Case Report:
Retroperitoneal Ancient Schwannoma

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Abstract: Schwannomas are rare tumors which arise from nerve sheath and are mostly benign in nature. They are usually located in the head, neck and flexor surfaces of extremities. Schwannomas are very rare in the retroperitoneal region. Amongst all schwannomas 0.7% of benign ones and 1.7% of malignant ones are located in the retroperitoneum. Preoperative diagnosis is difficult because of vague symptoms. We report a case of retroperitoneal schwannoma in a 70 years female patient because of its rarity and unusual location.

Key Words: Ancient schwannoma; Retroperitoneum

Introduction:
Schwannomas (Neurilemomas, Neurinomas or perineural fibroblastoma) are benign tumors arising from nerve sheath of Schwann cells. The etiology is still unknown.(1,2) They usually affect adult patients in the age group of 20-50 years and are more common in males.(3) The usual location is in head, neck and on flexor surfaces of extremities, retroperitoneum being the rare site with the incidence being 0.3-3.2%. Retroperitoneal schwannomas account for 1-10% of all primary retroperitoneal tumors.(1) Majority of retroperitoneal schwannomas are benign in nature. We describe a case of incidentally diagnosed retroperitoneal ancient type schwannoma in a female patient because of its rarity and unusual site.

Case Report
A 70 years female came to medicine department with complaints of epilepsy and vomiting. Ultrasonography revealed a round to oval heteroechoic mass near the lower pole of right kidney in the paracaval region. Other investigations were normal. Exploratory laparotomy was done, mass was excised and specimen was sent for histopathology.

Pathological examination: On gross examination it was a single, encapsulated, well-circumscribed, globular, grey white to grey brown mass measuring 7x6x4 cm. Cut section revealed a grey white to grey brown variegated appearance with cystic areas, hemorrhages and areas of calcification. Microscopically it was a well circumscribed, encapsulated tumor showing hypercellular and hypocellular areas. Elongated spindle shaped neoplastic cells showed tapering nuclei with nuclear palisading. Occasional area showed enlarged, hyperchromatic nuclei with very low mitotic activity. Areas of cystic degeneration, myxoid change and ossification were noted.

Figure 1: A well circumscribed tumour, cut section showing variegated appearance, cystic areas, hemorrhages and calcification.
Figure 2: Tumor with hyper and hypocellular areas, nuclear palisading, cystic change. Inset: Ossification. (400x, H&E)

Discussion:
Retroperitoneal schwannomas are rare tumors and are usually detected incidentally after radiological examination. Neurological symptoms are rare. Few cases present with abdominal and lower back pain and digestive disturbances. The tumor is usually detected with pre-operative cross sectional imaging which reveals a solid mass in the retroperitoneum with a few cystic areas. Though ultrasound and computed tomography can detect the tumor, MRI allows better visualization and involvement of other organs. These radiological findings are usually non-specific and fail to give accurate diagnosis on most of the times. Definitive diagnosis relies on histopathological examination of the tumor. CT guided biopsy can be attempted for preoperative diagnosis. Kapan M stated the limited role of this investigation due to the risk of hemorrhages, infection and tumour seeding.

Complete surgical excision is the only treatment for these tumors as they are not sensitive to radiotherapy and chemotherapy. Prognosis of benign schwannomas is good. Only 5-10% cases show recurrence because of incomplete excision. Grossly schwannomas are encapsulated, g loboid masses of variable size. Retroperitoneal schwannomas are usually large tumors. Cut section is glistening. Ancient schwannomas show infarct like necrosis, cystic changes, calcification and ossification related to degeneration. Microscopically conventional schwannomas are composed of neoplastic schwann cells showing hyper and hypocellular areas, elongated cells with nuclear palisading. Stroma shows thick walled dilated vessels. Ancient schwannomas show nuclear polymorphism including bizarre forms with nuclear inclusions and other secondary changes like calcification and ossification.

Differential diagnosis include paranglioma, pheochromocytoma, liposarcoma, malignant fibrous histiocytoma, retroperitoneal cystic masses such as hemangiom a and lymphangioma.

Conclusion:
Retroperitoneal ancient schwannomas are rare types of soft tissue nerve sheath tumors which are difficult to diagnose preoperatively due to vague symptoms. Definitive diagnosis relies on histopathology. CT guided FNAC should be tried to increase the accuracy of preoperative diagnosis. Treatment of choice is complete excision of mass. Recurrence and malignant transformation in retroperitoneal ancient schwannomas is rare.

References: