Autoimmune Hepatitis and Immunoglobulin G4-Associated Autoimmune Hepatitis

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Key Words
Autoimmune hepatitis · Autoimmune pancreatitis · Immunoglobulin G4

Abstract
Autoimmune hepatitis (AIH) is a disease that is characterized by the presence of autoantibodies and elevated levels of serum immunoglobulin G (IgG) and hepatic enzymes. Its characteristic findings in the liver include interface hepatitis, infiltration of lymphocytes and plasma cells, and rosette formation, and is treated with immunosuppressive drugs. Autoimmune pancreatitis, a pancreatic disease caused by an autoimmune mechanism, is associated with elevated levels of serum IgG4 and the infiltration of IgG4-positive cells into the pancreatic parenchyma, and it is occasionally accompanied by systemic features. This systemic inflammatory disease characterized by the infiltration of IgG4-positive plasma cells and elevated serum IgG4 levels was recently classified as an IgG4-related disease. A few studies have reported AIH cases with infiltrated IgG4-positive plasma cells in the liver, suggesting the involvement of IgG4 in the pathogenesis of AIH. This feature was called IgG4-associated AIH and only two studies have been published. However, the diagnostic criteria of IgG4-associated AIH has not been defined and the epidemiology and clinical features remain uncertain. The degree of IgG4-positive plasma cell infiltration in the liver was different in each article. The serum IgG4 level was not elevated in one study, whereas it was severely elevated in the other. Corticosteroid therapy normalized liver enzymes in both studies. Further studies are needed to define the epidemiological features or diagnostic criteria.

Introduction

The serological features of autoimmune hepatitis (AIH) are the presence of antinuclear antibody (ANA), anti-smooth muscle antibody (SMA) and anti-type 1 liver-kidney microsomal antibody (anti-LKM-1), and elevated levels of serum immunoglobulin G (IgG) and hepatic enzymes. Characteristic findings in the liver include interface hepatitis, infiltration of lymphocytes and plasma cells, and rosette formation. AIH is treated with steroids and immunosuppressive drugs.

Autoimmune pancreatitis (AIP), a pancreatic disease caused by an autoimmune mechanism, is associated with elevated levels of serum IgG4 and the infiltration of IgG4-positive cells into the pancreatic parenchyma, and it is occasionally accompanied by nonpancreatic features. AIP is known to be accompanied by multiple organ failure, and
systemic inflammatory diseases characterized by the infiltration of IgG4-positive plasma cells and elevated serum IgG4 levels were recently classified as IgG4-related diseases.

Recent studies have reported AIH cases with infiltrated IgG4-positive plasma cells in the liver, suggesting the involvement of IgG4 in the pathogenesis of AIH. Here, we review the characteristic features of IgG4-related disease and its association with AIH.

**Autoimmune Hepatitis**

**Clinical Features**

The term ‘autoimmune hepatitis’ was introduced in 1965 [1]. Upon the discovery of hepatitis C virus in 1989, chronic hepatitis C infection was separated from non-A, non-B hepatitis. Subsequently, AIH was classified as an independent disease category, and AIH diagnostic crite-
Serological findings are the predominant elevation of liver enzymes over biliary enzymes, elevated serum IgG levels, and the presence of antibodies such as ANA, SMA, and anti-LKM-1. Characteristic findings in the liver include interface hepatitis, infiltration of lymphocytes and plasma cells, and rosette formation. AIH is treated with steroids and immunosuppressive agents.

In Japan, AIH diagnostic criteria were established in 1996 by the Intractable Liver Disease Research Project Team of the Ministry of Health, Labor and Welfare (table 1) [4], which recommended the use of the IAIHG diagnostic criteria. The IAIHG diagnostic criteria were first established in 1993, revised in 1999, and are currently used as the international criteria (table 2) [3]. Characteristic AIH findings, including sex, clinical and serological features, liver histology and treatment response, are as-

### Table 1. Japanese diagnostic criteria for AIH in 1996

<table>
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<th>Concept</th>
<th>AIH is a disease that develops frequently in women past middle age and progresses to chronic hepatitis, and the pathogenesis of hepatocellular damage suggests the involvement of autoimmune mechanisms. For effective diagnosis, viral hepatitis, alcohol- and drug-induced hepatitis, and liver damage due to other autoimmune diseases should be eliminated. For treatment, immunosuppressants, particularly corticosteroids, are highly effective.</th>
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| Major findings | 1. Positive autoantibodies in the serum (particularly ANA and SMA)  
2. Elevated serum γ-globulin or immunoglobulin G levels (≥2 g/dl)  
3. Continuous or repetitive increases in serum transaminase levels  
4. Negative for viral hepatitis, in principle  
5. Histological chronic hepatitis or cirrhosis accompanied by hepatocellular necrosis and piecemeal necrosis, often with marked plasma cell infiltration and sometimes with acute hepatitis. |
| Diagnosis | If AIH is suspected based on the major findings 1–4 above, histological analysis is needed to make a diagnosis in accordance with the diagnostic criteria for AIH recommended by the IAIHG (see table 2) |
| Treatment guidelines | 1. In principle, immunosuppressive therapy (e.g., PSL) is recommended for cases with a definitive diagnosis of AIH.  
2. A sufficient dose of PSL (≥30 mg daily) is administered as the initial treatment, and efficacy is evaluated based on the improvement of serum transaminase levels. Maintenance doses are determined after the normalization of serum transaminase activity.  
3. For treating AIH with hepatitis C viremia:  
   a. Corticosteroid therapy is recommended for cases that score high on the international diagnostic scoring system.  
   b. Interferon therapy may be used in cases with a low score calculated in accordance with the international criteria. However, the indication for interferon therapy should be carefully determined based on a virological search before administration. Upon the initiation of interferon therapy, it is necessary to examine viral titers and liver function. If no improvement is observed, administration should be promptly terminated and the administration of immunosuppressive drugs should be considered. |

Modified from [4]. 1. Human leukocyte antigen-DR4-positive cases are more common in Japan. 2. Interferon therapy may be effective in some AIH cases with obvious hepatitis C infection. 3. Some AIH cases are accompanied by hepatitis C viremia in Japan.
sessed using the scoring system, and definitive and probable diagnoses are compared before and after treatment. The criteria are complex and purely meant for scientific purposes. In 2008, the IAIHG introduced the simplified version of the AIH diagnostic criteria (table 3) [5] for routine clinical practice. This has made diagnosis easier and early therapeutic intervention possible. However, the diagnosis of AIH can be difficult because of its various clinical presentations, the presence of atypical and acute cases, and the involvement of other autoimmune diseases.

### Treatment

Steroids and immunosuppressive drugs such as azathioprine are effective treatments for AIH, and ursodeoxycholic acid has shown good efficacy in many studies.

### IgG4-Related Diseases

AIP is known to be accompanied by multiple organ failure, including sclerosing cholangitis, liver failure, inflammation of the lacrimal and salivary glands, thyroiditis, interstitial pneumonia and interstitial nephritis. Systemic inflammatory diseases characterized by the infiltration of IgG4-positive plasma cells and elevated serum IgG4 levels were recently classified as IgG4-related disease. In 2011, the Ministry of Health, Labor and Welfare established the 2011 comprehensive diagnostic criteria for IgG4-related disease (table 4) [6].
Liver injury is observed in 60–70% of AIP cases; however, with the exception of obstructive jaundice, the cause of liver injury in AIH is not always clear. Umemura et al. [7] showed that AIP is accompanied by hepatocellular damage, such as the infiltration of IgG4-positive plasma cells near the portal vein, patterns of portal inflammation, large biliary duct damage, portal sclerosis, lobular hepatitis and cholestasis, and improvement in histological findings after steroid therapy. They called the disease IgG4 hepatopathy.

**AIH and IgG4-Associated AIH**

Recently, some AIH cases fulfilling the criteria of IgG4-related disease have been designated as IgG4-associated AIH [8, 9]. Chung et al. [9] reported that a group of AIH patients with infiltration of IgG4-positive plasma cells were successfully treated with prednisolone (PSL) therapy. Based on the IgG4 immunoreactivity of liver biopsy samples (≥5 IgG4-positive plasma cells/high-power field, HPF), they divided 26 patients with a definitive diagnosis of AIH into IgG4-positive (9 patients, 35%) and IgG4-negative (17 patients, 65%) groups (table 1). No pancreaticobiliary lesions were observed in the AIH patients. No IgG4-positive plasma cells were observed in 10 cases of primary biliary cirrhosis or 20 cases of chronic hepatitis C. The IgG4-positive group had a significantly higher level of serum IgG than the IgG4-negative group (p < 0.01), but no significant differences in IgG4 levels were observed between these two groups. In addition, there were no significant differences in alanine aminotransferase (ALT), alkaline phosphatase, γ-glutamyl transpeptidase or ANA. On the other hand, the severity of plasma cell infiltration and lobular hepatitis were significantly high in the IgG4-positive group. Although portal inflammation and interface hepatitis were similar in both groups, the severity of portal inflammation was significantly higher in the IgG4-positive group. Although portal inflammation and interface hepatitis were similar in both groups, the severity of portal inflammation was significantly higher in the IgG4-positive group. Although portal inflammation and interface hepatitis were similar in both groups, the severity of portal inflammation was significantly higher in the IgG4-positive group. The infiltration of B cells, T cells and plasma cells was also significantly higher in the IgG4-positive group than in the IgG4-negative group (p < 0.05). Furthermore, ALT levels at 4, 48, 72 and 96 weeks after the initiation of PSL therapy were significantly lower in the IgG4-positive group than in the IgG4-negative group (table 2). During the administration of PSL, hepatitis relapse was observed in 6 IgG4-negative patients (35%) but not in any IgG4-positive patients. Even with the definition of IgG4 positive as the

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**Table 4. Comprehensive diagnostic criteria for IgG4-related disease, 2011**

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<th>Concept</th>
<th>IgG-related disease is a disease with an unknown cause and is characterized by synchronous or asynchronous swelling, nodule formation and hypertrophic lesions in multiple organs due to tissue fibrosis and the marked infiltration of lymphocytes and IgG4-positive plasma cells. The pancreas, biliary duct, lacrimal and salivary glands, central nervous system, thyroid gland, lungs, liver, gastrointestinal tract, kidneys, prostate gland, retroperitoneum, arteries, lymph nodes, skin and mammary gland are known to be affected. In general, the disease produces systemic manifestations, with some cases of single organ involvement. Clinically, individual organs present different symptoms, with occasional severe complications such as enlarged organs, obstruction caused by hypertrophic lesions, compression, infiltration and organ dysfunction due to fibrosis. Corticosteroids are generally effective.</th>
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| Clinical diagnostic criteria | 1 Clinical findings of characteristic diffuse or focal swelling, nodule formation and hypertrophic lesions in single or multiple organs.  
2 Hematological findings of elevated IgG4 levels (≥135 mg/dl).  
3 Histopathological findings of the following:  
   a) Histologically marked infiltration of lymphocytes and plasma cells and tissue fibrosis.  
   b) Infiltration of IgG4-positive plasma cells: ≥40% IgG4/IgG-positive cells and >10 IgG4-positive plasma cells/HPF.  
1 + 2 + 3 Definite IgG4-related disease.  
1 + 3 Probable IgG4-related disease.  
1 + 2 Possible IgG4-related disease.  
However, by adding a histopathological diagnosis as possible, differential diagnosis of IgG4-related disease from similar diseases (Sjögren’s syndrome, primary sclerosing cholangitis, Castleman’s disease, secondary retroperitoneal fibrosis, Wegener similar disease) or malignant tumors (cancer, and malignant lymphoma) is important. Even if no definite diagnosis is made using the present criteria, diagnosis may be possible if the diagnostic criteria for each organ are used. |
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infiltration of ≥10 IgG4-positive plasma cells/HPF, the response to PSL therapy and the serum levels of IgG were significantly different between the groups.

Umemura et al. [8] also reported the pathology of IgG4-related AIH. In a study of 60 AIH patients, they defined IgG4-related AIH as the infiltration of IgG4-positive plasma cells (≥10 cells/HPF), IgG4-positive serum (≥135 mg/dl) and the ratio of IgG4 to IgG as 0.073. However, because only 2 cases fulfilled this definition, they concluded that the prevalence of IgG4-positive AIH is extremely low (2/60 cases, 3.3%).

The main difference between the studies by Chung et al. [9] and Umemura et al. [8] is that, in the latter study, the definition included the ratio of IgG4 to IgG in addition to high serum IgG4 concentration (≥135 mg/dl) and the infiltration of IgG4-positive plasma cells in liver tissue. Furthermore, while Chung et al. [9] examined cases that matched the definite diagnosis of AIH, the 60 AIH cases investigated by Umemura et al. [8] included 12 which were probable AIH. Moreover, the infiltration of IgG4-positive cells in the gallbladder and common bile duct was observed in 1 of the IgG4-associated AIH cases, indicating a case of IgG4-related sclerosing cholangitis rather than IgG4-associated AIH. Regardless of the differences, the two studies revealed that IgG4-related pathologies are associated with at least some AIH cases and that steroid treatment is effective in such IgG4-associated AIH, as in other IgG4-related diseases.

**Conclusion**

Some patients with AIH present symptoms of IgG4-related disease and respond effectively to steroid treatment. Although no unified diagnostic criteria are currently available, it is important to keep this disease in mind for definitive diagnosis and appropriate treatment. Further studies are needed to define the epidemiological features or diagnostic criteria.

**Disclosure Statement**

The authors have no conflicts of interest to declare.

**References**