Desmoplastic Small Round Cell Tumour (DSRCT) is a rare, highly aggressive, mesenchymal tumour that arises from the peritoneal cavity. It is commonly seen in adolescent and young adult males and its occurrence in females is uncommon. We are reporting here a rare case of DSRCT in a young woman, which clinically masqueraded as an ovarian malignancy, with metastasis to liver, lung, spleen and peritoneum. The cytologic findings, histomorphological and immunohistochemical features have been discussed, with a brief review of literature.

**CASE REPORT**

A 36-year-old woman presented with a progressive, abdominal distension of 1 month's duration, which was associated with severe pain in the abdomen since 15 days. On examination, she was found to have ascites, with a mass being present above umbilicus. Computed Tomography (CT) scan of the abdomen revealed a large heterogeneously enhancing, predominantly solid, abdominopelvic mass which was located posterior to the uterus; ovaries were not visualized separately. Multiple, large, omental, peritoneal, hepatic, splenic and lung deposits were seen (Table/Fig-1a). The omental deposits were found to infiltrate transverse colon at several places. Serum CA-125 level was elevated to 110.2 U/ml. All other parameters were within normal limits. A clinical diagnosis of a stage IV malignant ovarian tumour was made. Fine Needle Aspiration (FNA) which was obtained from the mass revealed a small round cell tumour; whose exact typing was not possible (Table/Fig-1b).

Exploratory laparotomy and cytoreduction surgery, with excision of tumour mass was performed and the specimens were received in the pathology laboratory for histopathological evaluation.

**Pathological Findings**

Grossly, tumour consisted of multiple grey white nodules, largest measuring 24x10x7cm. Cut section showed a grey white variegated tumour with fibrous, necrotic and cystic areas.

Microscopy revealed a tumour which was composed of well-defined nests of small round undifferentiated cells with round to oval hyperchromatic nuclei, inconspicuous nucleoli, scant eosinophilic cytoplasm, which were surrounded by abundant dense fibrous connective tissue with scattered fibroblasts. Occasional mitoses (0-1/hpf), multi-focal areas of tumour necrosis and interspersed thin walled blood vessels were seen (Table/Fig-2a). No ovarian tissue was seen in the sections which were studied.

On Immunohistochemistry (IHC), tumour cells showed membrane positivity for Epithelial Membrane Antigen (EMA) and a paranuclear dot positivity for desmin and vimentin (Table/Fig-2b,c,d).

Based on the clinical data, histopathological and IHC findings, a final diagnosis of Desmoplastic small round cell tumour arising from abdominopelvic peritoneum was made.

Patient received three cycles of chemotherapy (Cisplatin and Paclitaxal). Three years of follow up of the patient revealed one episode of increasing ascites, for which she was treated symptomatically. She was subsequently lost to follow up.
Desmoplastic small round cell tumour (DSRCT) is a rare mesenchymal malignancy that primarily affects adolescents and young adults. It was first described in 1989 by Gerald and Rosai [1]. DSRCT is characterized by the presence of small round cells that show positivity for epithelial and myogenic tumour markers. Cytogenetic studies have demonstrated a characteristic reciprocal chromosomal translocation, t(11;22) (p13;q12), which is different from the t(11;22) (q24;q12) translocation observed in Ewing's sarcoma/ PNET [3].

DSRCT mainly develops in adolescents and young adults, with a strong male predominance and a male to female ratio of 4:1 [4]. Clinically, the patients present with symptoms of abdominal sarcomatosis, such as ascites, abdominal pain, distension, constipation or bowel obstruction, vomiting, and weight loss [5].

CA125 and Neuron specific enolase are frequently raised in the sera of patients with intra-abdominal DSRCTs before therapy, but these are not reliable monitors of the course of the disease [6]. The present case also showed elevated serum levels of CA125. However, as the patient was a female, this finding led to a clinical misdiagnosis of a primary ovarian malignancy.

Abdominal imaging done by ultrasound, Computed Tomography (CT), Magnetic resonance Imaging (MRI) is more useful for staging purposes, to assess tumour burden. However, the presence of a single or multiple dominant masses within the diffuse intraperitoneal process is more characteristic of DSRCT, as compared to those seen in other lesions. Imaging evidences of tumour heterogeneity, calcification, or intratumoural degeneration are additional supportive evidences [7].

The cytologic findings of DSRCT have seldom been reported. They share cytomorphic features of other small round cell tumours. In the current case, FNAC reported mass as a small round cell tumour. Presence of collagenous stromal fragments, along with small round cells which show positivity for epithelial and myogenic markers, could prove useful to achieve a cytological diagnosis [8].

In conclusion, DSRCT is a rare and an aggressive mesenchymal tumour that has been described to occur at various anatomical sites. Its classical clinical presentation is as large bulky peritoneal masses with uncommon organ involvement. The present case, however, showed deposits in the liver, lung, spleen and omental deposits which infiltrated the transverse colon at several places. Diagnosis mainly rests on histopathology and immunohistochemistry. CA125 elevation can cause confusion with an ovarian malignancy, particularly so in female patients, as was observed in this case.

REFERENCES


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