A new conceptualization for Mikulicz’s disease as an IgG4-related plasmacytic disease

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Abstract Mikulicz’s disease (MD) has been included within the diagnosis of primary Sjögren’s syndrome (SS), but it represents a unique condition involving persistent enlargement of the lacrimal and salivary glands characterized by few autoimmune reactions and good responsiveness to glucocorticoids, leading to the recovery of gland function. Mikulicz’s disease was recently reported to be associated with elevated immunoglobulin G4 (IgG4) concentrations in the serum and prominent infiltration of plasmacytes expressing IgG4 into the lacrimal and salivary glands. The following features were used for diagnosis: (1) visual confirmation of symmetrical and persistent swelling in more than two lacrimal and major salivary glands; (2) prominent mononuclear cell infiltration of lacrimal and salivary glands; and (3) exclusion of other diseases that present with glandular swelling, such as sarcoidosis and lymphoproliferative disease. These features are not observed in most SS cases. The complications of MD include autoimmune pancreatitis, retroperitoneal fibrosis, tubulointerstitial nephritis, autoimmune hypophysitis, and Riedel’s thyroiditis, all of which show IgG4 involvement in their pathogenesis. Mikulicz’s disease thus differs from SS and may be a systemic IgG4-related plasmacytic disease.

Key words Autoimmune pancreatitis · Glucocorticoid · Immunoglobulin G4 · Mikulicz’s disease · Sjögren’s syndrome

Introduction to Mikulicz’s disease

Mikulicz’s disease (MD) refers to idiopathic, bilateral, painless, and symmetrical swelling of the lacrimal, parotid, and submandibular glands. Because MD and Sjögren’s syndrome (SS) are histologically similar, MD has been considered as a subtype of SS. However, there are some clinical differences between MD and typical SS. In MD, the enlargement of lacrimal and salivary glands is persistent and secretory dysfunction is either not detectable or slight. Further, MD shows good responsiveness to steroids. Serologically, MD patients exhibit normal or hypergamma-globulinemia and normal or hypocomplementemia, but lacks anti-SS-A and anti-SS-B antibodies. Recently, it has also been confirmed that MD patients show elevated serum immunoglobulin G4 (IgG4) concentrations and infiltration of plasmacytes expressing IgG4 into the lacrimal and salivary glands. Thus, MD apparently differs from SS. Attention has also been paid to the pathological similarities between MD and autoimmune pancreatitis (AIP), particularly the role of IgG4, and new developments are expected to clarify the pathogenesis of both diseases. Here, we summarize the results of recent studies and provide an overview of MD.

Historical details and the concept of Mikulicz’s disease

In 1888, Johann von Mikulicz-Radecki reported a case that exhibited bilateral, painless, and symmetrical swelling of the lacrimal, parotid, and submandibular glands. Napp later reported that these symptoms presented with leukemia, lymphoma, sarcoidosis, tuberculosis, and lues. To clarify this, in 1927 Schaffer linked cases with obvious diseases presenting these symptoms as Mikulicz’s syndrome and idiopathic cases as MD. On the other hand, in 1933, Sjögren analyzed 19 cases with keratoconjunctivitis sicca, 2 of which displayed swelling of the major salivary glands.

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The concept of SS was established after this report. In 1953, Morgan and Castleman examined specimens from 18 cases diagnosed with MD. After finding histological similarities between MD and SS, they announced that most cases reported as MD could be considered as SS.6,7 Subsequently, MD has been considered as a subtype of SS, and case reports of MD have disappeared.

However, when we resurveyed the thesis by Morgan and Castleman,5 we identified several problems, such as the inclusion of the many cases that lack swelling of the lacrimal glands or only show swelling of one side of the glands. This led us to question whether these cases were actually MD. The histological characteristics that Morgan reported as being common among the 18 cases were as follows: (1) degeneration and disappearance of acini due to severe mononuclear cell infiltration, (2) proliferation of ductal epithelial cells and duct stenosis, (3) formation of myoepithelial islands, and (4) cystic dilatation of peripheral ducts. However, it is reported that these characteristics were observed not only in SS but also in persistent and destructive pathogenesis such as sialolithiasis in the glands.7 Thus, the exclusion of MD from SS, which Morgan defined is under increasing scrutiny, and numerous studies on the relationships between MD and SS have been undertaken in Japan.

**Differences between Mikulicz’s disease and Sjögren’s syndrome**

Our classification of Mikulicz’s disease

We based our classification of MD on the following features: (1) visual confirmation of symmetrical and persistent swelling in more than two lacrimal and major salivary glands, (2) prominent mononuclear cell infiltration of lacrimal and salivary glands, and (3) exclusion of other diseases that present with glandular swelling such as sarcoidosis and lymphoproliferative disease.7 The aim of this classification is to establish disease independence and clarify the serological and histopathological differences between MD and SS.8

Analyzes were performed in 16 MD patients (4 men and 12 women) who consulted doctors at Sapporo Medical University and its related facilities between April 1997 and October 2005; these cases were compared with primary SS cases. Another 16 patients in whom SS was diagnosed on the basis of the revised Japanese criteria were also studied.9 We defined cases as MD when they satisfied both our classification and the revised Japanese criteria for SS.

Clinical characteristics of Mikulicz’s disease

Our MD patients were mainly middle-aged or elderly females. The average age was 58.6 years (range 25–73 years) with a sex ratio of approximately 3:1 in favor of females. However, male patients predominated when reports from other facilities were analyzed together with our data.10 The enlargement of the lacrimal and salivary glands in MD patients was found to be elastic, painless, and persistent (occurring for more than 3 months). Half of the MD patients did not exhibit keratoconjunctivitis sicca. With regard to salivary gland function, secretion by these glands in MD was normal or slightly decreased, and this improved with steroid treatment. On the other hand, the 16 patients with SS were all females in their forties and fifties. The sex ratio for SS is usually approximately 20:1 (female: male).11 Our results did not contradict the general consensus. The lacrimal and salivary swelling in SS presented repeatedly and disappeared without treatment. The sicca symptoms were severe in SS. We prescribed glucocorticoids to SS patients for treating the swelling of the lateral submandibular gland, but the amount of saliva did not increase. We do not expect improvements in glandular secretion in advanced stages of SS since SS is an autoimmune disease that irreversibly damages glands.12 There were a few reports on the efficacy of a low dose of prednisolone on saliva production in patients with primary SS; however, in the thesis, it was not clear how long these cases had suffered from SS.12,13 It was then mentioned that in authoritative textbooks, systemic corticosteroids were used for treating only severe extraglandular diseases, including diffuse interstitial pneumonia, glomerulonephritis, vasculitis, and peripheral neuropathy.14–16 The impact of steroids on the natural course of SS is not well established. The clinical figures of MD are thus quite different from those of typical SS.

Diagnostic imaging of Mikulicz’s disease

Computed tomography (CT) from the head to the neck in MD patients revealed severe enlargement of symmetrical lacrimal and submandibular or parotid glands. The size of the neck lymph nodes was less than 1 cm. Ultrasonography of the enlarged glands revealed that the foci were irregular, hypoechoic, and multilocular, while the power Doppler method showed the presence of hypervascular masses. Sialography was normal, and the “apple-tree sign,” which is typical of SS, was not observed in MD.

Serological tests in Mikulicz’s disease

Half of the MD patients exhibited hypergammaglobulinemia. A few patients had antinuclear antibodies in MD, and with the exception of one patient, did not have anti-SS-A or anti-SS-B antibodies. On the other hand, SS patients exhibited hypergammaglobulinemia and were largely positive for anti-SS-A (69%) and anti-SS-B (31%) antibodies.

**Mikulicz’s disease and IgG4**

We focused on hypergammaglobulinemia, which was observed in MD. Serum IgG subclasses in each group of MD and SS (each 16 patients) were measured by nephelometry. Mean total IgG levels were 3226.9 mg/dl and 2398.1 mg/dl in MD and SS, respectively. There were no significant differences between the groups. Serum IgG subclasses in MD
were 1256.4 mg/dl and 1111.0 mg/dl for IgG1 and IgG4, respectively, and those in SS were 1624.9 mg/dl and 88.8 mg/dl for IgG1 and IgG4 (P < 0.05, P < 0.0001), respectively. The IgG1/total IgG and IgG4/total IgG ratios were 41.5% and 28.6%, respectively, in MD and 73.0% (P < 0.0001) and 2.8% (P < 0.0001), respectively, in SS (Fig. 1). In healthy controls, these ratios for each IgG subclass were as follows: IgG1, 65%; IgG2, 25%; IgG3, 6%; and IgG4, 4%. IgG1 usually accounts for most of the total IgG. Generally, the amount of IgG4 does not vary with sex or age, and the quantity of IgG4 as well as the IgG4/total IgG ratio tends to remain constant. Our serological analysis revealed that all MD patients had elevated concentrations of IgG4, which has not been observed in any other connective tissue disease, including SS, systemic lupus erythematosus, rheumatoid arthritis, and polymyositis. Elevated serum IgG4 concentrations were very specific to MD patients.

As described above, Morgan and Castleman reported that MD and SS were histopathologically identical. Upon hematoxylin–eosin staining, lacrimal or salivary gland specimens from MD showed severe mononuclear cell infiltration and lymphoid follicles around the ductal and acinar cells. We were not able to identify any difference between MD and SS using typical histopathological analyses. However, Tsubota et al. reported that the frequency of apoptosis of lacrimal gland cells was significantly lower in MD. 

We then examined the lacrimal and salivary glands for anti-IgG4 antibody staining since elevated concentrations of IgG4 were detected in MD patients. The infiltration of numerous IgG4-positive plasmaocytes near acinar and ductal cells and around lymphoid follicles was confirmed in MD; however, the specimens from SS patients showed no plasma cells with IgG4 (Fig. 2). Thus, this finding clearly differentiates between MD and SS (Table 1). It is possible that MD is a systemic IgG4-related plasmacytic disease because in MD, we also detected abundant IgG4-bearing plasma cells in the stomach, colon, and kidney as well as in lymphoid tissues such as the cervical lymph nodes and bone marrow.

### Treatment of Mikulicz’s disease

Mikulicz’s disease is mainly treated by the administration of steroids. We initiate prednisolone at 30-40 mg/day against MD without organ failure. This leads to rapid improvement in glandular swelling as well as in lacrimal and salivary secretion. Prescription of glucocorticoids for 2 months led to an increase from 6.9 mm/5 min to 15.7 mm/5 min (P < 0.05) in Schirmer’s test (16 eyes), which reflects improved lacrimal secretion, and an increase from 1.98 g/2 min to 3.66 g/2 min (P < 0.05) in Saxon’s test (8 MD patients), which reflects improved salivary secretion. Glucocorticoid administration also improved hypergammaglobulinemia. A prescription for 2 months induced decreases in serum IgG levels (from 4270.0 mg/dl to 1440.1 mg/dl; P < 0.005) in 8 MD patients. The serum IgG1 levels decreased from 1632.9 mg/dl to 681.0 mg/dl (P < 0.005) and those of serum IgG4 decreased from 1556.4 mg/dl to 234.7 mg/dl (P <

| Table 1. Clinical characteristics of Mikulicz’s disease and Sjögren’s syndrome |
|-----------------------------|-----------------------------|
| **Mikulicz’s disease**      | **Sjögren’s syndrome**      |
| Age of disease onset        | From 40s to 50s              |
| Sex ratio                   | 20:1 in favor of females     |
| Gland swelling              | Recurrent                   |
| Keratoconjunctivitis sicca  | Mild to severe              |
| Dysfunction of salivary secretion | Mild to severe      |
| Response to steroid         | No change, or sometimes good||
However, the ratio of IgG1/total IgG increased from 38.3% to 46.8% ($P < 0.01$) while that of IgG4/total IgG decreased from 34.9% to 16.7% ($P < 0.005$). When steroids were discontinued, swelling of the lacrimal and salivary glands was observed, and the serum IgG4 levels increased. It is thus necessary to continue prednisolone administration at 5–10mg/day or combine it with an immunosuppressant such as azathioprine.

**Controversial points regarding Mikulicz's disease**

**Prognosis and relationship with lymphoproliferative diseases**

The long-term prognosis of MD is unknown. Among SS patients, the incidence of non-Hodgkin’s lymphoma is 43.8-fold higher than the prevailing incidence in the general population. Sugai classified the severity of SS into three stages: stage I, having only sicca symptoms; stage II, having systemic organ involvement; and stage III, is complicated by lymphoma. Glandular and extraglandular SS may thus be considered as an early stage of lymphoma. There have also been reports on malignant transformation in MD. Recently, Ihrler and Harrison speculated that the first MD case reported by Mikulicz in 1892 was MALT lymphoma based on Mikulicz’s sketch of the specimen. However, none of the present cases exhibited IgH gene rearrangement. It is necessary to continue the analysis of complicating lymphoma in MD as there is no data on its long-term prognosis.

**Interpretation of asymmetrical or individual lacrimal or salivary gland swelling**

In MD, enlargement of only the lacrimal or submandibular glands sometimes leads to other glandular swelling.

Küttner’s tumor is chronic sclerosing sialadenitis that presents with asymmetrical firm swelling of the submandibular glands. It is predominantly found in males. The histological characteristics of Küttner’s tumor are severe fibrosis, atrophic acinus, and mononuclear cell infiltration. Kitagawa et al. analyzed 12 patients with Küttner’s tumor and found that five cases were associated with sclerosing lesions in the extrasalivary glandular tissues. Immunohistochemically, the proportion of IgG4/IgG-positive plasmacytes was more than 45% in Küttner’s tumor, while it was less than 5% in SS. We experienced a case of Küttner’s tumor presenting with enlargement of only the right submandibular gland. We did not evaluate the gland by immunostaining but found elevated concentrations of serum IgG4 (33.1%). We also experienced a case of chronic dacryadenitis presenting with bilateral swelling of only the lacrimal glands. This case also exhibited elevated concentrations of serum IgG4 (26.0%) and prominent infiltration of IgG4-bearing plasma cells into the glands. It is difficult to assess whether Küttner’s tumor and chronic dacryadenitis will develop in MD or whether these differ substantially from MD. We cannot resolve this problem at this point because the opinions of otorhinolaryngologists, ophthalmologists, and pathologists must be considered. However, we have the impression that some of the pathogenic features diagnosed in Küttner’s tumor or chronic dacryadenitis are the same as those of MD. Thus, we must also analyze and resolve the relationships between MD and other diseases.

We regard symmetrical glandular swelling to be an indispensable characteristic for diagnosing MD. We use the name MD out of respect for the first reporter of this condition – Johann von Mikulicz-Radecki. When it is proved that Küttner’s tumor and chronic dacryadenitis are identical to MD, we will call them by another common name. It is possible that bilateral glandular swelling occurs in SS; however, it usually disappears within a few weeks without any treatment. Persistence of the condition for a period of more than 3 months is very rare.

![Specimens of labial salivary glands in patients with Mikulicz’s disease (MD) and Sjögren’s syndrome (anti-IgG4 monoclonal antibody staining, magnified 1:200).](image-url)

a Mikulicz’s disease, b Sjögren’s syndrome. The MD specimen exhibits abundant infiltration of plasmacytes with IgG4.
Autoimmune pancreatitis is an IgG4-related disease. It also exhibits hypergammaglobulinemia, particularly with regard to IgG4, and severe infiltration of IgG4-positive plasmacytes into the pancreas. Autoimmune pancreatitis was proposed by Yoshida et al. in 1995 because it was presumed that autoimmunity was a part of the mechanism of idiopathic chronic pancreatitis. The diagnostic criteria for AIP were designed by the Japan Pancreas Society. AIP include autoimmune hypophysitis, Riedel's thyroiditis, interstitial pneumonitis, sclerosing cholangitis, retroperitoneal fibrosis and interstitial tubular nephritis. In our analysis, the concentrations of serum IgG4, infiltration of IgG4-bearing plasma cells into the glands, and recovery of secretion by glucocorticoid treatment. Furthermore, AIP is sometimes complicated by MD. Thus, MD and AIP are considered to be related. Nakamura et al. reported an AIP case in which IgG4-positive mononuclear cell infiltration was observed not only in the pancreas but also in the salivary gland and kidney. Other complications associated with MD and AIP include autoimmune hypophysitis, Riedel's thyroiditis, interstitial pneumonitis, sclerosing cholangitis, retroperitoneal fibrosis and interstitial tubular nephritis. In our analysis, the concentrations of serum IgG4 tended to be proportional to the frequency of these complications. It is possible that serum IgG4 concentrations reflect disease activity in systemic IgG4-related plasmacytic diseases, which include MD and AIP.

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