Wunderlich syndrome - unusual complication of spontaneous renal haemorrhage

Keywords: wunderlich syndrome, subcapsular, neoplasms, renal, vasculitis, arteriosclerosis, rupture, renal, artery, aneurysm, hypovolemic

Abbreviations: WS, wunderlich syndrome; RAML, renal angiomyolipoma; MDCT, multidetector computed tomography

Introduction

Wunderlich syndrome (WS) is a rare condition characterized by non-traumatic spontaneous acute renal hemorrhage into the subcapsular and perirenal spaces, found and described in this case. The WS is characterized by Lenk’s triad: acute abdominal pain, mainly in the flank, palpable mass and hypovolemic shock. Renal neoplasms are common cause for WS, and renal angiomyolipoma (RAML) is the most prevalent. However, several other causes must be ruled out, including renal malignant neoplasm, vascular diseases (vasculitis, arteriosclerosis, rupture of the renal artery aneurysm), kidney infection, undiagnosed blood dyscrasia and anticoagulant therapy.

Case presentation

A 50-year-old man went to the emergency service with an unspecific abdominal pain complaint. We performed a multidetector computed tomography (MDCT) of the abdomen, with and without iodinated intravenous contrast, that showed a hypodense nodule formation with regular and well delimited outlines, few heterogeneous foci and fat density (-35 HU), sizing about 1.8 cm from the inferior pole of the left kidney (Figure 1). The patient was relieved of the pain and was discharged from hospital in the same day.

Figure 1

MDCT of left kidney without intravenous contrast, demonstrating nodular mass with 1.8 cm in the lower pole, with areas of fat density (-35 HU).

After a month, the patient returned to the emergency service with a worse abdominal pain and an altered hemodynamic state. We performed another MDCT of the abdomen which detected the same nodule spontaneously dense and larger, with 2.2 cm. We also detected an extend subcapsular renal and retroperitoneal hematoma, all aspects related to a spontaneous renal bleeding (Figure 2).

Figure 2A

MDCT of the abdomen without intravenous contrast. Note spontaneously dense nodule associated with left renal subcapsular hemorrhage, and densification of adjacent fat.

Figure 2B

MDCT during nephrographic contrast phase. Observe the low enhancement of the nodular area, similar to the first examination but slightly larger.

Discussion

The display of fat in a kidney tumor by MDCT, as described in the case, is very suggestive of a RAML. Other kidney tumors can also present high levels of fat, such as the clear cell renal cell carcinoma, liposarcomas, atypical Wilms’ tumor and teratomas. However, a cortical renal mass mainly composed of fat (less than -20 HU) can be diagnosed as a RAML.
The RAML is the most common benign renal tumor and typically consists of smooth muscle, blood vessels and adipose tissue. A tumor occurs as an isolated entity, sporadic in 80% of the cases, and generally manifests itself in middle aged women. The other 20% develops associated with tuberous sclerosis. The tendency for bleeding is multifactor and includes foci deficiency of the elastic tissue in abnormally rigid and thick blood vessels, hypervascularization and venous invasion.

The spontaneous rupture of an angiomiolipoma is an infrequent urological emergency, although presenting a potential risk to life. The risk of bleeding is associated with a large tumor diameter. Tumors with less than 4cm of diameter have less risk of bleeding. Our case did not have this usual aspect and was a relatively small tumor.

The proper choice of treatment depends on the clinical state of the patient, the laboratory results, the rupture degree of the kidney, and the size of the retroperitoneal bleeding. Many authors have argued that the WS can be treated in a conservative way if the hemorrhage is self-limited and the patient responsive to resuscitation fluids. Others recommend conservative treatment for RAML of less than 6cm suggesting that only 34% of the patients with tumors bigger than 4 cm require intervention. In our case, the conservative treatment was our choice and is under serial and outpatient monitoring, with benign development for 1 year.

MDCT is the best imaging test for WS, and the existence of RAML may be presumed and conservative therapy implemented. Arteriography with embolization is an important therapeutic method in the acute phase when necessary, as it controls the bleeding, and avoids surgery in a considerable number of patients.

Conclusion

Despite rare, the WS must be immediately recognized through its findings in MDCT, regardless of its cause, because it can develop complications that can put the patient’s life at risk. The radiologist has a fundamental role in diagnosis and in conservative therapeutic follow-up in these cases.

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Conflict of interest

Author declares that there is no conflict of interest.

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