Persistent left superior vena cava: clinical importance and differential diagnoses

Aynur Azizova, Omer Onder, Sevtap Arslan, Selin Ardali and Tuncay Hazirolan*

Abstract

Persistent left superior vena cava (PLSVC) is the most common thoracic venous anomaly and may be a component of the complex cardiac pathologies. While it is often asymptomatic, it can lead to significant problems such as arrhythmias and cyanosis. Besides, it can cause serious complications during vascular interventional procedures or the surgical treatment of cardiac anomalies (CA). The clinical significance of PLSVC depends on the drainage site and the accompanying CA. In this article, we will describe the epidemiology, embryology, and anatomic variations of PLSVC. Possible accompanying CA and heterotaxy spectrum will be reviewed with the help of multidetector computed tomography (MDCT) images. Radiological pitfalls, differential diagnoses, and the clinical importance of PLSVC will be highlighted.

Keywords: Persistent left superior vena cava, Cardiac anomalies, Clinical significance, Differential diagnoses, Computerized tomography

Key points

- Persistent left superior vena cava (PLSVC) may lead to significant clinical symptoms and may affect surgical management.
- PLSVC may accompany various congenital cardiac diseases as well as heterotaxy spectrum.
- To be aware of the differential diagnoses of PLSVC is essential for correctly interpreting left-sided mediastinal vascular structures.

Background

Persistent left superior vena cava (PLSVC) is a rare vascular anomaly that begins at the junction of the left subclavian and internal jugular veins, passes through the left side of the mediastinum adjacent to the aortic arch. It mostly drains into the right atrium via the coronary sinus (CS). Although PLSVC is infrequent among all vascular anomalies, it is the most common thoracic venous anomaly. Mostly, PLSVC is asymptomatic and detected incidentally in diagnostic and therapeutic examinations due to different reasons. However, it can be discovered as a component of the complex cardiac pathologies and may lead to significant problems such as arrhythmia [1–4].

There are different modalities for evaluation of PLSVC, such as perinatal echocardiography, multidetector computed tomography (MDCT), magnetic resonance imaging (MRI), and invasive angiography. The advantages, disadvantages of these modalities, and optimal techniques for imaging of PLSVC are shown in Table 1 [5–7].

In this article, we will describe the epidemiology, embryology, and anatomic variations of PLSVC. Possible accompanying cardiac anomalies (CA) and heterotaxy spectrum will be reviewed with the help of MDCT images. The radiological pitfalls with their CT imaging features that may help make the differential diagnosis, and the clinical importance of PLSVC will be highlighted.

Epidemiology

The exact frequency of PLSVC is not known because PLSVC is often asymptomatic and is detected incidentally. There is no significant difference in its prevalence between males and females. The prevalence of PLSVC ranges from 0.2 to 3% in the general healthy population. In patients
Table 1: The advantages, disadvantages, and techniques of different modalities for evaluation of PLSVC

| Imaging modality                  | Pros                          | Cons                                       | Techniques                                                                 |
|-----------------------------------|-------------------------------|--------------------------------------------|---------------------------------------------------------------------------|
| Echocardiography                  | ✓ Cheap                       | Difficult to interpret                     | * Coexistence of dilated coronary sinus without any evidence of right-sided congestion and positive “Bubble study” are diagnostic sonographic findings for PLSVC. |
|                                   | ✓ Widely available            | Operator-dependent                         | “Bubble study” is conducted with the injection of agitated saline from the left peripheral arm veins. If PLSVC is present, the agitated saline bubbles firstly are seen in the coronary sinus, before the right atrium. |
|                                   | ✓ No ionizing radiation       | Acoustic window dependent                  | In case of isolated PLSVC, positive “Bubble study” is observed after injection from right peripheral arm veins, as well. |
|                                   | ✓ Not affected by cardiac rhythm | The spatial resolution could be limited.   | * Contrast-enhanced echocardiography and transesophageal echocardiography are other useful modalities for the detection of PLSVC. |
|                                   | ✓ Portable (bedside assessment) |                                             |                                                                           |
|                                   | ✓ Real-time imaging           |                                             |                                                                           |
|                                   | ✓ Enables evaluation of flow direction |                                             |                                                                           |
| Multidetector computed tomography  | ✓ Accessible                  | Radiation exposure (Recently developed dose reduction methods have partially reduced concerns about radiation exposure) | * “ECG-gated CCTA with thin slices and multiplanar reformation” provides a detailed assessment. |
|                                   | ✓ Fast scanning speed         | An iodinated contrast agent (allergy, nephrotoxicity) | * Intravenous non-ionic iodinated contrast injection with a dose of 0.5-2 ml/kg at a rate of 1-2 ml/s is recommended. |
|                                   | ✓ The best spatial resolution | Cardiac rhythm changes may cause artifacts. | * The identification of PLSVC is usually independent of the contrast injection route (right or left, upper or lower extremities). The optimal contrast opacification of PLSVC is mostly seen in the delayed venous phase images. |
|                                   | ✓ Enables multiplanar imaging and reformatting | Need for sedation in the pediatric age group |                                                                           |
| Magnetic resonance imaging        | ✓ Radiation free              | High cost                                  | * Axial and coronal cine SSFP sequences are the best sequences for imaging of PLSVC. |
|                                   | ✓ High spatial resolution     | Less accessible                            | * The black blood TSE T2 images is also useful. |
|                                   | ✓ Enables multiplanar image acquisition | Slow scanning speed                         | * Contrast-enhanced MRA (± Dynamic imaging) and phase contrast angiography can be used as auxiliary modalities. |
|                                   | ✓ Enables assessment of flow direction | Contraindications such as the magnetic implant, claustrophobia |                                                                           |
|                                   | ✓ Depiction of PLSVC even without the administration of contrast media | Cardiac rhythm changes may cause artifacts. |                                                                           |
|                                   | ✓ Non-iodinated contrast      | Need for sedation in the pediatric age group |                                                                           |
| Invasive angiography              | ✓ Gold standard               | Invasive                                   | * The catheter angiography with water-soluble contrast agent is performed. Venograms are obtained after bolus contrast injection from the catheter. |
|                                   | ✓ Excellent morphologic information | Radiation exposure                         | * Invasive angiography is not a routine imaging modality for evaluation PLSVC. |
|                                   | ✓ Interventions can be made if necessary | An iodinated contrast agent (allergy, nephrotoxicity) | * PLSVC can be detected incidentally during procedures like central venous catheter insertion or pacemaker implantation. |

PLSVC: persistent left superior vena cava, ECG-gated CCTA: electrocardiogram-gated coronary computed tomography angiography, SSFP: steady-state free precession, TSE: turbo spin echo, MRA: magnetic resonance angiography
with congenital heart disease (CHD), its prevalence ranges between 1.3 and 11%. Additionally, the prevalence of PLSVC is thought to be higher in the prenatal period since the accompanying anatomic anomalies, including heart defects, may cause spontaneous abortions and premature deaths [1–3, 8, 9].

**Embryology**

The primitive venous system consists of three paired veins: vitelline veins (VV), umbilical veins (UV), cardinal veins (CV). Superior and inferior CVs are essential structures that allow the blood to return from the cranial and caudal parts of the embryo to the primitive heart. They combine to form common CVs (or the duct of Cuvier) draining into the double horned sinus venosus [2, 3, 9]. The caudal part of the right superior CV, together with the common CV, forms the right superior vena cava (RSVC). Generally, the left common CV and the caudal part of the left superior CV will regress. If these veins do not regress, then they will persist as PLSVC [2, 3, 8–11]. The detailed schematic anatomy of the developmental stages of the primitive venous system is shown in Fig. 1.

Different hypotheses regarding the development of PLSVC have been proposed. One of these is “low left atrial pressure theory.” According to this theory, in the presence of anomalies, which may cause reduced left atrial pressure and insufficient development of the left atrium, such as atrioventricular septal defect (AVSD), the left atrium will be smaller than expected. Thus, it will not be able to compress the CS and left CVs adequately. As such, the left common CV and caudal part of the left superior CV will not regress, and PLSVC will develop. Some hypotheses suggest the vice versa. According to the “obstructive theory” hypothesis, the presence of PLSVC, which may cause an increase in CS size, could lead to the formation of a left-sided obstructive lesion because of the space restriction [1].

**Drainage site and its impact on the anatomy**

PLSVC is responsible for approximately 20% of the total venous blood return from the left arm, left half of the head and neck. The right atrial drainage is seen in 80–90% of cases, while the left atrial drainage accounts for the remaining 10–20%. Generally, it joins into the right atrium through the CS and mostly has no hemodynamic impact.
effect. However, CS ostial atresia may accompany PLSVC. In that case, PLSVC becomes the major retrograde drainage pathway for coronary veins unless collateral drainage pathways develop between the coronary sinus and the heart chambers. The left atrial drainage, which is rare, occurs directly via the left atrial appendage or indirectly through the left pulmonary veins or the CS. In some sources, the latter is defined as an unroofed CS or CS atrial septal defect. The association of the atrial septal defect (ASD) and PLSVC draining into the left atrium via unroofed CS is called as Raghib syndrome [2–4, 9, 12].

Left atrial drainage is a cause of right-to-left shunt and is mostly accompanied by CA. However, it was reported that this condition might also be observed without any cardiac defects [2, 3, 12].

It has been reported that the drainage of more than the expected venous blood volume into the right atrium leads to some changes in the heart anatomy. Of those, the most well-known is the enlargement of the CS, which is a helpful clue indicating PLSVC existence. This enlargement may rarely reach the aneurysmatic level (Fig. 2g, h). There are also other anatomical changes reported in the literature, and they are described in Table 2, together with possible underlying mechanisms [2, 3, 12].

### Table 2

| Reported anatomical changes | Possible underlying mechanisms |
|----------------------------|-------------------------------|
| The decrease in RSVC dimensions | Reduction of blood volume drained through RSVC |
| Decrease in mitral valve area | Compression to the left atrium via dilated CS |
| Atrophy of the valves of cardiac veins such as Vleussens, Thebesian | Increased blood volume draining into the CS |
| The presence of a common left pulmonary vein trunk | Limited space caused by the dilated CS |
| Increase in heart weight | – |

PLSVC persistent left superior vena cava, RSVC right superior vena cava, CS coronary sinus

---

**Fig. 2** Clinical importance and different drainage sites of PLSVC and possible anatomical changes. **a, b** Posterior-anterior (PA) chest X-ray (a), and sagittal-oblique reformatted CT image (b) of different patients depict the course of central venous catheters inserted into the PLSVC (red arrows) draining into the right atrium via the coronary sinus (blue star). **c** CT imaging performed to evaluate the cardiac anatomy and possible variations of pulmonary veins draining to the left atrium (LA) before the radiofrequency catheter ablation in a patient with atrial fibrillation. The three-dimensional volume rendering technique (3D VRT) image shows that PLSVC (red arrow), which is detected incidentally, drains to the coronary sinus (CS). **d** Coronal-oblique maximum intensity projection CT image indicates PLSVC (red arrow), which indirectly drains into the left atrium via the left upper pulmonary vein (blue arrow). **e** Sagittal-oblique reformatted CT image shows PLSVC (red arrow) draining into the left atrium via the unroofed coronary sinus (blue arrows). Since ASD (not shown) is also present, the findings are compatible with Raghib syndrome. **f** Sagittal-oblique maximum intensity projection CT image demonstrates that the coronary sinus is connected to the left atrium with an aberrant vein as a collateral drainage pathway in a patient with coronary sinus ostial atresia (not shown). Thus, PLSVC (red arrow) drains into the left atrium through the coronary sinus (red star) and interatrial aberrant vein (blue arrow). **g, h** Axial CT (f) and 3D VRT images with different views (g, h) indicate the coronary sinus aneurysm (blue arrows) in a patient with PLSVC (red arrow) draining into the coronary sinus.

---

**Presence of RSVC and bridging vein**

In up to 90% of the cases, the right superior vena cava (RSVC) accompanies PLSVC, and this situation is known as double SVC (DSVC). If the caudal part of the right superior CV regresses in the intrauterine period, RSVC cannot develop, resulting in the presence of isolated PLSVC (IPLSVC). Mostly, IPLSVC is associated with CA and cardiac situs disorders. However, there are examples of IPLSVC without any accompanying apparent CA in the literature. In cases of DSVC, dimensions of RSVC may be larger or smaller than PLSVC (Fig. 3a–c) [2, 3, 9, 13].

In 65% of the cases, DSVC runs along each side of the mediastinum without interconnection. However, there could be the left brachiocephalic vein (LBCV)
connecting them, which is also called the bridging vein (BV) (Fig. 3d, e) [3].

PLSVC and accompanying cardiac anomalies

To date, many CA associated with PLSVC have been identified and grouped in different ways [1, 14, 15]. Shunt lesions (Figs. 4 and 5), conotruncal malformations (CTMs) (Figs. 6 and 7), left-sided obstructive lesions (LOLs) (Fig. 8), right-sided lesions, and single ventricular anomalies (Fig. 9) constitute the main CA groups. Aortic arch anomalies are also associated with PLSVC (Figs. 10 and 11). The subgroups of these anomalies are listed in Table 3. Besides, a summary of the literature about PLSVC and accompanying CAs is compiled in Table 4. Additionally, heterotaxy forms another disease spectrum associated with PLSVC and will be discussed under a separate title.

In the literature, there is a wide range of information about the frequency of cardiac anomalies accompanying PLSVC [1, 6, 14–22]. According to Lendzidan et al., the most common cardiac anomalies accompanying PLSVC are single ventricle, atrioventricular septal defect (AVSD), and tetralogy of Fallot (TOF). Cha et al. reported that the most frequent concomitant anomaly is ASD, whereas, according to Eldin et al., complete atrioventricular septal defect comes the first [17–19].

Moreover, attention has been drawn to the relationship of some specific cardiac anomalies with PLSVC in many publications. In addition to left-sided pathologies such as mitral atresia, cor triatriatum, and hypoplastic left heart, transposition of the great arteries (TGA) and tricuspid atresia are other rarer anomalies that have been reported to be closely related to PLSVC in the literature [6, 16, 20].

Different cardiac anomalies come to the fore in different situations such as type of accompanying cardiac anomaly (cyanotic or acyanotic), presence of heterotaxy, and drainage location of PLSVC [14, 15, 21]. Different parameters, such as odds ratio and PLSVC index, are calculated in the literature and used to determine the relationship between PLSVC and cardiac anomaly [1, 8].
In some publications, cardiac anomalies accompanying PLSVC were grouped and evaluated as in Table 3, and in others, they were examined separately [1, 17].

Association of PLSVC with aorta-related pathologies such as right-sided arcus aorta (RAA) and coarctation of the aorta (CoA) have also been emphasized in the literature. It was mentioned that the association of PLSVC with RAA is approximately 16% [14]. In another study, CoA was reported to be an independent and powerful factor for the existence of PLSVC [8]. Gustapane et al. underlined the coexistence of PLSVC with coarctation of the aorta (CoA) (21.3%) and suggested that fetuses with
Fig. 7 Conotruncal malformations accompanying PLSVC-2. a–d Axial (a), sagittal-oblique reformatted (c), and 3D VRT (b, d) CT images show the accompanying D-TGA in a patient with PLSVC (red arrows). The aorta (Ao) is located to the anterior and right of the pulmonary truncus (PA) (a, b). Please note the parallel course of the aorta and pulmonary truncus without “crossing over” (c, d). e–h Axial (e, h) and 3D VRT (f, g) CT images depict the accompanying L-TGA anomaly in a patient with PLSVC (red arrows). The aorta (Ao) is located to the left and anterior of the pulmonary truncus (PA) (e–g). The infundibular muscle around the aorta (blue arrow) indicates the right ventricular origin (e). The left-sided ventricle has the tricuspid valve (yellow arrows), which is the closer atrioventricular valve to the ventricular apex and indicates the right ventricular configuration (h).

Fig. 8 Left-sided obstructive lesions accompanying PLSVC. a–c Axial (a) and 3D VRT (b, c) CT images. The narrow segment compatible with aortic coarctation (blue arrows) is seen in a patient with PLSVC (red arrow) (a, b). Aortic coarctation treated by the endovascular intervention (blue circle) is depicted in another patient with double SVC (green and red arrows) (c). d–f Axial (d) and 3D VRT (e, f) CT images show the bicuspid aortic valve (blue arrows) (d) and the accompanying ascending aorta dilation (red stars) (e, f) in the patient with double SVC (green and red arrows). A bridging vein between RSVC and PLSVC (yellow arrows) is also seen (e, f).
PLSVC that are detected in the antenatal period should be followed during pregnancy in terms of CoA development [22].

**PLSVC and heterotaxy**

The term heterotaxy comprises situs inversus and situs ambiguous (right/left isomerism) (Fig. 12). DSVC or IPLSVC anomalies may be present in patients with heterotaxy. Meanwhile, in a study, “patients with both IPLSVC and situs inversus” were considered normal because of mirror image and excluded. In contrast, “patients with both isolated RSVC and situs inversus” were regarded as abnormal and accepted as SVC anomaly [1].

According to the literature, PLSVC-heterotaxy coexistence is frequently observed, and PLSVC is present in

---

*Fig. 9* Right-sided lesions and single ventricular anomalies accompanying PLSVC. a, b Axial-oblique CT images indicate unicuspid (blue arrow, a) and bicuspid (yellow arrows, b) pulmonary valve in different patients with PLSVC. c, d Axial (c) and coronal-oblique reformatted (d) CT images depict severe pulmonary stenosis (blue circles) in a patient with PLSVC (red arrow). e–g 3D VRT (e), axial (f), and sagittal-oblique reformatted (g) CT images depict complex cardiac anomaly in a patient with PLSVC. This patient has pulmonary atresia with confluent right and left pulmonary arteries connected to the aorta with a large caliber PDA (blue arrow) (e). There is a single ventricle (red star) and a single atrioventricular valve (orange arrows) (f). PLSVC (red arrow) and IVC (green arrow) are draining into the common atrium (blue star) (g).

---

*Fig. 10* Aortic arch anomalies accompanying PLSVC-1. a Axial CT image depicts azygos lobe (red star), and ARSA with Kommerell diverticulum (blue arrow) in a patient with PLSVC (red arrow). b Axial CT image indicates ARSA (blue arrow) in a patient with PLSVC (red arrow). c, d 3D VRT images with anterior and posterior views depict ARSA with Kommerell diverticulum (blue arrows) in a patient with double SVC (green and red arrows). e–h Coronal-oblique reformatted (e), axial (f), and 3D VRT CT images with different views (g, h) show the right aortic arch (RAA) and ALSA (blue arrows) in a patient with double SVC (green and red arrows).
50–70% of heterotaxy cases. Additionally, it is informed that 45% of patients with PLSVC in the antenatal period have accompanying heterotaxy [15]. In a study, DSVC is detected in nearly half of patients with heterotaxy [8]. Another study reported that 72% of heterotaxy patients with SVC anomaly have DSVC, while the remaining have IPLSVC [23].

According to a study, while right atrial isomerism in patients with PLSVC is about 7%, left atrial isomerism is about 9% [14]. In another study, those prevalences are nearly 15% and 30%, respectively. It was stated that the absence of inferior vena cava (IVC) is associated with left atrial isomerism, while the juxtaposition of IVC is observed in right atrial isomerism [15] (Fig. 13).

Complete atrioventricular septal defect, right ventricular outflow tract obstruction (RVOTO) (pulmonary stenosis and atresia), and double outlet right ventricle (DORV) are found as the most common accompanying anomalies of PLSVC in patients with heterotaxy [15] (Table 4, Fig. 14). The presence of concomitant heterotaxy and atrioventricular septal defect in patients with PLSVC during the antenatal period has been associated with poor prognosis.

Berg et al. reported that they never saw CS dilatation, which is a well-known sonographic finding supporting the presence of PLSVC, in the heterotaxy group. The absence of CS dilatation has been associated with unroofed CS, which is found in almost all heterotaxy cases. It should be kept in mind that the absence of CS dilatation does not exclude the presence of PLSVC in patients with heterotaxy during the antenatal period, and the possibility of concomitant unroofed CS anomaly is high [15].

**Clinical importance**

The clinical significance of PLSVC depends on the drainage site and the accompanying anomalies. PLSVC without CA is generally asymptomatic and is detected as an incidental finding. In the case of PLSVC with right atrial drainage, the CS often expands (Fig. 2g, h). This enlargement may cause compression of the

### Table 3 Main groups and subgroups of cardiac/aortic arch anomalies associated with PLSVC

| Main groups                        | Subgroups                                                                 |
|------------------------------------|---------------------------------------------------------------------------|
| Shunt lesions                      | ASD, VSD, AVSD, PDA, APVD                                                 |
| Conotruncal malformations          | TOF, PA with VSD, L/D-TGA, TA, DORV                                      |
| Left-sided obstructive lesions     | CoA, cor triatriatum, mitral stenosis, bicuspid aortic valve              |
| Right-sided lesions                | PS, PA, tricuspid atresia, bicuspid pulmonary valve, Ebstein anomaly      |
| Single ventricular anomalies       | None                                                                      |
| Aortic arch anomalies              | Cervical arch, RAA, ARSA, RAA + ALSA                                     |

PLSVC persistent left superior vena cava, ASD atrial septal defect, VSD ventricular septal defect, AVSD atrioventricular septal defect, PDA patent ductus arteriosus, APVD anomalous pulmonary venous drainage, TOF tetralogy of fallot, PA pulmonary atresia, L/D TGA-levo/dextro-transposition of the great arteries, TA truncus arteriosus, DORV double outlet right ventricle, CoA coarctation of the aorta, PS pulmonary stenosis, RAA right aortic arch, ARSA aberrant right subclavian artery, ALSA aberrant left subclavian artery.
Table 4 Summary of literature about PLSVC and accompanying CA

| Name of the author | Reported findings of PLSVC and associated CA |
|--------------------|-----------------------------------------------|
| Perles et al. [1]   | The most common groups of anomalies associated with PLSVC (Based on odds ratio) AVSD, CTMs, LOLs |
| Nagasawa et al. [6] | The highest incidence group of cardiac anomalies according to PLSVC index CoA and DORV |
| Lendzidan et al. [14]| The most common cardiac anomalies associated with PLSVC Single ventricle, AVSD, TOF |
| Ari et al. [12]     | The most common cyanotic heart diseases associated with PLSVC DORV and TOF |
| Berg et al. [13]    | The most common concomitant anomalies in patients with heterotaxy and PLSVC ASD and PDA |
| Perles et al. [1]   | The most common concomitant anomalies in patients with PLSVC, without heterotaxy AVSD, RVOTO, DORV |
| Oztunc et al. [16]  | The most common anomalies with PLSVC drained into the left atrium TOF and PS |
|                    | The most common anomalies with PLSVC drained into the left atrium Tricuspid atresia, TGA, situs anomalies |

PLSVC persistent left superior vena cava, CA cardiac anomaly, AVSD atrioventricular septal defect, CTMs conotruncal malformations, LOLs left-sided obstructive lesions, CoA coarctation of the aorta, DORV double outlet right ventricle, TOF tetralogy of Fallot, ASD atrial septal defect, PDA patent ductus arteriosus, RVOTO right ventricular outflow tract obstruction, VSD ventricular septal defect, PS pulmonary stenosis, TGA transposition of great arteries

atrioventricular node and His bundle. So, it can lead to cardiac arrhythmias, such as atrial/ventricular fibrillation. The compression of the left atrium and decreased cardiac output may occur due to this enlargement. Moreover, the presence of CS dilatation may complicate mitral valve surgery due to the close anatomic relationship [2, 9, 12, 19].

In a recent study by Yun Gi Kim et al. [24], it was demonstrated that PLSVC plays a considerable role in the induction and maintenance of atrial fibrillation (AF) in nearly half of the patients. So, pre-radiofrequency catheter ablation cardiac imaging in AF patients is useful and necessary for not only the evaluation of pulmonary venous anatomy but also for the detection of PLSVC existence. If PLSVC is detected as the trigger or driver of AF, it can be ablated (Fig. 2c).

It is crucial to know the PLSVC existence in advance in invasive procedures, such as central venous catheter (CVC) insertion (Fig. 2a, b), cardiac resynchronization therapy leads, or pacemaker implantation. It may complicate pacemaker implantation by causing fixation difficulties of the electrode due to the tortuous course. CVC insertion without fluoroscopy may cause angina, hypotension, and heart perforation. Furthermore, there may be constriction or atresia of the CS ostium. In this case, the catheterization will be challenging and may result in serious complications, such as dangerous arrhythmias, cardiogenic shock, and tamponade [2, 9, 12, 14].

The presence of CS ostial atresia is also critical in the operations that require PLSVC ligation. In this case, the CS still drains the blood from the coronary veins to the right atrium via the retrograde PLSVC-LBCV-RSVC pathway, instead of the atretic ostium. The ligation of PLSVC will be catastrophic due to the acute interruption of the cardiac venous drainage [12].

The left atrial drainage of PLSVC (Fig. 2d–f), sometimes, remains asymptomatic because it does not cause a right-to-left shunt at a significant level. In cases where the shunt is more pronounced, as a result of desaturation, the condition manifests itself with severe cyanosis, syncope, reduced exercise tolerance, and progressive fatigue. Thromboembolic events and even brain abscesses may develop in these patients. In this case, treatment can be done in two ways based on anatomy: PLSVC can be ligated if there is an adequate sized BV, and PLSVC can be re-anastomosed to the CS if the BV is not adequate in size or there is no RSVC [2, 9].

The knowledge of PLSVC is fundamental in some cardiac surgeries such as venous rerouting procedures, operations with cavo-pulmonary anastomosis (Glenn, Fontan), and heart transplantation. In heart transplantation surgery, if PLSVC without BV is present in the recipient’s heart, the bicaval anastomosis technique will be performed. It requires separation of the CS of the donor’s heart for the establishment of the recipient’s PLSVC anastomosis to the donor’s right atrium [1].
In the case of unknown PLSVC, retrograde cardioplegia, a common practice for cardiac surgeries for myocardial protection, will be ineffective. Clamping of PLSVC may be required for the prevention of retrograde flow. However, cardioplegia may fail even after clamping of PLSVC, due to the steal effect by the hemiazygos venous system linked to PLSVC [1, 3].

During cardiopulmonary bypass, not knowing PLSVC existence may result in both surplus blood return through the right atrium and insufficient venous return to the pump. This problem is mostly encountered in pathologies such as pulmonary atresia, tricuspid atresia, TOF, where increased systemic venous pressure gets over the level of left atrial pressure [19].

With the help of screening echocardiography, PLSVC can be detected as early as in the prenatal period. It can be used as a marker for cardiac or non-cardiac embryopathy. It may require extensive evaluation to exclude possible developmental anomalies. In cases with CHD, symptoms will be mainly due to these anomalies [1].

**Pitfalls and differential diagnoses**

In the presence of the vessel on the left side of the aorta in the mediastinum, other vascular structures apart from PLSVC should be considered in the differential diagnosis. They are vertical vein, levatoatriocardinal vein, left superior intercostal vein, aberrant left brachiocephalic vein, pericardiophrenic vein, and vascular structures secondary to surgery.

To make the definitive diagnosis, features which should be taken into consideration are as follows: “origin site,” “drainage site,” “orientation of the route between the origin and drainage site according to mediastinal structures,” “the expected direction of the blood flow,” and “characteristics of accompanying cardiac and non-cardiac diseases.” According to the above-mentioned features, a comprehensive summary of the differential diagnoses of PLSVC is depicted in Fig. 15.

Some masses on the expected course of PLSVC could be confusing at first look due to their location. For making the differential diagnosis, it is essential to follow all

---

**Fig. 13** PLSVC with heterotaxy-2 (right and left isomerisms). a–d Axial (a, c, d) and coronal-oblique minimum intensity projection (b) CT images depict right isomerism characterized by a bilateral broad-based triangular atrial appendages (blue arrows) (a), bilateral trilobed lungs (red circles) (b), asplenia (c), and juxtaposition of IVC (blue circle) in a patient with PLSVC (not shown). e–i Axial (e, f, h, i) and coronal-oblique reformatted (g) CT images depict left isomerism characterized by a bilateral narrow-based finger-like atrial appendages (blue arrows) (e, f), bilateral bilobed lungs (red circles) (g), polysplenia (blue circle) (h), dilated azygos vein due to the IVC absence (yellow arrow) (h), and right aortic arch (RAA) in a patient with double SVC (green and red arrows) (i).

**Fig. 14** PLSVC with heterotaxy-3 (the most common accompanying cardiac anomalies). Axial (a), coronal-oblique (b), and sagittal-oblique reformatted (c) CT images depict the three most common cardiac anomaly seen in the PLSVC and heterotaxy coexistence: AVSD (with dextrocardia) (a), pulmonary atresia (with VSD) (blue arrow, b) and DORV (c).
Fig. 15 A comprehensive summary of differential diagnoses of PLSVC. The course of the most frequent PLSVC variation (right atrial drainage via coronary sinus) and the courses of possible differential diagnoses are shown as columns with axial CT images from superior to inferior. For depicting the expected flow direction of vascular structures, the upstream zones are marked with darker shades, whereas the downstream zones with lighter shades. Red arrows represent pulmonary veins. Blue circles are used to depict systemic venous structures, while red circles are used to show the BT shunt, which is an interarterial structure. Orange circles represent the mass. In the column of Glenn shunt, the course of PLSVC is shown with solid blue circles while the course of RSVC is shown with hollow blue circles. Please note that (1) in the second row of the figure, all differential diagnoses of PLSVC are observed to be in a similar location in the mediastinum. (2) While two vascular structures are seen in front of the left main bronchus in the presence of PLSVC, no vascular structure is seen in this area in the presence of VV with PAPVD. (3) Expected flow directions for VV and LACV are caudocranial, unlike other vascular structures shown in the figure. Also, LACV is located in the posterior of the pulmonary artery, unlike PLSVC, which is located anterior to the pulmonary artery.

Fig. 16 Masses mimicking PLSVC. a–c Axial (a, b) and coronal-oblique reformated (c) CT images depict a neurofibroma (red arrows) arising from the left phrenic nerve, which has a parallel course with the pericardiophrenic vein, in a patient with neurofibromatosis type 1. If the beginning/end of the mass and relationship with vascular structures are not carefully evaluated, it can be confused with PLSVC due to its location. d–f Axial CT images depict multiple mediastinal hypervascular lymphadenopathies (blue stars) in the patient with renal cell carcinoma. Hypervascular lymphadenopathies in the left half of the mediastinum (red arrows) may mimic PLSVC. Green arrows show RSVC.
of the slices carefully and see the beginning and end of
the mass (Fig. 16) [25, 26].

Moreover, an interesting variant of PLSVC, which has
an intra-atrial course within the left atrium, has been
identified recently. If this pitfall variant is not known, it
may be misunderstood as left atrial cystic mass, may
cause patient anxiety, and may lead to unnecessary effort
for further investigations [27].

Vertical vein
The vertical vein (VV) is the vessel that drains the blood
from the pulmonary veins into the LBCV in the presence
of supracardiac type total or partial APVD (TAPVD or
PAPVD) (Fig. 17). It may be left- or right-sided. The left
APVD accounts for approximately 18% of all PAPVD
and left superior pulmonary veins are affected mostly.
The left-sided VV is one of the differential diagnoses of
PLSVC. The critical point in the distinction is the caudal
continuity of the vessel with atrial chambers. If there is
no continuity, it is compatible with the VV. However,
PLSVC may have a direct connection with the left pul-
monary veins. In this scenario, the pulmonary vein
drains into the left atrium after joining PLSVC [12, 28].

There are also some auxiliary features to differentiate
PLSVC and VV. The expected flow direction is craniocau-
dal in PLSVC, while it is caudocranial in the VV. In the
case of PLSVC, there are two vessels in the anterior aspect
of the left main bronchus: one of them is PLSVC, and the
other one is the left superior pulmonary vein. Ordinarily,
only the left superior pulmonary vein is expected to be at
this location. However, in the case of the VV with PAPVD,
no vessel is seen in the anterior aspect of the left main
bronchus. The size of the LBCV can also be helpful in
finding for differentiation. In the case of APVD, the LBCV
and RSVC may be of large caliber because the VV trans-
ports blood via these venous structures. On the other
hand, PLSVC, frequently, is associated with an absent or
small-sized LBCV [12, 28].

Levoatriocardinal vein
The levoatriocardinal vein (LACV) is the interatrial con-
nection that originates from the left atrium (68%) or pul-
monary vein (32%). It drains into one of the systemic
venous structures, mostly, into the LBCV (48%) (Figs. 18
and 19) [29, 30].

The differentiation of LACV from PLSVC with right
atrial drainage is straightforward. Because this drainage
site is unlikely for the levoatriocardinal vein. Similarly, in
cases where PLSVC drains into the left atrium via the
unroofed CS, unroofed CS and ASD facilitate the differ-
ential diagnosis in favor of PLSVC, since they are un-
usual for LACV [29–31].

However, PLSVC may drain directly into the left
atrium or pulmonary vein. In this situation, the expected
origin and drainage site of those two vessels will be the
same, and it is necessary to search other features for dis-
tinguishment. The anatomical feature that may help dis-
tinguish is their relative orientation according to the left
pulmonary artery. PLSVC is seen in the anterior aspect
of the left pulmonary artery, while the LACV is in the
posterior aspect. The evaluation of the flow direction
with echocardiography or velocity-encoded cine mag-
netic resonance imaging is another way to make differ-
ential diagnoses. The blood flows in the caudocranial

---

Fig. 17 Vertical vein. a-e Axial (a, b, c, e) and coronal-oblique reformed (d) CT images depict abnormal drainage of the left upper pulmonary vein (blue arrows) into the left brachiocephalic vein via the VV (red arrows). In the presence of VV with PAPVD (left-upper), the absence of vascular structure anterior to the left main bronchus (red circle) is an important clue for differential diagnosis (e). f-h 3D VRT (f), coronal-oblique maximum intensity projection (g), and axial (h) CT images depict the drainage of both right and left pulmonary veins (blue arrows) into the VV (red arrows), in a patient with supracardiac type TAPVD. The VV transports the whole pulmonary venous blood to the SVC through a large-caliber bridging vein (yellow arrows).
direction in the LACV while it flows craniocaudal direction in the PLSVC. However, the bidirectional flow could be seen in the levoatriocardinal vein [29–31].

Moreover, the caudocranial flow may be observed in PLSVC when there is atresia or stenosis of the CS ostium. Identification of accompanying CA may also help in the differential diagnosis. If LOLs without ASD are present, LACV should be considered in the differential diagnosis, firstly. It is hypothesized that, in the presence of in utero LOLs such as mitral stenosis, collaterals

![Fig. 18 Levoatriocardinal vein-1. a-f CT images of a patient with mitral valve stenosis due to acute rheumatic fever. The first four axial CT images (a-d) depict the venous structure (red arrows) coursing between the left brachiocephalic vein (blue arrow) and the left upper pulmonary vein (yellow arrow). The density difference between the cranial and caudal ends of the vessel suggests that flow direction is caudo-cranial. 3D VRT CT images with anterior and posterior views (e, f) show the course of the vessel. This vascular structure observed in the posterior of the pulmonary artery in the patient with mitral stenosis, which is a LOL, is compatible with LACV (red arrows).](image)

![Fig. 19 Levoatriocardinal vein-2. Axial (a, b, c, f) and 3D VRT (d, e) CT images of a patient with double SVC (green and red arrows, a) and complex cardiac anomaly, who underwent bivacaval Glenn shunt operation. Axial CT images (b, c) depict the thrombus extending from the right Glenn shunt to confluent pulmonary arteries (red stars). Due to the thrombus in the right Glenn shunt, the distribution of the contrast agent injected from the right arm into the mediastinal collaterals and the azygos system is observed. Axial (b, c) and 3D VRT reconstructed (d, e) CT images indicate bilateral vascular structures, which are compatible with LACV (blue arrows), originating from right Glenn shunt, coursing in the posterior of bilateral pulmonary arteries and draining into the right and left upper pulmonary veins. Please note that LACV may accompany LOLs (yellow arrow shows hypoplastic left heart, f), may be seen together with PLSVC, may be associated with any venous structure in the cardinal system not only left brachiocephalic vein, and in some cases, may have blood flow in the craniocaudal direction.](image)
between pulmonary and systemic circulations cannot re-
gress due to increased pressure in the left atrium and re-
maintain as LACV in the postnatal period. However, in the
presence of complex CA, the diagnosis of PLSVC should
be considered mainly [29–31].

LACV could be isolated without any CA, like PLSVC. Nevertheless, the frequency of this probability is very
low for LACV compared to PLSVC. Additionally, they
may be seen together, and the LACV may drain into
PLSVC [28, 30].

Pericardiophrenic vein
The pericardiophrenic veins (PCPV) are responsible for
pericardial and diaphragmatic venous drainage. They lie
along the lateral border of the heart and mediastinum,
accompany pericardiophrenic arteries/phrenic nerve and
drain into the internal thoracic, superior intercostal, or
BCV. Due to the connection with inferior phrenic veins,
dilated PCPVs could be observed as a collateral pathway
in cases of SVC or IVC occlusion. Besides, they can
serve as a collateral route via portosystemic shunting in
portal hypertension (Fig. 20) [32–34].

In the case of catheters located at the left parame-
diastinal region, the left PCPV is one of the possible
differential diagnoses. In posteroanterior chest X-ray,
left PCPV has a lateral course along the left heart
border, while PLSVC turns medially near the left
atrium. Although they both are located in the middle
mediastinum and connected with left brachiocephalic
vein cranially, their caudal courses differ in CT im-
gaging. While the caudal end of PLSVC is either the
coronary sinus or the left atrium, the left pericardioph-
renic vein moves toward the diaphragm lateral to
the heart when it is followed from top to bottom [35,
36].

Left superior intercostal vein
The left superior intercostal vein (L-SICV) drains the
blood from the second, third, and fourth left intercostal
veins into RSVC through the hemiazygos/azygos venous
systems. Ordinarily, it can be seen as a small aortic nip-
ple (1.4–5%) on the chest radiograph and is indistin-
guishable in CT. If its diameter exceeds 4.5 mm, it
should be considered as abnormal. In the case of occlu-
sion of SVC at the distal level of the azygos vein, the
connection between SVC and IVC becomes possible
with the dilation of L-SICV and other collateral vessels
(Fig. 21) [9, 37, 38].

Furthermore, L-SICV may dilate in congenital condi-
tions such as hypoplasia of LBCV, and diseases leading
to volume overload such as congestive heart failure. In
such cases, L-SICV might be confused with PLSVC.
However, knowing their courses and drainage sites will
facilitate the diagnosis [9, 37, 38].

Aberrant left brachiocephalic vein
Aberrant left brachiocephalic vein (ALBV) is a rare
anomaly (~1%) and is often associated with CAs,
such as TOF, septal defects, and right atrial isomer-
ism. Ordinarily, the LBCV passes through the anterior
of the arcus aorta and connects with the right BCV.
In the presence of an aberrant course, the LBCV be-
gins with the junction of the left subclavian and jugu-
lar veins, moves inferiorly along the left side of the
mediastinum, and joins to the right BCV passing

---

**Fig. 20** Left pericardiophrenic vein. **A–C** Axial CT images show a vascular structure compatible with PCPV (red arrows) in the left half of the mediastinum in a patient with SVC stenosis (green arrow). **D–F** Axial non-enhanced (**d**) and enhanced (**e, f**) CT images of a patient with portal hypertension secondary to Budd-Chiari syndrome depict varicose veins compatible with PCPV (red arrows) in the left half of the mediastinum. These vascular structures are connected with hepatic veins via a transdiaphragmatic course (blue arrows) (**f**).
behind the ascending aorta or esophagus. Retroesophageal ALBV is a more rare variation (Fig. 22) [10, 11, 39].

Vascular structures secondary to surgery
Vascular structures located on the left side of the mediastinum in patients with the history of cardiac surgery performed for complex CA may also be included in the differential list of PLSVC. The differential diagnosis could be made by knowing the performed surgery and demonstrating the drainage site of the vessel. Bicaval Glenn shunt, the left-sided Blalock-Taussig (BT) shunt, and collateral vessels after Fontan surgery are possible differentials (Fig. 23).

Bicaval Glenn shunt is an anastomosis of both SVCs to pulmonary arteries in the presence of PLSVC. The Glenn shunt allows the direct drainage of venous blood
into the pulmonary arteries via bypassing the right heart chambers [40, 41].

BT shunt is one of the surgical methods for complex CA. In this procedure, the connection of the subclavian artery and the pulmonary artery is enabled via the graft placement [40, 42].

In Fontan surgery, the SVC and IVC are anastomosed to the pulmonary artery. After surgery, collaterals, which may be seen as large vessels on the left side of the mediastinum, may develop and may be confused with PLSVC [43].

**Conclusion**

In conclusion, PLSVC is the most common thoracic venous anomaly known to be mostly asymptomatic. However, contrary to common misconception, it may cause a number of clinically significant symptoms, even in a heart with normal anatomy. Likewise, it may significantly affect the proper approaches to heart transplantations, effective surgical treatments for complex cardiac anomalies, and ablative procedures for cardiac arrhythmias. Thus, it should be recognized correctly and reported explicitly in radiological reports, even when it is an incidental finding. Besides, it is important to be aware of differential diagnoses of PLSVC and their radiological features to correctly interpret the vascular structures on the left side of the mediastinum.

**Abbreviations**

3D VRT: Three-dimensional volume rendering technique; AF: Atrial fibrillation; ALBV: Aberrant left brachiocephalic vein; ALSA: Aberrant left subclavian artery; APVD: Anomalous pulmonary venous drainage; ARSA: Aberrant right subclavian artery; ASD: Atrial septal defect; AVSD: Atrioventricular septal defect; AzV: Azygos vein; BT shunt: Blalock-Taussig shunt; BV: Bridging vein; CA: Cardiac anomalies; CCV: Common cardinal vein; CHD: Congenital heart disease; CoA: Coarctation of the aorta; CS: Coronary sinus; CTMs: Conotruncal malformations; CV: Cardinal vein; CVC: Central venous catheter; DORV: Double outlet right ventricle; DSVC: Double SVC; IPlSVC: Isolated PLSVC; ITVP: Inferior transverse venous plexus; IVC: Inferior vena cava; JV: Internal jugular vein; L/D-TGA: Levo/dextro-transposition of the great arteries; LA: Left atrium; LACV: Levoatriocardinal vein; LIBV: Left brachiocephalic vein; LIOV: Left inferior cardinal vein; LOLs: Left-sided obstructive lesions; LSCV: Left superior cardinal vein; LSVC: Left superior intercostal vein; MDCT: Multidetector computed tomography; MRA: Magnetic resonance angiography; MRI: Magnetic resonance imaging; OV: Oblique vein of the left atrium; PA: Pulmonary atresia; PAPVD: Partial PAPVD; PCPV: Pericardiophrenic vein; PDA: Patent ductus arteriosus; PLSVC: Persistent left superior vena cava; PS: Pulmonary stenosis; RA: Right atrial structure; RAA: Right aortic arch; RIOV: Right inferior cardinal vein; RSCV: Right superior cardinal vein; RSVC: Right superior vena cava; RVOTO: Right ventricular outflow tract obstruction; SCV: Subclavian vein; STVP: Superior transverse venous plexus; SV: Sinus venosus; SVC: Superior vena cava; TA: Truncus arteriosus; TAPVD: Total APVD; TOF: Tetralogy of Fallot; VSD: Ventricular septal defect; VV: Vertical vein

**Acknowledgements**

A part of this paper was submitted as an educational exhibit to RSNA 2020.

**Authors’ contributions**

AA and OO wrote the manuscript. SA contributed to the data collection. SA and TH edited the text. All of the authors read and approved the final manuscript.
to create a single caval vein. World J Pediatr Congenital Heart Surg 9(4): 446–450. https://doi.org/10.1177/2150135118765888

42. Kiran U, Aggarwal S, Choudhary A, Uma B, Kapoor PM (2017) The blalock and taussig shunt revisited. Ann Card Anaesth 20(3):323–330. https://doi.org/10.4103/aca.ACA_80_17

43. Lluri G, Levi DS, Aboulhosn J (2015) Systemic to pulmonary venous collaterals in adults with single ventricle physiology after cavopulmonary palliation. Int J Cardiol 189:159–163. https://doi.org/10.1016/j.ijcard.2015.04.065

Publisher’s Note
Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.