Insights Into Endovascular Management of Superior Vena Cava Obstructions

Alexandre Ponti*, Sarah Saltiel, David C. Rotzinger and Salah D. Qanadli

Department of Diagnostic Radiology and Interventional Radiology, Lausanne University Hospital (CHUV) and University of Lausanne (UNIL), Lausanne, Switzerland

Superior vena cava obstruction results from any limitation of blood flow through the superior vena cava. Circulation to the heart may persist through various collateral vessels whose development depends on the level of obstruction. Depending on the level and degree of occlusive disease, the severity of clinical symptoms may vary considerably, up to lethal. Etiologies have changed dramatically in recent years, mainly due to the increasing use of intravascular devices. However, guidelines for treatment are lacking, and various options are available. Endovascular therapies developed considerably in recent years, may offer a rapid improvement in symptoms and proved to be safe. However, knowledge and selection of appropriate techniques are essential to venous angioplasty, involving specific tools to guarantee satisfying outcomes. This review aims to discuss the particular venous anatomy of the upper body, the physiopathology of superior vena cava obstruction, and specificities of endovascular treatment compared with other management options.

Keywords: superior vena cava, venous obstruction, venous disease, endovascular therapies, angioplasty, stent

INTRODUCTION

William Hunter first described superior vena cava (SVC) syndrome in 1757 as a complication of compression from an aortic aneurysm (1). Superior vena cava obstruction (SVCO) may complicate many mediastinal disorders whose etiologies have changed considerably in recent years. Until about 60 years ago, infectious diseases accounted for the majority of cases (particularly tuberculosis and syphilitic aortic aneurysm). Since then, malignant conditions have become the major cause followed by non-malignant thrombosis, reflecting the increased use of intravascular catheters and pacemakers leads (2, 3).

Limitation of venous blood return from the head, arms, and upper torso to the heart through the SVC results in a constellation of signs and symptoms, constituting the SVC syndrome. As in other parts of the body, blood flows through collateral vessels in case of vessel obstruction. In SVCO, collateral flow is directed to the lower body and inferior vena cava (IVC) or the azygos vein. The severity of symptoms depends on several factors, including the speed of onset, degree of narrowing, and the quality of collateral supply.

Imaging studies play a crucial role in the diagnosis and choice of treatment. Management is guided by the underlying etiology and the severity of symptoms.

This article aims to review the causes and endovascular therapeutic options for SVCO.
SVCO PATHOPHYSIOLOGY

Relevant Anatomic Considerations

Vascular occlusion brings about collateral circulation. When the SVC blood flow is compromised, collaterals may form to shunt the flow back to the heart. Depending on the obstruction site, collaterals may be intra- or extrathoracic. The most important pathway is the azygos system—the latter consists of the azygos, hemiazygos, and accessory hemiazygos veins. Depending on the level of obstruction, the collateral network and, therefore, the symptoms, will vary (4).

SVC occlusion below the junction of the azygos vein arch will allow direct and indirect shunting by the azygos system. Direct shunting will pass countercurrent through the azygos system to the IVC. Indirect shunting will pass through the pericardophrenic veins and secondarily join the azygos system and the IVC (through esophageal, diaphragmatic, mediastinal, and bronchial veins). Transverse shunting also exists through communications from the azygos system to the intra- and extra-spinal venous plexuses. The superficial venous system also allows a cavo-caval anastomosis (through the internal thoracic veins and epigastric veins). The superficial venous system also allows a cavo-caval anastomosis (through the internal thoracic veins and epigastric veins to the external iliac veins or through the external thoracic veins and superficial epigastric veins to the internal saphenous veins). Parietal veins may also join the para-umbilical veins.

Other shunting may form a right-to-left venous shunt (from anastomosis between the azygos system to the bronchial and then pulmonary veins) or a shunt to the portal system (esophageal veins to the left gastric veins or from anastomoses to the para-umbilical veins and ligamentum teres) (5, 6).

SVC occlusion above the azygos vein arch or the occlusion of the innominate vein on one side will force the development of supra-ternal venous anastomoses, allowing mainly a transverse contralateral shunt (anterior and external jugular veins are directly and indirectly anastomosed by the thyroid veins). Posterior anastomoses also exist with the vertebral veins and the spinal plexuses. By these derivations, blood will flow through the superior intercostal veins to the azygos system and the SVC below the occlusion.

Knowledge of this particular anatomy explains, therefore, why clinical SVC syndrome doesn’t develop when only one brachiocephalic vein is occluded and also that the symptoms are usually less severe if the level of obstruction is below the azygos system.

A comprehensive classification of SVCO based on previously published works (7, 8) integrating anatomic patterns that may impact the treatment strategy is provided here:

- Type I—isolated stenosis of the superior vena cava
- Type II—stenosis of central veins (subclavian/brachiocephalic) with or without extension to SVC
- Type III—chronic total occlusion of the SVC
- Type IV—chronic total occlusion of one or more central veins with or without extension to SVC

- IVa: occlusion of the brachiocephalic veins
- IVb: occlusion of the brachiocephalic veins and SVC
- IVc: occlusion of brachiocephalic veins, SVC and jugular veins

Etiologies of the SVCO

SVCO can be subdivided into benign or malignant and extrinsic or intrinsic etiologies (Table 1). Malignant causes are the most common (85% of cases) with bronchopulmonary cancer, mainly small cell cancers, germ cell tumors, and lymphoma as the most frequent. They induce mainly extrinsic compression of the central veins (9). Benign extrinsic causes are much rarer, such as benign tumors (goiter, teratoma), cardiovascular causes (aneurysm, constrictive pericarditis, pericardial effusion), and acute or chronic mediastinitis.

On the other hand, intrinsic causes are less common (3–20% of cases) but are the leading benign etiologies (9). In the vast majority of cases, benign causes involve a thrombus around a foreign body such as a central catheter or, more rarely, pacemaker wires. Venous thrombosis is favored either by endothelial lesions from mechanical friction against the vessel wall (for example, due to a short catheter tip above the right atrium) or rheological modifications caused by flow turbulences. Other favoring factors are related to the toxicity of the administered treatment (for example, chemotherapy through an implanted venous access device) or hypercoaguability of cancer patients (10, 11).

CLINICAL PROFILE OF SVCO

The severity of symptoms depends on the degree of narrowing and speed of onset. History should always specify the duration of symptoms, previous diagnoses of malignant diseases, or previous intravascular interventions. Symptoms are usually progressive over a few weeks and may improve in some cases as collaterals develop.

Patients can present with a wide variety of symptoms, the most frequent being upper body swelling (face, neck, arms). Less commonly, chest pain, respiratory symptoms (edema

### Table 1 | SVCO etiologies.

| Etiologies | Malignant |
|------------|-----------|
| Benign     | Malignant |
| External compression | Lung cancer |
| - Cardiovascular diseases | Lymphoma |
| - Lung diseases | Sarcoma |
| - Trauma | Metastatic cancer |
| - Mediastinitis | Leiomyosarcoma |
| - Thyroid goiter | Plasmocytoma |
| - Thymoma | |
| - Cystic hygroma | |
| Intraluminal fibrosis | |
| - Idiopathic | |
| - Long-term catheters | |
| - Pacemakers/defibrillators leads | |
| - Hypercoagulable state | |
| - Infectious (tuberculosis, syphilis, and histoplasmosis) | |

Most frequent etiologies of SVCO classified as benign and malignant.

Abbreviations: SVC, superior vena cava; SVCO, superior vena cava obstruction; IVC, inferior vena cava; MRI, magnetic resonance imaging; CT, computed tomography; CTO, chronic total occlusion; CAJ, cavoatrial junction.
of the larynx or pharynx causing cough, stridor, dyspnea, dysphagia, hoarseness), or neurologic manifestations (cerebral edema causing headaches, confusion, and possibly coma) can be observed (2, 3, 12–15).

There are two scoring systems to stratify the severity of superior vena cava syndrome (SVCS). Neither is validated, but both provide a valuable approach to select patients and create a decision tree. In the classification system proposed by Yu et al. (16), symptoms are categorized by their severity ranging from asymptomatic patients (grade 0) to severe (grade 3) and even life-threatening (grade 4) manifestations (Table 2), the two latter requiring emergent management. Grade 5 SVCS is lethal. The Kishi score (Table 3) also allows quantifying the gravity of symptoms and guide therapy. A score above 4 was retained as an indication for superior vena cava endoprosthesis by the authors (17).

Depending on how rapidly the symptoms appeared, SVCO can also be classified as acute, subacute, and chronic with a direct repercussion on treatment:
- **Acute**: <2 weeks, management relies mainly on thrombolysis and/or thromboaspiration and anticoagulation.
- **Subacute and chronic**: Subacute (2 weeks-2 months) or chronic (>2 months) SVCO, on the other hand, are best approached by endovascular therapies (angioplasty and/or stenting).

### SVCO IMAGING

Diagnosis of venous obstruction requires two findings—first, reduced vessel opacification distal to the obstruction, intraluminal filling defects, or an occlusive lesion—second, the presence of collateral pathways. Phlebography, either catheter-based or by computed tomography (CT) provides the most expedient diagnosis (18). To guide the endovascular treatment, additional information provided only by CT, such as vein thickness and the vein wall's fibrotic pattern, is helpful.

Catheter-based phlebography provides the fastest path to both diagnosis and immediate treatment (endovascular recanalization), in case of severe or life-threatening symptoms (cerebral edema, laryngeal edema, and hemodynamic compromise).

On the other hand, CT phlebography may better determine the precise origin of SVCO and is suitable for treatment planning in patients with less severe symptoms. CT also helps define the level and extent of venous blockage as well as collateral pathways of venous drainage, but above all, it allows to identify the underlying cause of venous obstruction. CT phlebography must be performed with a specific protocol to reduce flow artifacts due to non-opacified veins (18).

As cross-sectional imaging, magnetic resonance imaging (MRI) may help in the same manner in patients with a known allergy to iodinated contrast or when venous access cannot be obtained for contrast-enhanced studies (19).

Duplex ultrasound can be used for peripheral analysis (subclavian, axillary, and brachiocephalic veins) and indirect findings that suggest SVCO (dampening of the venous waveforms in the upper body with loss of venous pulsatility and respiratory variation) but cannot image the SVC directly.

### TREATMENT OPTIONS

Guidelines for SVCS are lacking, but traditionally treatment approaches have included chemotherapy and/or radiation therapy, surgical bypass, or endovascular therapy. Case selection

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**TABLE 2 | Yu Grading system (18).**

| Grade | Category            | Estimated incidence (%) | Definition                                                                 |
|-------|---------------------|--------------------------|----------------------------------------------------------------------------|
| 0     | Asymptomatic        | 10                       | Radiographic superior vena cava obstruction in the absence of symptoms     |
| 1     | Mild                | 25                       | Edema in head or neck (vascular distention), cyanosis, plethora            |
| 2     | Moderate            | 50                       | Edema in the head or neck with functional impairment (mild dyspghagia, cough, mild or moderate impairment of head, jaw or eyelid movements, visual disturbances caused by ocular edema) |
| 3     | Severe              | 10                       | Mild or moderate cerebral edema (headache, dizziness) or mild/moderate laryngeal edema or diminished cardiac reserve (syncope after bending) |
| 4     | Life-threatening    | 5                        | Significant cerebral edema (confusion, obtundation) or significant laryngeal edema (stridor) or significant hemodynamic compromise (syncope without precipitating factors, hypotension, renal insufficiency) |
| 5     | Fatal               | <1                       | Death                                                                      |

Each sign or symptom must raise suspicion of superior vena cava obstruction and the effects of cerebral or laryngeal edema or impact on cardiac function. Symptoms caused by other factors (e.g., vocal cord paralysis, compromise of the tracheobronchial tree, or heart as a result of mass effect) should not be considered as they are due to mass effect on other organs and not superior vena cava obstruction.

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**TABLE 3 | Kishi Score (17).**

| Signs and symptoms                                      | Grade |
|---------------------------------------------------------|-------|
| **Neurologic symptoms**                                 |       |
| - Stupor, coma, or blackout                             | 4     |
| - Blurry vision, headache, dizziness, or amnesia        | 3     |
| - Changes in mentation                                  | 2     |
| - Uneasiness                                            | 1     |
| **Laryngopharyngeal or thoracic symptoms**              |       |
| - Orthopnea or laryngeal edema                          | 3     |
| - Stridor, hoarseness, dysphagia, glossal edema, or shortness of breath | 2     |
| - Cough or pleural effusions                            | 1     |
| **Nasal and facial signs or symptoms**                  |       |
| - Lip edema, nasal stiffness, epistaxis, or rhinorrhea   | 2     |
| - Facial swelling                                       | 1     |
| **Venous dilatation**                                   |       |
| - Neck vein or arm vein distention, upper extremity swelling, or upper body plethora | 1     |

The scoring system of Kishi et al. (17), is used to quantify the gravity of symptoms. A score above 4 was retained as an indication for superior vena cava endoprosthesis by the authors. The total score for signs and symptoms was calculated as the sum of the highest grades in each class.
Medical management aims to relieve symptoms, preferably through relief of the obstruction but not always. A pure symptomatic benefit may be obtained by reducing hydrostatic pressure when elevating the patient’s head or administering loop diuretics. The real effect of those manoeuvres is, however, debated. In the same way, glucocorticoids are often prescribed in SVCO of malignant origin, while their effect is also questionable (2). Still, in malignant causes, SVCO may be resolved with medical therapy if the tumor’s histology is predictive of good tumor response (small cell carcinoma, lymphoma, and germ cell tumors). Chemotherapy or radiotherapy may induce a rapid improvement of symptoms, however not immediate (2, 14, 20). Symptoms may even be transiently worse due to the inflammatory phase of treatment. In benign causes, most often due to intravascular devices, removing the catheter and anticoagulation therapy may be beneficial. Surgical bypass with venous graft may also retain
some indications, mostly in benign cases, but is often substituted by endovascular treatment, as surgery is associated with more complications and longer hospital stays than endovascular therapies (21, 22).

**STRATEGIES FOR ENDOVASCULAR THERAPIES**

Acute SVCO management relies mainly on thrombolysis and/or thrombo-aspiration combined with anticoagulation for inpatients. Conversely, subacute and chronic SVCO is best treated by angioplasty and/or stenting. The latter are also commonly performed for inpatients. However, depending on the complexity, underlying disease, and co-morbidities, some procedures might be done on a day care basis.

**Principle of Angioplasty**

Endovascular treatment is usually performed under fluoroscopic guidance. Ultrasound is used to guide the percutaneous puncture of veins. Rarely, additional techniques might be used, such as intravascular ultrasound, cone-beam CT, or transesophageal echography.

Procedures can be performed under local anesthesia in simple cases, but general anesthesia is preferred when multiple accesses are necessary (typically type III and IV requiring recanalization of the occlusion).

Venous access can be unique or combined (femoral, basilic, and/or internal jugular). The typical introducer sheath is 10F 60 cm to allow insertion of large diameter stents from the femoral veins. In the case of double access, an additional 6F is usually sufficient. 5,000 to 10,000 Units (UI) of heparin are administered. Perfusion of sheaths with heparin infusion is also performed.

The standard angioplasty technique using a balloon over a wire extensively described in the literature for different anatomic sites is also valid for the SVCO. Hydrophilic or hybrid guidewires are most commonly used to cross the stenosis or occluded veins. Stiffer guidewires are necessary for large diameter stents (>14 mm). In chronic total occlusions (CTO) or severe stenosis due to implanted metallic devices, guidewires are supported by dedicated catheters (CTO catheters) to give the necessary push for stenosis/CTO crossing. CTO wires on 0.018 are rarely used.

Venous CTO requires gradual balloon angioplasty with small balloons first and then larger ones to overcome venous stricture. High-pressure balloons are preferred. In a CTO with complete fibrotic stenosis, a scoring balloon might help to open the vessel.
After angioplasty, stents are used. Self-expandable stents are preferred—oversized by 10–20% in diameter and length adjusted to the obstruction span. Multiple overlapped stents can be delivered if necessary. In-stent remodeling, if necessary, is performed with shorter in-stent balloon dilatation, with caution paid to avoid stent displacement. A long introducer sheath (60 cm) helps secure balloon retrieval. To prevent secondary sent migration, the objective is to restore sufficient flow and not necessarily reach 0% residual stenosis. A mild non-hemodynamically significant stenosis in the stent is usually considered optimal.

Self-expanding (nitinol, elgiloy), balloon-expandable (cobalt chromium, stainless steel) as well as self-expanding covered (PTFE-coated nitinol) stents and stent-grafts have been used with similar results (23–33). However, self-expanding stents with greater flexibility may be advantageous for adaptability and ease of positioning and deployment (27). On the other hand, covered stents may offer better long-term patency but without any significant effect on clinical success rate or survival (30, 34). Covered stents should also be used with caution due to concerns of migration and coverage of venous collaterals.

Adjunctive therapies, such as thrombectomy or catheter-directed thrombolysis, are used in the case of associated thrombotic material.

Sometimes, thrombectomy must be applied to dissolve associated large clots above the obstruction.

Endpoints of SVO Revascularization
For type I and II, treatment can be limited to the site of stenosis. For type III and IV, the goal is to re-establish direct venous flow between the heart and the brain through the dominant jugular axis. If not possible, as is sometimes the case in type IV, the non-dominant jugular vein is treated. As a last resort, if no jugular axis can be re-vascularized, flow is re-established between the SVC and a subclavian vein. Figure 1 illustrates the goals of venous reconstructions in various cases of chronic total occlusion without and with central venous catheters.

Types of SVC Reconstruction
Different reconstructions are possible in the SVC system. One line reconstruction, defined as revascularization based on connecting the dominant jugular vein to the proximal SVC or even the right atrium, is the most appropriate strategy to resolve SVC syndrome. Two line reconstruction aims to reconstruct both sides (brachiocephalic or jugular veins) to the proximal SVC based on a Y reconstruction or kissing stents into the SVC. This reconstruction is useful for type IV CTO in patients who need a central venous catheter through the stent. In such cases, we prefer asymmetric reconstruction as previously reported (35). Bilateral symmetric reconstruction was reported to have no clinical benefit and is associated with higher complication rates and recurrence of SVCO (31).

Management of Implanted Central Venous Catheters During SVCO Revascularization
Preserving previously implanted central venous catheters can be challenging when treating SVCO. First of all, tip positioning must be precisely determined. According to Glauser et al., three types are distinguished (36). In type I positioning, the tip is located more than or <1 cm from the superior cavoatrial junction (CAJ). Endovascular reconstruction of the superior vena cava is usually possible without mobilization of the catheter. The catheter will be flattened between the stent wall and the vessel, but the tip will stay free at the cavoatrial junction, saving device function. On the contrary, in some type II or III positioning, the catheter tip is suboptimal or non-optimal. In type II positioning, the tip is located >1 cm above or below the cavoatrial junction. In type III, the tip is located >3 cm below the CAJ or not inside the SVC. Stenting at the tip level may render the catheter unusable (type II or III with the tip above the CAJ). In such cases, the catheter should be withdrawn from the venous segment treated before endovascular reconstruction and then repositioned in the stent lumen. This maneuver is typically performed by brachial access using a snare to catch and mobilize the tip (35, 37). Figures 2–4 show examples of treatments. Sometimes, a supplemental manoeuvre of mechanical adhesiolysis is required to allow catheter mobilization (38).

Post-operative Care
Patients with significant pre-existing heart disease (ejection fraction <30%) might benefit from intermediate care facilities for continuous monitoring during 24 h following the SVCO revascularization, to prevent pulmonary edema. This is specially indicated for patients with subacute severe type III/IV SVCO. Although no consensus exists, at least a short-term anticoagulation or antiplatelet therapy is indicated (39). We recommend long-term anticoagulation therapy in case of residual stenosis and low hemorrhagic risk.

COMPLICATIONS
The primary risk during venous angioplasty is vessel rupture, which may precipitate pericardial tamponade. Faged et al. found that using stents over 16 mm was the only significant predictive factor for complications (hemothage, acute pulmonary edema from SVC reopening, stent collapse, stent migration, stent infection, and death). The same authors found that the median time to recurrence was 30 days, and the risk factors were initial CTO of the SVC, initial thrombosis, and the use of steel stents.

Interestingly, the use of long-term anticoagulation therapy was not associated with changes in recurrence or complication rates (27). Finally, the risk of pulmonary edema can be prevented by close monitoring of patients with reduced cardiac function.

FOLLOW-UP
Generally, successful endovascular therapy ensues in more than 95% of patients, and over 90% of them report symptom relief (8, 24, 26, 39–41).

Restenosis (defined as occlusion or stent thrombosis) is infrequent, reported in about 13% of patients, and often, patency can be restored with re-intervention (39).
Independently of intervention success, the average life span in patients with malignant related SVCS is poor, about 6 months, but offering an immediate and sustained symptomatic relief (28). On the other hand, survival is expected to be normal in patients with benign causes of SCVS, and endovascular stenting was shown to offer durable mid-term relief, equal to surgical reconstruction (22).

CONCLUSIONS

Despite the absence of consensus, careful selection of patients with SCVS according to the etiology, severity, and length of symptoms may offer the best treatment with the highest likelihood of success. Endovascular treatment of SVCS is an effective and safe method to rapidly improve symptoms due to the obstruction of blood flow through the SVC.

AUTHOR CONTRIBUTIONS

AP wrote the first draft of the manuscript. AP, SS, DR, and SQ contributed to manuscript extension, revision, read, and approved the submitted version. All authors contributed to the article and approved the submitted version.

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