**Gastrointestinal Mantle Cell Lymphoma: incidental finding on routine CT scan following pneumothorax.**

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**ABSTRACT**

Gastric mantle cell lymphoma is a rare form of gastrointestinal tumor and represents 2.5–7% of all non-Hodgkin’s lymphomas. A 69-year-old male admitted with spontaneous pneumothorax was found to have a gastric mass on CT scan. Results of histopathological, immunohistochemical and genetic analysis were consistent with mantle cell lymphoma (MCL). Patient responded well to intensive chemotherapy and subsequent allogeneic hematopoietic stem cell transplant and remains in complete remission. We would like to emphasize that a gastric mass, as in this case can occur as a rare presentation of MCL involving the gastrointestinal tract without any other overt signs and symptoms.

**INTRODUCTION**

Mantle cell lymphoma (MCL) is a distinct subtype of B-cell lymphoma accounting approximately 3-10% of lymphoma diagnoses. 1 About 2.5-7% of all Non-Hodgkin lymphoma cases are mantle cell and despite being a low-grade lymphoma, it has the worst prognosis among all lymphomas. It tends to occur in older adults with a median age of approximately 60 years and is common in males [M: F-2-7:1]. 2-3 Primary gastric mantle cell lymphoma is an extremely rare form of gastrointestinal tumor and represents 2.5–7% of all non-Hodgkin’s lymphomas. 4 In the present report, we describe the clinicopathologic features of a case of early primary gastrointestinal mantle cell lymphoma found incidentally on routine Computed Tomography (CT) scan following pneumothorax.

**CASE REPORT**

A 69-year-old male with a past medical history of gastroesophageal reflux disease, atrial flutter, hypertension and anxiety presented in the emergency department with sudden chest pain and shortness of breath and was diagnosed with spontaneous pneumothorax. Physical examination did not reveal any palpable peripheral lymph nodes, liver or spleen enlargement or any pathologic abdominal mass. His Computed Tomography (CT) scan of the chest revealed a small right pneumothorax with subcutaneous emphysema and a 3 cm gastric fundal mass with irregular thickening of gastric wall. Numerous mildly enlarged para gastric lymph nodes were also noted. He underwent an esophagogastroduodenoscopy that revealed a normal esophagus and several gastric masses with two (large greater than 3 cm) in the fundus and a few satellite lesions in the antrum with thickened folds. Positron Emission Tomography (PET) /CT showed large hypermetabolic stomach mass from gastric antrum, intense hypermetabolic area in the cecum/ileoceleal valve with multiple enlarged mesenteric lymph nodes. Multiple biopsies taken showed lymphoma cells positive for CD20, CD5, BCL2 and negative for CD10 and CD 23. Initial diagnosis of mantle cell lymphoma involving intra-abdominal lymph nodes was made (ICD 200.43, working diagnosis). His LDH was normal and β2 microglobulin was mildly elevated at 2.53. Bone marrow evaluation revealed hypocellularity (20-25% cellular) with subcortical trephine core biopsy showing multiple crushed lymphoid aggregates. Marrow-aspirate smears showed approximately 13% lymphocytes with occasional small atypical forms, erythroid hyperplasia and adequate granulopoiesis and megakaryopoiesis without significant dyspoiesis or increased blasts (1%). Immunohistochemical stains performed on core biopsy showed the lymphoid aggregates to be predominantly CD20 and CD79a positive B-cells with aberrant CD5 expression occupying approximately 10-15% subcortical marrow space. Cyclin D1 (BCL-1) was negative within the neoplastic lymphocytes. KI-67 showed a proliferative activity of approximately 5-10%. Flow cytometry analysis of marrow aspirate showed CDS positive lambda-monoecional B cell population. Fluorescent In Situ Hybridization (FISH) study was positive for CCND1 (BCL1) rearrangement. Karyotypic studies showed a normal male karyotype. Marginal anemia, slight absolute neutrophilia without left-shift with no lymphocytosis was seen on peripheral blood smear. Reported findings were consistent with approximately 10-15% marrow involvement by mantle cell lymphoma.

Patient was started on an intensive chemotherapy with hyper-Cytosan, Vincristin, Doxorubicin (Adriamycin), Dexamethasone (CVAD) and Rituxan. He completed cycle#1 (part A and part B) and cycle#2 (part A and part B) along with Neulasta (Pegfilgrastim) support. He then underwent a successful peripheral blood stem cell mobilization (with Neupogen) and

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was then admitted for autologous stem cell transplantation during which she received BEAM-rituxan conditioning. He engrafted neutrophils and platelets. He had hypogammaglobulinemia secondary to rituxan and chemotherapy for which he received IVIG. Pentamidine prophylaxis was given for pneumocystis carinii pneumonia (PCP).

He subsequently recovered and follow up PET/CT showed no evidence of lymphoma and his bone marrow biopsy was negative for residual or recurrent disease.

**DISCUSSION**

The genetic hallmark of mantle cell lymphoma is the chromosomal translocation t(11;14)(q13;q32) which leads to the deregulation and upregulation of Cyclin D1 which is an important regulator of the G1 phase of the cell cycle. As per MIPI, Cyclin D1 plays an important role in the cell cycle regulation G1-S transition following mitotic growth factor signaling. Cyclin D1 binds to CDK4 and CDK6 to form a CDK/cyclin complex able to phosphorylate the tumor suppressor retinoblastoma (RB1) facilitating cell cycle progression. Thus, cyclin D1 overexpression would contribute to the lymphomagenesis in MCL by overcoming the suppressor effect that retinoblastoma performs in the G1/S transition.1

Majority of patients with mantle cell lymphoma present with advanced stage disease (70%). While 75% of the patients initially present with lymphadenopathy, extranodal disease is the primary presentation in the remaining 25%. Common sites of nodal involvement are the lymph nodes, spleen (45-60 percent), Waldeyer’s ring, bone marrow (>60 percent), blood (13-77 percent) and extranodal sites like the gastrointestinal tract.2

Mantle cell lymphoma can involve any region of the gastrointestinal tract, occasionally presenting as multiple intestinal polyposis. A prospective study on gastrointestinal involvement of mantle cell lymphoma showed the following results: stomach (57 percent), duodenum (52 percent), jejunum/ileum (87 percent), colon (90 percent) and rectum (69 percent).4 Up to a third of patients have systemic B symptoms, such as fever, weight loss and night sweats.

Initial diagnostic evaluation include a complete blood count, blood chemistries for renal and liver function; computed tomography scans of chest, abdomen and pelvis and an aspiration and biopsy of the bone marrow. Many patients have circulating lymphoma cells detectable by a peripheral blood smear or by flow cytometry.2,11 Immunohistochemistry can still be false negative, mainly due to the quality of the material.12

Cell proliferation (Ki-67) has been exploratively analyzed as an important biologic marker and has shown strong additional prognostic relevance. CCND1 expression, due to the presence of t(11;14)(q13;q32) translocation, is present in virtually all cases of MCL,23 however, a very few cases that do not express CCND1 might exist.15 Recently a new prognostic model has been introduced by the German Lymphoma Study Group called the MIPI (MCL International Prognostic Index) on the basis of age, performance status, serum lactate dehydrogenase and leukocyte count.16 As per MIPI, patients are classified into low risk (44% of patients, median overall survival not reached), intermediate risk (35%, 51 months), and high-risk groups (21% 29 months).

The treatment approach to newly diagnosed patients with MCL depends on the patient’s eligibility for stem cell transplantation (SCT). Those who are eligible are usually treated with either rituximab-CHOP (cyclophosphamide, doxorubicin, vincristine and prednisone) followed by SCT or rituximab-HyperCVAD (cyclophosphamide, vincristine, doxorubicin, decadron, cytarabine, and methotrexate) followed by observation.25

There are basically three potential induction regimens among several: 1) Standard doxorubicin-containing regimens such as R-CHOP 2) intensive combination chemotherapy regimens, including anti-metabolites such as R-HyperCVAD; and 3) purine analog-based regimens such as R-FCM (fludarabine, cyclophosphamide, and mitoxantrone). There have been limited studies using single-agent radioimmunotherapy (RIT) for patients with relapsed MCL.

The role of HSCT remains controversial and is not clearly defined despite the availability of newer therapies in MCL. Most of the studies that analyzed the role of HSCT in MCL were single-center trials with small number of subjects. Thus individualizing therapeutic approaches are essential to improve outcome in this aggressive disease.26 If the patient is <75 years old and in excellent physical condition, then autologous stem cell transplantation (SCT) can be considered a potential option but if the patient is elderly and not a SCT candidate, the use of rituximab-containing chemotherapy regimen is appropriate.25

Newer agents have been identified for relapsed MCL. Proteosome inhibitors like Bortezomib are inhibitors of the intracellular protein degradation pathway known as the proteosome and has shown antitumor activity in relapsed MCL in several studies. Thalidomide, used primarily for multiple myeloma is another agent that has been evaluated in MCL alone and in combination.27 Temsirolimus is an inhibitor of the mammalian target of rapamycin (mTOR) targeting the phosphatidyl inositol 3 kinase pathway.17 This mechanism inhibits the translation of mRNA, including cyclin D1 mRNA, making this agent a sound therapeutic agent in MCL.

**CONCLUSION**

MCL has the most aggressive clinical course thus necessitating the need for early diagnosis and management. Patients generally present at later stages of disease with extensive involvement of lymph nodes. Nevertheless, any gastric mass, as in this case can be a rare presentation of primary isolated gastrointestinal MCL without any other overt signs and symptoms. Cyclin D1 plays an important role in the pathogenesis of MCL and is a highly specific marker for diagnosis. No standard guidelines have been established till date for the treatment of MCL and its prognosis remains poor. Our patient responded well to intensive chemotherapy and subsequent allogeneic hematopoietic stem cell transplantation and remains in complete remission.

**REFERENCES**


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