Primary venous leiomyosarcoma: Cross-sectional imaging modalities, including CT, MRI, and F18-FDG PET/CT

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CASE SUMMARY
A 57-year-old man presented to the emergency department (ED) with 3 days of sudden onset left-sided flank pain radiating to his back. Past medical history was remarkable for pulmonary embolism with subsequent negative lower-extremity Doppler-ultrasound studies to rule out deep venous thrombosis. Other pertinent history included right anterior cruciate ligament tear, hypertension, hyperlipidemia, and depression. On review of systems the patient recalled bilateral leg swelling for several years, which he attributed to his knee problems.

IMAGING FINDINGS
A contrast-enhanced computed tomography (CT) revealed a large right-pelvic mass and adjacent right-iliac lymphadenopathy. Diffuse retroperitoneal inflammation and stranding surrounding the aorta and inferior vena cava (IVC) was noted. There appeared to be direct extension of the mass into the right common iliac vein and infra-renal IVC, which were dilated. A large tubular low-attenuation mass within the lumen of the IVC was concerning for tumor thrombus (Figure 1).

The patient was admitted for further workup, and a CT-guided core
biopsy of the pelvic mass was performed indicating high-grade sarcoma of muscle origin. Further imaging with gadolinium-enhanced magnetic resonance (MR) arteriography and venography revealed that the thrombus within the common iliac vein and IVC consisted mostly of enhancing tumor with some scattered areas of nonenhancing bland thrombus (Figure 2). A positron emission tomography/computed tomography (PET/CT) scan performed for staging prior to planned en-bloc resection demonstrated fluorodeoxyglucose (FDG) uptake within the IVC and right common iliac vein consistent with malignancy as well as metastasis to a right supraclavicular lymph node (Figure 3). Surgical removal of the pelvic mass was attempted, however, only partial removal could be achieved due to extensive intraoperative blood loss (> 6.5 L) from collaterals. A second procedure was performed to remove the involved IVC and thrombus one week later, following delayed primary closure with wound-vac device.

Final pathology revealed intravascular and extravascular components of high-grade sarcoma. Desmin IHC stain confirmed the smooth muscle origin of the tumor (Figure 4). The larger part of the leiomyosarcoma was found to be intravascular and tethered to the common iliac vein and IVC in the primary excision (Figure 5). A large part of the thromboembolus in the subsequent excision was found to be partially infarcted tumor thrombus (Figure 6). The expansile extravascular spread of the tumor and fragmentation of the specimen received from the OR did not allow for accurate determination of origin of tumor, however, the large tumor thrombus was more in keeping with an intravascular leiomyosarcoma.1

**DIAGNOSIS**
Primary venous leiomyosarcoma

**DISCUSSION**
Determining whether a sarcoma is of primary intravascular origin versus extravascular origin from a retroperitoneal mass that invades the vessel secondarily, is of paramount importance in guiding appropriate treatment and determining prognosis.2 The differential diagnosis of primary intravenous neoplasms includes retroperitoneal leiomyosarcoma, intravenous leiomyomatosis, endometrial stromal...
sarcoma, sarcomatoid renal cell carcinoma, angoisarcoma, and hemangioendothelioma. Treatment via surgical resection with or without adjuvant chemotherapy and radiation is not always curative, with local recurrence rates of 40% to 60%, and poor prognosis with median 3.5-year survival and 10-year survival of only 14%. Mean survival time in inoperable patients is about 1 month. The tumor arises from vascular smooth-muscle cells of the tunica media and may spread endoluminal, extraluminal or both, although extraluminal growth predominates. With extraluminal growth-pattern leiomyosarcoma is difficult to distinguish from other retroperitoneal masses.

Several signs on CT can help distinguish intravenous leiomyosarcomas from primary retroperitoneal tumors based on the plastic properties of vessel walls. A mass arising from the retroperitoneum will compress the vessel into a crescentic configuration, this is known as the negative-embedded organ sign. When the vessel is embedded in the periphery of the mass without being compressed the tumor likely arises from the vessel itself. Also known as the embedded organ sign, this is rarely seen in leiomyosarcoma. More commonly, the vessel itself is obliterated by the tumor so that the lumen becomes imperceptible. This phenomenon is likely due to the preponderance of extraluminal growth pattern of leiomyosarcomas and was found to have high specificity in detecting masses arising from the IVC in a recent study. A characteristic of intraluminal growth of tumor is that it expands the vessel. However, intraluminal tumor is not specific for intravenous leiomyosarcomas, as it can be seen with direct extension from retroperitoneal malignancy.

Another diagnostic challenge is that bland and tumor thrombi can coexist. The presence and extent of tumor thrombi influences staging and treatment, and may sometimes render patients unsuitable for surgical resection. The high rate of recurrence and dismal prognosis associated with tumor thrombosis make it all the more important to distinguish between bland thrombus. Tumor thrombus can be caused by primary tumor arising from the vessel wall as in leiomyosarcoma or by extension from an adjacent organ—most commonly in renal cell carcinoma, hepatocellular carcinoma, adrenocortical carcinoma, or retroperitoneal metastasis. The distinction between bland and tumor thrombus necessitates contrast enhancement whether in CT or MR cross-sectional imaging. By exploiting the inherent metabolic differences between tumor and bland thrombi, FDG-PET can be useful in this regard.

CT is the most accurate modality for detecting pulmonary embolism, a frequent complication of intravenous leiomyosarcoma. Neovascularity of tumor thrombus is visible on early arterial phase, which can also define extent into the right atrium. When evaluating the extent of thrombus delayed imaging should be used as typical portal-venous phase imaging is prone to artefactual filling defects due to venous admixture.

**FIGURE 4.** Desmin IHC stain demonstrating tumor (red arrow) and normal smooth muscle (blue arrow) in common iliac vein wall.

**FIGURE 5.** Intravascular leiomyosarcoma within the common iliac vein. Common iliac vein with high-grade leiomyosarcoma in tunica media (red arrow), adjacent normal smooth muscle (blue arrow) and normal tunica intima (green arrow) (H&E 20X).
MRI provides superior soft-tissue contrast resolution, allowing improved characterization of thrombus composition. A retrospective review of 10,908 oncologic patients using MRI identified a greater proportion of tumor thrombi than bland thrombi, 0.79% versus 0.5% respectively. Bland thrombus is typically isointense to muscle on T2, whereas leiomyosarcoma is isointense to hyperintense to muscle on T2 imaging. Enhancement by gadolinium is characteristic of tumor thrombi as with primary neoplasms. Intravascular leiomyosarcomas usually enhance, however, areas of necrosis can occur when tumors are large. In differentiating malignant versus bland portal-vein thrombosis in hepatocellular carcinoma, an increase of enhancement by >15% was established by Shah et al as characteristic for malignancy. Care must be taken not to confuse enhancement with collateral vessels around bland thrombi as is the case with cavernous transformation in portal vein thrombosis. When evaluating retroperitoneal pathology, it is helpful to acquire images with fat saturation.

Due to increased cellular density and altered nuclear-to-cytoplasmic ratio, neoplasms typically display restricted diffusion and decreased apparent diffusion coefficients. Recent studies have shown significantly higher apparent diffusion coefficient in bland thrombi versus thrombi in hepatocellular carcinoma on diffusion weighted images and even suggest DW imaging can be used as an alternative to contrast enhancement for thrombus characterization.

On PET/CT imaging, intense FDG uptake is usually seen in tumor thrombi due to high metabolic neoplastic activity and hypervascularity, while bland thrombus is typically less FDG-avid. PET has been useful in detecting neoplastic invasion of the IVC in cases of renal cell carcinoma, adrenal cortical carcinoma, colorectal cancers, and involvement of the superior mesenteric and portal veins in patients with neuroendocrine, primitive neuroectodermal tumors, hepatocellular carcinoma, and abdominal or pelvic malignancy. The value of adding contrast enhancement to the CT portion of PET/CT has been discussed in recent reports.

A recent study utilized a maximum standard uptake value (SUV) of 3.63 to differentiate tumor thrombosis from benign thrombus with a sensitivity of 71.4% and specificity of 90%. Our patient demonstrated intravascular maximum SUV of 5.6, however, one should be cautious of relying on absolute cut-off values of FDG uptake, which can vary significantly between patients and scans. More studies should be performed in this regard. Certainly the use of PET in this patient further defined the extent of vascular involvement, allowing for better surgical planning. In addition, distant metastatic disease was found in a right supraclavicular lymph node, which increased disease staging. Thus, in this particular patient, PET was instrumental in treatment planning.

**CONCLUSION** Although rare, intravascular neoplasms present a diagnostic challenge that can be met by utilizing multiple imaging modalities. Cross-sectional imaging with contrast-enhanced CT or MRI is paramount in diagnosing primary venous leiomyosarcoma, in differentiating tumor from bland thrombus, delineating intraluminal growth versus extravascular disease extension, and determining extent of disease, which influences surgical approach. FDG-PET with CT can confirm malignant involvement versus bland thrombus, and is instrumental in disease staging by detecting distant metastasis, thus influencing treatment planning. Using various modalities at
the diagnosticians disposal can lead to more accurate diagnosis and guide the most appropriate therapy.

REFERENCES