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Successful Management with CHOP for Pulmonary Lymphomatoid Granulomatosis in a Dog

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ABSTRACT. A 3-year-old, spayed female miniature dachshund was presented for vomiting and anorexia. Thoracic radiographs and CT scan revealed abnormal pulmonary opacities at bilateral caudal lobe. Cytological analysis of the pulmonary mass revealed the presence of large lymphohistiocytic cells and small lymphocytes with occasional neutrophils and plasma cells. An open lung biopsy was performed and a diagnosis of pulmonary lymphomatoid granulomatosis (LYG) was made. The dog was administered CHOP based therapy (modified UW-25), and it survived for 1,022 days after admission. Immunohistochemistry revealed pulmonary lesions consisted of many CD79a positive B cells aggregation and proliferation with prominent angiocentric pattern. This was the first case of canine pulmonary LYG managed by CHOP chemotherapy.

KEY WORDS: canine, CHOP, lymphomatoid granulomatosis, pulmonary.

Lymphomatoid granulomatosis (LYG) is a unique form of pulmonary angiitis and granulomatosis which was first described by Liebow et al. in 1972 [12]. Histologically it is a necrotizing angiocentric and angiodestructive infiltrative process composed of small lymphocytes, plasma cells, histiocytes and atypical lymphoreticular cells [11, 12]. In veterinary medicine, canine pulmonary LYG is a rare pulmonary lymphoproliferative disease; very few treatment studies have been conducted and no standard treatment is established [2, 17]. In this report, we describe clinical and pathological findings of a canine LYG, which results in long-term survive with CHOP therapy, modified version of the University of Wisconsin-Madison (UW-25) protocol for canine lymphoma [7].

A 3-year-old, spayed female miniature dachshund, weighing 3.5 kg, was presented to Osaka Prefecture University Veterinary Clinical Center for 4 week-history of vomiting and anorexia. Clinical examination revealed pyrexia and dyspnea. A peripheral lymphadenopathy was not recognized. There was no evidence of hepatomegaly or splenomegaly. A complete blood count showed abnormal leukocytosis (40,100 cell/µl; reference range 6,000 to 17,000/µl) with mature neutrophilia (23,258 cell/µl; reference range 3,000 to 11,500 /µl) and lymphocytosis (16,441 cell/µl reference range 1,000 to 4,800/µl). The blood biochemical findings included an increased total protein concentration (8.4 g/dl) with a high globulin concentration (A/G=0.5). Increased α-2 and γ fraction were identified via serum electrophoresis. The dog was negative for Dirofilaria immitis infestation. Thoracic radiographs revealed abnormal pulmonary opacities at bilateral caudal lobe (Fig. 1). Radiographic evidence of sterna lymphadenopathy was also noted. A chest computed tomographic (CT) scan of the thorax showed bilateral, wedge-shaped opacities (Fig. 2). Cytological analysis of the pulmonary mass, which was performed by fine-needle aspiration revealed the presence of large lymphohistiocytic cells and small lymphocytes with occasional neutrophils and plasma cells. The dog was treated with Enrofloxacin (5 mg/kg po SID) and Cephalexin (25 mg/kg po BID) from first admission. Although the number of neutrophil and lymphocyte was normalized, clinical signs did not resolved by antibiotics therapy. In an attempt to obtain a diagnosis, an open lung biopsy was performed in the left caudal lobe field on day 20 after admis-

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Fig. 1. Thoracic radiograph showing the presence of sterna lymphadenopathy (arrowhead) and abnormal pulmonary opacities (arrows) at bilateral caudal lobe (left lateral).
sion.  Enucleated firm, cream-colored nodular mass was found and that was 5 cm in diameter (Fig. 3A).

Histopathology revealed that an atypical angiocentric infiltration and proliferation of pleomorphic mononuclear cells that spread into the adjacent alveolar parenchyma (Fig. 3B and 3C).  The infiltrating mononuclear cells resembled large lymphoid cells (Fig. 3D).  Based on the histopathological findings, a diagnosis of pulmonary LYG was made.  The dog was started on a multiple drug protocol based on modified UW-25.  This protocol consisted of vincristine (0.5 mg/m² iv), cyclophosphamide (250 mg/m² iv), doxorubicin (30 mg/m² iv), and prednisone, without L-asparaginase.  Seven days after the first administration of vincristine, the pulmonary mass and sterna lymphadenopathy markedly decreased in size and almost disappeared (Fig. 4).  After treatment with chemotherapy protocol for 25 weeks, the condition of dog was very well and the dog was considered in complete remission. The dog was scheduled to be monitored at routine appointments including clinical examination, CBC, serum chemistry panel, thoracic radiographs and abdominal ultrasound.  On day 786 after presentation, a resurgence of clinical signs, pyrexia, dyspnea, diarrhea, and vomit, occurred. Ultrasound of the abdomen revealed abdominal lymphadenopathy and gastric wall thickening. The dog was tried to treat on a modified UW-25 again. However, because these treatments were not effective on this dog, administration of mitoxantrone (5 mg/m²) was started from day 830 once a month and clinical improvement and partial remission of abdominal lymphadenopathy was achieved. From day 1,014, dyspnea, diarrhea and severe vomit were observed. The dog finally died 1,022 days after the first presentation.  Complete necropsy was conducted.  Necropsy revealed lobar pneumonia with angiocentric infiltration of lymphocytic cells in the pulmonary parenchyma.  Clustered lymphocytes were also observed in lamina propria and submucosa of stomach.  Immunohistochemical examination of the lung revealed many CD79a positive B cells and a few CD3 positive T cells in the lymphoid cells around the blood vessels (Fig. 5A and 5B).  Those angiocentric cells were negative for macrophage/histiocyte markers, AM-3K and lysozyme (Fig. 5C and 5D).  Polymerase chain reaction (PCR) assays for the immunoglobulin and T-cell receptor genes were performed according to the protocol described by Burnett et al. [3] and Valli et al. [21].  The clonality for B lymphocyte was detected from lung tissues.

The typical character described in human and canine pulmonary LYG include multiple, bilaterally nodular masses [10, 17].  Moreover, LYG is an angiocentric and angiodestructive lymphoproliferative disease involving extranodal sites [2, 10].  In most patients with human LYG, similar cellular infiltrates are found in other organs such as skin, kidney, and central nervous system [10].  Canine LYG also often metastasizes to the regional lymph nodes, liver, and spleen [17].  Moreover, the case of human LYG affecting
that diffuse multinodular patterns of pulmonary lymphoma is rare in canine [1] and human [4]. Moreover, although one report of canine pulmonary lymphoma revealed that cytological findings were consistent with lymphoma [15], this case revealed the presence of large lymphoid cells and small lymphocytes with occasional neutrophils and plasma cells, that were consistent with LYG previously reported [2, 12]. Finally, histopathology in the present case was in agreement with histologic criteria for LYG, namely a nodular infiltrate of an angiocentric/angiodestructive lymphoid infiltrate within the pulmonary parenchyma [2, 12]. Those results supported that this case was diagnosed as a LYG.

Human LYG comprised of a usually small number of B cells admixed with prominent T cell infiltrates, and moreover, this is considered a form of T-cell rich B cell lymphoma on the basis of histopathological features and immunophenotypic analysis [8]. The phenotype of canine LYG is now controversial because immunohistochemical analyses of the canine LYG have been documented in only a few papers [14, 16, 20]. In one report, canine LYG has been associated with malignant T-cell type lymphocytes and is considered a T-cell lymphoma [20]. Moreover, the presence of clonally expanded T-cell was revealed by polymerase chain reaction (PCR) in a CD3-positive canine cutaneous LYG, recently [19]. The other report of a pulmonary case, however, showed that there were both CD3 positive T cells and CD79 positive B cells [16]. The immunophenotypic analysis showed that canine cutaneous LYG had both neoplastic T- and B-cells [14]. Same as later reports, the present pulmonary LYG had both T cell and B cell lineage. Moreover, B cell monoclonality was revealed by PCR in this case. Therefore, the present dog had features similar to those of canine LYG and human LYG.

Canine LYG is very difficult to diagnose, and treatment has not been attempted actively [17]. In human and veterinary medicine, the LYG was diagnosed by open biopsy and histological evaluation of tissues [6, 10]. Same as those reports, this case could not be diagnosed by cytological examination, due to the marked cytologic variation of sample collection, and thus open lung biopsy was needed. In human LYG, it has been shown that a good response and long-term survival may possibly be obtained through appropriate chemotherapy [9], and the present canine case also could survive for long-term by CHOP chemotherapy. Therefore, the open lung biopsy is necessary to diagnose canine LYG and to treat properly.

Because canine LYG is rare, only a few treatment studies including prednisone, COP and COAP have been conducted and there is no standard treatment [2, 17]. In human medicine, however, the patient with LYG was treated by a more aggressive combination chemotherapy regimen utilizing doxorubicin [5, 9]. To the best of our knowledge, this was the first case of canine pulmonary LYG treated by CHOP chemotherapy, and could have long-term survival. Therefore, CHOP therapy may be effective for canine LYG, as applied for human LYG.

Rescue protocols have not been reported in canine and human LYG. After CHOP was not effective on this dog, administration of mitoxantrone was started once a month and clinical improvement and partial remission was achieved. This drug was used as rescue protocols of canine lymphoma [13]. The results of this report demonstrate that mitoxantrone may be an effective rescue protocol for dogs with LYG.

Further studies including immunohistochemical analyses, PCR clonality analysis and clinical trials will be necessary to confirm the benefits of CHOP and mitoxantrone in the management of canine LYG.

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