Anatomy, Physiology &
Classification of Varicose Veins

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Varicose veins are dilated, tortuous, elongated superficial veins that are usually seen in the legs. It can occur in any age group but most common in mid-twenties. It is a progressive disease. It is more common in females than males.

ANATOMY
All veins in the body are either part of the superficial venous system or the deep venous system. The principal superficial veins of the lower extremity are the small saphenous vein (SSV), which usually runs from ankle to knee and the great saphenous vein (GSV), which usually runs from ankle to groin (Figure 1).

Superficial collecting veins deliver their blood into the great and small saphenous veins, which deliver most of their blood into the deep system. Superficial veins are also connected to a variable number of perforating veins (PV) that pass through openings in the deep fascia to join deep veins of the calf or thigh either directly or through smaller plexus of smaller veins (Figure 2).

All venous blood is eventually received by the deep venous system on its way back to the right atrium of the heart. The principal deep venous trunk of the leg is called the popliteal vein from below the knee until it passes upward into the distal thigh, where it is called the femoral vein (FV) for the remainder of its course in the thigh. This is the largest and longest deep vein of the lower extremity.

Unlike arteries with thick walls, most veins are very thin and easily distendable, so the peripheral venous system also functions as a reservoir to hold extra blood. You could say that the venous system is almost magical in its function when you are told that the entire cardiac output volume of 5–10 L/min is received into periphery venous system for eventual delivery back to the heart and lungs.

PATHOANATOMY
The veins have one-way valves to prevent them from backward flow. The correct functioning of the venous system depends on a complex series of valves. It has been known that varicose veins in the legs are caused by weakening of the veins and valves in the great saphenous veins and/or small saphenous veins. When the valves in these malfunction, blood begins to collect in the legs resulting in the buildup of pressure. The veins become enlarged and knotted and are visible near the surface of the skin as a varicose vein (Figure 3a and b).

Major valves which dysfunctions in the caution of varicose vein are saphenofemoral junction (SFJ) and saphenopopliteal junction (SPJ).

The termination point of the GSV into the common femoral vein, located proximally at the groin, is called the Saphenofemoral junction. The terminal valve of the GSV is located within the junction itself. In most cases, at least one additional sub terminal valve is present within the first few centimeters of the GSV. Most patients have
a single sub terminal valve that can be readily identified approximately 1 cm distal to the junctional valve.

**PATHOPHYSIOLOGY**

The pathophysiology behind their formation is complicated and involves the concept of ambulatory venous hypertension.

In healthy veins, the flow of venous blood is through the superficial system into the deep system and up the leg and toward the heart. One-way venous valves are found in both systems and the perforating veins. Incompetence in any of these valves can lead to a disruption in the unidirectional flow of blood toward the heart and result in ambulatory venous hypertension (AVH). 6

Incompetence in the superficial venous system alone usually results from failure at valves located at the SFJ and SPJ. The gravitational weight of the column of blood along the length of the vein creates hydrostatic pressure, which is worse at the more distal aspect of the length of the vein.

Reflux at or near the SFJ does not always come through the terminal valve of the GSV, nor does it always involve the entire trunk of the GSV. Reflux can enter the GSV below the sub terminal valve or even immediately below the junction, passing through a failed sub terminal valve to mimic true SFJ incompetence. Reflux can also pass directly into any of the other veins that join the GSV at that level, or it may pass a few centimeters along the GSV and then abandon the GSV for another branch vessel.

Incompetence of the perforating veins leads to hydrodynamic pressure. The calf pump mechanism helps to empty the deep venous system, but if perforating vein valves fail, then the pressure generated in the deep venous system by the calf pump mechanism are transmitted into the superficial system via the incompetent perforating veins.

Once venous hypertension is present, the venous dysfunction continues to worsen through a vicious circle. Pooled blood and venous hypertension leads to venous dilatation, which then causes greater valvular insufficiency. Over time, with more local dilatation, other adjacent valves sequentially fail, and after a series of valves has failed, the entire superficial venous system is incompetent. This can then cause subsequent perforator and deep venous valvular dysfunction.

The clinical findings of varicose veins, reticular veins, and telangiectasias are due to the hypertension in the superficial venous system that spreads to collateral veins and tributary veins, causing dilated tortuous structures. Treatment modalities are geared towards correcting the superficial venous hypertension.

In contrast to the superficial veins, the deep veins do not become excessively distended. They can withstand the increased pressure because of their construction and the confining fascia.

**THE CLASSIFICATION OF VENOUS DISEASE**

Venous disease of the legs can be classified according to the severity, cause, site and specific abnormality using the CEAP classification. The elements of the CEAP classification are:

- **C** (Clinical): Symptoms
- **E** (Examination): Physical signs
- **A** (Anatomy): Anatomical location
- **P** (Pathophysiology): Pathophysiological condition
- **I** (Intervention): Therapeutic modality

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**Fig. 3a: Showing normal veins and diseased veins**

**Fig. 3b: Showing normal veins and diseased veins**

**Fig. 4: Showing terminal and preterminal valve**

VF=femoral vein
GSV=Great saphenous vein
SSV=Supra-saphenic valve
TV=Terminal valve
PTV=Pre-terminal valve
For the initial assessment of a patient, the clinical severity is the most important and can be made by simple observation and does not need special tests. There are seven grades of increasing clinical severity:  

<table>
<thead>
<tr>
<th>Grade</th>
<th>Description</th>
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<tbody>
<tr>
<td>C 0</td>
<td>No evidence of venous disease.</td>
</tr>
<tr>
<td>C 1</td>
<td>Superficial spider veins (reticular veins) only</td>
</tr>
<tr>
<td>C 2</td>
<td>Simple varicose veins only</td>
</tr>
<tr>
<td>C 3</td>
<td>Ankle edema of venous origin (not foot edema)</td>
</tr>
<tr>
<td>C 4</td>
<td>Skin pigmentation in the gaiter area (lipodermatosclerosis)</td>
</tr>
<tr>
<td>C 5</td>
<td>A healed venous ulcer</td>
</tr>
<tr>
<td>C 6</td>
<td>An open venous ulcer</td>
</tr>
</tbody>
</table>

The majority of patients referred to the vascular surgical clinic have grade 2 diseases (simple varicose veins). Patients with C3-6 disease are demonstrating increase severity of chronic venous insufficiency, and all have a functional abnormality of the venous system. These patients are most at risk of chronic ulceration and require specialized tests such as venous duplex and ambulatory venous pressure measurement to diagnose and characterize the underlying venous abnormality. If we correct the venous abnormality in the disease process then the risk of complications associated with the venous disease are much lower.
REFERENCES


