Vascular malformations: an update on imaging and management

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ABSTRACT

Vascular malformations comprise a broad and heterogeneous range of lesions that often represent a diagnostic and therapeutic challenge for the pediatrician. For a long time, the use of an inaccurate nomenclature has led to confusion. Since management depends on the specific vascular malformation, a proper classification and identification is critical.

The objective of this article is to provide the necessary information about the current classification and terminology of vascular anomalies, including basic concepts about available imaging diagnostic and therapeutic tools for the management of such complex condition.

Key words: vascular malformations, diagnosis, lymphatic abnormalities, arteriovenous malformation, sclerotherapy, therapeutic embolization.

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INTRODUCTION

Vascular anomalies include vascular tumors and malformations. Frequently, they are incorrectly called “angiomas” or “hemangiomas”, and comprise a broad range of lesions, from a simple “birthmark” to life-threatening conditions. In general, they affect newborn, infants, young children and young adults. Vascular anomalies are diffuse or localized defects occurring during the embryonic phase and may involve any part of the body. They usually develop between the fourth and tenth week of gestation. These anomalies are present at birth, but may not be evident until weeks, months or years later.

CLASSIFICATION

In spite of multiple efforts for the diffusion and usage of a standard classification, nomenclature and terminology are still confusing for general pediatricians.

The terms “hemangioma” or “angioma” are frequently used in everyday clinical practice to refer to either vascular malformations or vascular tumors, although their etiology, natural course and treatment are different.

In 1982, Mulliken and Glowacki described the most widely accepted classification, which became the basis for naming and studying these lesions. It is a biological classification based on the presence of cell proliferation, histological features, and clinical findings.

This classification divides these anomalies into vascular tumors and malformations.

- Vascular tumors: hemangiomas are the most common type. They are benign tumors that may not be apparent at birth; they present increased cell proliferation and hyperplasia, and are characterized by an early proliferation phase and a subsequent involution stage.

- Vascular malformations: they are present at birth. They are characterized by dysplastic vascular channels with normal cell proliferation. They grow at the same rate as the child and present no spontaneous involution. Vascular malformations are further classified based on the vascular channel components involved: arterial, venous, lymphatic and capillary, or a combination of these. Their clinical features define their onset, follow-up and specific management.

In 1993, Jackson et al. proposed a radiological classification based on flow and hemodynamics. They classified vascular malformations as high-flow (arteriovenous fistula, arteriovenous malformation) or low-flow (capillary, venous, lymphatic malformations). Any vascular malformation involving an arterial channel...
will be considered “high-flow”, while any malformation without an arterial channel will be deemed “low-flow”.2,7-10

In 1996, the International Society for the Study of Vascular Anomalies (ISSVA) embraced and extended this classification, which has now become the most widely used system.

Conceptually, the ISSVA defined two types of anomalies, vascular malformations (which will be analyzed in this article) and vascular tumors (being hemangioma the most common type) (Table 1).

In 2014, the ISSVA expanded this classification based on ongoing research and updates on this topic.11 Vascular malformations are divided into simple and combined, and a section on the association with other anomalies and syndromes has been introduced (Table 2), among others. This new classification is available at www.issva.org.

**CLINICAL EXAMINATION**

In clinical practice, there are vascular malformations with a typical clinical course, clear characteristics on physical exam that require no imaging studies for diagnosis, while others are atypical, deep lesions covered by normal skin or involving vital structures for which clinical diagnosis is unclear or insufficient. Imaging studies are essential for an accurate diagnosis, to make a specific management decision and to discuss the prognosis with parents.

Recent scientific and technological advances in the field of imaging studies, surgery and interventional radiology offer multiple resources to diagnose and treat these malformations.

In this article we will focus on currently available tools used to diagnose and manage this complex condition.

**VASCULAR MALFORMATIONS DURING CHILDHOOD**

These are localized vascular defects occurring during development, and they are basically classified by the type of channel components involved (arterial, capillary, venous or lymphatic) and the malformation hemodynamics (high-flow or low-flow).1,3,8,11

They may grow at the same rate as the child, never going away or involuting. Many of these malformations become apparent during puberty because they have hormone receptors that make them sensitive to estrogen and testosterone variations.11,12

**Low-flow malformations**

**Venous malformations**

These are the most common type. They involve an abnormally developed vein wall that affects vein function. Involved channel

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<th>Table 1. Classification of vascular malformations/International Society for the Study of Vascular Anomalies, 1996</th>
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<td>Parkes-Weber: CM + AVF + limb overgrowth</td>
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<td>Servelle-Martorell: limb VM + bone undergrowth</td>
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<td>Sturge-Weber: facial + leptomeningeal CM + eye anomalies + bone and/or soft tissue overgrowth</td>
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<td>Maffucci: VM + spindle-cell hemangioma + enchondroma</td>
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<td>Macropelphalody:CM: CM + megalencephaly + polymicrogyria</td>
</tr>
<tr>
<td>CLOVES: CM + VM + LM +/- AVM + congenital lipomatous overgrowth</td>
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<td>Proteus: CM + VM and/or LM + asymmetrical somatic overgrowth</td>
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<tr>
<td>Bannayan-Riley-Ruvalcaba: AVM + VM + macrophalphy + lipomatous overgrowth</td>
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AVM: arteriovenous malformation; CLOVES: congenital lipomatous overgrowth, vascular malformations, epidermal nevi, skeletal/scoliosis and spinal abnormalities.
components are progressively dilated, which results in blood stagnation, leading to thrombosis, inflammation and pain. Venous malformations (VMs) frequently develop spontaneous localized thrombosis and thrombolysis. Persistent thrombi may calcify and phleboliths may be formed, typical of these lesions.\textsuperscript{1,3,7,11,12}

In general, they are isolated and mostly develop in the cervicofacial region, limbs and trunk.

**Clinical presentation**

Clinical presentation is highly variable. Superficial lesions are soft, bluish, compressible and non-pulsatile; they may be flat, slightly elevated or markedly exophytic (Figure 1).

Lesions may be localized, well-defined or extensive, diffuse and infiltrating. On palpation, a compressible, sometimes tender mass is often found, especially when there are recently-formed clots or phleboliths. Many times, spontaneous pain occurs, especially in relation to the onset of intralesional clots.\textsuperscript{3,4,12,13}

These malformations may grow in size depending on the patient’s position (e.g., with the head down), with the Valsalva maneuver, during infant crying, or with compression. They may grow suddenly after an incomplete surgical resection, a biopsy, a trauma and/or infections.\textsuperscript{13,16} Sometimes, pain is attributed to joint, tendon or muscle involvement. These lesions may even get worse during menstrual periods or puberty\textsuperscript{13,15} (Figure 2).

Extensive VMs may present localized intravascular clotting due to the above mentioned intralesional thrombosis; they occur accompanied by hypofibrinogenemia and elevated D-dimer levels. It is important to know the patient’s basic coagulation parameters before any intervention.\textsuperscript{12,14,17}

VMs may be associated with other clinical conditions. Klippel-Trenaunay syndrome typically has capillary malformations, venous and lymphatic (macro- and/or microcystic) malformations, and soft tissue and bone hypertrophy. Generally, these lesions affect the lower limbs, and sometimes they may be bilateral or involve the upper limbs.\textsuperscript{10,12,18}

**Imaging assessment**

For all vascular anomalies, Doppler ultrasound and magnetic resonance imaging (MRI) are generally the preferred methods for an adequate and complete assessment and follow-up of patients.
In relation to VMs, the initial assessment is a clinical exam. Small, superficial lesions that are clearly diagnosed may not require imaging studies; however, most patients will require at least a Doppler ultrasound.

**Ultrasound:** Two patterns may be identified. A cavitary pattern (the most common type), which shows heterogeneous images with mixed echogenicity. Phleboliths may be present, and thus reinforce the diagnosis of a venous malformation. The other is a dysplastic pattern with multiple tortuous dilated veins infiltrating subcutaneous fat, muscles, tendons and adjacent tissues.\(^{10,14,19}\) (Figure 1). These are low-flow lesions, so a Doppler ultrasound would be able to confirm a low-rate monophasic venous flow or the absence of flow within the malformation. Sometimes, flow may be observed when performing the Valsalva maneuver or compression (Figure 2).

**Magnetic resonance imaging:** An MRI allows to assess the extent of a lesion and its relationship to adjacent organs and structures. Recommended sequences are T1 (pre- and post-contrast) and T2 weighted images, with fat saturation. Typically, images may have an intermediate signal intensity in T1 and a hyperintense signal in T2 in relation to its content or the presence of hemorrhage or thrombosis. A gadolinium injection usually shows a diffuse enhancement of venous channels, unlike lymphatic malformations, which generally do not have this kind of enhancement. With T2 weighted images, phleboliths appear as focal areas of hypointense signal.\(^{13,16,20}\)

**Arteriogram:** It is not indicated.

**Percutaneous phlebography:** If treatment is required for the lesion, this is the imaging method of choice to observe the hemodynamics and angioarchitecture of the malformation. This invasive procedure is reserved as an immediate step prior to percutaneous management.

**Treatment**

Small, asymptomatic lesions can be managed conservatively. Compression stockings may be used by patients with limb involvement. The main indication for treatment is pain. In other cases, in spite of the absence of pain, treatment is indicated if there is joint involvement, functional impairment or for aesthetic reasons.\(^{4,13,17,21}\)

An effective medical treatment has not yet been discovered. In ongoing studies, it has been demonstrated that angiogenesis inhibitors have clinical effectiveness.\(^{22}\)

In patients with extensive venous malformations, coagulation should be assessed in depth. Alterations in the muscle wall of the affected vein leads to blood stagnation and thrombi formation. In some cases, localized intravascular coagulation (LIC) may occur, which

![Figure 2](image-url)

**Figure 2. Venous malformation**

A.

Thirteen-year-old female patient with an extensive venous malformation in the left lower limb (patient in ventral decubitus position).

B.

Doppler ultrasound of thigh region showing a cavitary pattern with no flow inside.

C.

Sagittal T2 MRI of the left knee showing a hyperintense signal caused by a venous malformation that diffusely involves the intra-articular and peri-articular region.
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is translated into elevated D-dimer levels and reduced fibrinogen.

At referral centers, LIC is managed with low-molecular-weight heparin. Sclerotherapy is contraindicated until LIC is resolved due to the risk of generalized intravascular coagulation.12-14

Sclerotherapy is, at present, the treatment that has shown the best results. It is performed at the Interventional Radiology Unit, usually under general anesthesia and with strict monitoring of vital signs. Percutaneous treatment consists of injecting sclerosing agents (sodium tetradecyl sulfate, ethanol and bleomycin, among other common agents) that cause endothelial damage and therefore lead to inflammation and fibrosis.2,7,13,14,17

Many times, treatment is combined with additional surgical procedures.

In children with extensive head and neck malformations that involve the airways, it is necessary to consider the possibility of performing a tracheostomy prior to sclerotherapy or to have the child in mechanical ventilation for 48/72 hours after the procedure because the inflammatory response may collapse the airways. For any of these cases, management is the result of a multidisciplinary approach.23-25

Lymphatic malformations

These are the second most common type of malformations. Lymphatic malformations are sequestated and dilated lymphatic sacs that lack connection with the peripheral lymph system. The rate of these lesions is similar in both sexes. They may classified as macrocystic, microcystic, combined and mixed.2,7,26

Although there are multiple ambiguous definitions, a lesion is considered macrocystic if the lesion can be successfully aspirated (and subsequently sclerosed), which reduces its size. This process is more complex in microcystic lesions.2,4,14

These malformations are more commonly located in the head and neck (70-80%), usually in the posterior cervical triangle13,26 (Figure 3).

The remaining 20% may be located in the axillary, superior mediastinal, mesenteric, retroperitoneal areas, the pelvis and the lower limbs. They may be associated with Turner syndrome, Noonan syndrome, multiple pterygium, and certain trisomies.18

Clinical presentation

Most cases are detected around two years old, although some may be apparent at birth and even get diagnosed during an antenatal ultrasound.2,3,7

These malformations appear as palpable tumors with a soft consistency and a temperature similar to the rest of the skin. Their clinical appearance is varied and depends on their size and, especially, their location and depth. Lymphatic malformations grow naturally at the same rate as the patient. During adolescence, the size and symptoms of these lesions may increase, probably in relation to hormone receptors present in them.26

In general terms, changes in skin color and/or small blisters may occur, which are characteristic of these lesions, or they may even have healthy skin.

A sudden increase in volume may be the result of an infection or an intracystic hemorrhage. Superinfected lesions are tense, warm and erythematous masses; an empiric antibiotic therapy is indicated in these cases. Intracystic hemorrhage is less common, although it may be secondary to a trauma or occur spontaneously. The lesion’s size increases suddenly, with a change in color and bruising, and less frequently, with an elevation in local temperature.

Imaging assessment

Ultrasound: Macrocystic lesions show multiple images of different size cysts, separated by echogenic septa. In case of bleeding, echogenic cysts or air-fluid levels are observed (Figure 3).

Microcystic lesions appear as echogenic images with ill-defined borders. Mixed lesions show cystic images inside an echogenic area. No flow is observed inside cysts, but vascularization may be evident in the septa separating them. Microcystic lesions lack blood flow.10,19,27

Magnetic resonance imaging: An MRI allows to assess the extent and depth of these lesions and their relationship to adjacent structures, as well as to observe their extension into the chest and airway involvement in the case of head and neck malformations. Macrocystic lesions show hyperintense signal in T2 images and low intensity signal in T1 images, with post-contrast enhancement of the septa. Microcystic lesions generally appear as T2 images with homogeneous hyperintense signal.4,10,16,20

Treatment

It has been demonstrated that medical treatment with antiangiogenic agents, such as rapamycin or sirolimus, is effective in very extensive, diffuse malformations, especially in newborn infants with extensive cervicofacial lesions. At present, there are different ongoing clinical studies with promising results for
these specific indications. In many cases, this treatment is usually associated with percutaneous management of macrocystic lesions or even an additional surgical resection.

Treatment with other drugs, such as sildenafil, has not proven to be safe and effective for this indication.

Sclerotherapy is usually indicated as a first-line option for patients with macrocystic lesions. It is performed at the Interventional Radiology Unit, usually under general anesthesia. The same precautions considered for VMs should be taken into account in the case of malformations involving the airways.

The most commonly used sclerosing agents are tetradecyl sulfate, doxycycline and alcohol for macrocystic lesions. Acceptable results have been observed with bleomycin for microcystic malformations.

In some cases, it is necessary to complete treatment with surgical resection of the remaining lesions.

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**Figure 3. Lymphatic malformation**

**A.**  
Eighteen-month-old male patient with an extensive mixed-type lymphatic malformation that involves the cervicofacial region, especially the tongue.  

**B.**  
Sagittal ultrasound image of the neck showing the typical pattern of a macrocystic lymphatic malformation. Large cysts with fluid content (lymph) and thin walls identified.  

**C.**  
Magnetic resonance images. Axial (C) and sagittal (D) T2 of the neck showing a hyperintense signal caused by the high protein content and multiple septa.  

**D.**  
Sagittal ultrasound image of the neck showing the typical pattern of a macrocystic lymphatic malformation. Large cysts with fluid content (lymph) and thin walls identified.
**Capillary malformations**

“Capillary malformation” (CM) is a generic term used to describe the presence of a “vascular stain” on the skin rather than to indicate a specific lesion. The most frequently recognized type of CM is a port-wine stain or nevus flammeus. This term also encompasses other types of vascular skin lesions, such as telangiectasias and hyperkeratosic macules.

CMs affect the capillary network vessels of the skin and mucous membranes, which are made up of ectatic dermal capillaries, and increase with age. These lesions are frequently associated with focal hypertrophy of the bones and underlying soft tissues.

CMs may be single and isolated, and may result in a severe aesthetic compromise. They may also be the expression of extracutaneous manifestation. CMs are present at birth, and no differences have been observed in their rate between both sexes. They are often located in the head and neck, and also in the trunk and limbs.

CMs are frequently combined with other malformations (capillary-venous, capillary-arteriovenous, or capillary-lymphatic malformations) and they may even be a component of more complex anomalies, such as Sturge-Weber syndrome, Parkes-Weber syndrome, the macrocephaly-CM complex, or hereditary hemorrhagic telangiectasia. In the Sturge-Weber syndrome, CMs affect mainly, but not exclusively, the upper lids and the supraorbital region, and are frequently associated with seizures.

Imaging studies for CMs are ultrasound and MRI, which only show a non-specific thickening of the dermis, occasionally extended to the subcutaneous cellular tissue. The study of choice in patients suspected to have Sturge-Weber syndrome is an MRI. A brain MRI with intravenous contrast would be able to show soft tissue and adjacent bone hypertrophy, focal brain atrophy, leptomeningeal enhancement and thickening and the typical intracranial calcifications, among other findings.

**Clinical presentation**

Symptoms are related to hypervascularization, the presence of abnormal arteriovenous shunts, venous hypertension and tissue ischemia. Symptoms may include pulsatile masses, generally with increased skin temperature and palpable thrill. During auscultation, superficial lesions usually indicate a murmur. They may be accompanied by bone and soft tissue overgrowth and/or be associated with osteolytic lesions. In advanced stages, AVMs may have trophic changes due to venous hypertension, and distal ischemia due to vascular steal syndrome. Some cases are associated with congestive heart failure due to right ventricular overload generated by the high flow of these lesions (Figure 4).

**Imaging assessment**

**Ultrasound:** An ultrasound shows a heterogeneous, ill-defined mass with vascular dilations in and around the mass. Doppler ultrasound recognizes numerous vessels with a high arterial flow, a high diastolic peak, turbulent flow and pulsatile venous flow.

**Magnetic resonance imaging:** An MRI allows to assess vascular anatomy, the extent of the lesion and its relationship to adjacent structures. With T1 images, it is possible to recognize tubular structures with hypointense signal (flow void phenomenon), which usually represent afferent arteries and the malformation nidus, which reflects a high flow. Angio-MRI images are very useful, especially in children, because they show the lesion’s afferent and efferent vessels and help to plan an eventual endovascular procedure. Additionally, an MRI is a very useful follow-up method after a procedure.

**Angiogram:** This invasive study shows the exact details of the AVM angioarchitecture and arteriovenous malformations and fistulas.

**Arteriovenous malformations**

This group mostly refers to two anomalies:
Transarterial occlusion of afferent vessels that supply the AVM should be particularly careful and occlusion of normal arterial vessels that feed other organs or tissues should be avoided. It should be an intra-nidus occlusion, without exception. If large-caliber draining veins are predominant in the AVM, transvenous endovascular embolization and subsequent occlusion of draining veins are an alternative approach because it has been shown that venous occlusion has a coadjuvant effect in nidus thrombosis. Another alternative would be to directly approach the malformation nidus by a direct percutaneous fine-needle puncture.

Treatment

The treatment of choice of AVMs is endovascular embolization. This procedure is a challenge for the treating team and requires a deep understanding of available techniques and of the vascular anatomy of these lesions.
This may be done in the case of superficial malformations which are easily accessed through direct puncture under angiographic endovascular control. The transarterial, transvenous and direct puncture techniques have also been frequently used in combination to manage AVMs.\(^7,17,40,41\)

Most commonly, liquid (ethanol) and semi-liquid (Onyx\(^8\), Histoacryl\(^19\)) embolic agents are used.\(^8,9,41\)

In short, a vascular malformation is a rare but severe condition that frequently requires a multidisciplinary approach.

The best results for these anomalies are achieved through the joint multidisciplinary management, control and follow-up of patients. In everyday practice, diagnostic and treatment decisions are commonly discussed among radiologists, general and plastic surgeons, dermatologists, pediatricians, trauma surgeons, hematologists, and ophthalmologists, just to name a few.

Imaging methods are the cornerstone of an adequate diagnosis and the basis of an appropriate management of these conditions. Pediatricians should be aware of these concepts to be able to guide children with such complex conditions and their families.

REFERENCES


