Rituximab therapy for primary glomerulonephritis: Report on two cases

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Abstract
The evidence in the medical literature on the efficacy and safety of rituximab therapy for primary glomerulonephritis is limited and controversial. We describe two male Caucasian patients with rapidly progressive kidney failure due to primary proliferative glomerulonephritis. Both of them received high-dose intravenous corticosteroids and oral cyclophosphamide with limited benefit. The first patient (hepatitis C virus-negative mixed cryoglobulinemia) underwent plasma-exchange with intravenous immunoglobulins; he showed significant benefit on kidney function (he became dialysis independent with serum creatinine going back to 1.6 mg/dL) after one rituximab pulse even if urinary abnormalities were still present. No improvement in renal function or urinary changes occurred in the second patient. Both these individuals developed sepsis over the follow-up, the first patient died two months after rituximab therapy. This report is in keeping with the occurrence of severe infections after rituximab therapy in patients with renal impairment at baseline and concomitant high-dose steroids.

Key words: Chronic kidney disease; Cryoglobulinemic vasculitis; Membranoproliferative glomerulonephritis; Rituximab

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Core tip: A small but growing body of evidence is emerging on the efficacy and safety of rituximab therapy for primary glomerulonephritis. Various authors have claimed that rituximab for glomerular diseases is effective and has minimal adverse effects. We report on two male Caucasian patients who were refractory to conventional immunosuppressive therapy; each of them received one rituximab pulse and developed sepsis over the follow-up, the first patient died two months after rituximab therapy. The risks (and the predictive...
Primary glomerulonephritis (GN) remains an important cause of end-stage kidney disease. Preliminary trials have recently shown the efficacy of rituximab for adult-onset primary GN\(^{[1]}\); rituximab being a genetically chimeric monoclonal antibody directed to CD20 antigen, a B-cell-specific transmembrane found on immature and mature cells, as well as on malignant B cells. Following treatment with rituximab (RTX), B-cells are prevented from proliferating, and undergo apoptosis and lysis through complement-dependent and -independent mechanisms. B-cell depletion usually persists for 6-9 mo in around 80% of patients, but the degree of depletion is greatly variable. Rituximab is currently approved for treating various malignancies including B cell non-Hodgkin's lymphoma and chronic lymphocytic leukemia; also, it has been licensed for refractory rheumatoid arthritis, granulomatosis with polyangiitis (Wegener's granulomatosis) and microscopic polyangiitis. Rituximab was expected to inhibit the production of autoantibodies involved in the pathogenesis of the disease without the toxicity of nonspecific immunosuppression. It has been the first monoclonal employed for the treatment of glomerular diseases and has been initially used for patients with membranous nephropathy but its use has rapidly spread to other glomerular diseases\(^{[2]}\). Membranous nephropathy and membranoproliferative GN are characterized by glomerular deposition of immune complexes; a crucial role of B cells in membranous (MN) and membranoproliferative glomerulonephritis pathogenesis through autoantibody production and antigen presentation has been mentioned. A small but growing body of literature is emerging on the benefits of rituximab in MN and membranoproliferative glomerulonephritis as primary treatment or as treatment of lesions refractory to other immunomodulatory regimens. In this setting, the drug appears to be well tolerated with small adverse events (Table 1)\(^{[3-8]}\).

We report here our experience on rituximab use in two patients with progressive kidney failure due to primary proliferative GN. Both of them received conventional immunosuppressive therapy with limited benefit on urinary and biochemical abnormalities; then, they underwent one RTX pulse but developed sepsis over the follow-up. A brief review on the safety and efficacy of rituximab for primary GN has been also added.

### CASE REPORT

**Patient 1**

A 51-year-old Caucasian male patient was admitted to hospital for two-week's duration of abdominal pain with vomiting and diarrhoea. His medical history included arterial hypertension and symptomatic hepatitis C virus (HCV)-negative mixed cryoglobulinemia (since three years) with recurrent purpura and peripheral neuropathy at the lower extremities. Skin biopsy had shown leukocytoclastic vasculitis whereas neurological evaluation had revealed mono-neuritis at the left foot with axonal ischemic damage, probably related to cryoprecipitable immune complexes in the vasa nervorum. He had received low dose oral corticosteroids and azathioprine with partial control of cutaneous and neurological abnormalities. A bone marrow biopsy had reported no evidence of malignant lymphoma, and a small expansion of B lymphocytes (10%-15%).

A physical examination showed bilateral edema and hypertension (180/100 mmHg), purpuric rash with ulcers at the legs (Figure 1); no bowel movements were apparent from the clinical standpoint, this being confirmed by an abdomen X-ray. An ultrasound scan of the abdomen showed normal sized kidneys bilaterally, with normal echotexture. At presentation (Table 1), abnormal laboratory results included serum creatinine level of 2.5 mg/dL, proteinuria, 3.6 g/24 h, and hypoalbuminemia (2 g/L). Other pertinent chemistries were: positive cryoglobulins, with a cryocrit of 3% (polyclonal IgG and monoclonal IgM), elevated rheumatoid factor (148 IU/mL) and hypocomplementemia. Serology was negative for hepatitis B virus (HBV), HCV and human immuno-deficiency virus (HIV) markers, polymerase chain reaction tested negative for HCV RNA. Repeat urine sediment, analyzed by phase-contrast microscopy, showed severe microscopic hematuria (> 50 erythrocytes/microscopic field), many dysmorphic erythrocytes and casts (jailine, granular and red cell casts). Bence Jones proteinuria (kappa type) was positive. The search for anti-neutrophil cytoplasmic antibody (proteinase 3 and myeloperoxidase), anti-glomerular basement membrane antibody, extractable nuclear antigen antibody, antinuclear and anti-double stranded DNA tested negative.

Renal biopsy was not performed due to anatomic reasons, and a diagnosis of essential MC with rapidly progressive renal failure due to nephritic/nephrotic syndrome was made. Treatment was initiated with intravenous methylprednisolone (600 mg/d for three days), oral prednisone (50 mg/d on taper), and oral cyclophosphamide (100 mg daily). The progressive deterioration of kidney function (serum creatinine raised to 6.3 mg/dL, blood urea nitrogen to 279 mg/L) led us to make sequential plasma-exchange (nine sessions) and high-dose intravenous immunoglobulins (five procedures); in addition, hemodialysis was started. We...
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Table 1  Blood chemistries at presentation and over follow-up (patient 1)

|                       | Admission | Discharge | Middle follow-up | Final follow-up |
|-----------------------|-----------|-----------|------------------|-----------------|
| Creatinine (0.5-1.2, mg/dL) | 2.8       | 1.59      | 1.76             | 0.8             |
| Blood urea nitrogen (8-20, mg/dL) | 147      | 104       | 86              | 85              |
| AST (5-32, IU/L)       | 12        | 24        | 27              | 23              |
| ALT (5-31, IU/L)       | 9         | 43        | 39              | 7               |
| γGT (5-36, IU/L)       | 12        | 44        | 41              | 69              |
| Cholinesterase (5300-12900, IU/L) | 3833   | 2160      | 2920            | 3173            |
| Total bilirubin (0.2-1.1, mg/dL) | 0.2      | 0.25      | 0.22            | 0.23            |
| Direct bilirubin (0.0-0.3, mg/dL) | 0.09  | 0.07      | 0.08            | 0.16            |
| Total protein (6.6-8.7, g/dL) | 3.9      | 4.6       | 4.6             | 4.5             |
| Albumin (3.4-4.8, g/dL) | 2.4       | 3         | 2.5             | 3               |
| Prothrombin time (0.88-1.16) | 1.08    | 1.07      | 1.06            | 1.08            |
| Partial thromboplastin time (0.85-1.18) | 1.01  | 1         | 1.03            | 1.19            |
| C1 (90-180)           | 20        | 56        | 59              | 99              |
| C1 (10-40)            | 0         | 1         | 3               | 2               |
| Cryoglobulins         | Present   | Absent    | Absent          | Present         |
| Leucocytes (4.8-10.8, 10^3/mmc) | 10670 | 3440      | 3400            | 3730            |
| Hemoglobin (12-16, g/dL) | 10.8      | 9.7       | 10.2            | 10.5            |
| Platelets (130-400, 10^3/mmc) | 313000 | 159000    | 153000          | 73000           |

AST: Aspartate aminotransferase; ALT: Alanine aminotransferase; γGT: Gamma-glutamyl transpeptidase.

Figure 1  Purpuric rash with ulcers at the leg (patient 1).

Figure 2  Biochemical/serological response to therapy (patient 1). HD: Haemodialysis; MP: High-dose methylprednisolone; PEX: Plasma-exchange; RTX: Rituximab.

observed healing of skin ulcers and improvement of neuropathic pain; GI disorders disappeared but immunosuppressive therapy was complicated by Clostridium Difficile-positive diarrhea, which was successfully treated with oral vancocine. Due to the persistence of severe renal failure (serum creatinine of 4.2 mg/dL), he received one infusion of RTX (375 mg/m²) off-label (Figure 2); by day 6 of the RTX dose, an improvement of urine output occurred. Since then, serum creatinine went back to 1.6 mg/dL, and blood urea nitrogen to 104 mg/dL. At discharge from the hospital, his medications included oral steroids, rabeprazole, amlodipine, furosemide, calcium carbonate, gabapentine, and darbopeoetin.

Two weeks after hospital discharge, serum creatinine was 1.76 mg/dL, blood urea nitrogen, 86 mg/dL, serum albumin 2.5 g/dL; severe and dysmorphic hematuria and non-nephrotic proteinuria (1.43 g/L) being still present. Low white blood cell count after RTX administration (3.4 × 10^3/mm³) occurred. According to flow cytometry, CD20⁺ B cells were 19% of total peripheral blood lymphocytes (before RTX) and fell to 1% (after RTX, since day 6) with no increase over the following weeks.

One month later, he became pyrexial with a temperature around 38℃ and was admitted again to the hospital. Pertinent biochemistry included serum creatinine (0.8 mg/dL), cryoglobulins and rheumatoid factor tested positive. Complete blood count gave the following features: white blood cells 2.67 × 10^3/mm³, erythrocytes 3.57 × 10^6/mm³, platelets 209 × 10^3/mm³. Flow cytometry: lymphocytes 169/mm³, CD3⁺ cells 142 (83.9%), CD19⁺ 0 (0%), natural killer cells 25 (14.9%). Gamma globulins were 0.16 g/L (3.9%, 11%-18.8%). IgA < 4 mg/dL (70-400), IgG 84 mg/dL (700-1600), IgM 108 mg/dL (40-230). Monoclonal component by serum electrophoresis (IgMk + oligoclonal Ig) was again detected. An active urinary sediment with non-nephrotic proteinuria (1.21 g/d) was still present. Sepsis from Enterococcus Spp. was identified whereas the chest radiograph reported multiple pneumonias. The culture of the bronchoalveolar lavage fluid was positive for Candida albicans, thus, we initiated intravenous imipenem and antifungal
medications. Medical and supportive therapy was unsuccessful and the patient ultimately expired due to septic shock (two months after RTX pulse).

**Patient 2**

A 49-year-old Caucasian male underwent kidney biopsy for evaluation of serum creatinine 1.35 mg/dL (estimated glomerular filtration rate (eGFR) 56 mL/min per 1.73 m² by MDRD equation), 3.2 g of proteinuria on 24-h urine collection and active urinary sediment (severe microscopic hematuria with red blood cell casts). Renal biopsy showed global ilinosis in some glomeruli (5 out of 14); the others had intense glomerular hypercellularity (mainly due to mesangial proliferation), a limited number of mesangial immune deposits and segmental thickening of glomerular basement membrane were also present. Final diagnosis was mesangial proliferative GN with immune deposits of unclear significance. Other pertinent chemistries were: negative cryoglobulins, normal rheumatoid factor and complement fractions; serum protein electrophoresis in the normal range. Serology tested negative for HBV, HCV and HIV markers, polymerase chain reaction was negative for HCV RNA. The search for anti-neutrophil cytoplasmic antibody (proteinase 3 and myeloperoxidase), anti-glomerular basement membrane antibody, extractable nuclear antigen antibody, antinuclear and anti-double stranded DNA tested negative. At discharge from the hospital, his medications included oral steroids, rabeprazole, allopurinol, doxazosine, furosemide, calcium carbonate, and darboepoetin. Partial remission of nephritic/nephrotic syndrome with improvement of kidney function (serum creatinine going back to 1.1 mg/dL) was obtained with intravenous methylprednisolone pulses, oral cyclophosphamide, azathioprine, and mycophenolate mofetil in variable associations. Eight years later, he was again admitted to our unit, showing bilateral lower-extremity edema, arterial hypertension and serum protein electrophoresis in the normal range. Pertinent chemistries were: negative cryoglobulins, normal rheumatoid factor and complement fractions; serum protein electrophoresis in the normal range. Glomerular filtration rate (eGFR) 56 mL/min per 1.73 m² by MDRD equation). Nephrotic proteinuria was demonstrated (proteinuria of 9.2 g/d) with active urinary sediment. A repeat kidney biopsy revealed intracapillary/extracapillary glomerular proliferation with several crescents and fibrinoid necrosis, diffuse arteriolosclerosis, in addition to uniform and diffuse thickening of the glomerular basement membrane. Immunofluorescence demonstrated sporadic and granular deposition of C1q/C3 in the mesangium and capillary walls; fibrinogen in the Bowman space. Renal biopsy was complicated by perirenal hematoma and a few units of red packed cells were given. He received oral antibiotics and adequate hydration. One month after rituximab administration he was again hospitalized (acute pulmonary insufficiency with septic shock); serum creatinine of 4.56 mg/dL (eGFR, 11 mL/min per 1.73 m²). Active urinary sediment and nephrotic proteinuria persisted. Pulmonary aspergillosis was documented- medical plus supportive therapy was initiated and the patient recovered in a few weeks; however, he developed irreversible kidney failure and initiated dialysis acutely. He is currently doing well on maintenance hemodialysis (thrice weekly) treatment.

**DISCUSSION**

We report here on two patients with rapidly progressive renal failure due to idiopathic proliferative GN who were resistant to conventional immunosuppressive therapy. Both the patients underwent rituximab treatment in off-label condition, RTX infusion was well tolerated by both the patients but sepsis developed over the follow-up, fatal course occurring in patient 1. Numerous case reports and case series have suggested that the addition of rituximab to standard chemotherapy for malignant lymphoma increases the risk of viral infections such as varicella zoster [10], cytomegalovirus [11], HBV [12], parvovirus [12], and enteroviral encephalitis [13]. The risk of HBV reactivation has been added to the existing Boxed Warning of the rituximab label by the Food and Drug Administration in 2013 [14]. Impaired immunity against non-viral pathogen agents such as Pneumocystis jirovecii [15] or cryptococcus after rituximab therapy has been also noted.

A recent systematic review and meta-analysis has shown that rituximab plus standard chemotherapy for malignant lymphoma increases the incidence of severe leucopenia (RR = 1.24; 95%CI: 1.12-1.37) and granulocytopenia (RR = 1.07; 95%CI: 1.02-1.12) even if the overall risk of severe infections has not been increased (RR = 1.0; 95%CI: 0.87-1.14) [17]. We have already reported on a case of cholestatic hepatitis C after rituximab therapy for gastric cancer in a renal transplant recipient [18]. On the other hand, various authors have claimed that rituximab use for glomerular diseases is effective and has minimal adverse effects (Table 2) [11,19,20].

Our first patient presented idiopathic cryoglobulinemic vasculitis which has undefined therapeutic management [21]. There is some evidence on the efficacy and tolerance of RTX in patients with HCV-associated mixed cryoglobulinemia vasculitis who were naïve, resistant or intolerant to antiviral therapy [22-25]. Two randomized controlled trials have compared RTX with conventional immunosuppressive therapy for HCV-related mixed cryoglobulinemia vasculitis [26-27]. As listed in Table 3, evidence in the medical literature on RTX use among patients with non-infectious cryoglobulinemia vasculitis targeting kidneys is extremely limited, and a total of 16 cases were retrieved [28-37]. Patient 1 gives emphasis on the efficacy of RTX, as one RTX pulse made possible the control of renal disease: kidney function normalized, nephrotic syndrome disappeared and only nephritic urinary changes persisted. However, RTX use was complicated by sepsis a few weeks after...
RTX pulse. On the basis of the evidence reported in Table 3, severe infections after RTX treatment are not uncommon [35% (6/17)].

RTX therapy in patients with nonviral cryoglobulinemia vasculitis or membranoproliferative GN raises various questions such as the role of RTX as first-line or rescue therapy, the efficacy/safety of maintenance therapy with RTX, and the tolerance to RTX. In the absence of randomized controlled trials, such questions remain unanswered; as an example, the poor tolerance of our patients after RTX administration remains unclear. The French multicenter CRYOvas survey retrospectively evaluated 242 patients with non-infectious mixed cryoglobulinemia vasculitis, RTX plus corticosteroids had greater therapeutic efficacy compared with corticosteroids alone and corticosteroids plus alkylating agents [38]. However, RTX plus corticosteroids was associated with more frequent infections than corticosteroids alone (HR = 9; 95%CI: 3.1-20, P < 0.001). Prospective data from the AIR (AutoImmunity and Rituximab) registry, which includes data on patients treated with rituximab off-label, have shown that among patients (n = 23) with nonviral cryoglobulinemia vasculitis on RTX, side-effects occurred in almost half of the patients (n = 11), including severe infections [34]. Both our patients had important kidney impairment at baseline and concomitant therapy with intravenous high-dose corticosteroids, among other immunosuppressive agents.

The current study calls for further research on the RTX-based treatment of essential cryoglobulinemic vasculitis or membranoproliferative GN but the low frequency of patients in individual centers would make

| Ref.      | n | Age (yr)/gender | Treatment prior to RTX | Features | Response to RTX | Side-effects after RTX |
|-----------|---|-----------------|------------------------|----------|-----------------|------------------------|
| Arzoo et al [35] | 1 | 71/F | CS | C, N, R | Remission | None |
| Ghijels et al [34] | 1 | 48/M | CS, CPH, CHL | C, Ca, R | Remission | None |
| Koukoulaki et al [36] | 1 | 48/F | CS, CPH | GI, P, R | Partial remission | None |
| Bryce et al [37] | 1 | NA | NA | R | No response | None |
| Rich et al [38] | 1 | 64/M | CS | R | Remission | Cold agglutinin disease, sepsis |
| Annear et al [39] | 1 | 42/F | CS | C, R | Remission | None |
| Terrier et al [40] | 7 | 73 ± 5/M (n = 4) | CS (n = 4) | C (n = 6), N (n = 2), A (n = 2), R (n = 3) | Remission (n = 3), partial remission (n = 1), NA (n = 3) | Severe infections (n = 4) |
| Wink et al [41] | 1 | 72/F | CS, Aza | C, P, R | Remission | None |
| Choudhry et al [42] | 1 | 61/F | CS, CPH | C, P, R | Remission | None |
| Kemel et al [43] | 1 | 77/F | CS | C, A, R | Remission | None |
| Own case | 1 | 51/M | CS, Aza | C, GI, N, R | Remission | Severe infection |

A: Arthralgias; Aza: Azathioprine; C: Cutaneous; Ca: Cardiac; CHL: Chlorambucil; CPH: Cyclophosphamide; CS: Corticosteroids; GI: Gastrointestinal; N: Neurological; NA: Not available; P: Pulmonary; R: Renal; RTX: Rituximab.

Table 2: Literature review: Adverse events during rituximab therapy for primary membranous and membranoproliferative glomerulonephritis

| Ref.      | n | Rituximab treatment dose | Follow-up period | Concomitant therapy | Response to RTX | Side-effects after RTX |
|-----------|---|--------------------------|-----------------|---------------------|-----------------|------------------------|
| Fervenza et al [44] | 15 | 1 g × 2, on days 1 and 15 | 12 mo | ACE-I + ARB | Complete (n = 2) or partial remission (n = 6) | Nonserious transient AE (n = 10) pneumonia (n = 1) |
| Segarra et al [45] | 13 | 375 mg/m² once weekly × 4 | 30 mo | Tac (n = 10), CyA (n = 3), CCS (n = 3) | Partial remission (n = 13) | Nonserious transient AE (n = 11) pneumonia (n = 1) |
| Fervenza et al [46] | 20 | 375 mg/m² once weekly × 4 | 24 mo | ACE-I + ARB | Complete (n = 4) or partial remission (n = 12) | Nonserious transient AE (n = 11) pneumonia (n = 1) |
| Michel et al [47] | 28 | 375 mg/m² once weekly × 2 or 3 or 4 (n = 27) | 12 mo | ACE-I + ARB, CCS (n = 1), Tac (n = 1) | Complete (n = 6) or partial remission (n = 13) | Nonserious transient AE (few) |
| Ruggenenti et al [48] | 100 | 375 mg/m² once weekly × 4 | 29 mo | CCS | Complete (n = 27) or partial remission (n = 38) | AE (n = 28) |
| Dillon et al [49] | 6 | 1 g × 2, on days 1 and 15 | 12 mo | ACE-I + ARB | Complete (n = 2) or partial remission (n = 3) | Nonserious transient AE (n = 6) Pneumonia (n = 1) |
| Kong et al [50] | 13 | 500 mg × 1 (n = 6) | 31.5 mo | CCS (os) (n = 9) | Remission (n = 19) | Nonserious transient AE (n = 6) |
| Kong et al [51] | 13 | 500 mg × 2 (n = 3) 500 mg × 4 (n = 4) | | CyA (n = 2) CCS (iv) (n = 2) | | |

ACE-I: Angiotensin converting enzyme inhibitors; AE: Adverse events; ARB: Angiotensin receptor blockers; CCS: Corticosteroids [by intravenous (iv) or oral (os) route]; CyA: Cyclosporine; MN: Membranous nephropathy; MPGN: Membranoproliferative glomerulonephritis; Tac: Tacrolimus; RTX: Rituximab.

Table 3: Overview of cases with non-viral hepatitis mixed cryoglobulinemia (and kidney involvement) on rituximab
randomised controlled trials extremely difficult. Rituximab has surfaced as potential treatment option for some primary glomerular diseases and the HCV KDIGO Study Group[39] had already included rituximab among the recommended drugs (steroids, and cyclophosphamide) for the immunosuppressive treatment of HCV-associated kidney disease. The risks (and the predictive factors) of infections in kidney patients on RTX-therapy are not yet understood and are an area of active research. These patients should be monitored over the follow-up to avoid the occurrence of infectious episodes.

**COMMENTS**

**Case characteristics**
Two male Caucasian patients with progressive kidney failure.

**Clinical diagnosis**
Arterial hypertension, bilateral lower-extremity edema.

**Differential diagnosis**
Progressive kidney failure due to secondary glomerular disease.

**Laboratory diagnosis**
At presentation serum creatinine ranged between 2.5 and 2.9 mg/dL, proteinuria 3.6 and 9.2 g/dL, microscopic haematuria with dysmorphic erythrocytes and red cell casts.

**Imaging diagnosis**
Computed tomography scan revealed normal sized kidneys bilaterally with normal echotexture in both the patients.

**Pathological diagnosis**
Renal biopsy (patient 2) showed intracapillary/extracapillary glomerular proliferation with several crescents and fibrinoid necrosis, in addition to uniform diffuse thickening of the glomerular basement membrane.

**Treatment**
Both the patients received one infusion of rituximab (375 mg/m²) off-label.

**Related reports**
Various authors have claimed that rituximab use for glomerular diseases is effective and has minimal adverse effects.

**Term explanation**
Phase-contrast microscopy is a microscopy technique to analyze the morphology of urine erythrocytes.

**Experiences and lessons**
The risks and the predictive factors of severe infections in kidney patients on rituximab therapy are still unclear and appear an area of active research.

**Peer-review**
It is a good article.

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