Leiomyosarcoma Presenting as Abdominal Pain in a 50 Years Old Woman: A Case Report

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ABSTRACT

Leiomyosarcoma of uterus is a comparatively rare tumor we come across in daily histopathological examination. Clinical presentation of a leiomyosarcoma is varied and includes dysfunctional uterine bleeding, abdominal distention, abdominal pain and metastatic symptoms depending on the organ involved like Lung, bone and brain. Here we report a case of leiomyosarcoma in a 50 year old woman having a single complain of persistent abdominal pain for last 1 year or so. Clinically and by ultrasonogram it was diagnosed as a case of large fibroid. Subsequent histopathological diagnosis was leiomyosarcoma. Any post menopausal woman having persistent abdominal pain and a clinical and USG diagnosis of fibroid a differential of leiomyosarcoma should be borne in mind of the physicians.

Key Words: Leiomyosarcoma, Abdominal Pain

Introduction

Leiomyosarcoma comprises of about 10-20% of soft tissue sarcomas. They occur in adults and most commonly women. The common sites are skin1, soft tissue of extremities and retroperitoneum and uterus, cervix of women2,3. Some cases has also been reported in intramural blood vessels, renal vein and other vessels4,5,6. These leads to early metastasis. Extrauterine leiomyosarcomas present as painless firm masses where as uterine leiomyosarcomas present as abdominal swelling, pain, dysfunctional uterine bleeding and metastatic manifestations in the organs commonly involved like lungs, bone and brain. Uterine leiomyosarcoma is found equally before and after menopause and have a peak incidence at 40-60 years of age. The tumor has striking tendency to recur after removal and the 5 year survival is 40% in average. Leiomyosarcomas grow within uterus in two distinctive pattern a bulky, fleshy mass projected into the uterine wall or polypoidal mass projecting in to the uterine lumen. This uncommon neoplasm arise denovo from the myometrium or endometrial stromal precursor cells. It has linked to complex variable karyotype that include frequent deletions. It is a resistant cancer, meaning generally not very responsive to chemotherapy or radiation. The best outcomes occur when it can be removed surgically with wide margins early, while small and still in situ. Other treatment options include radiation therapy, hormone therapy, and chemotherapy – used either alone or in combination with another therapy6.

Treatment recommendations depend on the specific type of sarcoma, the stage of the cancer, and the patient's sarcoma history. A novel chemotherapy regimen that has demonstrated excellent efficacy in patients with leiomyosarcoma that has metastasized and cannot be removed surgically. The success of this regimen – a combination of gemcitabine and docetaxel – to treat leiomyosarcomas has led to additional research efforts, including two larger studies of our
treatment regimen by the Gynecologic Oncology Group, which proved the efficacy of this regimen for metastatic uterine leiomyosarcoma. This study shown that this regimen is effective in treating patients with other types of soft tissue sarcoma. For patients with early-stage leiomyosarcoma of the uterus that has been completely removed surgically, the role of adjuvant chemotherapy to reduce recurrence in women with high-risk types of the disease is also very important.

Case history

This case is a woman of 50 years of age hailing from Mirpur of Dhaka city, a house wife by profession. She is married and have a para of 3 child, the age of the last child is 33 years. She complains of persistant pain abdomen and severe weakness for last one year. She took some analgesic and antispasmodic as per doctors advice and get cured temporarily but the pain comes again. She then consulted a surgeon and on routine blood examination her hemoglobin was 6.2gm/dl and ESR was 120mm in first hour. Per vaginal examination was nothing contributory except some early prolapse of cervix. Ultrasonogram showed enlarged uterus with a large fibroid. A clinical diagnosis of leiomyoma was made by the physician and she was advised abdominal hysterectomy which she underwent subsequently on 22.02.2011 at a local clinic. Total abdominal hysterectomy with bilateral salpingo oophorectomy was done by spinal anaesthesia. After all aseptic precautions abdomen was opened by pfennelsteil incision. After opening a large uterus was found measuring 20X20cm in size. After successive clumping total abdominal hysterectomy with bilateral salpingo oophorectomy was done There was no adhesion. After proper haemostasis abdomen was closed in layers. Bleeding was average. She was discharged with advice with medication and rest for three months and periodic follow-up. Her uterus and both sided adnexa were sent for histopathological examination.

Gross and histopathological findings

Specimen consists of a resected uterus with cervix and both sided adnexa. The measurement of uterus with cervix is 15X12x10cm. Cut surface of uterus showed a large fibroid which occupies the whole uterine cavity measuring about 9.0cm in the maximum diameter. Cut sections of the fibroid show cystic spaces containing dirty fluid and it showed focal areas of calcification, hyalinization and hemorrhage. Endometrium was not clearly discernible. The measurement of ovaries were 2.5cm and 2.2cm in the maximum diameter. Cut section of the both ovaries were unremarkable. Blocks are embedded from cervix, endometrium, ovaries and fibroid.

Microscopic features

Sections of cervix show hyperkeratosis of ectocervix consistent with prolapse. Some endocervical glands are cystically dilated containing secretions. Sections of both ovaries are unremarkable, only shows corpora albicancia. Multiple sections from grossly described fibroid show a highly cellular tumor of smooth muscle cell origin with marked nuclear atypia, increased mitotic activity (1-8/HPF). Many tumour giant cells, infarction, myxoid and hyaline degeneration and foci of dystrophic calcification are also seen. The atypical tumour cells are arranged in nodules, irregular sheets infiltrating
the underline stroma. None of the section show any endometrial tissue. (Figure 2). A diagnosis of leiomyosarcoma was made.

Figure: 2

![Microphotograph of leiomyosarcoma, H&E stain X 40 showing cellular atypia, pleomorphism and mitosis.](image)

**Review of literatures and discussion**

Less than 4 percent of all uterine cancers are uterine sarcomas. Because uterine sarcomas are so rare and the behavior of these tumors is so variable, expert pathological review is critical. It is important for women diagnosed with these cancers to receive treatment at a center with experience caring for such patients. Uterine leiomyosarcoma is an aggressive tumor. In one retrospective series of 27 patients, there were 25 with nonmyxoid high-grade leiomyosarcoma of the uterus.

Uterine sarcomas (US) are rare and carry a poor prognosis characterized by high rates of local recurrence and metastasis. The three major histologic criteria: tumor differentiation, mitotic count, and tumor necrosis as well as other histologic and clinical factors were important and had an influence on survival and relapse.

Our case also showed infarction. Necrosis increased mitotic count and poor tumor cell differentiation.

Forty-nine patients were diagnosed with uterine leiomyosarcoma (25 stage I, 4 stage II, 8 stage III, and 12 stage IV). Of the 49 patients, 35 (71%) died of disease and 2 died of intercurrent disease. The 5-year survival rate was 33%. In a multivariate analysis with survival as the end point, stage was found to be the most important factor. All p53 immunopositive stage I-II tumors recurred within 28 months from diagnosis. This study found that stage represents the most important prognostic factor for uterine leiomyosarcomas. In stage I patients, tumor grade gives significant information regarding clinical outcome. In addition, p53 overexpression may predict a higher risk of recurrence in early stage leiomyosarcomas. In other study the overall survival rate was 66% at 2 years and 57% at 5 years with a disease free survival of 54% at 2 years and 50% at 5 years. Seven patients relapsed locally and 8 developed metastases. Another retrospective review from 1967 to 1995 of clinical and pathological characteristics of 93 patients with primary uterine sarcoma was done. The overall three-year survival rate was 67.9% and the overall five-year survival rate was 64.5%. It found statistical differences (p < 0.001) between the stage I survival rate and other stage survival rates.

A retrospective review of patients treated for a uterine sarcoma in Edinburgh from 1974 to 1992 has been performed. Clinical details at presentation, tumour pathology, treatment and the outcome of treatment were all recorded. 82 patients' case notes were reviewed. 54 patients had died and 28 were alive (mean follow-up period 80.3 months). 39 patients had a malignant mixed mesodermal tumour, 12 patients had an endometrial stromal sarcoma, and 27 had a leiomyosarcoma. The overall median survival is 15 months and the 5-year actuarial survival is 31%. 25 of the 26 surviving patients had Stage I disease at presentation. Post-operative pelvic radiotherapy did not influence either survival or local tumour control. 51 of the 54 patients who relapsed had evidence of distant metastases. The conclusion that total abdominal hysterectomy and bilateral salpingo-oophorectomy remains the treatment of choice for uterine sarcomas.

Three hundred and ten cases of uterine sarcomas, among them, according to pathology: LMS (Leiomyosarcoma)–117, MMS (Mixed mesodermal sarcoma)–62, ESS (Endometrial stromal sarcoma)–56, CS (Carcinosarcoma)–27, RBMS (Rhabdomyosarcoma)–18 and other sarcomas–30, were retrospectively evaluated at the Maria Sklodowska-Curie
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Memorial Cancer Center between 1950 and 1985. Overall 5-year survival was 34.5% and in particular subgroups the survivals were as follows: the best, 52% in LMS; lower, 30% in ESS; poor survival about 20% in CS and MMS.

A case of an epithelioid leiomyosarcoma of the uterine cervix in a 42 year-old woman was reported. This is a very rare tumor. Usually, the presenting symptoms are vaginal bleeding and abdominal pain. Two problems have to be solved by the microscopy: to prove the smooth muscle differentiation of the tumor and to assert the malignancy. Surgery remains the basis of therapy.

Conclusions
Any postmenopausal woman having persistent abdominal pain, discomfort apart from irregular vaginal bleeding suspected of having fibroid should undergo clinical ultrasonographic and histopathological examination of the specimen after surgery to exclude leiomyosarcoma. Because early diagnosis bears a good prognosis and late diagnosis carries a poor prognosis to the patient in spite of surgery, chemo and radiotherapy.

References