PARAGANGLIOMA OF URINARY BLADDER - A RARE CASE

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ABSTRACT: A 27 year old male presented with hematuria & dysuria since 9 months. Hematuria was transient at times, pain less & terminal. History of passing clots occasionally. Ultrasound revealed echogenic isolated mass measuring 5x4.6 cm arising from anterolateral wall of bladder with evidence of small foci of calcification at the periphery. Partial cystectomy was done—but marked fluctuation in blood pressure and tachycardia was observed during the procedure. The histopathological findings were diagnostic of paraganglioma of the urinary bladder. The diagnosis was confirmed by means of immuno histochemical studies.

KEY WORDS: Paraganglioma, extraadrenal pheochromocytoma, bladder neoplasms.

INTRODUCTION: Paragangliomas originating in urinary bladder are extremely rare, arising from chromaffin tissue of the sympathetic nervous system. It is usually benign. The tumor usually develops in young adult women. Common symptoms and signs are dysuria, hematuria, and hypertension. We report a single case of non-functional paraganglioma of the urinary bladder clinically diagnosed as transitional cell carcinoma.

CASE REPORT: A 27 years old male presented to outpatient clinic of urology department with a 9 month history of hematuria and dysuria. He had no significant past medical history. His family history was non contributory. Physical examination was unremarkable, no organomegaly was observed. At admission, his blood pressure was 120/80mm of Hg, pulse rate was normal. Both ultrasound and CECT demonstrated isolated, enhancing sessile mass measuring 5x4.6 cm arising from right anterolateral wall of urinary bladder with specks of calcification. Perivesical fat planes were preserved. Wall thickness appeared normal.

On the basis of the clinical diagnosis of primary bladder tumor, patient was admitted for partial cystectomy. During surgery, marked fluctuation in blood pressure with rise of blood pressure upto 230/80 mm of Hg and pulse rate of 200/minute was observed. Patient developed intraoperative cardiac arrest and was revived after DC shocks and was kept on ionotropic...
infusion and mechanical ventilator. He attained conscious state on 2nd day of surgery. On follow up, his blood pressure and pulse rate are within normal limits.

MACROSCOPIC AND MICROSCOPIC FINDINGS: Grossly specimen received consisted of two soft tissue masses showing brown to yellow color. Largest mass measured 5x2.5x1 cm and smaller mass 3x2x1 cm. On cut section, the both masses displayed uniform/homogenous yellowish brown appearance with no evidence of hemorrhage or necrosis.

Histopathological examination revealed a neoplasm composed of tumor cells arranged in nesting pattern andzell ballen pattern with delicate fibrovascular stroma. Individual cells are large round to polygonal with distinct cell border, abundant eosinophilic granular cytoplasm and vesicular nucleus. Mitoses are absent. A provisional histopathological diagnosis of paraganglioma of urinary bladder was made.

On immunohistochemistry, this bladder tumor was negative for epithelial marker cytokeratin and positive for the neuroendocrine markers – chromogranin, synaptophysin. Sustentacular cells showed positivity for S-100, thereby confirming the diagnosis of Paraganglioma of urinary bladder.

DISCUSSION: Paragangliomas are extra adrenal neoplasms arising from chromaffin tissue of the sympathetic nervous system. According to the latest edition Campbell Walsh Urology 1, trigone and posterior wall of urinary bladder have been stated as the most common sites for paraganglioma, while in our case, the tumor arised from the anterolateral wall. The first case of paraganglioma of the urinary bladder was reported by Zimmerman2 in 1953. Paraganglioma of the urinary bladder accounts for <1% of all bladder tumors and 0.6% of extra adrenal pheohromocytomas 3. In most cases, paragangliomas of the urinary bladder often causes micturitional attacks, headache, palpitation, fainting and visual disturbances. No such symptomatology was seen in our case. During surgery, marked fluctuation in blood pressure with tachycardia was observed 4.

The tumor usually show females preponderance in (F: M = 3:1), occurring in 20-40 years of age, while this case was 27 years old male.

In most series patients may present with headache, palpitation, paroxymal HTN due to catecholamine excess especially during micturition. As the patient in our study had none of these symptoms, hence no endocrine tests were performed 5. As many as 50% of the paragangliomas are hereditary and may be associated with familial paragangliomas, neurofibromatosis type1, Von Hippel Lindau disease and the Carney triad 6.

Macroscopically, these tumors are red-brown lobulated solid, submucosal or intramural masses covered by intact epithelium. They are as large as 10 cm but most are only a few centimeters in greatest dimensions. They usually appear as fungiform rounded or pedunculated masses that bulge into the lumen with variable ulceration.

On microscopic examination: tumor shows nests of cells arranged in zellballen pattern or trabecular architecture or a mixture of the two, diffuse or solid architecture can also be seen7. Individually, tumor cells are large with abundant granular and basophilic to amphophilic cytoplasm with cellular and nuclear pleomorphism. Sometimes prominent nuclear pseudoinclusions are present in some cases. Spindle cells are present in about 2% of cases.

A small cell variant is also described. Intracytoplasmic hyaline globules occur commonly. These are PAS positive and diastase resistant. Hemorrhage and hemosiderin deposits are common. Oncocytic variants have also been reported.
IHC paragliomas are positive for chromogranin-A, synaptophysin and NSE. S-100 protein demonstrates sustentacular cells.

**CONCLUSION:** Bladder paragangliomas may be misdiagnosed when characteristic symptoms are absent. Although it shows female preponderance, our case is a 27 year male. It should be considered as a differential diagnosis in neoplasm of urinary bladder. Laparoscopic partial cystectomy may be the first choice in treating paraganglioma of the urinary bladder. Recurrence and metastasis though infrequent, have been reported in the literature; therefore, long term follow up is required.

**GROSS:** Both masses displayed uniform/homogenous yellowish brown appearance with no hemorrhage or necrosis. Largest measuring 5x2.5x1 cm and smaller 3x2x1 cm.

**HPE:** LP (10x) view: neoplasm composed of tumor cells arranged in nesting and zell ballen pattern with delicate fibrovascular stroma. HP view (40x): Individual cells are large round to polygonal with distinct cell border, abundant eosinophilic granular cytoplasm and vesicular nucleus. Mitoses are absent.
I H C: Tumor showing positivity for neuroendocrine markers – Chromogranin A

REFERENCES: