Ovarian dysgerminoma: a case report

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INTRODUCTION

Ovarian germ cell tumours are derived from primordial germ cells of the ovary. They can be malignant or benign and they comprise about 30% of ovarian neoplasms and 3% of all malignant ovarian neoplasms.2 It arises by malignant transformation of the primordial germ cells and in some cases may occur on anomalies of genital tract such as gonadal dysplasia.3 They usually arise in young women between 10 and 30 years of age. These tumors grow rapidly, and present as abdominal pain and large tumour mass in the abdomen.4

Malignant GCT’s are being classified into subgroups, the most common of it is dysgerminina. Most common mode of spread is through lymphatic system into paraaortic lymph node. In cases of unilateral encapsulated unruptured tumour conservative surgery is the treatment of choice. In advanced stages of the tumour, radical surgical procedure followed by chemotherapy will be done. The five years survival rate of tumour in stage 1 is 80-90%.5,6

CASE REPORT

A 22 years old female with paral live1 had come to the ANC OPD with amenorrhea and lump in the abdomen since last 6 month with chief complaint of pain in the abdomen requesting for routine antenatal sonography for fetal well-being. History of significant weight loss was present. She was pale and cachexic and on clinical examination a 36 weeks size huge abdominal mass. Subsequent computed tomography revealed 22×23×32.4 cm mixed density lesion in pelvi-abdominal region and multiple paraaortic and mesenteric lymph node with gross pleural effusion. On further evaluation raised beta-hcG and LDH were noted and hence dysgerminoma was suspected. Total abdominal hysterectomy with right salpingo-ophorectomy with resection of tumour mass with partial omentectomy and lymphadenectomy was performed. Histopathology reports were suggestive of dysgerminoma.

ABSTRACT

An ovarian dysgerminoma is a rare, malignant tumour occurring in young women, accounting for 1% to 2% of all primary ovarian neoplasms. A 22 years old female presented with 6 months of amenorrhea and lump in the abdomen. Her physical examination was remarkable with 36 weeks sized huge abdominal mass. Subsequent computed tomography revealed 22×23×32.4 cm mixed density lesion in pelvi-abdominal region and multiple paraaortic and mesenteric lymph node with gross pleural effusion. On further evaluation raised beta-hcG and LDH were noted and hence dysgerminoma was suspected. Total abdominal hysterectomy with right salpingo-ophorectomy with resection of tumour mass with partial omentectomy and lymphadenectomy was performed. Histopathology reports were suggestive of dysgerminoma.

Keywords: Dysgerminoma, Malignant ovarian tumor, Surgical intervention
tapping showed elevated protein in the pleural fluid and with no LDH activity or bacilli. After correction of anaemia with 2 pint blood transfusion and surgical fitness she underwent exploratory laparotomy with right salpingo-oopherectomy with resection of tumour mass with partial omentectomy with lymphadenectomy was performed. The pathological diagnosis was turned as dysgerminoma. The tumour was encapsulated. Tumour was weighing 7.5 kg with maximum gross dimension of the tumour was 30 cm. The removed tumour was sent for histopathological analysis. A peritoneal washing was collected for cytological analysis. seen and no macroscopically visible secondary deposits were visible vascular infiltration was not found.

Peritoneal washing for malignant cells was negative. The consulting team decided to carry out adjuvant chemotherapy and hence referred to higher centre for the same. Postoperative CT scan suggestive of no obvious lesion in the pelvic bed but metastatic retroperitoneal lymphadenopathy were noted. Post-operative tumour markers were done and elevated (LDH-294U/L, beta-HCG-2.95mIU/mL) but were reduced as compare to preoperative values. According to the protocol patient was planned for 3 cycles of BEP (Bleomycin, etoposide, cisplatin) at higher centre.

DISCUSSION

Dysgerminoma is the most common type of malignant germ cell tumour, comprising of 1-2% of all malignant ovarian tumours. In most cases, it occurs at the ages of 20 and 30 as our patient was 22 years old. The tumour most often occurs unilaterally in about 80-85% of all cases, as was the case in our patient. Dysgerminomas tend to spread by the perirectal lymphatic system to lymph nodes near the aorta as our patient who had paraaortic lymphnode involvement. The main clinical feature is rapid growth of the tumour and symptoms may persist from one to six months prior to diagnosis. The first symptom is pain in the abdomen followed by abdominal mass. Most commonly it spreads through the lymphatic para aortic in the lymph nodes, and haematogenous spread may take place in the advanced stages of the disease. Tumour marker may aid in the diagnosis and postoperative management of dysgerminoma. Elevated beta-HCG (10mIU/mL) and LDH (600U/L) were documented in this case report as seen in most of the cases of dysgerminoma. There were no pathological findings on the liver, stomach, small and large intestine, omentum or parietal peritoneum on naked eye examination in our patient. The cytological finding of the ascitic fluid taken from the abdomen was negative for tumour cells. Patient management is focussed on treating the tumour while preserving the fertility. While in
advanced stages of the disease such as Ib and above, radical surgical approach can be done which includes hysterectomy with bilateral salpingo-oopherectomy, lymphadenectomy with omentectomy as done at our centre.\textsuperscript{13} Postoperative chemotherapy consisting of BEP were applied to this patient as it is considered as gold standard and widely accepted.\textsuperscript{14,15}

**CONCLUSION**

Reproductive aged group women presenting with lower abdominal pain and rapidly growing mass with recurrent culture negative UTIs ovarian tumours should be considered and should be evaluated aggressively to rule out malignancies at the earliest. Dysgerminoma represent a distinct category with excellent response to chemotherapy and radiotherapy. Post surgical clinical surveillance can be planned for patient at low risk to prevent secondary malignancies, steriliry, and gonadal dysfunction associated with pelvic radiotherapy but careful followup is required to ensure early detection of recurrence. Patients education on compliance with treatment plan and specialist follow-up will definitely improve their outcome in postoperative period.

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**REFERENCES**
