A 58-year-old woman was seen for follow-up in the urology clinic 18 months after undergoing resection of 2 renal cell carcinomas from her left kidney (partial nephrectomies). The pathology from the resections showed pT1a clear-cell renal cell carcinoma, with negative surgical margins. The patient was clinically well, with no flank pain or hematuria. In addition to her most recent cancer diagnosis, she had previously undergone a right radical nephrectomy 5 years prior for renal cell carcinoma (pT3a clear-cell renal cell carcinoma) and a distal pancreatectomy and splenectomy for an invasive pancreatic ductal adenocarcinoma 1 year prior. A genetic work-up to explain her multiple malignancies at a young age was negative. As part of her routine cancer surveillance, computed tomography (CT) with intravenous contrast showed new, enhancing, nodular tissue abutting the left perirenal fascia adjacent to the site of her previous partial nephrectomy (Figure 1). This was reported as most in keeping with recurrence of renal cell carcinoma. Blood tests (complete blood count, electrolytes, creatinine, calcium, phosphate, albumin, urea) done before the appointment were within normal limits and were all stable compared to earlier results.

The CT images were reviewed by her urologist (R.H.B.), who had performed the partial nephrectomies. Taking into consideration our patient’s negative surgical margins, the positioning of the nodules on her CT scan, and her history of multiple abdominal surgeries (including an open splenectomy), we thought it likely that the new soft tissue lesion could represent benign splenic regrowth. Local recurrence of renal cell carcinoma after partial nephrectomy is typically seen as an enhancing lesion in the parenchyma of the kidney at the base of the prior surgical resection, and we did not see this in the CT scan.

**Figure 1:** Axial (A) and coronal (B) contrast-enhanced computed tomography images in a 58-year-old woman, with a history of multiple renal cell carcinomas resected from left kidney, showing nodular soft tissue growth (red arrows) abutting the left perirenal fascia adjacent to site of previous partial nephrectomy.
Box 1: Differential diagnosis of enhancing soft tissue mass after abdominal surgery for cancer

- Local recurrence of previously resected malignancy
- Metastatic deposit of renal cell carcinoma or pancreatic adenocarcinoma
- Metastatic deposit of new primary malignancy
- Splenosis
- Peritoneal carcinomatosis
- Peritoneal mesothelioma
- Abdominal lymphoma

An accessory spleen (splenule) is a separate entity from splenosis. Accessory spleens form during embryologic development as a consequence of failed fusion and migration of the splenic buds. They are present in 10%–30% of the population and are typically located adjacent to the spleen (common sites include the splenic hilum, adjacent to the pancreatic tail and along the splenic artery) although various ectopic locations have been reported.5–7

Most patients with splenosis or accessory spleens are asymptomatic, and the splenic abnormality is found incidentally on imaging performed for another reason. When symptoms do occur, they are generally a result of the splenic tissue causing blockage of another organ (e.g., bowel obstruction or hydro- 

Figure 2: Positive heat-damaged red blood cell scan. Scintigraphy scan showing intense uptake in nodular tissue consistent with splenic tissue (red arrow). Expected hepatic uptake in the background is seen.

Discussion

Splenosis is the ectopic growth of benign splenic tissue often seen after splenectomy or splenic trauma.1 The presumed etiology is spillage of splenic cells at the time of the trauma or surgery to the spleen. These cells then grow over time and can exhibit contrast enhancement on cross-sectional imaging.2 Splenosis, which does not require resection or treatment, can mimic concerning processes such as metastatic malignancy, peritoneal carcinomatosis, peritoneal mesothelioma and lymphoma (Box 1).1,2

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On conventional imaging of a lesion, specific characteristics may raise suspicion of splenic tissue. On ultrasonography, splenosis or accessory spleens are often round with well-defined margins, uniformly hypoechoic centrally with a hyperechoic rim.5,7 On CT, these lesions are homogeneous and hypodense, and have similar contrast enhancement as normal splenic tissue.6 Magnetic resonance imaging (MRI) can also be used to assess a lesion suspected to represent splenosis or an accessory spleen. On MRI, splenosis appears dark on T1 and T2 series and bright on diffusion-weighted imaging.6 The contrast enhancement on MRI is similar to what is seen on CT.6 Because of its high sensitivity and specificity for splenosis, heat-damaged RBC scanning is considered the gold-standard imaging modality when this diagnosis is contemplated.8

Heat-damaged RBC scans are more specific than MRI studies for assessing splenosis.9 Before the test, a patient’s blood is drawn. Red blood cells are labelled with technetium-99m and heated with a specific protocol to elicit damage before being infused back into the patient.5 The test relies on the normal function of the spleen to filter and destroy damaged RBCs. Thirty minutes after the tracer is infused, the patient undergoes imaging. If splenic tissue is present, the heat-damaged RBCs will accumulate in this area and the tracer will be detected on imaging.

The ability to differentiate splenosis from growing malignant lesions is important. For our patient, confidence in the diagnosis of splenosis prevented the risk of invasive testing and the anxiety that goes along with uncertainty. Still, there are limitations to this technology. Use of radiotracers and scintigraphy requires the institutional expertise to produce reliable results. Imprecise technique may result in false-negative results.10 Thus, these scans are not available in many centres and may require referral to tertiary care facilities.

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