Reconstruction of the superior vena cava with an autologous pericardial patch for a giant superior vena cava aneurysm

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A 54-year-old man was transported to our hospital as the result of a loss of consciousness. Emergent enhanced computed tomography (CT) revealed a large superior vena cava (SVC) aneurysm (62 × 92 × 72 mm) and pulmonary thromboembolism. As the patient was hemodynamically stable, prudent follow-up with anticoagulant therapy was performed. During the follow-up, thromboembolism of the pulmonary artery improved; however, chest radiography performed 6 months after admission showed an enlarged SVC shadow (Figure 1, A and B). CT revealed rapid expansion of the aneurysm (65 × 117 × 88 mm) into the right upper thoracic cavity, and surgery was planned. The orifice of the aneurysm (approximately 2 cm in size on enhanced CT) was located immediately above the junction of the azygos vein; patch repair with autologous pericardium was planned. Aortography was performed to exclude arteriovenous fistula, and no communication was detected.

Surgery was performed with the patient under general anesthesia. Under full sternotomy, the SVC aneurysm was confirmed between the pericardium and right-side parietal pleura (Figure 2, A). The wall of the aneurysm was extremely thin and was depressed and dilated on breathing. To enable safe and further dissection, cardiopulmonary bypass was initiated to stop ventilation during dissection and reduce the risk of rupture. A venous drainage tube was inserted in the right and left brachiocephalic veins and inferior vena cava, and an arterial cannula was inserted in the ascending aorta. After the veins were clamped, the SVC was opened longitudinally (Figure 2, B). A thrombus was not observed inside the aneurysm cavity. The anterior, right, and posterior SVC wall, constituting 75% of the SVC, appeared damaged, thin, and dilated. The wall structure of the lesions differed from the structure of the remainder of the venous wall. Reconstruction of the SVC wall with pericardium is a good treatment option.
in accordance with the patient’s wish. Cardiopulmonary bypass weaning was uneventful, and the SVC pressure was normal (Video 1). No complications developed, and the postoperative course was uneventful. Pathologic examination of the aneurysm revealed normal venous wall structure with no localized wall structure abnormalities; however, the wall was thinner at the aneurysm site than at the other sites. CT at 6 months postoperatively revealed a well-reconstructed SVC (Figure 2, D). Informed consent for the publication of this paper was obtained from the patient.

**DISCUSSION**

SVC aneurysm is a rare venous pathology. Abbott et al. first described congenital aneurysm of the SVC in 1950 and with Leigh reported the classification in 1963. Congenital etiologies, trauma, previous surgery, and other factors may lead to the development of an SVC aneurysm; however, this patient presented with no specific history, such as high-energy trauma of the chest wall, surgery, obstruction, neoplasms, and arteriovenous aneurysms.

The management of SVC aneurysms after diagnosis is also controversial. Some studies report that conservative treatment is beneficial, particularly for primary, small (maximum diameter of <40 mm), and fusiform aneurysms and asymptomatic cases. These studies indicate that the treatment strategy should be based on the type and size of the aneurysm, but identification of the aneurysm type is challenging. If conservative treatment is selected, CT follow-up is essential.

In this patient, SVC aneurysm was apparent on CT at admission but was difficult to identify on chest radiographs. Chest radiography is not suitable for prudent follow-up.

Another treatment option is endovascular therapy. Griviau and colleagues reported a case of successful endovascular therapy for SVC aneurysms. This less-invasive method is also a good treatment option for small aneurysms, but large aneurysms are more difficult to treat.

**FIGURE 1.** A and B, SVC aneurysm (white arrows) observed on chest radiographs recorded at admission and 6 months later. C, Thromboembolism of the right pulmonary artery. D and E, Multilocular large SVC aneurysm (65 × 117 × 88 mm). The estimated SVC diameter was approximately 2 cm on preoperative computed tomography.
Another aspect of this disease is that a thrombus may form inside the aneurysm and cause pulmonary thromboembolism. In this patient with a large SVC aneurysm, bilateral pulmonary thromboembolism was detected. In patients with rapidly expanding SVC aneurysms, surgery may be the best treatment option, and with large aneurysms, the saphenous vein was too small to reconstruct the SVC. SVC reconstruction with an autologous pericardial patch is a good option for avoiding anticoagulant therapy.3,5

References
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FIGURE 2. A, Intraoperative image of the SVC aneurysm. B, Deep aneurysmal cavity after longitudinal incision. C, Image obtained after SVC reconstruction with the pericardial patch. D, Computed tomography image of the reconstructed SVC at 1-year postoperatively. Rt, right; BCV, brachiocephalic vein; Lt, left; SVC, superior vena cava.

VIDEO 1. Large superior vena cava aneurysm and reconstruction of the superior vena cava with a fresh pericardial patch. Video available at: https://www.jtcvs.org/article/S2666-2507(20)30582-4/fulltext.

VIDEO 1. Opening the anterior wall
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