A 55-year-old African American man presented to the emergency room with 3 days of new-onset right-sided flank and abdominal pains. His abdominal pains were dull in nature, constant, moderate in severity (approximately 6 out of 10) and radiated to his back and right flank regions. There were no aggravating factors but the patient reported temporary relief with over-the-counter acetaminophen. The pains were associated with mild nausea and vomiting but he denied diarrhea, constipation, weight loss, fevers or other associated symptoms. His only known medical history was hypertension, controlled with lisinopril. Surgical, social and family history were unremarkable. Other than abdominal pains, a complete review of systems was negative.

His blood pressure was 117/78 mm Hg and his pulse was 82 bpm. His physical examination revealed right flank and right lower quadrant abdominal pain without rebound or guarding.

He underwent an abdominal CT scan which showed a 5.2 cm×4.2 cm hypodense solid right adrenal mass measuring 22 Hounsfield units (figure 1). Subsequent abdominal MRI demonstrated a 5.2 cm×4.8 cm×3.6 cm heterogeneous right adrenal mass with mass effect on liver and inferior vena cava (figure 2A). Axial diffusion-weighted image of the MRI showed a heterogeneous, significantly T2 hyperintense right adrenal mass (figure 2B). The patient’s blood work showed markedly elevated plasma fractionated metanephrines 805 pg/mL (<57 pg/mL) and elevated plasma normetanephrines 1938 pg/mL (<148 pg/mL) consistent with pheochromocytoma. He was started on phenoxybenzamine and metoprolol and underwent successful right adrenalectomy (figure 3). Histopathology was notable for chromaffin cells (figure 4A) with chromogranin A stain positive (100×), synaptophysin stain positive (100×).

Figure 1 Non-contrast CT abdomen axial view shows a hypodense solid 5.2 cm×4.2 cm right adrenal mass measuring 22 Hounsfield units, compressing the inferior vena cava.

Figure 2 (A) MRI abdomen axial T2 view shows a heterogeneous right adrenal mass measuring 5.2 cm×4.8 cm×3.6 cm displacing the inferior vena cava and the liver. (B) MRI abdomen axial diffusion-weighted image shows a T2 hyperintense right adrenal mass.

Figure 3 A 6 cm yellow–brown solid adrenal pheochromocytoma tumour, confined within the capsule.

Figure 4 (A) H&E stain showing nested tumour cells (200×), (B) chromogranin A stain positive (100×), (C) synaptophysin stain positive (100×).
positive (figure 4B) and synaptophysin stain positive (figure 4C), consistent with pheochromocytoma.

Pheochromocytoma is a catecholamine-producing tumour which usually arises from the adrenal medulla. If left untreated, it is potentially fatal because patients may develop hypertensive crisis, cardiac arrhythmias or myocardial infarction. It is extremely rare, with an annual incidence of two cases per million per year. Patients usually present with symptoms of excessive catecholamine production, such as palpitations, diaphoresis, headache and paroxysmal hypertension. Patients may also be asymptomatic or present with a variety of other symptoms mimicking other diseases, which may make diagnosis difficult.

Given the large size of his tumour, our patient presumably presented with flank and abdominal pains secondary to localised mass effects. Patients with pheochromocytoma presenting as our patient did is quite unusual and seldom reported.

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Learning points
► Pheochromocytoma is a rare disease in which patients typically present with palpitations, diaphoresis, headache and paroxysmal hypertension.
► Patients with pheochromocytoma may also present with atypical signs and symptoms which can delay diagnosis or make diagnosis difficult.
► It is crucial in patients with adrenal tumours, even when found incidentally, be evaluated for pheochromocytoma because it is a potentially fatal disease.