Male breast cancer and mantle cell lymphoma in a single patient
A case report and literature review
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Abstract
Rationale: Although still relatively rare, multiple primary malignant neoplasms (MPMNs) have been increasingly reported in recent years.

Patient concerns and diagnoses: A 65-year-old man was referred to our hospital for a painless, incidental left axillary lump. Ultrasound showed enlarged left axillary lymph nodes. An excisional biopsy was conducted on 3 lymph nodes. The pathological diagnosis was determined to be metastatic adenocarcinoma and mantle cell lymphoma (MCL) in the lymph nodes. Further physical examination of the patient yielded a 1.5-cm hard, left subareolar mass.

Interventions and outcomes: The patient underwent modified radical mastectomy. The diagnosis was grade II invasive ductal carcinoma (stage IIA). The axillary lymph node showed MCL (stage I, group A), but not metastatic ductal carcinoma. The patient received chemotherapy, including 6 courses of CHOP (A chemotherapy protocol consists of cyclophosphamide 1.2 g day 1, doxorubicin 80 mg day 1, vindesine 4 mg day 1, and prednisone 90 mg from day 1 to 5) for lymphoma and breast cancer. The patient was also administered endocrine therapy. After a 54-month follow-up, the patient was well with no evidence of disease.

Lessons: MPMNs are easily misdiagnosed as a primary and metastatic tumor, leading to delayed or erroneous treatment. Male breast cancer in a patient with MCL is rare. Early diagnosis and proper therapy are necessary for an optimal prognosis. Further studies are required to define the mechanisms and risk factors of MPMNs.

Abbreviations: IHC = immunohistochemistry, MBC = male breast cancer, MCL = mantle cell lymphoma, MPMNs = multiple primary malignant neoplasms.

Keywords: male breast cancer, mantle cell lymphoma, multiple primary malignant neoplasms

1. Introduction
Billroth was the first to report a case of dual malignancies in 1889.[1] Multiple primary malignant neoplasms (MPMNs) are rarely reported and are defined as the diagnosis of ≥2 independent, primary malignancies of different histologies/origins in a single individual. The prevalence of MPMNs varies from 0.73% to 11.70% of all patients diagnosed with carcinomas in western countries.[1] With improved health care provision, an increasing number of patients survive cancer for longer intervals, which in turn increases the risk of developing a second tumor. In this study, we report a patient with male breast cancer (MBC) and coexisting mantle cell lymphoma (MCL).

2. Patient information
A 65-year-old man was referred to our hospital for a painless, incidental left axillary lump.

2.1. Clinical findings
Examination by palpation revealed a well-defined, rounded 4-cm mass of soft texture and distinct boundary under the left armpit. There was no fixation to the skin.

2.2. Diagnostic assessment
Ultrasound showed enlarged left axillary lymph nodes. An excisional biopsy was conducted on 3 lymph nodes, measuring 5.0, 4.0, and 0.8 cm, respectively. These nodes were white to tan and fleshy with areas of hemorrhage and necrosis. The resection specimens were fixed in formalin, routinely processed, and 3-μm sections were stained with hematoxylin and eosin. Immunohistochemistry (IHC) was also performed.
Microscopic examination of 2 of the 3 lymph nodes showed effaced architecture, being replaced by small- to medium-sized atypical lymphocytes (Fig. 1AA), with a mantle zone and partly nodular growth. On immunohistochemical examination, the neoplastic lymphoid cells were found to be positive for CD5 (Fig. 1B), CD20 (Fig. 1C), cyclin D1 (Fig. 1D), CD79α, Bcl-2, p53, and Ki-67 (5%), and negative for CD3, CD45RO, CD10, Bcl-6, Mum-1, CD21, and CD23. Therefore, a diagnosis of MCL was rendered.

Examination of 1 of the 3 lymph nodes revealed the presence of malignant cells, with enlarged hyperchromatic nuclei containing duct/tubule formations. IHC showed that the malignant cells were positive for CK7, CK19, CA153, CEA, cyclooxygenase-2, and synaptophysin, and negative for thyroglobulin, thyroid transcription factor-1, prostate-specific antigen, chromogranin A, CA19-9, CK20, AFP, and CD117. The pathological diagnosis was determined to be metastatic adenocarcinoma.

Further physical examination of the patient yielded a 1.5-cm hard, left subareolar mass with a smooth surface and ill-defined margin that was fixed to the underlying tissue. A computed tomography scan showed the patient’s subareolar mass with axillary lymph node enlargement (Fig. 2). A lumpectomy was performed and frozen sections showed adenocarcinoma with duct/tubule formations.

The patient underwent modified radical mastectomy. The diagnosis was invasive ductal carcinoma (grade II) (Fig. 3AA). IHC showed that the malignant cells were strongly positive for estrogen receptor (ER) (Fig. 3B) and progesterone receptor (PR) (Fig. 3C), positive for E-cadherin, p53, and Ki-67 (50%), and negative for human epidermal growth factor receptor 2 (HER2) (Fig. 3D), gross cystic disease fluid protein-15, synaptophysin, chromogranin A, and high-molecular-weight cytokeratin. The resected margins were negative. A total of 30 axillary lymph nodes showed no metastatic ductal carcinoma, but 28 of 30 axillary lymph nodes showed MCL.

The final diagnosis was grade II infiltrating ductal carcinoma of the left breast (stage IIA) and MCL of the left axillary lymph node (stage I, group A).

2.3. Therapeutic intervention

The patient received chemotherapy, including 6 courses of CHOP (cyclophosphamide 1.2 g day 1, doxorubicin 80 mg day 1, vindesine 4 mg day 1, and prednisone 90 mg from day 1 to 5) for lymphoma and breast cancer. The patient was also administered endocrine therapy.

2.4. Follow-up and outcomes

The patient underwent ultrasonography every 6 months after the operation. He was well with no evidence of recurrence or distant metastasis after 54 months of follow-up.

3. Discussion

The diagnosis of multiple primary malignancies in this study was made according to the criteria developed by Warren and Gates\(^2\): each tumor must be malignant; each tumor must have its own unique pathological features; and metastasis or recurrence must be excluded. Depending on the time of diagnosis, the dual malignancies can be synchronous or metachronous. This case showed a combination of MBC and MCL, 2 completely different
cancer types, diagnosed at the same time, and therefore met the
criteria for simultaneous MPMNs.

The pathogenesis of MPMNs remains unclear. Risk factors
include age, genetic predisposition, alcohol, smoking, and
chemoradiotherapy.[3] The association between MBC and
MCL is unknown and should be further investigated. This
patient denied smoking or alcohol intake, had no history of
exposure to radioactive substances and had no family history of
cancer. One possible mechanism of the formation of the MPMNs
in this patient may be immune defects. In addition, the polycentric
theory of tumorigenesis[4] could also be a possible mechanism.
Based on this theory, there are multiple tumor-susceptible centers
in the human body. Eventual tumor formation is subject to
different carcinogenic factors, locations, and types, resulting in
multiple primary tumors. The incidence rates of both MBC and
MCL are very low, underpinning the rarity of the current case.

MBC is rare, accounting for ~0.6% of all breast cancers and
<1% of all cancers in men.[5] The pathogenesis of MBC is not
clear. Known risk factors include genetic defects (BRCA2),
estrogen–androgen imbalance, radioactive injury, and testicular
disease (cryptorchidism, orchitis, or orchiectomy).[6] There is no
direct evidence of an association between gynecomastia and
MBC. MBC frequently displays skin, nipple, and chest involve-
ment at the early stages, with the periareolar lymphatic network
facilitating tumor metastasis. Moreover, MBC has a poor
prognosis. Based on 2537 cases and 38,316 cases of breast
cancer in men and women, respectively, Giordano et al
retrospectively found that, when compared to female breast
cancer, MBC had a later age of onset, demonstrated a higher rate
of lymph node metastasis, was diagnosed at later clinical stages,
and had a higher tendency to be ER/PR-positive.[5] The median
age of onset has been reported to be 68 years, with half of these
patients being diagnosed with stage III or IV disease.[7] All
pathological types of breast cancer found in women have been
reported in men, with invasive ductal carcinoma accounting for
the majority of cases (93.7%), followed by papillary carcinoma
(2.6%), medullary carcinoma, tubular carcinoma, mucinous
carcinoma, metaplastic carcinoma, inflammatory breast cancer,

| Event number | Event                                                                 |
|--------------|----------------------------------------------------------------------|
| 1            | A 65-year-old man was referred to our hospital for a painless, incidental left axillary lump |
| 2            | Ultrasound showed enlarged left axillary lymph nodes                  |
| 3            | An excisional biopsy was conducted on 3 lymph nodes. The pathological diagnosis was determined to be metastatic adenocarcinoma and mantle cell lymphoma in the lymph nodes |
| 4            | Further physical examination of the patient yielded a 1.5-cm-hard, left subareolar mass |
| 5            | The patient underwent modified radical mastectomy                     |
| 6            | The diagnosis was invasive ductal carcinoma grade II. The axillary lymph node showed MCL (stage 1, group A), but not metastatic ductal carcinoma (grade II) |
| 7            | The patient received chemotherapy, including 6 courses of CHOP for lymphoma and breast cancer. The patient was also administered endocrine therapy |
| 8            | After a 54-month follow-up, the patient was well with no evidence of disease. |

MCL = mantle cell lymphoma.
The current case showed tubular structures and was compatible with infiltrating ductal carcinoma. Most MBCs have been reported to be hormone receptor positive and HER2 negative, which is similar to the findings in this case (Table 1).

MCL is a subtype of B-cell non-Hodgkin’s lymphoma with a unique immune phenotype, genetic features, and clinical characteristics. MCL is relatively rare, accounting for <10% of all lymphomas. In addition, this disease is more common in older men, often involving the gastrointestinal tract, liver, spleen, bone marrow, and peripheral blood, with diagnosis at later clinical stages (III or IV). The patient in this report had lymphadenopathy, but not splenomegaly, and the bone marrow was spared. The MCL originated from the mantle zone, and cytogenetically showed the characteristic t (11;14) (q13;q32) translocation involving the BCL-1 and immunoglobulin heavy chain genes; the enhanced BCL-1 gene expression resulted in cyclin D1 overexpression. In this patient’s case, the lymph node structure was effaced by single small- and medium-sized lymphocytes, with a partial mantle zone and nodular-like growths and atrophy of the non-neoplastic follicular centers. Immunotyping showed a cell profile that confirmed the diagnosis of MCL. Because of different tumor pathological types and clinical stages, MPMNs have different treatment principles. Therefore, the appropriate treatment should be based on the specific characteristics of the primary tumors. MPMNs are easily misdiagnosed as a primary and metastatic tumor, or the second primary tumor is undetected altogether, leading to delayed or erroneous treatment. Although MBC is rare, the incidence of this tumor is increasing. Current treatment principles are based on protocols for postmenopausal female breast cancer patients. However, whether this represents the optimal therapy for male patients is uncertain, as biological and epidemiological differences exist. MCL runs an aggressive clinical course and is not sensitive to radiotherapy or chemotherapy. The prognosis is poor, with an average survival period of 2 to 5 years.

To the best of our knowledge, this is the first report of MBC in a patient with MCL. This diagnosis is consistent with MPMNs, which have an increasing incidence and an unknown underlying mechanism. Early diagnosis and treatment are needed for optimal prognosis. Further studies are required to define the mechanisms and risk factors of MPMNs. Molecular genetics could be used in the early detection and prevention of MPMNs, potentially increasing patient survival.

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