Localized biphasic malignant mesothelioma presenting as a giant pelvic wall mass: a rare case report and literature review

Yunsong Liu¹, Jingjun Wu¹, Ying Zhao¹, Pengxin Zhang², Zhengyu Hua², Wan Dong¹, Tao Lin¹ and Ailian Liu¹*

Abstract

Background: Localized biphasic MPeM is rare in clinical practice, we reviewed 8 cases of localized biphasic MPeM (including our present case), and summarized the clinical and imaging features of the disease.

Case presentation: We reported a 79-year-old man with chief complaint of a narrowing in the caliber of the stool for one year. A soft tissue shadow was occasionally found by CT examination in the right pelvic wall, and it was diagnosed as localized biphasic malignant peritoneal mesothelioma (MPeM) by postoperative pathology. Radical excision was performed and no radio-chemotherapy was applied. Nearly six years after surgery, the mass was significantly enlarged, and the neighboring tissues including rectum, prostate, seminal vesicle, and right ischial ramus were all infiltrated. The patient was in the end stage of cancer with poor prognosis.

Conclusions: The localized biphasic MPeM may show following characteristics: (1) with heterogeneous low-density and obscure margin; (2) with low incidence rate of ascites; (3) with few central hemorrhage and necrosis; (4) with few calcified structures; (5) with mild to moderate heterogeneous delayed enhancement on contrast-enhanced CT. The imaging characteristics can provide further information for the diagnosis of localized biphasic MPeM in the future.

Keywords: Biphasic, Malignant peritoneal mesothelioma, Pelvic, Computed tomography (CT)

Background

Mesothelioma is defined as the transformation of mesothelial cells from the lining of any human cavity into a tumor. The malignant mesothelioma (MM) is relatively rare in clinical practice, and has a highly invasive form. The most common site of MM was the pleura, and MM arising from the peritoneum of the pelvic wall was rarely reported [1]. The distribution of MM is diffuse or localized in two ways. The former is more common and presents as diffuse nodule or mass, the latter is relatively rare and presents as localized mass, which is usually large in size [2]. Histologic classification of MM includes epithelial, sarcomatoid, and biphasic subtypes according to World Health Organization (WHO) [3]. Simple epithelioid MM is the most common histologic type of the disease. Sarcomatoid and biphasic MM are relatively rare. To the best of our knowledge, only seven similar cases [2, 4–9] of localized biphasic malignant peritoneal mesothelioma (MPeM) have been reported.

The present study reported a localized epithelial and sarcomatoid mixed mesothelioma derived from pelvic wall, of which diagnostic and therapeutic experience remain limited. We reviewed the patient’s clinical, imaging, pathological, therapeutic and prognostic information in order to provide more clues for this disease.
Case presentation
A 79-year-old man came to our gastrointestinal outpatient with complaint of a narrowing in the caliber of the stool without obvious cause for about 1 year. Otherwise, he had no history of asbestos exposure, hematochezia, diarrhea, constipation, abdominal pain and distention. He had a history of prostatectomy due to benign prostatic hyperplasia and he denied recent weight loss. Digital examination of rectum showed the lower boundary of a mass in the rectum was 2 cm from the anal margin, and the upper boundary could not be palpated. Therefore, the patient underwent colonoscopy endoscopic electrocoagulation resection.

A soft tissue shadow was occasionally found by pelvis CT examination in the right pelvic wall. The mass had well-defined boundary and was oval shape with size of 7.3 × 5.3 cm. No calcification was found within the mass. On unenhanced CT images, the mass was heterogeneous with suspected necrotic area (Fig. 1a). In arterial, venous and delayed phases, CT values were 32–61 Hu, 65–90 Hu, and 46–77 Hu, respectively. The lesion showed mild heterogeneous on delayed enhancement CT images (Fig. 1b-d). Otherwise, the rectum and prostate were pressed, and no sign of destruction was observed in the adjacent bone. The CT examination suggested that the mass may originate from striated muscles with malignant transformation, and it may belong to neurogenic benign tumor. The source of blood supply to the mass was identified by pelvic computed tomography angiography (CTA), which showed the mass was mainly supplied by the right internal iliac artery (Fig. 4b). Unfortunately, the local dissection of the right common iliac artery was observed by CTA, and delayed the treatment of pelvic mass. Then, the right common iliac artery dissection was treated in other hospital more than a month later.

Six month after the discovery of the pelvic mass, the mass was slightly larger observed by CT examination (Fig. 4a). The patient underwent radical excision of pelvic mass. Intraoperatively, a solid mass with complete capsule was disclosed at the right obturator site of the pelvic wall. The lesion was 8 × 6 cm in size, with nodular surface and well-defined boundary. The surgeons removed the mass completely. The patient did not receive radio-chemotherapy and was in good condition after surgery.

The postoperative pathological examination showed that the mass was biphasic differentiated to both epithelium and mesenchyma (Fig. 2a-b). Immunohistochemistry is also important for the diagnosis of the mass. The ki-67 was less than 10%, which suggested that tumor cell proliferation is relatively inactive (Fig. 2c). Mesothelioma cells were positive for CD34, calretinin, EMA, MC and Vimentin, and negative for CD99, CD117, CK5/6, CK7, CK20, HMB45 and S-100 (Fig. 3a-l). In summary, the tumor was considered as biphasic malignancy mesothelioma.

One and a half years after surgery, the patient underwent pelvic CT reexamination and no sign of tumor recurrence was found. Four years after surgery, the patient attended a local hospital due to progressive dysuria and...
lower abdominal pain. Then pelvic CT showed a soft tissue mass in the preexisting position and the mass was measured as 7.89 × 10.41 cm with oval shape and well-defined boundary (Fig. 4c). We still saw that the mass had heterogeneous density in unenhanced CT scan. In the arterial, venous and delayed phases, CT values were 36 Hu, 58 Hu, and 60 Hu, respectively. The mass showed mild to moderate heterogeneous delayed enhancement. The right ischium was destroyed. Neighboring tissues including rectum, prostate and seminal vesicle were all infiltrated and pressed by the mass. Then, the patient underwent three times chemoembolization successively, and he was significantly relieved of symptoms after treatment. Nearly 6 years after surgery,
the patient came to our hospital for further palliative treatment of the tumor. CT reexamination revealed the mass had unclear boundary and significantly enlarged. The bone destruction of right ischial ramus was observed. Scattered and irregularly distributed patchy calcifications were observed in the mass, which may be a hypertrophic response caused by bone destruction. The walls of bladder, descending colon, and sigmoid colon were thickened due to tumor invasion (Fig. 4d). In addition, there was a peritoneal effusion. Finally, the patient was in the last stage with cachexia, and his prognosis was poor.

Discussion and conclusion

Our present case was a localized biphasic MM originated from the peritoneum of the pelvic wall. We made a detailed analysis about relevant publications, and our present case was also included (Table 1). The cases consisted of five males and three females (1.67: 1). In industrialized countries, the prevalence rates of MPeM ranged from 0.5 to 3 cases per million in men and from 0.2 to 2 cases per million in women [10]. So men may have a higher incidence than women. In Kawai et al.’s study [11], the ratio of men to women was 4.5 to 1, which is similar to our study. Patients were aged from 41 to 79 years, and the median age was 69 years old. The tumors appear to affect mainly the older population. Localized tumor in the liver was observed in 5/8 cases (62%), 2/8 (25%) in the abdominal wall, and 1/8 (13%) in the transverse colon. At present, the epidemiology of malignant peritoneal mesothelioma is more ambiguous than that of pleural mesothelioma. The epidemiology of MPeM varies with various factors. Asbestos exposure is the main cause of MPeM. But patients with peritoneal mesothelioma are less likely to have a well-defined history of asbestos exposure than patients with pleural mesothelioma [12]. Only 2 patients (25%) had a history of asbestos exposure. Typical initial symptoms of MPeM were abdominal pain, abdominal distention, or weight loss [13]. Initial symptom of 3 patients (38%) was abdominal pain. Four patients (50%) had no obvious symptoms. One (12%) patient noticed enlarging lump in right abdominal wall. Most patients showed normal results about biochemistry examinations. However, 4 patients (50%) had anemia on hematologic examinations, which may be due to bleeding from lumps. Tumor markers such as carcino-embryonic antigen (CEA), alpha fetoprotein (AFP), CA12–5, CA19–9 were normal in the 8 cases. Most of the cases reported as the localized tumors were with a median size of 11.2 cm (range 4–24 cm). One case had a very large mass, which occupied the right abdominal cavity and bilateral pelvic cavity.

Next, we reviewed the immunohistochemical data of all present cases. In Table 2, mesothelioma cells were positive for calretinin in 8/8 (100%) cases, vimentin in 7/7 (100%), CK5/6 in 5/6 (83%), WT-1 in 3/5 (60%) and negative for CK20 in 3/3 (100%), HMB-45 in 3/3 (100%), S-100 in 4/4 (100%). So we found that the most
dominant group of positive markers were calretinin, vimentin, CK5/6. Meanwhile, the most significant group of negative markers were CK20, HMB-45, S-100.

At present, immunohistochemistry has been commonly used to diagnose malignant mesothelioma. However, CT, a commonly used diagnostic method for abdominal lesions, shows no specific manifestation in the diagnosis of MPeM. In the study of 244 MPeM cases, Tandon et al. found that the most sensitive immunohistochemical markers were calretinin (100%), WT1 (94%), and CK5/6 (89%) [14], which was similar to our study. Saito et al. believed that calretinin, CK 5/6, mesothelin, vimentin, epithelial membrane, and WT-1 were specific markers of tumor mesothelial origin [15].

Firstly, MPeM should be distinguished from similar benign lesions, such as reactive mesothelioma and mesenteritis. Kawai et al. found that EMA, P53, desmin and p-glycoproteins were 100% expressed in malignant pleural mesothelioma, but no positive marker was found in the cases of reactive mesothelioma [11]. One of the most effective methods to distinguish MPeM and reactive mesothelial hyperplasia was fluorescence in situ hybridization (FISH), which could be used to analyze the homozygous deletion at site 9p21, which was positive in 67% of pleural mesothelioma, but the positive rate of peritoneal mesothelioma was low, only 25% [16]. Therefore, this method was not applied in our case. Liang et al. [17] found that the pattern of peritoneal thickening and contrast-enhanced imaging were effective markers for differentiating MPeM and peritoneal carcinomatosis, but their case was DMPeM (Diffuse MPeM), so it was of little help in differentiating this case. Liang et al. also found that

| Author/year | Age | Sex | Asbestos Exposure | Location | Size (cm) | Anemia | Initial symptom |
|-------------|-----|-----|-------------------|----------|----------|--------|-----------------|
| Sasaki et al [4]/2009 | 66 | Male | Yes | Liver | 4 | No | No obvious symptoms |
| Shao et al [2]/2011 | 77 | Female | No | Right abdominal wall | Very large | No | Notice enlarging lump |
| Kohno et al [5]/2012 | 69 | Male | Yes | Left abdominal wall | 10.7 | No | No obvious symptoms |
| Takehara et al [6]/2014 | 72 | Male | No | Transverse colon | 10 | Yes | Abdominal pain |
| Serter et al [7]/2015 | 66 | Male | No | Liver | 20 | Yes | Abdominal pain |
| Ali et al [8]/2016 | 41 | Female | No | Liver | 24 | Yes | No obvious symptoms |
| Dalal et al [9]/2018 | 69 | Female | No | Liver | 9 | Yes | Abdominal pain |
| Present case | 79 | Male | No | Liver | 8 | No | No obvious symptoms |

| Author/year | Tumor marker | Treatment | Follow-up |
|-------------|--------------|-----------|-----------|
| Sasaki et al [4]/2009 | Normal | Radical excision | No recurrence or metastasis 6 months after surgery |
| Shao et al [2]/2011 | Normal | Symptomatic treatment | Died 6 months after discovery |
| Kohno et al [5]/2012 | Normal | Radical excision | No recurrence more than 7 months after operation |
| Takehara et al [6]/2014 | Normal | Radical excision | Died 6 months after operation |
| Serter et al [7]/2015 | Normal | Radical excision | Unknown |
| Ali et al [8]/2016 | Normal | Radical excision | Unknown |
| Dalal et al [9]/2018 | Normal | Radical excision and adjuvant chemotherapy | Recurrence and progression during follow-up |
| Present case | Normal | Radical excision | Recurrence 4 years after surgery |

| Author/year | Central hemorrhage and necrosis | Calcification | Heterogeneous low-density | Enhanced mode | poorly-defined margins | Ascite |
|-------------|---------------------------------|--------------|--------------------------|---------------|---------------------|-------|
| Sasaki et al [4]/2009 | Yes | No | Yes | peripheral staining | No | No |
| Shao et al [2]/2011 | Yes | No | Yes | mild to moderate heterogeneous delayed enhancement | Yes | Yes |
| Kohno et al [5]/2012 | Yes | No | Yes | peripheral staining | Yes | No |
| Takehara et al [6]/2014 | No | No | No | peripheral staining | Yes | No |
| Serter et al [7]/2015 | Yes | No | Yes | peripheral staining | Yes | No |
| Ali et al [8]/2016 | Yes | Yes | Yes | Unknown | No | No |
| Dalal et al [9]/2018 | Yes | No | Yes | Unknown | Yes | No |
| Present case | No | No | Yes | mild to moderate heterogeneous delayed enhancement | No | Yes |
on CT images, the mesenteric lipomatosis showed soft tissue nodules, perivascular fatty halo and nodules, which may be helpful to distinguish MPeM from mesenteric lipomatosis. Malignant diseases include metastatic peritoneal adenocarcinoma and rhabdomyosarcoma were similar to MPeM. Peritoneal carcinoma was a metastatic feature of many organ malignancies, especially of the gastrointestinal tract and ovary, and must be considered as a first possibility even in the absence of a clear primary focus [17]. The most common malignancy reported by Walkey et al. was ovarian cancer [18]. Metastatic peritoneal adenocarcinoma is histologically difficult to distinguish from MPeM. Kawai et al. found that the best negative mesothelioma markers to distinguish epithelioid mesothelioma from serous carcinoma were be-ep4 and moc-31, and the best positive mesothelioma markers were d2–40 and calretinin [11]. The primary lesion of peritoneal adenocarcinoma found on CT is also a strong evidence for the diagnosis of metastatic peritoneal adenocarcinoma. Arora et al. found that a specific myogen, a muscle-derived marker, could rule out rhabdomyosarcoma if it was negative [19]. However, biphasic MPeM contained sarcoma components, so it was difficult to exclude rhabdomyosarcoma by relying on it alone. It required a combination of specific markers of epithelial and mesenchymal origin for a comprehensive analysis. In a word, immunohistochemical diagnosis of MPeM is progressing well, but there are still many problems. Due to the small number of cases, few specific imaging findings were found.

Therefore, we collected radiological data from all the biphasic MPeM cases of restricted growth patterns available at present. Radiological studies play an important role in the diagnosis, staging and prognosis of biphasic MPeM. Among them, the contrast-enhanced CT is the major imaging modality for MM [20]. CT has an advantage in distinguishing between biphasic MPeM and its surrounding tissues in order to observe whether there is pathological infiltration. In addition, CT has the ability to display images over a wide range and to clearly show lumps in different areas, which is helpful in finding the origin of biphasic MPeM. Because biphasic MPeM is extremely rare, there is currently few imaging description of localized biphasic MPeM. A review of eight cases was summed up about radiological data, and our present case was also included in Table 2. The masses presented as heterogeneous low-density lesion on non-contrast CT scan in 7/8 cases (88%). One case presented as homogeneous low density tumor. On dynamic enhanced CT scan, the lesions presented as peripheral staining in 4/6 (66.7%) and mild to moderate heterogeneous delayed enhancement in 2/6 (33.3%). Tumors with obscure margin were seen in 5/8 (63%) cases. Only one case had few small calcifications, and 75% cases (6/8) developed hemorrhage and necrosis in the center. In the present case, CT showed no obvious hemorrhage and necrosis in the center of the mass as shown in Fig. 1a, showing only a slightly heterogeneous density within the lesion. Although the histopathological image showed a small amount of extravasation of red blood cells in Fig. 2a, this didn’t definitively demonstrate significant bleeding in the central area of the lesion. Only two (25%) cases had ascites. We found no other significant imaging features.

Currently, no standard treatment of malignant peritoneal mesothelioma had been established, and localized MPeM has been usually treated with radical resection. In Table 1, all cases presented as localized tumor in the peritoneum at initial diagnosis. Radical excision was performed in seven cases, and only one patient undertook symptomatic treatment because the lesion was too large. In addition to radical resection, only one patient underwent postoperative adjuvant chemotherapy. According to the analysis of follow-up data, we perceived that the prognosis was variable: two cases had no recurrence less than a year after surgery, two cases were recurrent postoperatively, and two cases were dead less than a year after surgery. Our patient experienced a period of nearly seven years from biphasic MM discovery to the last time follow-up, which demonstrated a relatively good prognosis. We suspected that the prognosis of localized biphasic MPeM was generally poor, and early treatment was urgently necessary.

At present, the diagnosis of MM is still difficult, and the diagnostic standards are usually pathological examination including immunohistochemistry. Although imaging examination has only made little progress in the diagnosis of MM, it can still show the spatial or temporal features of mass with a non-invasive way compared with pathology. The diagnostic efficacy of radiological examination for MM is improving by reviewing more cases of MM. Histology and immunohistochemistry also have limitations in the classification of subtypes of MM. The classification of subtypes of MM by imaging has been explored recently [21–23]. For example, Escalon et al. [21]. found the calcified pleural plaques and local invasion were more common in non-epithelioid subtypes of malignant pleural mesothelioma. Similar studies in the abdomen and pelvic need be further carried out in the future.

In summary, the present case and literature review suggest that the localized biphasic MPeM may show following characteristics: (1) with heterogeneous low-density and obscure margin; (2) with low incidence rate of ascites; (3) with few central hemorrhage and necrosis; (4) with few calcified structures; (5) with mild to moderate heterogeneous delayed enhancement on contrast-enhanced CT. We hope that our report on localized biphasic MPeM will
provide further information for the diagnosis, classification and treatment of the disease in the future.

Abbreviations
AFP: Alpha fetoprotein; CEA: Carcino-embryonic antigen; CTA: Computed tomography angiography; MPeM: Malignant peritoneal mesothelioma; MM: Malignant mesothelioma; WHO: World health organization

Acknowledgements
Not applicable.

Authors’ contributions
AL, ZH, and PZ collected data during the study. YL and JW contributed to the study design. YL, JW, AL and YZ developed the first draft of the manuscript which was then reviewed and intensively revised by JW, AL, WD and TL. All authors read and approved the final manuscript.

Funding
This work was supported by the Program for Training Capital Science and Technology Leading Talents [grant number Z181100006318003]. The funding body contributes to the design of the study and analysis of data.

Availability of data and materials
All data generated or analyzed during this study are included in this published article.

Ethics approval and consent to participate
Not applicable.

Consent for publication
The patient provided written informed consent for publication of this case report and accompanying images.

Competing interests
The authors declare that they have no competing interests.

Author details
1Department of Pathology, The First Affiliated Hospital of Dalian Medical University, Xigang district, Zhongshan road, No.222, Dalian, China.
2Department of Radiology, The First Affiliated Hospital of Dalian Medical University, Xigang district, Zhongshan road, No.222, Dalian, China.

Received: 13 December 2019 Accepted: 15 April 2020

Published online: 06 May 2020

References
1. Muta H, Sugita Y, Oshima K, et al. Primary malignant pericardial sarcomatoid mesothelioma: an autopsy report.[J]. Pathol Int. 2017;67:311–5.
2. Shao ZH, Gao XL, Yi XH, et al. Malignant mesothelioma presenting as a giant chest, abdominal and pelvic wall mass.[J]. Korean J Radiol. 2011;12:750–3.
3. Gibbi AR, Thunnissen FB. Histological typing of lung and pleural tumours: third edition.[J]. J. Clin. Pathol. 2001;54:498–9.
4. Sasaki M, Araki I, Yayui T, et al. Primary localized malignant biphasic mesothelioma of the liver in a patient with asbestosis.[J]. World J. Gastroenterol. 2009;15:615–21.
5. Kohno M, Manuyama R, Kitagawa D, et al. Localized biphasic type malignant mesothelioma arising in the peritoneum: report of a case.[J]. Thorac Cancer. 2014;5:74–7.
6. Takehara Y, Endo S, Mori Y, et al. Malignant peritoneal mesothelioma with lymph node metastasis that originated in the transverse colon.[J]. World J Surg Oncol. 2014;12:112.
7. Serter A, Buyukpinarbasili N, Karatepe O, et al. An unusual liver mass: primary malignant mesothelioma of the liver - CT and MRI findings and literature review.[J]. Jpn J Radiol. 2015;33:102–6.
8. Haji Ali R, Khalife M, El Nounou G, et al. Giant primary malignant mesothelioma of the liver: A case report.[J]. Int J Surg Case Rep. 2017;30:58–61.
9. Hassan D, Ligato S. Localized biphasic malignant peritoneal mesothelioma with rhabdoid features involving the liver: case report and review of the literature.[J]. Case Rep Pathol. 2019;2019:2732674.
10. Boffetta P. Epidemiology of peritoneal mesothelioma: a review.[J]. Ann Oncol. 2007;18:985–90.
11. Kawai T, Tominaga S, Hiroi S, et al. Peritoneal malignant mesothelioma (PMM), and primary peritoneal serous carcinoma (PPSC) and reactive mesothelial hyperplasia (RMH) of the peritoneum. Immunohistochemical and fluorescence in situ hybridisation (FISH) analyses.[J]. J Clin Pathol. 2016;69:706–12.
12. Welch LS, Acherman YI, Haile E, et al. Asbestos and peritoneal mesothelioma among college-educated men. Int J Occup Environ Health. 2005;11:254–8.
13. Mohamed F, Sugarbaker PH. Peritoneal mesothelioma. Curr Treat Options in Oncol. 2002;3:375–86.
14. Tandon RT, Jimenez-Cortez Y, Taub R, et al. Immunohistochemistry in peritoneal mesothelioma: a single-center experience of 244 cases. Arch Pathol Lab Med. 2018;142:236–42.
15. Saito H, Hasuda S, Nau J, et al. A case of malignant peritoneal mesothelioma suggesting the utility of combining double-contrast radiography and endoscopy with computed tomography for diagnosis.[J]. Clin J Gastroenterol. 2017;10:271–6.
16. Ilie PB, Rusch Vw, Zaworski MF, et al. Homozygous deletion of CDKN2A and codeletion of the methylthioadenosine phosphorylase gene in the majority of pleuralmesotheliomas. Clin Cancer Res. 2003;9:2108–13.
17. Liang YF, Zheng QQ, Chen YF, et al. CT differentiation of diffuse malignant peritoneal mesothelioma and peritoneal carcinomatosis.[J]. J Gastroenterol Hepatol. 2016;31:709–15.
18. Walkey MM, Friedman AC, Sohota P. CT manifestations of peritoneal carcinomatosis. AJR Am J Roentgenol. 1988;150:1035–41.
19. Arora SK, Srinivasan R, Nijhawan R, et al. Malignant biphasic peritoneal mesothelioma in a child: fine-needle aspiration cytology, histopathology, and immunohistochemical features along with review of literature.[J]. Diagn Cytopathol. 2012;40:112–5.
20. Moore AJ, Parker RJ, Wiggins J. Malignant mesothelioma. Orphanet J Rare Dis. 2008;3:34.
21. Escalon JG, Harrington KA, Ploidkowski AJ, et al. Malignant pleural mesothelioma: are there imaging characteristics associated with different histologic subtypes on computed tomography?[J]. J Comput Assist Tomogr. 2018;42:601–6.
22. Senyigit A, Bayram H, Babayigit C, et al. Malignant pleural mesothelioma caused by environmental exposure to asbestos in the southeast of Turkey: CT findings in 117 patients. Respir; Int Rev Thorac Dis. 2000;67:615–22.
23. Seely JM, Nguyen ET, Churg AM, et al. Malignant pleural mesothelioma: computed tomography and correlation with histology. Eur J Radiol. 2009;70:485–91.

Publisher’s Note
Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.