Dear Editor,

A 44-year-old female presented with a 7-year history of paroxysmal episodes of dyspnea, headache, palpitation, tremors, and hypertension. In each episode, there had been a sudden onset of symptoms, with no triggering factors, and spontaneous improvement after approximately 15 min. On physical examination, she presented no relevant findings or comorbidities. At hospital admission, she reported having had episodes of palpitation, tachycardia, and profuse post-micturition sweating, remaining asymptomatic between episodes. She was submitted to computed tomography (CT) and magnetic resonance imaging (MRI), as shown in Figures 1A and 1B, respectively. The CT scan, with intravenous administration of contrast medium, revealed a nodular lesion, measuring 3.5 × 3.0 cm, with lobulated contours and increased density in its soft parts, showing intense, heterogeneous enhancement, in the anteroinferior wall of the bladder. On MRI, the lesion presented a lobular pattern, with a heterogeneous signal on T2-weighted sequences, a predominance of isointense signals, and foci of hyperintense signals in its center. Surgical resection of the lesion (partial cystectomy) was performed. Examination of the surgical specimen, retrieved from the right anterior wall of the bladder, showed a yellowish tumor measuring 3.0 × 3.0 cm, with a macroscopic appearance similar to that of adrenal tissue (Figure 1C). The pathological examination of the specimen revealed extra-adrenal paraganglioma and tumor-free margins (Figure 1D). In the postoperative period and during the remainder of the hospital stay, the patient did not present any of the adrenergic symptoms previously reported.

Pheochromocytomas are tumors of the sympathetic nervous system and can be functioning or nonfunctioning, sometimes secreting catecholamines, thus causing paroxysmal hypertension, palpitations, headache, and syncope\(^1\). They are most common between the fourth and sixth decades of life. Approximately 10% are bilateral, 10% are malignant, 10% occur in children, and 10% are extra-adrenal. More than 90% are located in the adrenal gland, and 98% are intra-abdominal. Pheochromocytomas can occur anywhere from the base of the skull to the bladder; when located outside the adrenal gland, they are known as paragangliomas\(^2\). Pheochromocytoma of the urinary bladder is a rare tumor, originating from chromaffin cells of the sympathetic nervous system and located within the bladder wall, accounting for 0.06% of all bladder tumors and 6% of all paragangliomas\(^3\). In the bladder, it can produce symptoms typical of pheochromocytoma, including hematuria and micturition syncope resulting from the release of catecholamines by bladder contraction. In 10–15% of cases, paragangliomas of the bladder are nonfunctioning; another 10% show hormonal activity without clinical expression\(^4\).

Recent studies have discussed the role of imaging examinations in the investigation of pelvic lesions\(^5\)–\(^10\). The diagnostic imaging methods used in the investigation of pheochromocytomas include ultrasound, CT, MRI, and scintigraphy. For the detection of adrenal pheochromocytomas > 1.0 cm in diameter, CT and MRI have a sensitivity of nearly 95% and 100%, respectively, and MRI has greater specificity than does CT\(^11\). On MRI, pheochromocytoma typically manifests as an expansive lesion with low signal intensity on T1-weighted sequences and high signal intensity on T2-weighted sequences, with intense impregnation after contrast administration. However, in rare cases, pheochromocytoma can present low signal intensity on T2-weighted sequences\(^2\). The treatment of choice for paraganglioma is surgical resection, because most are benign and can be completely resected\(^12\).
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Carpal boss syndrome: os styloideum fused to the trapezoid

Dear Editor,

A 29-year-old White female presented with chronic pain on dorsiflexion of the right hand and a hard prominence, which was painful on palpation, at the base of the second and third metacarpal muscles. An X-ray of the hand (Figure 1A) revealed a bony prominence in the region identified as palpable in the physical examination, as well as showing that there was lack of definition of the joint space between the trapezoid and the capitate. In multiplanar and three-dimensional computed tomography reconstructions, which provided greater detail (Figures 1B and 1C), an os styloideum was seen to be fused to the trapezoid bone and in neoarticulation with the base of the third metacarpal. Magnetic resonance imaging showed a hypointense signal on a T1-weighted image (Figure 1D) and increased intensity in a T2-weighted short-tau inversion-recovery sequence, with bone edema adjacent to the neoarticulation, which is indicative of apophysitis.

Os styloideum is an anatomical variation characterized by an accessory ossicle on the dorsum of the wrist, between the trapezoid and capitate, at the base of the second and third metacarpal bones. When it produces symptoms, mainly local pain, it is known as a carpal boss. The true incidence of carpal boss syndrome is unknown; it is probably underestimated and often confused, clinically, with other causes of tumor in the dorsum of the carpus.

Although a carpal boss can be classified as acquired (osteophytic), congenital (secondary to os styloideum), or of mixed etiology, the clinical presentations appear to be similar across the...