Coexistence of Acute Crescent Glomerulonephritis and IgG4-Related Kidney Disease

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Abstract

Introduction: IgG4-related disease (IgG4-RD) is a fibroinflammatory disorder that may involve almost each organ or system. IgG4-related kidney disease (IgG4-RKD) refers to renal lesions associated with IgG4-RD. The most frequent morphological type of renal lesions is IgG4-related tubulointerstitial nephritis (IgG4-TIN) which is associated with increased IgG4-positive plasma cell infiltration and interstitial fibrosis. Case Report: Herein, we present a rare case with coexisting IgG4-RKD and acute crescent glomerulonephritis with concomitant severe tubulointerstitial lesions instead of classic IgG4-TIN. Conclusion: IgG4-RKD and acute crescent glomerulonephritis can occur in the same patient. This case may give us a clearer viewpoint of the disease.
Introduction

IgG4-related disease (IgG4-RD) is a potentially multiorgan disorder which is characterized by increased IgG4-positive plasma cell infiltration, tissue fibrosis, and elevated serum levels of IgG4 [1, 2]. IgG4-RD can affect nearly every organ or system, including the pancreas, aorta, lung, salivary and lacrimal glands, thyroid, and kidney. Prior studies have reported that renal lesions seem to occur in approximately 15% of patients. Now, IgG4-related kidney disease (IgG4-RKD) is used as a comprehensive term for renal lesions associated with IgG4-RD.

IgG4-RKD is a recently recognized autoimmune renal disease which often, but not always, presents with high levels of serum IgG4, proteinuria, and acute or chronic renal dysfunction. Many characteristic abnormalities can be observed by radiological examinations which may resemble malignant tumors. The most frequent histological finding is IgG4-related tubulointerstitial nephritis (IgG4-TIN) which is associated with increased IgG4-positive plasma cell infiltration and interstitial fibrosis [3, 4]. Although TIN is the predominant lesion type in IgG4-RKD, herein, we present a rare case with coexisting IgG4-RKD and acute crescent glomerulonephritis with concomitant severe tubulointerstitial lesions instead of classic IgG4-TIN.

Case Report

A 61-year-old Chinese woman was hospitalized due to elevated serum creatinine (Scr). Four months previously, without any predisposing factors, she had suffered from sudden dryness, burning, and paroxysmal pinprick-like pains in her right eye, accompanied by a headache in the right hemisphere. Her discomfort was obvious in the morning, while it was slightly relieved in the afternoon without any treatment. Computed tomography (CT) did not reveal any abnormality in the brain. Iritis and scleritis were considered by an ophthalmologist, but the symptoms were not alleviated after dexamethasone eyedrops. One day previously, laboratory tests had shown elevated Scr of 324 μmol/l.

At physical examination, her body temperature was 36.6°C, her pulse rate 100 beats per minute, and her blood pressure 125/68 mm Hg. She had no skin lesions or swollen lymph nodes. There were no rales on bilateral lungs and no murmurs in the heart. Her abdomen was soft, and there were no signs of swollen liver, spleen, or edema.

Lab test results were as follows: white blood cells 7.3 × 10⁹/l, red blood cells 2.89 × 10¹²/l, hemoglobin 76 g/l, and platelets 183 × 10⁹/l; urinary red blood cells 140/μl, urinary occult blood (++), urinary white blood cells 6/μl, and urinary protein 1.93 g/24 h. Blood biochemistry analysis provided the following results: total protein 66 g/l, albumin 31 g/l, alanine aminotransferase 32 U/l, total bilirubin 9 μmol/l, serum iron 7 μmol/l, blood nitrogen urea 13.1 mmol/l, Scr 324 μmol/l, uric acid 385 μmol/l, sodium 129 mmol/l, potassium 3.8 mmol/l, chloride 93 mmol/l, and CO₂ 18.3 mmol/l. Clinical immunology tests revealed the following: anti-nuclear antibody (−), anti-neutrophil cytoplasmic antibodies (−), IgG 16.8 g/l, IgG 3.21 g/l, C3 0.84 g/l, C4 0.24 g/l, C-reactive protein 46 mg/l, and erythrocyte sedimentation rate 58 mm/h. Serum immune electrophoresis was normal. Hepatitis virus screening was as follows: hepatitis B surface antigen (−), anti-hepatitis B surface antibody (+), and anti-hepatitis C antibody (−). Blood levels of lipid series, glucose, thyroid function, and tumor markers were all normal.
Renal ultrasonography showed that the size of the right kidney was 98 × 34 mm, while the left one was 95 × 40 mm. The renal cortical echo was slightly enhanced, and no renal calculus was found. An enhanced CT scan indicated multiple low-density lesions under both of the renal capsules (fig. 1). Emission CT revealed chronic sialadenitis in the bilateral parotid glands.

Then, the patient underwent a renal biopsy. The immunofluorescence report found negative IgG, IgA, IgM, and C3. The pathological findings were as follows: there were 19 glomeruli in total, global sclerosis in 7 glomeruli, crescents forming in 10 glomeruli, and middle to severe mesangial cellular hyperplasia along with increased mesangial matrix in 2 glomeruli as well. There were diffuse tubular atrophy and extensively dense mononuclear cells among the tubulointerstitial tissues. Protein casts were also observed in some tubules. No obvious vascular lesions were found. Immunohistochemistry showed IgG4-positive plasma cells >10/HPF, and the ratio of IgG4+ cells to IgG+ cells was >40% (fig. 2). A pathological diagnosis of acute crescent glomerulonephritis with severe renal TIN damage was achieved.

Based on the physical examination, lab tests, and renal biopsy, the patient was diagnosed as IgG4-RKD combined with acute crescent glomerulonephritis after excluding vasculitis, Sjögren's syndrome, and plasmacytoma. The patient was treated with methylprednisolone (1 mg/kg every day) and cyclophosphamide (1.0 g per month). In addition, tablets of calcium, iron, and proton pump inhibitors were used at the same time. Four weeks later, the Scr dropped from 324 to 213 μmol/l, and symptoms were relieved. Furthermore, the Scr dropped to 130 μmol/l at the 16-week follow-up (fig. 3). At the same time, the serum IgG4 concentrations dropped to 1.21 g/l.

Discussion

IgG4-RKD is a recently defined term referring to renal lesions associated with IgG4-RD [5, 6]. In this case of an aged female patient, presenting renal insufficiency, hematuria, albuminuria, low serum C3, and elevated serum IgG4, renal biopsy showed acute crescent glomerulonephritis combined with severe renal tubulointerstitial lesions. Moreover, the immunohistochemistry revealed IgG4-positive plasma cells >10/HPF and the ratio of IgG4+ cells to IgG+ cells was >40%. Irregular fibrosis located around the infiltrated lymphocytes could also be observed. In addition, the eye and the parotid gland were involved. Diagnostic criteria for IgG4-RKD have been proposed by Kawano et al. [7]. By using these criteria, the results can be separated into definite, probable, and possible. According to the diagnostic criteria for IgG4-RKD, the patient was diagnosed with definite IgG4-RKD [1, 7]. In IgG4-RKD, steroids are commonly effective in ameliorating kidney lesions and abnormalities of radiology or serology [8–10]. Although a high-dosed steroid treatment works rapidly in IgG4-RKD, a long-term maintenance of low-dosed therapy remains essential for the recovery of renal function [11]. In this case, after the treatment with steroid and cyclophosphamide pulse therapy, the patient’s symptoms were relieved and the Scr became almost normal 16 weeks later.

This is the first report of IgG-RKD presenting with a pathological pattern of acute crescentic glomerulonephritis with severe renal tubulointerstitial injury. In earlier reports, IgG4-RKD usually resulted in IgG4-TIN without specific signs of immunofluorescent staining. Furthermore, IgG4-RKD presented with increased plasma cell infiltration in the swelling interstitial tissues, and slight fibrosis could be observed in the early stage. At the late stage, multi-
focal tubular atrophy in laminate structures could appear with a high number of IgG4-positive plasma cells, and storiform fibrosis was also apparent [12–14]. Obliterative phlebitis is also a critical pathological feature of IgG4-RD, although it is rarely seen in IgG4-TIN [3]. However, apart from TIN, other pathology types induced by IgG4-RKD have also been discovered, such as membranous nephropathy, pyelonephritis, or glomerulonephritis as recently reported [13, 15–17]. Acute crescent glomerulonephritis combined with IgG4-RKD has never been reported yet.

IgG4-RKD often occurs in old male patients with hyperglobulinemia and high serum levels of IgG [4, 18]. However, this case was an old female patient with anemia, maybe due to a long-term kidney injury. Another novelty of this case is that the histological type of the renal lesions was acute crescent glomerulonephritis with concomitant TIN. Also, TIN-uveitis syndrome and an atypical form of Cogan’s syndrome should be differentiated from IgG4-RD in this case, since the patient presented with combined iritis and kidney damage [19, 20].

As a recently clarified type of renal injury, IgG4-RKD is different from classic acute kidney injury or chronic kidney disease [21–23]. Usually, the renal pathological pattern of IgG4-RKD is TIN. This case of acute crescent glomerulonephritis combined with IgG4-RKD may give us a clearer viewpoint of this disease. In the long run, the mechanisms of the crescent glomerulonephritis in the present case and the proportion of IgG4-RKD patients affected need further investigation and discussion.

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Statement of Ethics

This case report was approved by the ethics committee of Shanghai Jiao Tong University Affiliated People’s Hospital, and the patient provided informed consent.

Disclosure Statement

The authors have no conflicts of interest to declare.

References

Z.L., J.Y. and H.B. contributed equally to this work.
Fig. 1. Characteristic CT of the kidney. Multiple low-density lesions can be seen on enhanced CT.

Fig. 2. Pathological findings of the renal biopsy. The major finding of the renal biopsy was acute crescent glomerulonephritis combined with severe TIN. a, b Extensive glomerular crescents have formed (a HE staining, ×400; b periodic acid-Schiff staining, ×400). c Dense mononuclear cells have infiltrated the interstitium (HE staining, ×400). d Tubular degeneration, atrophy, necrosis, inflammatory cell infiltration, and diffuse interstitial fibrosis were observed (trichrome staining, ×400). e There were plenty of IgG4-positive plasma cells in the infiltrate (IgG4 staining, ×400). f Infiltrated plasma cells or lymphocytes in the infiltrate (electron microscopy, ×5,000).
Fig. 3. Scr changes during follow-up. Scr levels decreased with the treatment of methylprednisolone and cyclophosphamide. W = Weeks.