Multiple Jejunal Myeloid Sarcomas Presenting with Intestinal Obstruction in a Non-leukemic Patient: A Case Report with Ultrastructural Observations

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Myeloid sarcoma is a rare extramedullary myeloid tumor, which is frequently misdiagnosed when no evidence of leukemia is initially observed. Here, we report on a peculiar case of a 49-year-old man afflicted with multiple masses in the jejunum, the superior mesentery, and the serosa of the transverse colon, without leukemic manifestation. The tumor was composed of undifferentiated small round cells containing eosinophilic cytoplasm, which were negative for myeloperoxidase, nonspecific esterase, lysozyme, terminal deoxynucleotidyl transferase, leukocyte common antigen, CD3, CD4, CD15, CD20, CD30, CD43, CD56, CD68/PG-M1, CD79a, human melanoma black-45, c-kit, and CD34 with positivity only for CD68/KP1, CD99, and vimentin. Under electron microscopy, those cells had abundant membrane-bound cytoplasmic granules that measured 200 to 300 nm in diameter, which were consistent with granulocytic azurophilic granules. The tumor was finally diagnosed as a myeloid sarcoma. The presence of non-leukemic myeloid sarcomas showing immunonegativity for conventional myeloid-leukemic markers necessitated a diagnosis by ultrastructural observation.

Key Words: Sarcoma, myeloid; Jejunum; CD68/KP1; CD99; Microscopy, electron

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The World Health Organization defines myeloid sarcoma as a tumor of myeloblasts or immature to maturing myeloid cells involving an extramedullary anatomic site. The neoplasm is also known by various names such as granulocytic sarcoma, chloroma, monocytic sarcoma, extramedullary myeloid cell tumor, and myeloblastoma. Myeloid sarcomas can occur in any site of the body. Most gastrointestinal myeloid sarcomas occur in the small intestine, mainly in the ileum. Only a few case reports have documented synchronous myeloid sarcomas involving multiple portions of the small intestine. Under light microscopy, myeloid sarcomas are frequently mistaken as other undifferentiated high grade tumors, and up to 56% of these tumors are initially misinterpreted as malignant lymphoma, Ewing’s sarcoma/primitive neuroectodermal tumor (PNET), rhabdomyosarcoma, medulloblastoma, or neuroblastoma. A confirmatory diagnosis is necessary using ancillary methods including immunohistochemistry, especially for those cases in which a previous diagnosis of leukemia or myelodysplastic syndrome has not been made.

We herein present a very rare case of multiple jejunal non-leukemic myeloid sarcomas that presented as an intestinal obstruction and were confirmatively diagnosed by electron microscopic identification of granulocytic granules.

CASE REPORT

A 49-year-old man presented with recurrent central colicky abdominal pain, which had commenced 2 weeks earlier. Abdominopelvic computed tomography (CT) showed a mass-like lesion at the left upper abdominal region (Fig. 1A). Segmental resection of the jejunum 30 cm below the Treitz ligament was performed. Intraoperatively, the obstructed portion was con-
glomerated and adhered to other segments of the distal small intestine. The mass-like portion demonstrated thick adhesion to the transverse colon and superior mesentery.

The resected segment measured 106 cm in length. The cut surface revealed two portions of circumferential stricture by two separate ulceroinfiltrating masses. The masses measured $15 \times 10$ cm and $12 \times 10$ cm each. The masses were separated from each other by 21 cm. Grossly, the masses showed transmural involvement and had invaded through the proper muscle layer of the adhered segment of the jejunum. The cut surface had a brown-colored homogeneous and fish-fleshy appearance (Fig. 1B). The mucosal surfaces of the masses were coated by necrotic exudates. Histologically, each tumor was diffusely infiltrated by round small to moderately sized cells containing eosinophilic granular

Fig. 1. (A) Abdominopelvic computed tomography reveals multifocal enhancing concentric wall thickening (arrow) with a skipped lesion in the jejunal loop. (B) A gross picture of the bisected jejunal segment reveals two separate encircling masses. (C) Primitive, undifferentiated, round-shaped tumor cells infiltrate the bowel wall. The inset indicates plump eosinophilic granular cytoplasm. (D, E) The tumor cells are cytoplasmically stained with CD68/KP1 (D) and CD99 (E).
cytoplasm (Fig. 1C). The tumor cells were irregularly shaped with angulated nuclei, irregular nuclear contours, and fine chromatin. Immunohistochemically, the tumor cells were cytoplasmically stained with CD68 (1:100, KP1, Lab Vision Co., Cheshire, UK) (Fig. 1D), CD99 (prediluted, 12E7, Dako, Glostrup, Denmark) (Fig. 1E) and vimentin (prediluted, V9, Dako). The cells produced negative results for myeloperoxidase (MPO), toluidine blue, lysozyme, terminal deoxynucleotidyl transferase (TdT), nonspecific esterase, leukocyte common antigen (prediluted, Dako), CD3 (prediluted, polyclonal, Dako), CD4 (4B12, prediluted, Dako), CD15 (1:50, Carb-3, Dako), CD20 (prediluted, polyclonal, L26, Dako), CD30 (prediluted, Ber-H2, Dako), CD43 (prediluted, Dako), CD56 (1:50, PG-M1, Dako), CD79a (1:50, JCB117, Dako), human melanoma black-45 (prediluted, HMB-45, Dako), CD117 (prediluted, c-kit, Dako), and CD34 (prediluted, QBEnd10, Dako). To exclude the remote possibility of a gastrointestinal stromal tumor (GIST), platelet-derived growth factor receptor-alpha (PDGFRA) and c-kit mutation studies were performed using polymerase chain reaction sequencing. No PDGFRA or c-kit mutations were observed.

Upon examination of their ultrastructure, the ovoid-shaped tumor cells possessed peripherally condensed chromatin with occasional involuted configurations (Fig. 2A, B). The tumor cells were packed with abundant intracytoplasmic membrane-bound granules that measured from 200 to 300 nm in diameter. The rectangular inclusions revealed a fine lamellar substructure, and the periodicity of the filamentous striations was about 10 nm (Fig. 2C, D). The peripheral blood smears failed to demonstrate any abnormal hematopoietic cells or blasts. Myeloid sarcoma was diagnosed. A bone marrow study was not performed due to the patient’s request. After the operation, a chest CT showed a 7 cm heterogeneous enhancing cavitary mass within low density in the right upper lobe. At that time, two enhancing small nodules were found at the liver. After 7 months, the size of the masses of the lung and liver had enlarged slightly. The peripheral blood smears indicated that no abnormal myeloid cells or blasts remained. Biopsies were not taken from these organs. The patient experienced severe recurrent hemoptysis. His condition continued to decline and then he died. An autopsy was not performed.

DISCUSSION

Non-leukemic myeloid sarcoma can manifest initially as an isolated intestinal mass, and chemotherapy is the treatment of
choice. Surgery is indicated only if there are complications such as bowel obstruction, bleeding, or perforation.\textsuperscript{7} Because of frequent misdiagnosis as other undifferentiated high grade tumors, an accurate diagnosis may be made only after acute myeloid leukemia (AML) has developed (mean, 9 months).\textsuperscript{3} Therefore, an early and accurate diagnosis will enable proper treatment, which has the potential to be successful despite the aggressive clinical course that is typical of myeloid sarcomas. Microscopically, a dense infiltrate of relatively small, undifferentiated high grade tumor cells showing an unfamiliar histology and uncertain origin are the first presentation of a hematologic malignancy.\textsuperscript{7} The diagnosis can be challenging to pathologists when there is a non-leukemic phase or in the absence of a known hematological disorder. From light microscopic morphology, the differential diagnoses of myeloid sarcoma include a GIST, malignant lymphoma, melanoma, and an undifferentiated tumor such as Ewing’s sarcoma/PNET. Once the possibility of a myeloid sarcoma is considered, immunohistochemistry can reliably make this distinction in nearly all cases. Immunohistochemical panels for myeloid markers such as MPO, lysozyme, c-kit, CD43, and CD68/KP1 are all sensitive and helpful, although markers for CD43, CD68, and c-kit are nonspecific, particularly CD43.\textsuperscript{11,13} Among them, CD68/KP1 is a good marker and even appears more sensitive than MPO for granulocytic precursors, while CD68/PG-M1 is more sensitive in monocytic differentiation.\textsuperscript{11,12} CD30 and CD56 are seldom recorded in myeloid sarcoma cases.\textsuperscript{15} Our case showed negativity for conventional immunohistochemical markers such as chloroacetate esterase, MPO, and lysozyme, and showed positivity only for CD68/KP1 and CD99. These results had the potential to be misdiagnosed as Ewing’s sarcoma/PNET because CD99 is a transmembrane glycoprotein p30/32\textsuperscript{MIC2}, a product of the MIC2 gene, and is positive in Ewing’s sarcoma/PNET and lymphoblastic lymphoma.\textsuperscript{14} However, CD99 has also been expressed by monococytes, B cells, and granulocyte-lineage cells such as those from myeloid sarcoma, lymphoblastic lymphoma, or TdT-positive AML.\textsuperscript{15} Immunoreactivity for CD99 has been reported in more than 50% of the myeloid sarcomas, which is in line with its not infrequent expression in hematopoietic tumors.\textsuperscript{11-13} TdT is expressed in approximately 90% of acute lymphoblastic lymphoma cases, which represents a small subset of AML cases. If the disease develops in the gastrointestinal tract, differential diagnoses of GIST and gastrointestinal lymphoma should be considered. Although immunopositivity for c-kit and CD34 are important diagnostic markers, shared features exist between GIST and myeloid sarcoma.\textsuperscript{16} In confusing cases showing negativity for a myeloid marker including MPO, a mutation analysis for PDGFR and c-kit, coupled with electron microscopic investigation, may be required.

In the present case, the presence of membrane-bound cytoplasmic granules in the ultrastructural analysis offered a diagnostic clue. Differential diagnoses from the various types of secretory granules included membrane-bound granulocytic granules, dense-core granules, zymogen granules, mucin granules, Birbeck granules, melanosomes, surfactant granules, laminated granules of the mast cells, lipofuscin granules, and Weibel-Palade bodies.\textsuperscript{17} Lipofuscin granules are aggregates of osmiophilic granules and appear as an electron-dense layer. Although there have been rare reports of granular variants of some tumors such as mammary lobular carcinoma, which may contain such cytoplasmic granules, granules are not commonly found even in the myeloid metaplasia of tumors such as those secondary to carcinoma, leukoerythroblastosis, or tuberculosis.\textsuperscript{18} Investigation of the nuclear morphology of the tumors cells is helpful for distinguishing their differentiation and maturation. As granulocytes mature, their nuclei become flattened, indented, and then lobulated.\textsuperscript{19} Cytoplasmic granules appear and then lose cytoplasmic basophilia. Ultrastructural changes of the granule content, with observed loss of electron density, indicates secretory activity. Giant granules up to 3 µm that are commonly observed in diseases such as Chediak-Higashi syndrome are of rare occurrence in myeloid sarcoma.\textsuperscript{20} Identification of the cytoplasmic granules offered a diagnostic clue in the present case.

In conclusion, the application of a screening panel for high grade undifferentiated tumors should be broad in order to avoid a diagnostic misinterpretation due to occasionally aberrant or unexpected antigen expression. Even in cases of negative clinical results and unusual immunohistochemical observations, a high index of suspicion is required to avoid overlooking the diagnostic findings, and in these cases, the use of electron microscopic examination is vital for the diagnosis of myeloid sarcomas.

**Conflicts of Interest**

No potential conflict of interest relevant to this article was reported.

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