Multicystic peritoneal mesothelioma treated with complete cytoreductive surgery, peritonectomy and hyperthermic intra-peritoneal chemotherapy—A case report

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ABSTRACT

BACKGROUND: Multicystic Peritoneal mesothelioma is a rare and distinct variety of peritoneal mesothelioma with borderline malignant potential. Conventional Tumor bulking has been associated with recurrence of 45–50%. Hence a comprehensive treatment with Complete cytoreductive surgery with involved field peritonectomy (CRS) and Hyperthermic Intra-peritoneal chemotherapy (HIPEC) is being increasingly adopted for MCPM.

CASE PRESENTATION: A 47-year-old lady evaluated for peri-menopausal disturbance was diagnosed to have a multicystic lesion in the pelvis. With a preoperative suspicion of diagnosis of pseudomyxoma peritonei, CRS with HIPEC was planned. On exploration a diffuse multicystic mass was found in omentum and pouch of douglas with typical morphological features of MCPM. Complete cytoreduction was achieved with anterolateral and sub-diaphragmatic peritonectomy, omentectomy and panhysterectomy. HIPEC was performed with cisplatin 50 mg/m² for 40 min. Pathological examination revealed MCPM of omentum and uterine surface with focal clusters of mesothelial proliferation. However there was low proliferative activity 1–2%.

DISCUSSION: MCPM presents with wide spread peritoneal spread but with relative sparing of visceral invasion. Literature review suggests the disease spread is similar to PMP and treatment with CCRS and HIPEC has yielded long term survivals in MCPM.

CONCLUSIONS: This patient with voluminous disease burden in abdomen required surgical management and HIPEC for her condition. Whether CCRS alone without HIPEC can be an alternative for limited disease will be interesting research for future clinical reports.

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1. Introduction

Mesothelioma is a rare tumor arising from the serous membrane of pleural, peritoneal or pericardial cavities. Peritoneal mesothelioma (PM) accounts for about one third of all mesotheliomas and has varied biological aggressiveness depending on the subtype.

Peritoneal adenomatoid mesothelioma is usually localized and has low biological aggressiveness whereas at the other end of the spectrum is diffuse malignant mesothelioma involving multiple quadrants of parietal and visceral peritoneal surfaces and has a high potential for local recurrence.

Multicystic Peritoneal mesothelioma (MCPM) is rare and distinct variety of PM with low to intermediate malignant potential with reported long-term survival after aggressive therapy. It is common in women of reproductive age group and is unrelated to asbestos exposure. It is thought to arise from chronic peritoneal inflammation secondary to prior surgery, endometriosis or pelvic inflammatory disease, though conclusive evidence is unavailable [1,2]. Evidence suggests that it arises from fluid secretion of ovary due to peritoneal irritation and consequent trapping of this fluid in peritoneal adhesions [3].

Several authors insist that this type of tumor is not benign but a borderline malignancy that can rarely coexist with diffuse malignant mesothelioma [4,5]. Hence a comprehensive treatment with Complete cytoreductive surgery with involved field peritonectomy (CRS) and Hyperthermic Intra-peritoneal chemotherapy (HIPEC) has been advocated for MCPM [2,6]. We present our experience

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of managing a 47 year old lady with MPCM in a teaching hospital. This case is reported in line with the SCARE criteria [7] and informed consent was obtained from patient for publication of case report.

2. Presentation of case

Our patient, a 47 year lady reported to her gynecologist with complaints of peri-menopausal menorrhagia, dysmenorrhea and pallor in December 2019. Her past medical, drug and social history was unremarkable. Lab evaluation showed anemia and Magnetic Resonance Imaging (MRI) of pelvis revealed fibroma of the uterus. In addition, MRI showed multiple cystic areas in the pelvis and right iliac fossa (Fig. 1). Tumor markers levels were marginally elevated- Serum CA 125 – 55 IU/mL (Normal -25-35) and CA 19-9 – 6 IU/mL (Normal -<25). Provisional diagnosis of Pseudomyxoma Peritonei (PMP) originating from Ovary was made by her gynecologist and she was referred to us further therapy.

After evaluating her, our decision was to perform Exploratory laparotomy, cytoreduction and HIPEC as she had good performance status and limited spread of disease amenable for complete cytoreduction (CCR –0).

In March 2019, she underwent open surgical exploration through a Xiphopubic laparotomy by the senior author (YY). A large multicystic lesion was found in omentum and right paracolic gutter. Similar cystic lesion was found in pouch of douglas adjacent to the uterus (Fig. 2). Cysts were filled with clear to turbid fluid and were non-invasive in nature. The surgical Peritoneal carcinomatosis Index (PCI) was 6 (Lesion Size score (LS) - 3 in omentum and LS-2 in pouch of douglas, LS-1 in right paracolic gutter) and the adjacent structures- bowel, rectum and bladder were free of tumor. Extensive Intraperitoneal lavage (EIPL) was performed with ten liters of physiological saline and a complete cytoreduction was obtained by performing total anterolateral peritotomy, Pelvic peritotomy with en bloc extrafascial hysterectomy (bladder and rectum preserving), total omentectomy and excision of falciform and umbilical round ligament. She then received HIPEC with Cisplatin 50 mg/m² for 40 min.

Pathological examination showed multiple cysts lined with a layer of cuboidal mesothelial cells with no atypia (Fig. 3). There
are focal areas of proliferation of mesothelial cells in the omentum (Fig. 3). Sections of uterus showed a benign fibroma. Immunohistochemically the cells were positive for mesothelin, calretinin and negative CEA, CD-31, p 53 and D-2-40 (Fig. 3c–d). Ki-67 was low 1–2\%.

She recovered well after surgery and her postoperative course was uneventful. No further adjuvant therapy was indicated.

3. Discussion

MCPM (variably called as Peritoneal inclusion cysts, Inflammatory cysts of peritoneum, postoperative peritoneal cysts and benign papillary cystosis) is extremely rare tumor with about 200 cases reported in literature hitherto. It commonly affects young and middle age women and is associated with a prior history of abdominal surgery, endometriosis or pelvic inflammatory [8]. The disease process mainly involves omentum peritoneal surfaces, pelvic visceral surfaces like uterus, fallopian tubes, ovaries and pouch of douglas.

Preoperative diagnosis can be challenging as the symptoms are non-specific like diffuse abdominal pain, abdominal mass or pressure symptoms like dyspareunia, dysuria and disturbance of bowel habits. Imaging shows a multicystic lesion but there is no characteristic sign to clinch the diagnosis of MCPM. However MRI can characterize the nature of cyst fluid and extent of the mass. Presence of serous fluid in MCPM appears hypointense on T1 and hyperintense on T2 sequences in contrast hemorrhagic cysts appear hyperintense on T1 and hypointense on T2 sequences [9,10].

The differential diagnosis (DD) of MCPM includes cystic lesions of abdomen such as adenomatoid tumors, lymphangiomas, loculated ascites, pseudomyxoma peritonei, Diffuse malignant peritoneal mesothelioma (DMPM). Lymphangiomas commonly occurs in children and cyst is filled with chylous fluid and immunostaining of cyst wall with factor VIII is specific. DMPMs are extensive and patients present with weight loss, abdominal distension due to ascites, multiple bulky intraabdominal masses and invasion of adjacent structures like bowel wall, pancreas and spleen [11]. MCPM has characteristic gross appearance with multiple grape like cystic lesions filled with clear to yellowish fluid attached to peritoneal surfaces and omentum. Eventually definitive diagnosis of MCPM is made by the pathologist after histopathological examination of excised specimen.

Though MCPM is reported to be a benign etiology by some, several others have reported instances of malignant transformation and co-existent DMPM in these patients [4,12–14]. In general, MCPM pathologically has multicystic or unilocular cystic lesions lined by single layer of cuboidal or flat mesothelial cells with no atypia. But in agreement to the above theory we noticed found focal areas of mesothelial proliferation which indicates there are areas of increased cellular activity (Fig. 3). Ki-67 staining was increased at this area compared to others. Similar findings has been noted by others and these areas may suggest points of ongoing transformation with higher proliferative activity [3].

Another interesting pathological finding we observed was the presence of Peritoneal free floating cysts (PFFC) in the peritoneal cavity. Yonemura et al. has reported these Peritoneal free floating cysts (PFFC) in 8/9 MCPM patients in their series [13]. Some of the cysts attached to the main mass through a thin stalk and can disintegrate easily (Fig. 4). Disintegrated cysts can be free floating and migrate to other areas in peritoneal cavity and continue to deposit on other areas to establish multicystic lesions. Due care should be taken to thoroughly examine the peritoneal cavity and remove free floating cysts. EIPL with ten liters of saline assists in mechanical washing and removal of PFFC.

Regarding the management of MCPM, considerable debate exists as there is conflicting theories regarding the nature of disease.
Some authors have advocated surveillance or non-operative methods in small and asymptomatic lesions while most others regard surgery as the mainstay of treatment in MCPM. With 45–50% recurrence and potential for future transformation to invasive phenomenon, optimal surgery should include cytoreduction to remove gross disease and HIPEC for the treatment of microscopic residue [15–17].

Several centers have reported their successful results with peritoneectomy, complete cytoreduction and HIPEC for MCPMs and summary of these studies are in the Table 1 [1,2,18]. Recently group
Table 1
Summary of studies of MCPM treated with Cytoreductive surgery and HIPEC.

| Center and Year | Patients | PCI | CC-0 | HIPEC        | Survival | Recurrence |
|-----------------|----------|-----|------|--------------|----------|------------|
| Sugarbaker PH et al. [20] | 5        | –   | 4    | Cis + Dox    | 4 (NED)  | 1<sup>a</sup> |
| Lyon Fr [21]    | 3        | –   | 3    | Cis + Mitomycin | 3 (NED)  |            |
| Multi-institutional, Chua 2010 [2] | 26       | 14 (6–39) | CC–0 – 19 patients | Cis + Dox | 25 (NED) |            |
| NCI, Milan      | 19 (20 procedures) | 15.5 (6–24) | CC–0 – 17 patients | Cis + Dox | 1 (AWD)  | 1          |
| Nizri et al. [1] |          |     | CC–1 – 3 patients |           | 15 (NED) |            |

AWD - Alive with Disease; NED - No evidence of disease; RFS - Recurrence free survival.
<sup>a</sup> One patient had transformation to malignant mesothelioma.

from Milan, Italy reported an impressive mean recurrence free survival (RFS) of 159 months (13.2 years) and 79 % RFS at 10 years with CRS and HIPEC. Such robust long term data is unavailable in studies with conservative measures or debulking surgery.

Our patient had diffuse multi cystic disease involving the omentum, pelvic and right paracolic gutter peritoneum. Her surgical PCI was 6 whereas most studies report a median PCI of 10–15 [1,2]. We performed cytoreduction and involved field peritoneectomy as described above. Due to the presence of giant fibroid in the uterus a panhysterectomy was performed. However in young women, tumor excision with uterus and ovary conservation for purpose of future pregnancy has been reported [19]. These findings reflect the borderline invasive nature of the disease and support visceral preservation in appropriate cases. HIPEC was performed to eradicate the microscopic residue in the peritoneum.

In addition to existing literature, we have observed two distinctive pathological features in this case. One is the presence of free floating cysts in peritoneum with potential to deposit in other areas and second areas of increased proliferation in the background of benign mesothelial cells.

4. Conclusion

MCPM is borderline tumor with potential for recurrence and transformation to invasive mesothelioma. Complete cytoreduction with HIPEC appears to be the optimal treatment to eradicate the disease process. Uterus and ovarian preservation can be offered with caution in women desirous of fertility preservation.

Conflicts of interest

No Conflicts of Interest.

Sources of funding

No Funding received.

Ethical approval

The procedure-specific consent, patient data and material of this study had reviewed and approved by the ethical review bodies of Kishiwada Tokushokai hospital with ethic number 19–35 dated 11th November 2019.

Consent

Written informed consent was obtained from the patient for publication of this case report and accompanying images. A copy of the written consent is available for review by the Editor-in-Chief of this journal on request.

Author contribution

Naveen Padmanabhan – Study conceptualization, Data collection, drafting manuscript.
Haruki Ishibashi- Data collection and analysis.
Kazoru Nishihara- Data collection and analysis.
Shouzou Sako- Data collection and analysis.
Kanji Katayama- Data collection and analysis.
Satoshi Wakama- Data collection and analysis.
Yasuyuki Kamada- Data collection and analysis.
Yutaka Yonemura Study conceptualization, Data collection and review and finalization of manuscript.

All members have been reviewed the final version of manuscript.

Registration of research studies

Not required as it is single case report and not first in human study.

Guarantor

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Provenance and peer review

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