INTRODUCTION
Spindle cell tumours are low grade fibromyxoid soft tissue sarcoma (LGFMS). The name is descriptive of its microscopic appearance with features of both mesenchymal and neuroendocrine differentiation. Cytogenetic studies of spindle cell tumour show two cell lines containing balanced translocation between chromosomes 7 and 16. A small subset of both LGFMS and hyalinizing spindle tumour (HSCT) displays areas of increased cellularity and atypia which qualify as intermediate to high grade sarcoma. Spindle cell carcinomas are rare presentation in the esophagus as well. These tumours rarely metastasized, but pulmonary and lymph nodes metastases are reported in the literature.

To the best of our knowledge, this case has not been reported earlier from Pakistan. This case is the description of LGFMS presented as a huge enlargement of abdomen.

CASE REPORT
A case of 60-year-old male, labourer by occupation, presented through the outpatient department with 4 months history of abdominal swelling and distention, weight loss and intermittent constipation. His vitals were within normal limits. On examination, abdomen was distended with everted umbilicus, dilated veins on right lower quadrant of abdomen with a hard non-tender mass almost occupying the whole of the abdomen reaching up to the pelvis with smooth surface. Left inguinal hernia was also present.

Ultrasound diagnosis was a retroperitoneal mass. CT scan showed a heterogeneous mass occupying the whole of abdomen and pelvis 23 x 18 cm in size. Posteriorly the mass was abutting the aorta, inferior vena cava, psoas muscle, ureter and inferiorly abutting the urinary bladder (Figure 1). The mass was reported most likely to be a neoplastic lesion. Mid line exploratory laparotomy showed a jelly like mass in the abdominal cavity and pelvis reaching up to the spermatic cord. This mass was approximately 20 x 20 cm in size smooth in surface and rounded in shape (Figure 2). Mass was separated from the surrounding structure and excised completely. Left side inguinal hernia repaired and orchidectomy was done through a separate left inguinal incision. Weight of the mass was 18 kg, measured after the operation. Postoperative follow-up in the ward was uneventful except draining about 500-600 ml blood stained fluid daily through the drain up to 10 days.

Histopathology confirmed the diagnosis of spindle cell tumour grade I-II. Patient presented again with recurrence after 4 months with the history of similar complain. CT scan confirmed a huge abdominal mass compressing the surrounding structures. This was reported as recurrence and patient was referred to oncology for further treatment.
DISCUSSION

Spindle cell sarcoma is a rare cause of large abdominal mass. Cytological diagnosis of spindle cell tumour is difficult and has a high potential for false positive and false negative diagnosis compared to other soft tissue tumours as seen in this case, where cytological diagnosis was inconclusive. Spindle cell tumors are usually diagnosed on histopathology based on the presence of two architectural components first giant rosettes composed of hyalinized acellular islands surrounded by round to oval plump cells in a palisading like pattern, second spindle cells, and hyper cellular areas with plump elongated cells with highly vascular background. Occasionally ghost cells in the centre and multinucleated giant cell may present, grading is suggested in soft tissue sarcoma in order to improve the management, prognosis and to prevent the recurrence, based on nuclear atypia, nuclear overlap, mitotic figures, and necrosis these tumours are grouped into grades. Treatment of spindle cell tumour is its complete excision.

To conclude spindle cell tumours are low-grade fibromyxoid sarcoma LGFMS with giant rosettes. It may present as a huge enlargement of abdomen. Cytological diagnosis is difficult. Wide excision of the tumour is the best treatment option, but high grade tumour may present with recurrence.

REFERENCES