CASE REPORT

Pulmonary Mucosa-associated Lymphoid Tissue Lymphoma with Spontaneous Regression after Computed Tomography-guided Needle Biopsy: A Case Report and Summary of 8 Reported Cases

Kazuaki Fukushima¹, Susumu Hirosako¹, Yuki Tenjin¹, Yosuke Mukasa², Keisuke Kojima¹, Sho Saeki¹, Shinichiro Okamoto¹, Hidenori Ichiyasu¹, Kazuhiko Fujii¹, Yoshitaka Kikukawa³, Koichi Kawanaka⁴ and Hirotsugu Kohrogi¹

Abstract

A 72-year-old woman was admitted to our hospital with a solitary right lung nodule. She had no symptoms and no abnormal physical findings except for bladder cancer. Tumor markers were mildly elevated but no other abnormal laboratory data were found. The nodule was diagnosed to be pulmonary mucosa-associated lymphoid tissue lymphoma on computed tomography-guided needle biopsy. Thereafter, she first underwent surgery for bladder cancer. The lung nodule was found to have slightly increased at three months and then disappeared at 15 months after the biopsy. The notable clinical course of this rare disease suggests the effectiveness of a non-interventional treatment strategy.

Key words: mucosa-associated lymphoid tissue lymphoma, spontaneous regression, computed tomography-guided needle biopsy


Introduction

Mucosa-associated lymphoid tissue (MALT) lymphomas occur frequently in the stomach and rarely in the lungs. The prevalence rate of pulmonary MALT lymphoma accounts for approximately 0.3-0.45% of all lung malignant tumors (1). Prospective trials have not so far been conducted for analyzing the appropriate treatment of pulmonary MALT lymphomas because the number of cases is extremely small. Therefore, no optimal treatment modality for this disease has yet been established (2). In general, localized tumors have been treated by surgical resection or radiation therapy and extended tumors have been treated by systemic chemotherapy (3). Therefore, little is known about the progressive course of untreated pulmonary MALT lymphomas. In general, pulmonary MALT lymphomas tend to progress indolently and have favorable outcomes (2); however, the spontaneous regression of this disease is very rare.

We herein report an unusual case of a localized pulmonary MALT lymphoma that disappeared spontaneously after the patient underwent a percutaneous needle biopsy and bladder cancer surgery.

Although conducting future clinical trials for the treatment of this disease will be difficult, if similar series of case reports are eventually accumulated, then the observation of pulmonary MALT lymphoma may be one of the recommended therapeutic choices.

¹Department of Respiratory Medicine, Kumamoto University Hospital, Faculty of Life Sciences, Kumamoto University, Japan, ²Department of Respiratory Medicine, Kumamoto Social Insurance General Hospital, Japan, ³Department of Hematology, Kumamoto University Hospital, Faculty of Life Sciences, Kumamoto University, Japan and ⁴Department of Diagnostic Radiology, Kumamoto University Hospital, Faculty of Life Sciences, Kumamoto University, Japan

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Correspondence to Dr. Susumu Hirosako, hirosako@kumamoto-u.ac.jp
Case Report

A 72-year-old woman with an abnormal shadow of the right lung and no symptoms presented to our hospital. The patient had a past history of chronic gastritis, surgery for a uterine myoma, and fever and diarrhea after the ingestion of sho-saiko-to. She had a 20-pack-year history of smoking and had no unusual family history. She had repeatedly undergone transurethral resections of bladder tumors (TUR-BT) for 15 years, and a total cystectomy with an ileal conduit was planned because a cancerous tumor had progressed. Before the operation, chest computed tomography (CT) was taken, and it revealed a solitary nodule in the right lower lobe of the lung. The lung nodule had remained undiagnosed by bronchofiberoscopy at her previous hospital, and she was thus referred to our hospital for diagnosis.

Physical examination showed no findings except for a surgical scar on the abdomen. Laboratory investigation revealed that carcinoembryonic antigen (CEA) was 5.8 ng/mL (normal: ≤3.4 ng/mL) and cytokeratin 19 fragment (CYFRA) was 3.8 ng/mL (normal: ≤3.5 ng/mL). Lactate dehydrogenase (LDH) and soluble interleukin-2 receptor were within the normal ranges. β-D glucan and fungal antigens were normal. A chest X-ray and chest CT showed a nodule measuring 12-mm diameter localized in the right lower lobe (Fig. 1A, 2A). We performed a CT-guided needle biopsy of the nodule.

Histological examination with Hematoxylin and Eosin (H&E) staining showed the proliferation of small lymphocytes with little atypia (Fig. 3A and B). Immunohistochemical staining proved that most of the cells in the lymphatic follicles were clusters of differentiation (CD) 20 positive (Fig. 3C). Immunostaining of cytokeratin confirmed the presence of a lympho-epithelial lesion (LEL), which indicated the infiltration of lymphocytes into the epithelium of the gland duct (Fig. 3D). In addition, we performed molecular analyses to evaluate the clonality of the tumor using paraffin-embedded tissue specimens. We detected the gene rearrangement of the immunoglobulin heavy chain (IGH)
Figure 3. A: Histological examinations showed an accumulation of small lymphoid cells [Hematoxylin and Eosin (H&E) staining, ×40]. B: Lymphoid cells consist of atypical small lymphocytes (H&E staining, ×200). C: Cells in the lymphatic follicle include clusters of differentiation (CD) 20 positive (CD20 stain, ×40). D: Infiltration of lymphoid cells into the epithelium, defined as lympho-epithelial lesion (LEL) (Cytokeratin stain, ×200).

DH1-6-JH by polymerase chain reaction (Fig. 4), although we did not detect the API2/MALT1 fusion gene by fluorescence in situ hybridization (FISH) and the monoclonal distribution of kappa/lambda light chains by in situ hybridization (ISH). According to these findings, we diagnosed the nodule to be pulmonary MALT lymphoma.

As the lymphoma was localized and the patient was asymptomatic, bladder cancer surgery was initially performed, and we observed the progress of the lymphoma by imaging until the patient had recovered from surgery. The solitary lung nodule increased slightly on the CT image at three months after the biopsy (Fig. 2B), but subsequently became unclear on a chest X-ray nine months after the biopsy (Fig. 1B), and then it completely disappeared on the CT image 15 months after the biopsy (Fig. 2C). To date, no recurrence has been found either in the lung or at any other sites as of 40 months after the initial biopsy.

Discussion

MALT lymphomas were initially suggested by Isaacson and Wright in 1983 (4) and described in the World Health Organization classification of tumors of hematopoietic and lymphoid tissues as an extranodal marginal zone B-cell lymphoma of the MALT type, which is a type of low-grade extranodal lymphoma (5). MALT lymphoma most frequently arises from the stomach, and it can involve the small intestine, salivary gland, lung, head and neck, ocular adnexa, skin, thyroid, and breast (5). Helicobacter pylori infection has been proved to be a potential cause of gastric MALT lymphoma, and the eradication of the pathogen could resolve the tumor (6). However, the etiology of pulmonary MALT lymphomas remains to be elucidated.

The prevalence of pulmonary MALT lymphoma is very small; Kurtin reported that among the 10,046 new diagnoses of non-Hogkin lymphoma, 106 (1.1%) originated from the lung and 50 (0.5%) of such cases were pulmonary MALT lymphomas (7). Pulmonary MALT lymphomas present with the following pathologic features: intraparenchymal masses composed of centrocyte-like cells, plasmacytoid lymphocytes and plasma cells that form LEL, the infiltration of neoplastic lymphocytes into the bronchial or bronchiolar epithelium, and a lymphangitic growth pattern (7). Because pulmonary MALT lymphoma is difficult to diagnose from small lung specimens, the diagnosis is often made by lung surgery that also serves as therapy for the localized disease (1). In fact, the proportion of patients who were diagnosed as having lymphoma by transbronchial biopsy or radiologically-guided transthoracic core-needle biopsy is only one-third, while the remaining two-thirds required video-assisted thoracoscopic surgery or thoracotomy to achieve definitive diagnoses (3). In the present case, using needle-biopsied specimens, pul-
Figure 4. GeneScan results of polymerase chain reaction products of the biopsied sample showing the gene rearrangement of the immunoglobulin heavy chain DH1-6-JH. The x-axes represent the molecular size (base pairs), and the y-axes represent the fluorescence intensity. A: Positive control of the peak of DH1-6-JH. B: DH1-6-JH gene products of the tumor (arrow) showing a higher peak than the positive control (A). C: Polyclonal control.

Pulmonary MALT lymphoma was diagnosed based on the histological proliferation of small lymphocytes with little atypia, the infiltration of CD20-positive cells in the lymphatic follicles, LEL, and the identification of the gene rearrangement of IGH DH1-6-JH.

Everson and Cole defined the spontaneous regression of cancer as the partial or complete disappearance of a malignant tumor in the absence of therapy which is considered to exert an influence on neoplastic diseases (8), and indicated the frequency of occurrence was one in 60,000-100,000 cases of cancer. Among a total of 741 cases of spontaneous cancer regression originating from various organs, the patients with cancer originating in the lung consisted of 23 cases [3.1% (8-10)]. The mechanism of spontaneous regression is unknown, however, several of these regressions occurred after invasive procedures such as biopsies, or after complications with bacterial infection (11, 12); some may have activated immune reactions, which might have contributed to the resolution of the disease. Cole showed that 71 (41%) cases of 176 spontaneous regressions were associated with surgery or surgical trauma (11). In the present case, stimulation by needle biopsy may have directly triggered a regional immune reaction or the trauma due to the bladder surgery may have activated the patient’s systemic immunity, which could have led to tumor remission.

The first report of spontaneous regression of pulmonary
MALT lymphomas was presented by Troch et al (13). In that report, six of 11 patients with non-treated pulmonary MALT lymphomas showed spontaneous regression, but not complete remission, during an observation period with a median of 28 months (13). In some patients from that study, this phenomenon was only seen in the lung, while extrapulmonary lesions did not change significantly. These facts indicate that the circumstances of lung tissue surrounding the tumor or the nature of the lymphoma originating from lung might be distinct from those of other sites. In addition, three out of six patients with spontaneous regression have at least one episode of tumor increase before regression (13). These courses were similar to that of the present case. A possible mechanism of the change in tumor size was the fact that the tumor initially increased in size after the invasive biopsy and healed after activation of the immune system. Sato et al. described a pulmonary MALT lymphoma of an 82-year-old man that regressed spontaneously after a bronchoscopic biopsy and had not relapsed during a 20-month period (14). Recently, Kang et al. described a 57-year-old woman with spontaneously disappeared pulmonary MALT lymphoma after thoracoscopic lung biopsy, although the lymphoma remained in her bone marrow (15). We summarized the clinical characteristics of these eight cases of pulmonary MALT lymphomas with spontaneous regression as well as the present case (Table). All cases were diagnosed by invasive procedure: endobronchial biopsy, CT-guided needle biopsy, or surgical biopsy (details of all patients are not known). The present case was a patient with a higher age and localized disease in a unilateral lung, which completely disappeared after the trauma induced by the CT-guided needle biopsy and bladder cancer surgery.

The prognostic factors of pulmonary MALT lymphoma other than lymphoma-specific therapy were indicated in the past studies. Patients whose lymphomas with LEL had better outcomes than those without LEL, and patients with amyloid-containing tumors showed a poorer prognosis than those without amyloid (7). The present case identified LEL, but not amyloid deposition in the biopsied lung specimens.

We herein reported a rare case of pulmonary MALT lymphoma that showed spontaneous regression, which completely disappeared after a CT-guided needle biopsy. When pulmonary MALT lymphoma is asymptomatic and not progressive, observation without immediate therapy should therefore be considered as a treatment option until the lymphoma size increases. In order to determine the appropriate management for pulmonary MALT lymphoma, more information concerning the progressive course of this disease should be obtained.

The authors state that they have no Conflict of Interest (COI).

References

Table. Summary of 8 Reported Cases and the Present Case of Pulmonary Mucosa-associated Lymphoid Tissue Lymphoma with Spontaneously Regression.

| Age (years) | Median 55 (range 42–82) | 72 |
| Gender | Male 2, Female 6 | Female |
| Bilateral/unilateral lung lesion | Bilateral 3, Unilateral 5 | Unilateral |
| Extrapulmonary organ involvement | Yes 4, No 4 | No |
| Autoimmune disease coincidence | Sjögren’s syndrome 2, None 6 | None |
| Degree of regression | Complete 2, Incomplete 6 | Complete |

* References (13–15)


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