IgA Nephropathy: A European Perspective in the Corticosteroid Treatment

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Keywords
Corticosteroid therapy · Europe · IgA nephropathy · Renal pathology · Risk factors for progression

Abstract
Background: IgA nephropathy (IgAN) is detected in Europe in 22% of glomerular diseases diagnosed by biopsy. The frequency of IgAN as cause of ESRD in Europe has increased in the last decades, accounting for 35% of young and adult transplanted patients. These data justify the interest for risk factors and a possible therapeutic approach. Summary: Insight into a European perspective of IgAN was allowed by the multicenter study VALIGA, on 1,147 patients, almost all Caucasians, with follow-up of 4.7 years. The predictive value of mesangial hypercellularity (M), segmental sclerosis (S), tubular atrophy interstitial fibrosis (T) as independent biomarkers of progression was validated. Endocapillary hypercellularity was predictive of increased follow-up proteinuria. Two groups of patients selected by a propensity score to perfectly match for histologic features (MEST) and clinical data treated with renin-angiotensin system blockers (RASBs) and corticosteroids, or RASBs alone were compared and a beneficial effect of corticosteroids in addition to RASB was found in patients with proteinuria > 1 g/day, with an initial eGFR < 50 mL/min/1.73 m². On the contrary, the STOP-IgAN RCT found that immunosuppressive therapy in addition to optimized supportive care did not provide substantial kidney-related benefits in European patients with IgAN, because there was no difference in the rate of decrease in eGFR, although corticosteroid/immunosuppressive therapy induced complete remission of proteinuria more frequently than supportive care alone. The NEFIGAN trial evaluated a targeted release formulation of budesonide (TRF budesonide) delivering the drug in the distal ileum. TRF budesonide, additionally to optimized RAS blockade, reduced proteinuria and maintained eGFR in IgAN patients, suggesting a reduced risk of future progression to ESRD. Key Messages: In Europe, there is a reasoned search of a balanced approach to corticosteroid therapy for patients with IgAN, with particular attention to selecting the patients at risk of progression while limiting the unwanted systemic adverse events.

Introduction

IgAN is the most frequent glomerular disease in Europe, with an incidence of 8 to 25 new cases/year/1mpmp (per million of age-related population) in adults and 3 to 5 new cases/year/1mpmp in children [1]. In Europe, the frequency of IgAN ranges from 19 to 51% of renal biopsies performed in glomerular diseases in various countries [2–4]. From the results of the International Kidney...
Biopsy Survey (IKSB) on glomerular disease frequencies, analyzing 42,603 renal biopsies in four continents, IgAN has a high frequency in Europe, though less than in Asia, accounting for 22 and 39%, respectively, of all glomerular diseases diagnosed by renal biopsies [5]. A genetic background is likely to play a role, since the increase in incidence of IgAN with eastward and northward distance from Africa is correlated with an increase of the 7-SNP genetic risk score [6]. However, according to the IKSB survey, IgAN frequencies differed by continent, even among patients of the same race/ethnicity, being lower in whites in North America in respect to those in Europe (14 vs. 25%, \( p < 0.001 \)) or in Asians in North America to those in Asia (27 vs. 40%, \( p < 0.001 \)). Regional dietetic and microbiota environmental differences and lifestyle factors may influence the different incidence of IgAN in the same ethnic groups, but most of all, local biopsy policies influence IgAN epidemiology, as reported by the Scottish registry almost 20 years ago [7]. The frequency is even more strongly influenced by the present renal biopsy policy first adopted in the US and nowadays followed in some (though not all) European countries as well, which has a trend to not biopsy a patient with chronic urinary abnormalities before a trial with renin-angiotensin system blockers (RASB) and the manifestation of persistent relevant proteinuria.

In Europe, IgAN is a relevant disease ending in end-stage renal failure and needing renal replacement treatment (RRT). The ERA-EDTA Registry reported from countries with a long-term follow-up since 1990 (including Denmark, Finland, Iceland, Norway, the Netherlands, Sweden, and UK/Scotland) showed an increase in incidence of RRT in patients with primary IgAN from 4 to 6% of all subjects entering the program/year, while IgA vasculitis (Henoch-Schoenlein purpura nephritis) maintained the same incidence of 1% over the last decades. According to the ERA-EDTA Registry, IgAN is the original disease in 35% of 20- to 44-year-old transplanted patients in Europe. These data justify the interest of European nephrology for IgAN and particularly for the risk factors for progression and a possible therapeutic approach.

Presentation of IgAN in Europe Focusing on Pathology Risk Factors Derived from the VALIGA Study on 1,147 Patients with IgAN

The Validation Study of the Oxford Classification of IgAN (VALIGA) [8] created a large European database of patients with IgAN followed over a prolonged time frame, which included clinical, laboratory, and histological data. The study was initially aimed to validate the Oxford Classification of IgAN in a European cohort. The MEST lesions – mesangial (M) or endocapillary (E) hypercellularity, segmental glomerulosclerosis (S), and tubular atrophy/interstitial fibrosis (T) – predicted renal outcome independently of clinical data at renal biopsy and during follow-up and were valid in children as well as in adults [9, 10]. However, the Oxford cohort was rather limited including 265 adults and children only from 15 centers in 11 countries [11]. Several validation studies have been performed in the following years [12]. The VALIGA study validated in Europe the Oxford classification in cohorts with different presentations and treatments expanding the enrollment criteria of the original study, including patients with mild proteinuria <0.5 g/day and those with follow-up shorter than 1 year, if ending in ESRD [8].

The multicenter VALIGA study involved 1,147 European patients, almost all Caucasians, with a follow-up of 4.7 years. In this large cohort, the value of M, S, and T as independent histological biomarkers of progression was confirmed. The strongest predictive value was the presence of irreversible extensive T lesions in >25% of the renal biopsy tissue (T1 and T2), but several data indicated also the value of reversible pathology risk factors.

In this European validation study, the independent value of endocapillary hypercellularity (E1) was not detected, but there was an important bias of a greater use of corticosteroid/immunosuppressive drugs in patients with E1. However, in patients with mild proteinuria (<0.5 g/day), E1 lesions were associated with significantly lower survival from a 50% reduction of e-GFR/ESRD (combined end point) and were risk factors for developing higher proteinuria levels (>1 g/day) [8]. In a single-center European study investigating patients with IgAN without corticosteroid or immunosuppressive treatment irrespective of clinical features at renal biopsy and followed for 6 years, E1 was the best predictor of renal survival [13]. This was indicated also by a Chinese study of repeated renal biopsies after treatment with various combinations of immunosuppressive drugs, in which a significant decrease in E1 lesions was reported [14]. In a recent European study, E1 was found to be reduced by mycophenolate moftetil treatment in repeated biopsies [15]. In conclusion, in Europe, the value of endocapillary hypercellularity as risk factor for IgAN progression and the possible benefit of corticosteroid/immunosuppressive treatment is taken into account, while waiting for the results of a future specific RCT.
Mesangial hypercellularity M1 was found in the VALIGA study to be predictive of outcome also in the subgroup of patients with eGFR < 30 mL/min [8]. The presence of M1 was a histological marker predicting benefits of steroid therapy [16]. In 261 young subjects aged < 23 years enrolled in VALIGA, the presence of M1 was a significant risk for survival from the combined end point and for high time-averaged proteinuria during the subsequent follow-up [17]. The remission of proteinuria to values < 0.5 g/day/1.73 m² was significantly predicted by the absence of mesangial proliferation, preserved GFR at renal biopsy (> 90 mL/min/1.73 m²), and use of corticosteroid/immunosuppressive therapy. These data indicate that mesangial hypercellularity in VALIGA children and young subjects is a risk factor for progression and may be reversed by steroid therapy. Almost 20 years ago, a Japanese RCT [18] enrolling children with mesangial hypercellularity in > 80% of glomeruli demonstrated a significant benefit of 2 years of prednisone and azathioprine therapy in addition to anticoagulation and antiplatelet treatment in comparison to anticoagulation and antiplatelet therapy only. The mesangial proliferation significantly decreased in children treated by corticosteroids. Ten years after the end of the treatment, the renal survival was significantly better in children previously treated with corticosteroids/immunosuppressors [19]. In Europe, the present tendency is to consider the presence of M1 as a possible marker of the need of corticosteroid therapy particularly in children and young subjects even with mild proteinuria, as suggested also from the collaborative study which pooled data from VALIGA, Chinese, Japanese, and North American cohorts [16]. This study proved that the presence of M1 was a significant risk factor for progression even in patients with proteinuria < 1 g/day at renal biopsy, which showed a decline similar to that of patients with persistent time-averaged proteinuria of 1–2 g/day for 2 years, hence fully eligible for corticosteroid treatment according to KDIGO 2012 [20].

Segmental glomerular sclerosis (S1) In VALIGA European patients was a risk factor for progression [8]. In a recent European study, podocyte hypertrophy or sclerosis at the tubular pole (tip lesion), features typically associated with podocytopathies, was correlated with more proteinuria at presentation and more rapid decline in renal function if not treated with corticosteroids/immunosuppressors [21]. The revised Oxford Classification [22] recommends reporting segmental sclerosis with/without podocyte hypertrophy/tip lesions. In the VALIGA study, propensity score-matched patients with S1 had a significant benefit from corticosteroid/immunosuppressive therapy [23]. These results are of interest, while in Europe, we are waiting for a RCT demonstrating the potential benefits of corticosteroids/immunosuppressors in IgAN with S1 podocytopathic lesions.

Crescents lesions were not found to have a prognostic value at multivariate analysis in the VALIGA study, but there was a bias of a great use of immunosuppressive treatment in these patients. A multicenter study analyzing pooled data from the Oxford, VALIGA, Chinese, and Japanese cohorts (3,096 patients, one-third with crescents) produced an interesting new analysis of crescents as a risk factor for IgAN [24]. Crescents were predictive of the combined event of 50% decline in GFR or ESRD but only in patients not receiving immunosuppressors. Patients with > 25% of crescents were at risk even if treated with immunosuppressive therapy. This study indicates the need to add crescent scores (C0, C1, and C2) to the Oxford MEST scores [22]. A new MEST-C score has been proposed and largely adopted in Europe. In Europe, patients with crescents in > 25% of glomeruli are most often treated with corticosteroid/immunosuppressive therapy.

**Indications for Corticosteroid Therapy for IgAN in Europe Derived from the Observational VALIGA Study, the STOP-IgAN, and the NEFIGAN RCTs**

**VALIGA Study**

The large cohort of European patients with IgAN observed in the VALIGA study allowed the identification of two groups of 184 patients, each selected by a propensity score which perfectly matched for histologic features (MEST scores) and clinical data: one group was treated with RASBs and corticosteroids, and the other with RASBs alone [23]. Patients treated with corticosteroids and RASBs had better outcomes compared with matched patients with RASBs alone [23]. Patients treated with corticosteroids and RASBs had better outcomes compared with matched patients with RASBs alone for survival to the combined end point (p = 0.004), decrease in proteinuria during the follow-up (p = 0.001), showing a significantly higher frequency of reducing proteinuria < 1 g/day. The benefits of corticosteroids increased according to pretreatment-persistent levels of proteinuria. The greatest protective effect was observed when proteinuria was > 3 g/day (p = 0.001) or > 1 g/day (p = 0.03), whereas no difference between treatments was found when proteinuria was < 1 g/day. A lower rate of renal function decline and a greater reduction in proteinuria were observed when patients with M1, S1, and T1 scores were treated with corticosteroids compared with those treated with RASB alone.

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This analysis of the European VALIGA patients led to the conclusion of a beneficial effect of corticosteroids in addition to RASB in IgAN patients with proteinuria >1 g/day, even with an initial eGFR <50 mL/min/1.73 m². However, this was a retrospective cohort study and the results of prospective RCT were expected in Europe as in the whole nephrologist community. The results of the STOP-IgAN RCT were published after a few months in 2015 [25].

STOP-IgAN RCT

The STOP-IgAN trial [25] tested the hypothesis that in patients with IgAN, corticosteroid/immunosuppressive therapy added to comprehensive supportive care would be superior to supportive care alone. The primary end point, after 3 years, was full clinical remission (<0.2 mg/mg proteinuria/cr with stable eGFR) or decrease in the eGFR >15 mL/min/1.73 m². Eligible patients had proteinuria >0.75 and <3.5 g/day and eGFR >30 mL/min/1.73 m². The randomization excluded 34% of responders to the 6-month run-in phase with supportive care who showed a decline in proteinuria to values <0.75 g/day) and 12% of high-risk patients with higher proteinuria levels and fast eGFR decline. Patients were randomized to continue supportive care or to receive corticosteroid/immunosuppressive therapy; 55 patients with eGFR of >60 mL/min received corticosteroid monotherapy for 6 months (9 methylprednisolone pulses and oral prednisolone at 0.5 mg/kg on alternate days), and 27 patients with eGFR >30 and <59 mL/min received, for 36 months, 1.5 mg/kg/day cyclophosphamide followed by 1.5 mg/kg/day azathioprine plus 40 mg/day oral prednisolone, waning over 36 months. At the end of the 3 years, 5% of patients in the supportive care group had full clinical remission (p = 0.001) versus 17% in the immunosuppression group (p = 0.001); however, no difference between the two groups was found in the decrease of eGFR >15 mL/min/1.73 m² (28 vs. 26%, respectively). Patients in the corticosteroid monotherapy group reached significantly lower mean proteinuria than those in the supportive care alone, but data were pooled together with those in patients with advanced CKD receiving the combination therapy, and no difference in proteinuria or GFR decline was found at 3 years. Microscopic hematuria disappeared more frequently in the treatment group at the end of the RCT.

The STOP-IgAN RCT found that immunosuppressive therapy in addition to optimal supportive care did not provide substantial kidney-related benefits in European patients with IgAN, because there was no difference in the rate of decrease in eGFR, although corticosteroid/immunosuppressive therapy induced complete remission of proteinuria more frequently than supportive care alone.

The results of this RCT were of great relevance, though some flaws were pointed out by the nephrologist community (in Europe as well as in other continents) [26]. The patients enrolled had a very limited renal function decline of ~1.6 mL/min/year, rendering it difficult to prove the benefits of any additional treatment after a relatively short follow-up [26]. The STOP-IgAN trial was primarily powered to detect a difference in clinical remission, which it did in favor of immunosuppression [27], showing a reduction in proteinuria which has been validated by a recent meta-analysis [28]. The trial was too short and underpowered to detect the changes in renal function which have been validated as surrogate markers of ESRD such as a 50% decline in eGFR. The two groups, receiving two immunosuppressive regimens, had different modification of proteinuria, which was good in the group with relatively preserved eGFR and receiving corticosteroid monotherapy: the effects on eGFR decline on the long-term course are expected to be different [26]. In the VALIGA, the protective effect of proteinuria >0.5 g/day and <1 g/day on the combined end point was detected after 10 years [8].

The authors of the STOP-IgAN RCT concluded that immunosuppressive therapy in addition to optimal supportive care would not provide substantial kidney-related benefits in European patients with high-risk IgAN, because there was no difference in the rate of decrease in eGFR, although corticosteroid/immunosuppressive therapy induced complete remission of proteinuria more frequently than supportive care alone. However, there were many flaws as detailed above, rendering the absolutely negative conclusion about the ineffectiveness of corticosteroid/immunosuppressive therapy in IgAN not fully acceptable.

Notably, the STOP-IgAN provided relevant information, demonstrating the strong benefit of aggressive optimization of supportive measures that result in a sustained reduction in proteinuria and a low short-term risk of renal function decline. However, the issue of side effects was raised (one death due to pneumogenic sepsis and two neoplasms in the immunosuppressive group), which was further claimed by the results of the TESTING RCT in mostly Chinese patients [29]. The latter trial was prematurely discontinued because of an imbalance of adverse events, mostly infectious, in the corticosteroid arm, in spite of greater protection against renal failure progression. Hence, great interest is focused in Europe and ev-
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Safety of Treatment of IgAN with Immunosuppressors

In a meta-analysis on 245 patients with IgAN from 9 RCTs treated with corticosteroids, the incidence of severe adverse events was 6.9% (mostly hypertension and diabetes and rarely, gastrointestinal bleeding) [32]. In a recent meta-analysis on 325 patients treated with intravenous steroid pulses alone or in combination with azathioprine, the risk of major adverse events was low in patients with normal renal function (2.9%) but increased in those with impaired renal function (15.4%), particularly when adding azathioprine (30%).

In the STOP trial, adverse events were more common than in prior trials. Serious adverse events had similar frequencies in the two study groups, but patients treated with immunosuppressive therapy reported more cases of infection, though not significantly different from control patients (36 vs. 40%), impaired glucose metabolism, and body weight gain. Moreover, two patients had malignant neoplasms during combined immunosuppression. In the TESTING trial, the frequency of adverse events was higher in patients treated with high FGR than in previous RCTs. Moreover, the frequency of side effects is also increased in protocols with exposure to corticosteroids/immunosuppressive drugs longer than 6 months: the frequency of adverse events was high in the placebo group (84%), similar to what was observed in the STOP trial (36% of serious adverse events in the placebo group). The frequency of any side effect was high in the treatment as well as in the placebo groups in recent RCTs in which adverse event reports were solicited.

Conclusion from Recent European Studies on Corticosteroid Treatment of Patients with IgAN

- A protracted and rigorous supportive care, targeting RAS for BP and proteinuria, metabolic and lifestyle targets can be of benefit in one-third of patients with proteinuria <0.75–3.5 g/day and not at high risk of rapid progression [25].
Addition of corticosteroid therapy needed? The STOP-IgAN trial provided a negative answer, but a personalized approach may be proposed: no when proteinuria 0.75–1.5 g/day and negative MEST scores [16]; yes when IgAN is in progression with rapid loss of GFR or when crescents are present in >25% of glomeruli (C1) [24]; probably yes when risk factors are present, with persistent proteinuria >1 g/day [8] or when proteinuria <1 g/day with M1 or E1 or S1 with podocytopathy [22].

Addition of corticosteroids to supportive care induces reduction in proteinuria and possible renal-protective effects on the long term with increase in adverse events but mostly in cases with impaired renal function [8, 28, 33].

The enteric formulation of budesonide seems to provide similar favorable results without serious side effects of steroids [31].

Addition of alkylating agents/antimetabolites to corticosteroids is not indicated in non-vasculitis-like progressive forms, particularly when GFR <50 mL/min/1.73 m² [25, 33].

Conflict of Interest Statement

The author has no conflict of interest to declare.

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