Case Report

A Rare Case of Retroperitoneal Liposarcoma and Castleman’s Disease

HARESH KUMAR GOBU1, SAMEER BABU UMMER8, GANESH CP BABU3, NILAKANTAN ANANTHAKRISHNAN

ABSTRACT

Retroperitoneal liposarcoma is a malignant mesenchymal tumour. The incidence is rare and early diagnosis is difficult as they present with non-specific symptoms. A 55-year-old male patient, presented to General Surgery Department with complaints of abdominal swelling since two months. Patient also complained of discomfort for three to four months duration followed by the swelling. With recent onset of dull, intermittent and non-radiating pain in the right upper abdomen since two weeks. The pain had no specific aggravating or relieving factors. There was associated nausea and vomiting. Vomiting was non-projectile in nature, containing partially digested food particles, non-foul smelling, non-bilious and was not blood stained. During the same two weeks, patient also complained of decreased appetite and generalised fatigue. No similar episodes or complaints in the past were noted.

On physical examination, the vitals recorded were pulse rate 82/min, blood pressure 130/80 mmHg and respiratory rate 21/min. Per abdomen examination showed fullness along right abdominal quadrants and a mass of 15×12 cm with no clear margins was palpated in the right quadrants extending to midline at the level of umbilicus. The mass was soft to firm in consistency, did not move with respiration or fall forwards. Mass was dull on percussion with no evidence of free fluid. Laboratory investigations showed low Haemoglobin-8.2 g/dL, total leucocyte count of 5000 cells/µm and other parameters (renal and liver function tests) were within normal limits. On performing Ultrasonography (USG) abdomen, an ill-defined heterogeneous mass occupying right hypochondrium, right lumbar and right iliac fossa was reported [Table/Fig-1].

With origin being uncertain, thoracic-abdominopelvic contrast enhanced computed tomography (CECT) performed showed a large well-circumscribed low attenuated mass lesion with epicentre in right anterior pararenal space measuring 26×22.9×14.5 cm crossing the midline to left with compression over right kidney and renal pelvis posteriorly. Mixed areas with minimal calcification and areas of necrosis were noted. Renal pelvis stretched and the ureter displaced anteromedially with resultant Grade 2 hydronephrosis [1]. The mass was visualised abutting the inferior surface of liver, gallbladder and inferior vena cava. The small bowel was displaced to opposite side. The CECT imaging showed no local nodal or distant metastases [Table/Fig-2]. No differential diagnosis was made in view of its retroperitoneal origin and clinical correlation.

Preanaesthetic evaluation showed no contraindication except for low haemoglobin which was optimised using two units of Packed Red Blood Cells (PRBC). With adequate cardio pulmonary status, reserved additional PRBC, patient underwent laparotomy three weeks from time of presentation with the aim of en-bloc resection.

In view of stretched right ureter, Double J (DJ) stenting was done. This was followed by transverse incision two to three cms below the umbilicus extending between both anterior superior iliac spine. A large lipomatous tumour measuring 29×28×17 cm and weighing 4.8 kg was extracted from the retroperitoneal space with meticulous dissection without solid organ or visceral organ involvement [Table/Fig-3,4]. Although the large mass displaced bowel, it was returned to normal position. No secondary lesions in the liver or abdominal nodes was identified.

The histopathological examination showed circumscribed tumour composed of adipocytes separated by fibrous septa. Individual adipocytes showing nuclear vacuolation with occasional lipoblasts corresponding to low-grade, well-differentiated liposarcoma and according to American Joint Committee on Cancer eighth edition [2] for soft tissue sarcomas arising from retroperitoneum, the staging was T4N0M0 with Grade 1 differentiation. Also, a section adjacent to the lesion showed dense lymphoid aggregates with germinal centres separated by wide band of collagens and plasma cell resembling Castleman’s disease. There was no evidence of lymphoma [Table/Fig-5a,b]. However, the resected margins were free of tumour infiltration. No specific treatment for castleman
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A wide spectrum of rare tumours arises from retroperitoneum that includes both benign and malignant tumours. Symptoms and imaging are considered non specific and histological diagnosis is confirmatory. Soft tissue sarcomas are relatively rare, among which retroperitoneal sarcomas constitute 10-15% of all soft tissue sarcomas. They are mesenchymal tumours and malignant tendency is less than 1% in adults. The reversible posterior leukoencephalopathy syndrome (RPLS) was first described by Giovanni Battista Morgagni in 1761 while performing autopsy of a 60-year-old woman. They are uncommon malignant tumours and a low incidence of approximately 4-5 per 100,000 individuals. It can affect both genders and age groups of 40-60 years [3].

The prognosis depends highly on tumour margins and grading than tumour size. According to World Health Organisation (WHO) classification, they were categorised as well differentiated, dedifferentiated, myxoid, pleomorphic and mixed type. The well differentiated liposarcoma are identified as five histological variants [4]. The clinical manifestations vary from vague abdominal pain or mass to mass effect based on tumour growth. The treatment of choice is complete surgical excision and it is reported that no survival benefits of adding adjuvant chemotherapy or radiotherapy [5-7].

Another rare retroperitoneal fibrotic disease namely the Castleman's disease also has been reported with similar incidence in age and tumour location [8]. This condition presents as lymphoproliferative disease of lymph nodes. The pathogenesis is unclear but possible immunodeficiency and human herpes virus (HHV) has been studied. Although commonly involves the mediastinal nodes, retroperitoneum has also been reported [9]. This condition also involves the same age group 40-60 years and equal gender distribution. The reported case here, showed similar age and the presentation as vague constitutional symptoms with abdominal mass [10]. However, the differentiating features further depends on centricity where unicentric castleman tends to be asymptomatic and multicentric castleman disease can present with constitutional symptoms and carries poor prognosis. This disease is not malignant condition however it increases risk of other tumours especially B-cell lymphomas and follicular dendritic cell sarcomas [11].

Retroperitoneal liposarcomas are diagnosed based on non specific or constitutional symptoms due to the mass or incidentally on imaging. The investigation of choice is CECT allowing an adequate staging and preoperative evaluation. Pelvic magnetic resonance is used for ruling out muscular or vascular invasion, satellite lesions or recurrences [12]. The possible differential diagnosis mainly includes tumours arising from retroperitoneal visceral (renal or adrenal, pancreas) structure, a lymphoma or metastatic lesion. There are instances where collision tumours, atypical breast liposarcoma and synovial liposarcoma have been reported [13-15]. While the diagnosis of castleman disease is mostly a postoperative or histopathological diagnosis [8]. Based on the prognostic factor established by WHO, the optimal treatment for retroperitoneal liposarcoma is en-bloc resection with or without adjacent viscera [4,5,16]. Although tumours are insensitive to chemo/or radiotherapy, adjuvant radiotherapy for tumours >5 cm with positive surgical margins has shown to reduce recurrence rate without increasing survival. Thus, complete resection with free margins is the most important prognostic factor. Extensive resection for tumours in contact with adjacent organs tends to increase overall morbidity [7,12,17,18]. The case reported here shows retroperitoneal liposarcomas could grow faster to large size and may not involve the adjacent structures. This case report shows a rarity in co-existence of well-differentiated tumour and a premalignant disease.

CONCLUSION(S)

Retroperitoneal tumours are rare and presents with non specific symptoms due to its location and are often diagnosed by imaging.
However, this requires confirmation by histopathology and the prognosis depends on en-mass surgical resection which was achieved in present case report.

REFERENCES


PARTICULARS OF CONTRIBUTORS:

1. Postgraduate, Department of General Surgery, Mahatma Gandhi Medical College and Research Institute, Puducherry, India.

2. Senior Resident, Department of General Surgery, Mahatma Gandhi Medical College and Research Institute, Puducherry, India.

3. Professor, Department of General Surgery, Mahatma Gandhi Medical College and Research Institute, Puducherry, India.

4. Professor, Department of General Surgery, Mahatma Gandhi Medical College and Research Institute, Puducherry, India.

NAME, ADDRESS, E-MAIL ID OF THE CORRESPONDING AUTHOR:

Haresh Kumar Gobu, 2/37, VIP Nagar, Vijlankurich Road, Combatorare-641035, Tamil Nadu, India.
E-mail: hareshg06@gmail.com

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