eviCore healthcare Clinical Decision Support Tool Diagnostic Strategies: This tool addresses common symptoms and symptom complexes. Imaging requests for individuals with atypical symptoms or clinical presentations that are not specifically addressed will require physician review. Consultation with the referring physician, specialist and/or individual’s Primary Care Physician (PCP) may provide additional insight.

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## Pediatric Peripheral Vascular Disease (PVD) Imaging Guidelines

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**PEDPVD-1.0: General Guidelines**

- A pertinent clinical evaluation including a detailed history, physical examination, appropriate laboratory studies and basic imaging such as plain radiography or ultrasound should be performed prior to considering advanced imaging (CT, MR, Nuclear Medicine), unless the patient is undergoing guideline-supported scheduled imaging evaluation. A meaningful technological contact (telehealth visit, telephone call, electronic mail or messaging) can serve as a pertinent clinical evaluation.

- Unless otherwise stated in a specific guideline section, the use of advanced imaging to screen asymptomatic individuals for disorders involving the peripheral vascular system is not supported. Advanced imaging of the peripheral vascular system should only be approved in individuals who have documented active clinical signs or symptoms of disease involving the peripheral vascular system.

- Unless otherwise stated in a specific guideline section, repeat imaging studies of the peripheral vascular system are not necessary unless there is evidence for progression of disease, new onset of disease, and/or documentation of how repeat imaging will affect patient management or treatment decisions.

**PEDPVD-1.1: Age Considerations**

- Many conditions affecting the peripheral vascular system in the pediatric population are different diagnoses than those occurring in the adult population. For those diseases which occur in both pediatric and adult populations, differences may exist in management due to individual’s age, comorbidities, and differences in disease natural history between children and adults.

- Individuals who are < 18 years old should be imaged according to the pediatric peripheral vascular disease imaging guidelines if discussed. Any conditions not specifically discussed in the pediatric peripheral vascular disease imaging guidelines should be imaged according to the general peripheral vascular disease imaging guidelines. Individuals who are ≥ 18 years old should be imaged according to the general peripheral vascular disease imaging guidelines, except where directed otherwise by a specific guideline section.

**PEDPVD-1.2: Imaging Appropriate Clinical Evaluation**

- See **PEDPVD-1.0: General Guidelines**
PEDPVD-1.3: Modality General Considerations

MRI
- MRI is generally performed without and with contrast unless the individual has a documented contraindication to gadolinium or otherwise stated in a specific guideline section.
- Due to the length of time required for MRI acquisition and the need to minimize patient movement, anesthesia is usually required for almost all infants (except neonates) and young children (age < 7 years), as well as older children with delays in development or maturity. This anesthesia may be administered via oral or intravenous routes. In this patient population, MRI imaging sessions should be planned with a goal of minimizing anesthesia exposure adhering to the following considerations:
  - MRI procedures can be performed without and/or with contrast use as supported by these condition-based guidelines. If intravenous access will already be present for anesthesia administration and there is no contraindication for using contrast, imaging without and with contrast may be appropriate if requested. By doing so, the requesting provider may avoid repetitive anesthesia administration to perform an MRI with contrast if the initial study without contrast is inconclusive.
  - Recent evidence-based literature demonstrates the potential for gadolinium deposition in various organs including the brain after the use of MRI contrast.
  - The U.S. Food and Drug Administration (FDA) has noted that there is currently no evidence to suggest that gadolinium retention in the brain is harmful and restricting gadolinium-based contrast agents (GBCAs) use is not warranted at this time. It has been recommended that GBCA use should be limited to circumstances in which additional information provided by the contrast agent is necessary and the necessity of repetitive MRIs with GBCAs should be assessed.
  - If multiple body areas are supported by eviCore guidelines for the clinical condition being evaluated, MRI of all necessary body areas should be obtained concurrently in the same anesthesia session.
- The presence of surgical hardware or implanted devices may preclude MRI.
- The selection of best examination may require coordination between the provider and the imaging service.

CT
- CT or CTA may be appropriate for further evaluation of abnormalities suggested on prior US or MRI Procedures.
- CT may be appropriate without prior MR or US, especially in the following (non-exhaustive list of) settings:
  - Lymphatic malformations
  - Vascular abnormalities including vasculitis, thrombosis, narrowing, aneurysm, dissection, and varices.
  - For preoperative planning or assessment of post-operative complications.
In some cases, especially in follow-up of a known finding, it may be appropriate to limit the exam to the region of concern to reduce radiation exposure.

CT should not be used to replace MRI in an attempt to avoid sedation unless listed as a recommended study in a specific guideline section.

The selection of best examination may require coordination between the provider and the imaging service.

- **Ultrasound**
  - Ultrasound can be helpful in evaluating arterial, venous, and lymphatic malformations.
  - Ultrasound can be limited by the imaging window and the patient body type.
  - CPT® codes vary by body area and presence or absence of Doppler imaging and are included in the table at the beginning of this guideline.

- **3D Rendering**
  - 3D Rendering indications in pediatric imaging are identical to those in the general imaging guidelines. See **Preface-4.1: 3D Rendering** in the Preface Imaging Guidelines

- **Nuclear Medicine**
  - Nuclear medicine studies are rarely used in the evaluation of peripheral vascular disorders, but are indicated in the following circumstances:
    - Lymphoscintigraphy (CPT® 78195) is indicated for evaluation of lower extremity lymphedema when a recent Doppler ultrasound is negative for valvular insufficiency.
    - Vascular flow imaging (CPT® 78445) is an obsolete study that has been replaced by MRA, CTA, or Duplex ultrasonography, and is not supported for any indication at this time.
    - Venous thrombosis imaging (CPT® 78456, CPT® 78457, and CPT® 78458) are obsolete studies that have been replaced by MRA, CTA, or Duplex ultrasonography, and are not supported for any indication at this time.
    - Radiopharmaceutical nuclear medicine studies (CPT® 78800, CPT® 78801, CPT® 78802 or CPT® 78803) can be approved for evaluation of the following:
      - Mycotic aneurysms
      - Vascular graft infection
      - Infection of central venous catheter or other indwelling device

The guidelines listed in this section for certain specific indications are not intended to be all-inclusive; clinical judgment remains paramount and variance from these guidelines may be appropriate and warranted for specific clinical situations.
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PEDPVD-2.1: General Information

Vascular and lymphatic malformations encompass a broad variety of conditions and have very heterogeneous natural history and treatment approaches. Lesions can be divided into low flow lesions (lymphatic, capillary and venous malformations), and high flow lesions (arteriovenous malformations and fistulas).

- Individuals with aggressive lesions being treated with systemic therapy can have imaging (see specific sections for details regarding modality and contrast level) approved for treatment response every 3 months during active treatment.
- Annual surveillance imaging of known vascular or lymphatic malformations can be approved for body areas where growth could cause significant organ dysfunction or functional impairment.

PEDPVD-2.2: Lymphatic Malformations

Lymphatic malformations are composed of dilated lymphatic channels filled with proteinaceous fluid and do not connect to normal lymphatic channels. They are typically soft, non-pulsatile masses with normal overlying skin.

- Ultrasound is indicated as an initial examination for superficial lesions.
  - Large lesion characterization may be limited by ultrasound imaging window.
  - Ultrasound is also limited in evaluating malformation relationship to airway or bony structures.
- MRI without contrast or without and with contrast of the affected body part is indicated for:
  - Lymphatic malformations involving deep tissues
  - Malformations too large to be completely imaged with ultrasound
  - Inconclusive ultrasound findings
  - Preoperative planning
- CT is of limited value in evaluating lymphatic malformations
  - CT with contrast of the affected body part can be approved for lesions with acute enlargement and concerns for compression when MRI is contraindicated.
PEDPVD-2.3: Venous Malformations

Venous malformations are slow-flow lesions characterized by dilated venous spaces and a normal arterial component. They are soft, compressible, non-pulsatile lesions that are usually blue to deep purple in color. Lesions can range from very small to large infiltrating ones. Some may change size with Valsalva.

Venous malformations are usually isolated, but they may be seen in multiple syndromes including Klippel-Trenaunay (KT) syndrome, Blue Rubber Bleb Nevus syndrome (BRBN), Maffucci syndrome, Proteus syndrome, Bannayan-Riley-Ruvalcaba syndrome, Parkes-Weber syndrome and congenital lipomatous overgrowth, vascular malformations, epidermal nevi and scoliosis/skeletal/spinal anomalies (CLOVES) syndrome.

- Ultrasound with Doppler is indicated as an initial examination for superficial lesions.
  - Large lesion characterization may be limited by ultrasound imaging window.
  - Ultrasound is also limited in evaluating malformation relationship to airway or bony structures.
- MRI without contrast or without and with contrast of the affected body part can be approved for venous malformations for preoperative assessment to evaluate the extent of malformation and their relationship to normal structures.
- MRA or CTA has a limited role in evaluating most venous malformations, but may be approved (contrast as requested of the affected body part) if MRI or CT are equivocal and the results will impact acute management decisions.
- CT can also be used to characterize venous malformations and their relationship to normal structures but is generally not as accurate as MRI.
  - CT with contrast of the affected body part can be approved when MRI is inconclusive or contraindicated
  - Both Klippel-Trénaunay syndrome and CLOVES syndrome have been found to have increased risk of venous thrombosis and pulmonary embolism, particularly after surgery or sclerotherapy. When pulmonary embolism is suspected in such patients, CT Chest with contrast with PE protocol (CPT® 71260) or CTA Chest (CPT® 71275) is indicated.

PEDPVD-2.4: Capillary Malformations

- Capillary malformations also known as port wine stains are characterized by a collection of small vascular channels in the dermis and generally do not require advanced imaging because the diagnosis is made clinically. However, MR imaging (without contrast or without and with contrast) may be approved to evaluate occult underlying neurologic structures, since these malformations are associated with encephalocele, spinal dysraphism, or Sturge-Weber syndrome.
**PEDPVD-2.5: Arteriovenous Malformations (AVMs) and Fistulas**

Arteriovenous malformations are characterized by a network of multiple abnormal vascular channels interposed between enlarged feeding arteries and draining veins. The arteriovenous fistula has a single communication interposed between a feeding artery and a draining vein. The normal capillary bed is absent in both lesions. Both lesions may have an aggressive clinical course and are characterized by a reddish pulsatile mass which has a thrill or bruit. Though often recognized at birth, these lesions may grow and present near adolescence.

- Ultrasound with Doppler is indicated as an initial examination for superficial lesions.
  - Large lesion characterization may be limited by ultrasound imaging window.
  - Ultrasound is also limited in evaluating AVM relationship to airway or bony structures.
- MRI without contrast or without and with contrast of the affected body part is also indicated for evaluation of AVMs, and is useful in evaluating the extent of AVMs and their relationship to normal structures.
- MRA (contrast as requested) of the affected body part can be approved for evaluation and surveillance of known AVMs.
- It is unusual for both MRI and MRA to be necessary for routine treatment response or surveillance imaging of AVMs, but both may be approved for preoperative planning.
- CT and CTA can also be used to characterize AVMs and their relationship to normal structures, but is generally not better than MRI and has associated radiation risks.
  - CT with contrast and/or CTA (contrast as requested) of the affected body part can be approved when MRI and/or MRA is inconclusive or contraindicated.

**PEDPVD-2.6: Vascular Tumors**

Vascular tumors include a variety of benign, borderline, and malignant tumors, which have variable clinical courses, including but not limited to Infantile Hemangiomas see PEDPVD-5: Infantile Hemangiomases, Epithelioid hemangioma, Kaposiform hemangioendothelioma, Kaposi sarcoma, Epithelioid hemangioendothelioma, and Angiosarcoma of soft tissue.

- Ultrasound with Doppler is indicated as an initial examination for vascular tumors.
  - Large lesion characterization may be limited by ultrasound imaging window.
  - Ultrasound is also limited in evaluating malformation relationship to airway or bony structures.
- MRI without contrast or without and with contrast of the affected body part is also indicated for evaluation of vascular tumors, and is useful in evaluating the extent of arteriovenous malformations and their relationship to normal structures, as well as response to therapy.
- MRA (contrast as requested) of the affected body part can be approved for evaluation and surveillance of known vascular tumors.
It is unusual for both MRI and MRA to be necessary for routine treatment response or surveillance imaging of vascular tumors, but both may be approved for preoperative planning.

CT and CTA can also be used to characterize vascular tumors and their relationship to normal structures, but is generally not better than MRI and has associated radiation risks.

- CT with contrast and/or CTA (contrast as requested) of the affected body part can be approved when MRI and/or MRA is inconclusive or contraindicated.

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PEDPVD-3.1: General Information
Systemic vasculitis is much less common in children than in adults, although the diagnostic pathways and treatment options are similar.

- PET/CT is considered investigational for management of pediatric vasculitis at this time.
  - There are limited data suggesting PET may have similar accuracy to MRA in the initial diagnosis of Takayasu arteritis but is not helpful in assessing treatment response and has not been shown to improve patient outcomes to date.

PEDPVD-3.2: Large Vessel Vasculitis
Takayasu arteritis is the predominant large vessel vasculitis occurring in children.

- Any of the following modalities may be indicated for evaluation of Takayasu arteritis:
  - MRA of the affected body area(s) (contrast as requested)
  - CTA of the affected body area(s) (contrast as requested)
  - Ultrasound with Doppler of the affected body area(s)

- Imaging is indicated at the following intervals:
  - Every 3 months for treatment response during active treatment in individuals being treated with systemic therapy.
    - See specific sections for details regarding modality and contrast level.
  - Annually for surveillance of known involved body areas to detect progressive vascular damage that may require intervention.

PEDPVD-3.3: Medium Vessel Vasculitis
Polyarteritis nodosa and Kawasaki Disease are the primary medium vessel vasculitides occurring in children.

- Imaging guidelines for Kawasaki Disease- see PEDCD-6: Kawasaki Disease in the pediatric cardiac imaging guideline.

- For evaluation of polyarteritis nodosa:
  - Any of the following modalities may be indicated:
    - MRA of the affected body area(s) (contrast as requested)
    - CTA of the affected body area(s) (contrast as requested)
    - Ultrasound with Doppler of the affected body area(s)
  - Imaging is indicated at the following intervals:
    - Every 3 months during active treatment with systemic therapy for treatment response.
      - For details regarding modality and contrast level see PEDPVD-1.3: Modality General Considerations
    - Annually for surveillance of known involved body areas to detect progressive vascular damage that may require intervention.
PEDPVD-3.4: Small Vessel Vasculitis

- Advanced imaging is not sensitive enough to detect changes in small vessels, and is not indicated for primary assessment of any small vessel vasculitis.

- End-organ damage occurs with several of the small vessel vasculitides. Advanced imaging is indicated for the following:
  - Henoch-Schönlein Purpura (HSP) is the most common vasculitis of childhood, mainly involving small blood vessels. Ultrasound abdomen (CPT® 76700) is commonly used to evaluate possible gastrointestinal complications (including bowel wall edema and hemorrhage, and intussusception) in known or suspected HSP, and should be approved when requested for that indication.
  - Granulomatosis with polyangiitis (GPA, formerly known as Wegener’s granulomatosis):
    - CT Sinuses (CPT® 70486) and/or CT Chest without contrast (CPT® 71250) or with contrast (CPT® 71260) is indicated for the following:
      - New or worsening clinical symptoms affecting the body area requested
      - To assess response to medical therapy when a change in treatment regimen is being considered
      - Annually-to evaluate the extent of disease
  - Eosinophilic granulomatosis with polyangiitis (EGPA, formerly known as Churg-Strauss Syndrome):
    - CT Chest without contrast (CPT® 71250) or with contrast (CPT® 71260) is indicated in the following circumstances:
      - New or worsening clinical symptoms affecting the body area requested
      - To assess response to medical therapy when a change in treatment regimen is being considered
      - Annually-to evaluate the extent of disease
  - Immune complex associated small-vessel vasculitis [immunoglobulin A–associated vasculitis (IgAV)]:
    - Doppler ultrasound of the affected body part (most commonly abdomen) is indicated in the following circumstances:
      - New or worsening clinical symptoms affecting the body area requested
      - To assess response to medical therapy when a change in treatment regimen is being considered
      - Annually-to evaluate the extent of disease
References


# PEDPVD-4: Disorders of the Aorta and Visceral Arteries

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PEDPVD-4.1: Thoracic Aortic Disease

**Familial Aortopathies**

- For Aortopathies such as the following:
  - Marfan
  - Ehlers-Danlos (EDS) - a genetic mutation known to predispose to aortic aneurysms/dissections (TGFBR1, TGFBR2, FBN1, ACTA2, or MYH11)
  - Loeys-Dietz
  - Familial thoracic aneurysm and dissections

- Screening: for Family history with first degree relative of aortopathy
  - Asymptomatic Individuals with no signs or symptoms of disease, whose first degree relative has no definitive gene defect, can have screening.
    - Echo (TTE) annually.

- Initial workup: Individuals with suspected aortopathies (gene positive, physical exam positive, or other findings) or definite disease associated with aortopathy
  - Echocardiogram (TTE) at the time of evaluation.
  - If the consideration is for Loeys-Dietz any of the following may be indicated in addition to the TTE at the time of work up:
    - MRA or CTA head
    - MRA or CTA neck
    - MRA or CTA chest
    - MRA or CTA Abdomen/pelvis
    - MRA or CTA of area of concern when there is an incidental finding on other imaging

- Surveillance: Suspected or known disease, but normal aortic imaging:
  - Individuals with suspected genetic aortopathies, but no disease, can have an echocardiogram to assess for change:
    - at 6 months
    - then annually
  - Individuals with Loeys-Dietz can be imaged with any of the following:
    - Echocardiogram
    - MRA or CTA of (any or all):
      - Head
      - Neck
      - Chest
      - abdomen
      - pelvis
  - Individuals with Loyes-Dietz can be imaged with the above at the following intervals:
    - at 6 months
    - then annually

- Surveillance: Suspected disease, and previous abnormal imaging
  - Individuals with abnormal thoracic imaging can be imaged with (both):
    - Echocardiogram
CTA or MRA of (any):
- chest
- abdomen
- pelvis
- head (Loyes-Dietz)
- neck (Loyes-Dietz)

The above imaging is indicated as follows:
- at the time of diagnosis
- in 6 months after diagnosis (if older than 2 years)
- then as follows based on the individual's age:
  - Individual's age 0 to 2 years:
    - every 3 months
  - Individual's age 3 to 12 years:
    - every 6 months
  - Individual's age 13 years and older:
    - every 12 months (if <4.5 or < 0.5 cm growth per year)
    - Every 6 months if ≥4.5 or ≥0.5 cm growth per year, or any Loyes-Dietz patient

If the diameter z score is increased, then a repeat study can be done prior to the next allowed study, to assess for rate of change.

If there are symptoms of dissection any or all of the following are indicated:
- Echo
- CTA or MRA of (any or all):
  - chest
  - abdomen
  - pelvis

For pediatric individual with dissection, imaging per vascular surgery and cardiology or any provider in consultation with vascular surgery at any interval.

Miscellaneous syndromes with potential aortopathy as major feature of congenital heart disease:
- Individuals with Turner syndrome see section CD-11.2.10: Aortic disease in Turner Syndrome in the Cardiac imaging guideline
- Williams syndrome See section PEDCD-2.4.10 in the pediatric cardiology imaging guideline
- Individuals with congenital heart disease would be managed based on PEDCD-2: Congenital Heart Disease in the pediatric cardiology imaging guideline

Miscellaneous disorders that can affect aorta, Osteogenesis imperfecta, Homocystinuria, polycystic kidney disease, Pseudo xanthoma elasticum, Hurler syndrome.
- Screening echocardiogram yearly.
- If positive findings, follow protocol for aortic root dilatation.
Follow-up of thoracic aortic abnormalities for other conditions please see discussions indicated elsewhere in the guidelines

- Coarctation of the Aorta- See PEDCD-2.3: Congenital Heart Disease
- Congenital rubella syndrome- See PEDCD-2.3: Congenital Heart Disease
- Kawasaki Syndrome- See PEDCD-6: Kawasaki Disease
- Neurofibromatosis- See PEDCD-1.2: Pediatric Cardiac Imaging Appropriate Clinical Evaluation

**PEDPVD-4.2: Aortic Congenital Vascular Malformations**

- Cardiac MRI without contrast (CPT® 75557) or without and with contrast (CPT® 75561), MRA Chest (CPT® 71555), CT Chest with contrast (CPT® 71260), or CTA Chest (CPT® 71275) may be indicated for evaluation.
- Vascular rings may impact both the esophagus and trachea. See PEDNECK-7: Esophagus and/or PEDNECK-8: Trachea for additional guidelines.

**PEDPVD-4.3: Visceral Artery Aneurysms**

- Visceral artery imaging indications in pediatric individuals are identical to those for adult individuals. See PVD-6: Aortic Disorders and Renal Vascular Disorders and Visceral Artery Aneurysms in the peripheral vascular disease imaging guidelines.

**References**


## PEDPVD-5: Infantile Hemangiomas

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**PEDPVD-5.1: Infantile Hemangiomas – General Considerations**

Infantile Hemangiomas are the most common benign tumor of childhood, occurring in close to 5% of infants. Infantile Hemangiomas typically have a phase of rapid and significant growth between 1 and 3 months of age; growth is usually completed by 5 months of age. Gradual involution then occurs, completed in 90% by age of 4 years, but with residual skin changes frequently persisting. Though usually not needed for diagnosis, biopsy can be done when needed to identify unique markers not found on other vascular tumors.

Most infantile hemangiomas do not require any imaging. Ultrasound with Doppler can be used when the diagnosis is uncertain, or with high risk clinical considerations. Other general imaging considerations for other vascular neoplasms regarding MRI, MRA, CT, and CTA also apply to infantile hemangiomas. See **PEDPVD-2.6: Vascular Tumors**.

- Multiple (5 or more) infantile hemangiomas can be associated with hepatic hemangiomas, with risk potential for high-output cardiac failure, and other risks see **PEDPVD 5.2: Multiple Infantile Hemangiomas**.
- High-output cardiac failure can also be caused rarely by large cutaneous infantile hemangiomas. Affected infants may present with “failure-to-thrive”, a hyperdynamic precordium, tachycardia, bounding pulses with a widened pulse pressure, and a palpable thrill and/or audible bruit over the hemangioma. This is an indication for cardiac evaluation, including echocardiography (CPT® 93303 ordered with CPT® 93320 and CPT® 93325).
- Life threatening risk of airway obstruction is associated with infantile hemangiomas of the lower face (“beard distribution”), or of the anterior neck, or of the oral and/or pharyngeal mucosa.
- Location-associated functional impairment can be found with periocular infantile hemangiomas larger than 1 cm (impairing vision), or infantile hemangiomas involving lip(s) or oral cavity (impairing feeding)
- Ulceration can occur with profuse bleeding that can be life threatening.
- Disfigurement risk is increased with large (5 cm or larger) infantile hemangiomas, facial or scalp infantile hemangiomas, and breast infantile hemangiomas in female infants.
- An infantile hemangioma at least 2.5 cm in diameter overlying the lumbar spine or sacrum is an indication to do a spinal ultrasound (under 6 months of age) and/or MRI Lumbar Spine without contrast (CPT® 72148) or MRI Lumbar Spine without and with contrast (CPT® 72158).
- Infantile hemangiomas 5 cm or larger in size have an increased risk of extracutaneous structural abnormalities.
- Other high risk indications include Syndromes or Associations with extracutaneous structural changes: for "PHACE(S) syndrome" See **PEDPVD-5.3: PHACE(S) Syndrome**, and for "LUMBAR syndrome" See **PEDPVD-5.4: LUMBAR Syndrome**.
When treatment is needed, imaging may be used to monitor response; consultation with a Hemangioma specialist may be useful in guiding evaluation, treatment, and follow up. The 2019 Clinical Practice Guideline of the American Academy of Pediatrics states “Unlike many diseases, management of IHs is not limited to 1 medical or surgical specialty. A hemangioma specialist may have expertise in dermatology, hematology-oncology, pediatrics, facial plastic and reconstructive surgery, ophthalmology, otolaryngology, pediatric surgery, and/or plastic surgery, and his or her practice is often focused primarily or exclusively on the pediatric age group.”

**PEDPVD-5.2: Multiple Infantile Hemangiomas**

Multiple (5 or more) hemangiomas- though hepatic hemangiomas can be asymptomatic, they rarely can cause a high flow rate that can cause high-output cardiac failure and can be potentially fatal. “Diffuse” hepatic infantile hemangiomas are a rare subset of hepatic hemangiomas at high risk for morbidity and mortality; affected infants usually present before 4 months of age with severe hepatomegaly, which can lead to lethal abdominal compartment syndrome with compromised ventilation, renal failure caused by renal vein compression, or compromise of inferior vena cava blood flow to the heart. Hepatic hemangiomas can also inactivate (via deiodination) thyroid hormones, causing risk of severe hypothyroidism.

- Multiple (5 or more) hemangiomas is an indication for Ultrasound with Doppler exam of the liver (CPT® 76700):
  - Initial imaging to look for hepatic hemangiomas
  - Repeat doppler ultrasound abdomen:
    - to monitor hepatic hemangiomas for progression
    - to monitor response to treatment.
**PEDPVD-5.3: PHACE(S) Syndrome**

"PHACE" (Posterior fossa malformations, Hemangiomas, Arterial anomalies, Coarctation of the aorta and Cardiac defects, and Eye abnormalities) syndrome or association (or "PHACE(S)" syndrome when also associated with Sternal cleft and/or Supraumbilical raphe) is frequently suspected when an infant has a large (5 cm in diameter or larger) infantile hemangioma of the face, scalp, or neck (risk of PHACE(S) Syndrome is then approximately 30%).

In rare cases, the face or scalp is not involved, with a large infantile hemangioma located on the torso and/or upper extremity instead. Cerebrovascular anomalies, present in more than 90% of individuals with PHACE(S) syndrome, are the most common extracutaneous feature of the syndrome, followed by cardiac anomalies (67%) and structural brain anomalies (about 50%).

- **Indications for imaging a young child for suspected PHACE(S) syndrome include the following:**
  - A large (5 or more cm in diameter) infantile hemangioma of the face, scalp, and/or neck.
  - Infantile hemangioma on face, scalp, or neck that is smaller than 5 cm in diameter, but with at least one major anomaly found in PHACE(S) syndrome, such as coarctation of the aorta or midline ventral defect.
  - Without any visible facial infantile hemangioma, PHACE(S) syndrome can also reasonably be suspected with the following:
    - an infantile hemangioma on upper chest or proximal upper extremity that is 5 cm or larger in size, with also major anomalies found in PHACE(S) syndrome
    - a large intraorbital infantile hemangioma.

- **When PHACE(S) syndrome is reasonably suspected, initial imaging would include the following:**
  - MRI Brain without contrast (CPT® 70551) or MRI Brain without and with contrast (CPT® 70553)
  - MRI Orbits without contrast (CPT® 70540) or MRI Orbits without and with contrast (CPT® 70543)
  - MRA Head without contrast (CPT® 70544) or MRA Head without and with contrast, (CPT® 70546)
  - MRA Neck may be done either without contrast (CPT® 70547), with contrast (CPT® 70548), or without and with contrast (CPT® 70549)
  - MRA Chest (CPT® 71555).
  - A screening transthoracic echocardiogram, CPT® 93303 (CPT® 93320 and CPT® 93325 are also indicated if ordered with CPT® 93303). If abnormalities are identified on echocardiogram, a cardiac MRI (CPT® 75557 or CPT® 75561) is then indicated.
  - If other clinical information or imaging shows involvement of the aorta, then MRI Chest without contrast (CPT® 71550) or MRI Chest without and with contrast (CPT® 71552) is also indicated.
Need for follow up or surveillance imaging is dictated by the results of the initial clinical and imaging assessment, and any subsequent clinical change. The most frequent follow up will be needed for those deemed at highest risk, including when the following has been found:

- Evidence of past arterial stroke
- Arterial stenosis or occlusions, with or without moyamoya-like vascular changes
- Structural brain changes, with neurosurgical evaluation clarifying the need for follow up.
- Changes in the aortic arch, coarctation of the aorta, and congenital cardiac anomalies, with pediatric cardiology evaluation clarifying the need for follow up. See PEDCD-2.5: Imaging and Surveillance per Congenital lesion

PEDPVD-5.4: LUMBAR Syndrome

The acronym “LUMBAR syndrome” refers to the association of Lower body infantile hemangiomas at least 5 cm in size (and other cutaneous defects), Urogenital anomalies and ulceration, “Myelopathy” (lipomyelocoele/lipo-myelomeningocele and/or tethered spinal cord), Bony deformities, Anorectal malformations and Arterial anomalies, and Renal anomalies. Though not exclusively true, there is a general regional correlation between the location of the cutaneous large infantile hemangioma(s) with underlying structural anomalies.

“LUMBAR syndrome” is reasonably suspected in a child with a large (5 or more cm in diameter) infantile hemangioma of any lumbosacral or perineal region or lower extremity. The following imaging is then indicated:

- Ultrasound spine (CPT® 76800) in infants up to 6 months of age, abdomen (CPT® 76700), and pelvis (CPT® 76856), with color Doppler.
- MRI Lumbar Spine without contrast (CPT® 72148) or without and with contrast (CPT® 72158) at 3 to 6 months of age, or earlier when either findings on an Ultrasound exam are inadequate or when requested by a hemangioma specialist or any provider in consultation with a hemangioma specialist.
- MRI of other relevant spinal level (relevance based on proximity of observed infantile hemangiomas larger than 5 cm) without contrast or MRI of the relevant spinal level without and with contrast.
- When ultrasound findings are inadequate and/or when recommended by a hemangioma specialist or any provider in consultation with a hemangioma specialist:
  - MRI Pelvis without contrast (CPT® 72195) or without and with contrast (CPT® 72197) and/or
  - MRI Abdomen without contrast (CPT® 74181) or without and with contrast (CPT® 74183).
- MRA Abdomen CPT® 74185 and/or Pelvis CPT® 72198, is indicated based on proximity of infantile hemangioma(s) at least 5 cm in diameter and/or other clinical evidence of vascular involvement, and/or when recommended by a hemangioma specialist or any provider in consultation with a hemangioma specialist.
- Infantile hemangioma of the lower extremity that is at least 5 cm in diameter is an indication for MRI of the relevant portion of the lower extremity without contrast.
(CPT® 73718) or lower extremity without and with contrast (CPT® 73720) and/or lower extremity joint without contrast (CPT® 73721) or lower extremity joint without and with contrast (CPT® 73723).

- When there is extensive lower extremity involvement with infantile hemangiomas the following are all indicated:
  - MRA (for both arterial and venous phase imaging)  Abdomen
  - MRA Pelvis
  - MRA Lower extremities
- Note: this should be reported as CPT® 74185 and CPT® 73725; the CPT® code for MRA Pelvis (CPT® 72198) should not be included in this circumstance.

References