Rituximab in Membranous Nephropathy: Not All Studies Are Created Equal

Paolo Cravedi
Renal Division, Department of Medicine, Icahn School of Medicine at Mount Sinai, New York, N.Y., USA

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Abstract
Many prospective studies and a recent randomized controlled trial have shown that the B-cell-depleting monoclonal antibody, rituximab, safely promotes the remission of nephrotic syndrome in approximately 65% of patients with membranous nephropathy (MN). Mechanistic studies have indicated that rituximab-induced proteinuria reduction is associated with clearance of anti-podocyte antigens phospholipase 2 receptor autoantibodies and subepithelial immune complexes, the hallmarks of the disease. A recently published study reported results which, at first sight, looked less favorable and implied that, due to a publication bias against negative results, the efficacy of rituximab in MN might be overestimated. Since patients received only one or 2 rituximab administrations, the authors suggest that when rituximab is used, higher doses and longer treatments should be considered. In this study, we highlight limitations of the study and warn against an oversimplified interpretation of the data. Though information on the optimal dose of rituximab to use in MN is still limited, available data from studies with predefined rituximab administration protocols collectively support the concept of titrating rituximab to the number of circulating B-cells that are invariably depleted after the first or second administration. Additional doses may increase the risk of adverse effects and related costs without augmenting efficacy. Importantly, underpowered studies with inconclusive results should not be confused with negative studies formally proving a neutral effect of a treatment. Until data from ad hoc designed clinical trials are available, the B-cell-driven protocol should be the preferred regimen, since it is similarly effective, but safer and more cost effective than other protocols employing multiple rituximab administrations.

The 2001 Kidney Disease: Improving Global Outcomes guidelines suggest the use of alkylating agents plus steroids as first-line therapy for membranous nephropathy (MN), while calcineurin inhibitors, either alone or in combination with steroids, are recommended for patients who do not tolerate or refuse alkylating agents [1]. These indications are largely based on studies performed in the 1980s and 1990s, when disease pathogenesis was only partially understood and treatment options were limited. Better understanding of disease mechanisms, including the identification of the main target podocyte an-
Evidence that B-cells play a central pathogenic role in MN, both as antigen-presenting cells [5] and as autoantibody producing cells [4], provided the background for explorative studies testing the role of B-cell-depletion therapy with the monoclonal antibody rituximab. The first report in 2002 showing that rituximab safely ameliorated nephrotic syndrome (NS) in 8 patients with primary MN [6] fueled a series of observational studies that uniformly confirmed the excellent safety/efficacy profile of rituximab in this glomerular disease. Evidence accumulated so far from prospective studies [7–9], including a series of 100 consecutive patients [10], collectively indicates that rituximab therapy safely induces complete or partial remission in approximately 65% of MN patients with NS (fig. 1). Proteinuria remission generally occurs within 1 year after therapy, even though late responses have been observed as well. Response to therapy is similar between patients who receive rituximab as first-line therapy or as a rescue treatment after other treatments have failed [11]. Importantly, mechanistic studies have shown that rituximab-induced depletion of circulating B-cells is followed by a decline in anti-PLA2R antibodies that invariably anticipates a decline in proteinuria [12, 13]. These data, along with the finding observed in patients with repeated renal biopsies that rituximab-induced proteinuria remission is associated with the disappearance of subepithelial immune complexes [14], the hallmark of MN indicates that rituximab targets a crucial pathogenic mechanism of the disease.

Recently, the randomized-controlled Evaluate Rituximab Treatment for Idiopathic Membranous Nephropathy (GEMRITUX) trial randomized 75 MN patients with NS to rituximab (two 375 mg/m² doses) versus no immunosuppression and evaluated the rate of complete or partial remission at 6 months after therapy (primary endpoint) [15]. Although the study failed in detecting a significant difference between the 2 groups in the primary 6-month endpoint (35 vs. 21% in rituximab-treated vs. control patients; p = 0.21), after an extended median follow-up of 17 months, 64.9% of rituximab-treated patients versus 34.2% of the untreated subjects achieved remission (p < 0.01). Treatment was very well tolerated.

Altogether, these results appear similar or even superior to those reported in trials testing the efficacy of alkylating agents and steroids both in terms of remission rates and time to remission. Despite similar efficacy, however, rituximab therapy is devoid of the serious toxicities of such therapies. Therefore, within the limitations of comparisons across different studies, available data suggest that rituximab is at least as effective as alkylating agents plus steroids but is associated with fewer adverse events [16].

A recent prospective cohort study by Moroni et al. [17] reported less favorable results. These authors evaluated the rate of partial and complete remission after one or 2 rituximab administrations (375 mg/m² each) in 34 patients with MN and NS. No predefined protocol to decide on single or dual rituximab administration is provided. At 6 months after therapy, 15 patients (44%) achieved remission, which is consistent with the data reported in the GEMRITUX trial [15], but in contrast to the GEMRITUX trial, no additional patient achieved remission between 6 and 12 months after therapy. There was no difference in response between patients who received one or 2 rituximab doses, nor between patients who received rituximab as first-line or second-line therapy. Only 24 (70%) of the initial cohort of patients had a follow-up longer than 12 months. Among these patients, 13 (54%) reached remission at 1 year after treatment (2 had a relapse of proteinuria), which is below what has been previously reported in larger studies with predefined rituximab administration protocols (fig. 1). The authors ascribed this result to the lower than commonly used doses of rituximab.

The issue of optimal rituximab dosing in MN is still a matter of debate. Rituximab doses used across the various studies in MN patients differ significantly, ranging from a single dose of 375 mg/m² to a repeated course of four 375 mg/m² weekly doses 6 months apart. The initially used 4-dose regimen was adopted from rituximab dosing in Hodgkin’s lymphoma, the only indication for rituximab therapy at that time [18]. However, as the number of CD20 cells in patients with MN is significantly lower than in patients with lymphoproliferative diseases, the need for anti-CD20 antibody to induce a complete lymphocyte depletion might be consequently lower, consistent with the evidence that CD20+ B-cells are fully depleted from the circulation after the first rituximab administration in patients with MN or lupus. To address this issue, a prospective, matched-cohort study compared the safety/efficacy profile of a B-cell-driven rituximab treatment with the standard four 375 mg/m² dose protocol in 36 MN patients with long-lasting nephrotic range proteinuria refractory to conventional therapy [19]. Patients allocated to the B-cell-driven protocol received a second infusion only if they had more than 5 B-cells/mm³ of pe-
ripheral blood after the first rituximab administration, which occurred in only 1 of the 12 patients in this group. Prompt and persistent B-cell depletion was achieved in all patients. Time-dependent changes in proteinuria and the other components of NS were similar in the 2 groups, but the B-cell-driven approach was associated with fewer adverse events and less hospitalizations, and was fourfold less expensive. These findings were confirmed by a large prospective cohort study including 100 MN patients showing that a B-cell-driven rituximab protocol provides similar efficacy than the 4-dose regimen [10]. Thus, B-cell titrated dosing seems as effective as a 4-dose regimen but is safer and cost-saving. Due to the excellent relationship between the levels of circulating anti-PLA2R antibodies and disease activity, this biomarker could be tested in the future as an alternative tool to titrate rituximab therapy [12].

Consistent with the aforementioned reports, the study by Moroni et al. [17] showed that B-cells were fully depleted in all the patients, regardless of the use of single or repeated rituximab administrations. Unfortunately, lack of serial B-cell measurements prevents any comparison in B-cell recovery between the 2 rituximab treatment regimens. Moreover, the absence of a control group of subjects receiving a 4-dose treatment precludes any conclusion on the impact of rituximab dosing on the proteinuria reduction.

Importantly, previous data have clearly indicated that patients with tubulointerstitial lesions at renal biopsy and impaired renal function have milder and slower response than patients with normal renal function [20]. Since about one third of patients enrolled in the study by Moroni et al. [17] had an estimated glomerular filtration rate <60 ml/min/1.73 m2, a longer follow-up period would have been important to adequately detect the rituximab effect. Lack of serial measurements of anti-PLA2R antibodies also prevents a full understanding of rituximab efficacy in this cohort of patients. Despite these limitations, the authors conclude that ‘since negative studies are seldom reported, the efficacy of rituximab in MN might be overestimated’.

The issue of publication bias against negative findings is of course very important. Statistically significant results are more likely to be published than papers with null results [21]. This bias may seriously distort the literature, drain scarce resources by undertaking research in futile quests, and lead to misguided research and clinical practices. However, the risk of publication bias may change direction over time. The publication cycle also clearly illustrates that significant findings are published ahead of

Fig. 1. Forest plot of the proportion of remission with rituximab after 12–24 months (left) and table reporting numbers of patients included in the studies and rituximab dosing (right). The estimated proportions and their 95% CIs were calculated with a random-effects model using the method of DerSimonian and Laird. The 8 patients included in the series by Ruggenenti et al. [10] and reported in 2013 were removed from the 132 patients reported by the same group in 2015. Even though the forest plot displays equal weights for the individual studies, weighting was indeed done using an iterative procedure. The dashed vertical line represents the overall estimate. Studies not crossing the vertical dashed line are significantly different from the overall average. ES = Estimated proportion. Rituximab doses: * 375 mg/m2; ** 1 g.
nonsignificant findings, and that significant findings seem to provide an incentive to publish nonsignificant studies [22]. Since studies have extensively shown that rituximab safely promotes remission in approximately 65% of patients with MN, there may be now the risk of a publication bias favoring unexpected negative results, which is of course as worrisome as the opposite bias.

More importantly, similar to studies with positive results, negative studies can be conclusive, exploratory, or inconclusive based on their design and statistical power [23]. Is the report by Moroni et al. [17] a true negative study or just an inconclusive collection of clinical data? Unfortunately, lack of a formal protocol or rationale for the choice of the rituximab doses, absence of sample size estimation and a follow-up inadequate to the already available knowledge on the timing for rituximab response make this study far from being a conclusive one.

The evidence supporting the use of rituximab in the treatment of MN represents a valuable advancement in new therapies for other glomerulopathies as well. Experimental studies investigating disease pathogenesis provided the background for this hypothesis-driven approach that due to its selective mechanism of action allowed further understanding of disease mechanisms. Data from small, mechanistic clinical studies led to the GEMRITUX trial and the currently ongoing larger trials (NCT01955187, NCT01180036) that, altogether, will formally define the place of rituximab in the treatment of MN. At the present time, even considering data from Moroni et al. [17], the response rate to rituximab is 65% (fig. 1), which is similar to what has been previously reported with more toxic treatments such as alkylating agents [24]. Importantly, published studies on rituximab therapy employed, in the vast majority of cases, doses similar to the ones employed by Moroni et al. [17]. However, the impact of different reports on the therapeutic decisions should always be grounded on critical appraisal of the quality of the study. Only a randomized controlled trial will definitively answer the question about the optimal dose of rituximab to use in MN. Based on available data, since no evidence supports a relationship between rituximab dose and efficacy in promoting proteinuria remission, while higher doses associate with more adverse events and costs, the B-cell-driven regimen should be the approach employed in everyday clinical practice.

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The author has no conflicts of interest to declare.

References


