, and right coronary artery had 50% discrete stenosis on the proximal. The patient was diagnosed with a moderate 3-vessel disease of the coronary artery, and it was decided to be treated conservatively.

The median sternotomy approach was chosen and we found that SVC was 2 times bigger than the aorta. Under guiding TEE the mass found to be occupied the SVC, then we decided to canulate the internal jugular vein and IVC. Under a total bypass with a cardiopulmonary bypass machine, RA was opened and we found a yellowish mass inside the RA that originate from SVC. The mass has a stalk that origin from the cranial of SVC. The mass was yellowish with the size of 15x5x4 cm, had a lobulated surface, mobile, and had rubbery consistency (Fig. 2). We pulled the mass and extirpated it on the highest part of the stalk that could be reached. RA was closed and cardiopulmonary bypass was quickly discontinued without any problem, and the surgery was done smoothly. The patient was stable with normal sinus rhythm on ECG post-surgery and was discharged four days after.

Pathology examination shows mature white adipose cells with no-centrally located nuclei dominated this mass, thin fibrous septa in some parts, and a few blood vessels. These histologic findings confirmed the mass as lipoma. The patient underwent cardiac rehabilitation program after being discharged and remains to show no symptoms in 2 years follow-up. There is also no evidence of residual mass or stalk on the SVC on Cardiac MRI with four chamber and right ventricle two chamber stack 10 slices, slice thickness 6 mm that was performed extended to the cervical region on 2 years follow-up (Fig. 3).

**Discussion And Conclusions**

Lipomas are benign tumors that rarely occur intraluminal in major vessels, which most prevalent in people between 40–60 years. It usually shows no symptoms, but when present it usually shows obstructive symptoms of cardiovascular like congestiveness and edema. We only found eight cases of SVC lipoma from a literature search in PubMed (Table 1.). Four cases described that patients showed obstructive symptoms. In our case, the patient shows symptoms of periodical arrhythmia which has never been described in other cases even though the one that extended to the right atrium like ours. We assumed that the symptom was due to its position in RA and its gigantic size, therefore we decided not to do any invasive intervention to it. It was confirmed so that the arrhythmia was disappeared after the resection.
| Author (Year) | Gender/Age | Clinical Presentation | Prediagnostic modalities | Tumor Size | Surgical Approach |
|--------------|------------|-----------------------|--------------------------|------------|------------------|
| Vinnicombe S (1994)<sup>10</sup> | F, 42 y.o | Fatigue, edema face and right hand | **CT scan**: rounded mass of fat compressing proximal right brachiocephalic vein and SVC  
**Venogram**: large lobulated filling defect up to 3.5cm diameter in SVC | 10x5x5cm | not well described |
| Thorogood SV (1996)<sup>11</sup> | M, 73 y.o | Asymptomatic | **CT scan**: mass of fat density in SVC and the right brachiocephalic vein | not specified | no surgical intervention |
| Mordant P. (2010)<sup>12</sup> | F, 55 y.o | Asymptomatic | **CT scan**: intraluminal nonenhancing tumor occluding the distal right subclavian vein, the right brachiocephalic vein, and the SVC up to the right atrium  
**Venogram**: total occlusion of the right subclavian and brachiocephalic veins and of the SVC to the level of the azygos vein  
**MRI**: fatty intravascular lesion | 9x6cm | median sternotomy with right transclavicular cervicotomy. Transverse venotomy in SVC. En bloc resection, end-to-end anastomosis left innominate vein - SVC |
| Bravi MC (2011)<sup>4</sup> | M, 63 y.o | Abdominal, right shoulder, and lumbar pain | **CT scan**: superior vena caval (SVC) filling defect with a subtotal occlusion that extended into the right atrium. MRI: uniform signal drop on fat-suppressed sequences | not specified | not well described |
| Tanyeli O (2015)<sup>1</sup> | M, 48 y.o | Right arm edema and paresthesia | **CT scan and MRI**: fat density within SVC | 5x2cm | mini J sternotomy, venotomy |
| Author          | Gender/Age | Clinical Presentation | Prediagnostic modalities | Tumor Size | Surgical Approach                     |
|-----------------|------------|-----------------------|--------------------------|------------|---------------------------------------|
| Concatto NH     | M, 58 y.o  | Asymptomatic          | CT scan: a hypodense elongated lesion with fat density within the superior vena cava | 11 x 3 cm  | not well described                    |
| (2015)          |            |                       | MRI: confirmed the fatty nature of the lesion |            |                                       |
| Wahab A         | F, 70 y.o  | Asymptomatic          | TEE: 2.6x1.6x1.6 cm partially obstructing round, echogenic mass at SVC and RA junction | 2–3 cm     | No surgical intervention              |
| (2017)          |            |                       |                           |            |                                       |
| Sundaram N      | M, 58 y.o  | Asymptomatic          | CT scan: intraluminal 5 cm mass in the right innominate vein extending into SVC | 5 cm       | median sternotomy with right cervical extension, venotomy in SVC, counter incision in right mid-jugular vein |
| (2020)          |            |                       | Venous duplex: large pedunculated 5 cm hyperechoic mass at the junction of the right internal jugular and subclavian veins |            |                                       |
| Soetisna TW. Et al | M, 54 y.o | Episodes of SVT and atrial flutter | CT scan: elongated lesion with low density from SVC to RA | 15x5x4cm    | conventional median sternotomy        |
| (2021)          |            |                       | MRI: big capsulated mass from SVC to RA (fat-rich content) |            |                                       |

None of those eight cases underwent biopsy before the intervention. There were only a few articles about intravascular lipoma and there was no literature that shows the incidents of intravascular lipoma or liposarcoma. Despite it, there were data about the incidence of lipoma and liposarcoma originated from the heart that shows the rarity of the case (lipoma 0.07%-8.4%; liposarcoma 0.19%-0.5%). Nevertheless the rarity of malignancy incidence in the cardiovascular tumor, we still cannot exclude the possibility of malignancy, in this case, due to its size (the biggest lipoma ever been reported in SVC) and the age of the patient. Studies have shown cardiac MRI to be the gold standard diagnostic imaging modality for cardiac lipoma, but it has limited sensitivity that could only distinguish 69% of cases in the setting of well-differentiated liposarcoma. Given that malignant tumor originated from cardiovascular required different
consideration in treatment options, therefore we still encourage to do the biopsy before intervention to better weigh the risks and benefit of the surgical treatment.

In our case, the cardiac CT and cardiac MRI didn't specify the origin of the lipoma's stalk; it was fully described by Elen, et al.\(^9\) Given the uncertainty of the tumor origin, we decided to not performed any extensive manipulation due to its probability of increasing surgical technique difficulty and postoperative morbidity or mortality. Two years after, the patient remains to shows no symptoms, and Cardiac MRI also shows no evidence of recurrence of the tumor or the stalk. This evidence certifies that it is not necessary to do any extensive manipulation or other surgical approaches to reach the origin of the stalk since lipoma is a very slow-growing tumor. Nevertheless, we still encourage to do throughout diagnostic approach before the procedure to define the whole mass’ precise location. Extension of cardiac MRI to the cervical region or venography should be considered in any similar cases.

The surgical approach in excising lipoma in SVC should be considered wisely with the support of adequate preoperative diagnostic. Extensive manipulation that could increase surgical technique difficulty or postoperative morbidity and mortality is not necessary since lipoma is a very slow-growing tumor.

**Abbreviations**

SVC  
Superior vena cava; TTE: Transthoracic echocardiography; RA: Right Atrium; CT: Computed tomography; MRI: Magnetic resonance imaging; TEE: Transesophageal echocardiography; IVC: Inferior vena cava; ECG: Electrocardiogram

**Declarations**

**Ethics approval and consent to participate**

**Not applicable.**

**Consent for publication**

**Informed consent was obtained from the patient.**

**Availability of data and materials**

The datasets used and/or analysed during the current study are available from the corresponding author on reasonable request.

**Competing interest**

The authors declare that they have no competing interests.
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Author's contributions

Tri Wisesa Soetisna: tricts2000@yahoo.com: conceptualized, wrote the paper and reviewed the literature. Lisca Namretta: liscanamretta@gmail.com: wrote the manuscript and edited the paper. Bagus Ronidipta: ronipradana@rocketmail.com: reviewed the literature. Elen Elen: elensahara@gmail.com: validated the data and reviewed the paper. Sunu Budhi Raharjo: sunu.b.raharjo@gmail.com: reviewed and edited the paper. Amin Tjubandi: amintjubandi@yahoo.com: supervised, reviewed the literature and edited the paper.

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**Figures**

![Figure 1](image)

**Figure 1**

Preoperative radiology. (a) CT scan coronal plane; (b) Cardiac MRI T1-weighted image axial plane.
Figure 2

Giant lipoma after surgically resected
Figure 3

year post operatif cardiac MRI (cMRI). (a) Cine cMRI coronal plane; (b) Cine cMRI axial plane.