Objective
We present here a case of retroperitoneal ancient schwannoma and review this rare entity under the guidance of recent literature.

Case
An 66-year-old men admitted to our clinic with a retroperitoneal mass in plain radiography without any symptoms. F-18 FDG PET- CT scan revealed 26 x 19 mm regularly contoured hypodense hypermetabolic nodular lesion in right inferior quadrant leaning on ileopsoas muscle. Histopathology from a tru-cut biopsy and was reported as mixoid spindle cell tumor revealing mixoid liposarcoma. The patient underwent total excision of the retroperitoneal mass. Postoperative pathologic diagnosis was reported as ancient schwannoma. The patient was recommended observation without any adjuvant treatment and is free of recurrence in postoperative 10th month.

Conclusion
Ancient Schwannomas are originating from perineural Schwann cells. They are accepted as benign, but some may be mistakenly termed malignant. As they are well encapsulated, recurrences following complete surgical excision are uncommon.

Anahtar kelimeler: Ancient, Retroperiton, Schwannom

Introduction
Schwannomas are benign tumors of peripheral nerve sheath. They usually occur as solitary encapsulated mass. It has a slight female predominance mostly seen between the second and fifth decades of life(1). The frequently seen clinical sites are head and neck and extremities. Retroperitoneal schwannoma is rare except the cases in associated with von Recklinghausen’s disease (2). Only 0.5% to 12% of retroperitoneal tumors and 0.5% to 5% of all schwannomas are retroperitoneal schwannomas. Malignant transformation is infrequent. The term “ancient schwannoma” refers to the degenerative changes, which develops with increasing duration (age) (3) Nuclear atypia may also be present in these tumors and may be mistaken for malignant change. Complete surgical resection results in cure and recurrences are uncommon. We describe a retroperitoneal ancient schwannoma case treated at our Institute.

Case Report
A retroperitoneal mass was realised in plain radiography of 66-year-old men without any symptoms. Clinical examination was normal.

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underwent a tru-cut biopsy and histopathology was reported as mixoid spindle cell tumor revealing mixoid liposarcoma. Keeping this report in view; the patient was recommended total excision of the retroperitoneal mass. In the intraoperative observation the close relation of the tumor with the adjacent peripheral nerve was remarked.

Postoperative macroscopic outlook was a mixed solid and cystic mass with hemorrhagic surface. Microscopic slices revealed a well capsulated lesion including cystic and hemorrhagic areas. Pathologic diagnosis was reported as ancient schwannoma (Figure 2a - 2b).

Treatment: The patient was recommended observation without any adjuvant treatment. She is free of recurrence in postoperative 4th month.

Discussion
Epidemiology-Clinical Features
Schwannomas or neurilemmomas are benign soft-tissue tumors, which arise from Schwann cells of the peripheral nerve sheaths. They are commonly originated from head and neck and the extremities (4) Retroperitoneal schwannomas account for 0.3% to 3.2% of these tumors (5). Ancient schwannoma is a rare variant of schwannoma first described by Ackerman and Taylor in 1951. The term ‘Ancient’ verbalize characteristic hypocellular and degenerative areas which is believed to occur with the long term progression of the tumor. They constitute 0.8% of all soft-tissue tumors (6). Patients are mostly diagnosed between ages 40 and 60 years and the ratio between men and women is 2:3(7).

Most cases are asymptomatic and incidentally discovered during medical examination or imaging. Infrequently they can cause pressure effects on surrounding large nerves (8). A recent study on clinical features of retroperitoneal schwannoma also confirmed that symptoms were nonspecific, and neurologic symptoms were rare (9). Those nonspecific symptoms can be hypertension, abdominal pain, flank pain,
renal colic pain, hematuria and headache (7). Mostly, due to the large loose areolar space, the clinical course is slow and and malignant change is uncommon (10).

**Radiologic Findings-Diagnosis**

Although ancient Schwannoma may exhibit some certain radiologic features, due to absence of pathognomonic ones 13 preoperative diagnosis is challenging in these cases. In ultrasonography lesion is seen as a well-defined, solitary and oval or sphere-shaped mass with regular contours that is hypoechoic or of mixed echogenicity. (5).

CT scans can reveal low or mixed attenuation. Cystic changes due to central cystic necrosis, are common in retroperitoneal schwannomas according to other retroperitoneal tumors. Density in CT-scan depends on the lesional rearrangements which can be cystic, necrotic, haemorrhagic or in the form of calcification. Pure cystic forms are also reported (11). CT-scan provides a better analysis of the bone extension. Besides, the opportunity of wide field examination with the adjacent vital organs enabling spatial analysis is the major advantage of CT scan in terms of pre-surgical evaluation of patient (5).

MRI provides better visualization in terms of tumor’s origin, vascular architecture, and involvement of surrounding organs in large retroperitoneal tumors (9), in other words, the tissue analysis is better with MRI (5). In MRI, the association of a T1 isosignal at the adjacent skeletal muscle, with a T2 hypersignal is reported frequently however it is non-specific. T2 signal intensity is inversely proportional to the cellularity of the tumour. So, Antoni type B which is hypocellular is associated with intense T2 hypersignal. The contrast enhancement shows tissue portions, partitions, septa and walls of the lesion. High intensity of rearrangements reveals more heterogeneous contrast enhancement (12).

Some authors advocate angiography for assessment of vascularisation in the tumour and/or evaluate presurgical embolisation (13). CT-guided biopsy and fine-needle aspiration are reliable methods of diagnosis for retroperitoneal schwannomas (8). Presence of heterogeneity and areas of degeneration on CT or MRI, cause a suspicion of malignancy. Therefore obtaining tissue diagnosis before embarking on surgical resection is recommended (14).

CT-guided biopsy works if sufficient Schwann cells for microscopic visualization is obtained. This procedure is not supported as a diagnostic tool by many investigators who based their contest on the risks of hemorrhage, infection, and tumor seeding (2). Duodeno-pancreatic sphere technically provides a good and efficient alternative, especially for the small lesions (under 3 cm) (12).

Efficacy of fine needle aspiration cytology(FNA)is not well established, while cellular pleomorphism leads to misinterpretation as malignancy (2) In more complicated cases above mentioned biopsy techniques fail, preoperative incision biopsy was advocated as an alternative by some authors. However these preoperative diagnostic procedures can be omitted in some cases while the tumor mostly has an intact smooth capsule, without any adjacent organ involvement despite the radiologically demonstrated heterogeneity raising the probability of malignancy (15). The differential diagnosis of retroperitoneal schwannomas includes neurofibroma, paraganglioma, pheochromocytoma, liposarcoma, malignant fibrous histiocyotma, lymphangioma, and hematoma (9).

**Histopathologic Features**

These tumors are macroscopically encapsulated and histologically includes dense, highly cellular Antoni A areas, and loose Antoni B areas of myxoid stroma (10). Predominat degenerative findings such as cystic degeneration, fibrosis, stromal edema, xanthomatosue change, and perivascular hyalinization are some of the histologic patognomonic features (7, 10). Pleomorphism, lobulation, or hyperchromasia due to degenerative nuclear changes are also common (6) Plenty of theories raised in an attempt to explain this degenerative process. One theory is degeneration of Antoni B areas leading to cyst formation while growing, a second one advocates central ischemic necrosis due to increasing tumor causing cysts within the tumor (16).

**Treatment Considerations-Prognosis**

Primary treatment of choice for retroperitoneal schwannoma is complete excision in healthy patients. If not feasible, partial or subtotal resection can be done to protect adjacent organs or neurovascular structures, although tumour recurrence may be inevitable in such cases (17). Controversy on this issue is proceeding especially when adjacent tissue or viscera is to be sacrificed for clear margins. Some investigators impose complete excision, if necessary, taking the risk of sacrifice of adjacent tissue (2). Others advocate more conservative surgical procedures like simple enucleation or partial excision accepting they are sufficient. After intralesional enucleation reported local recurrence rates range from 16% to 54% (7). Some other studies reported recurrence rates ranging from nonexistent to greater than 50% with simple enucleation (18). Complete surgical resection is recommended because frozen section analysis is not adequate to exclude malignancy (7). Considering the frequent concern for malignancy with ancient schwannoma due to the heterogeneous appearance on imaging, radical excision is usually the recommended treatment in these situations (19).

Laparoscopy is also safe and effective alternative modality for schwannomas, offering better visualization in narrow anatomic spaces. Although successful laparoscopic resection even near the great vessels has been described there may be an increased risk of hemorrhage, leading to unsuccessful resection or intraoperative mortality. Laparoscopic resection of larger tumors (> 5 cm) may entail longer hospital stays than even with open resection (2).

Prognostic behaviour of ancient schwannomas are similar to other schwannomas. Absence of mitoses and preservation of spindle shaped cells can be helpful for differentiation from malignancy. Complete surgical resection with preservation of the surrounding structures mostly ensures cure in these patients (19).

**Conclusion**

We presented a case of an unusual pathological entity, retroperitoneal ancient schwannoma. These are slow growing, clinically indolent behaving benign tumors, which
may present with pressure symptoms in some cases. Surgical resection provides cure.
Retroperitoneal schwannomas are usually identified incidentally on tomographic images. Diagnosis is based on histopathological examination after surgery and immunohistochemical examination. Total excision of the tumor had the best prognosis.

References