INVITED REVIEW ARTICLE

Novel emerging treatments for NMOSD

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ABSTRACT
Neuromyelitis optica spectrum disorders (NMOSD) are inflammatory demyelinating diseases of the central nervous system (CNS) that cause optic neuritis, transverse myelitis, and some other CNS syndromes.

Recently, diagnosis and understanding of these diseases has been markedly enhanced by the discovery that serum autoantibodies that target aquaporin-4 (AQP4) are strongly associated with the disease. This spectrum includes also a potential subset of patients with a phenotype of NMOSD who have anti-myelin oligodendrocyte glycoprotein (MOG) antibody. Although steroids and immunosuppressive drugs have been widely used for NMOSD treatment, until recently there was no approved therapy for these diseases. With improved understanding of the pathophysiology of NMOSD, numerous new therapeutic strategies have recently been evaluated. The results of these studies, involving monoclonal antibodies (mAbs) inhibiting terminal complement protein cleavage interfering with interleukin-6 receptor (IL-6 R) signaling and depleting CD19-positive B cells, have been published in recent months. All of these new therapeutics have shown a high degree of efficacy in diminishing NMOSD activity and inhibiting disability progression. At the same time, all these mAbs have demonstrated favorable safety and tolerability profiles, with a limited rate of adverse events. The first of these new drugs, eculizumab, have been approved in USA and Europe for NMOSD treatment within the last couple of months and it is expected that the other novel, effective and safe treatments for NMOSD will be approved in the near future.

Key words: neuromyelitis optica spectrum disorders, monoclonal antibodies, demyelinating diseases


Introduction
Neuromyelitis optica spectrum disorders (NMOSD) are autoimmune inflammatory disorders of the central nervous system (CNS) characterised by bilateral or rapidly sequential optic neuritis and/or transverse myelitis. Other suggestive presentations include episodes of brainstem symptoms, in particular area postrema clinical syndrome. The disease can result in severe muscle weakness and paralysis, loss of vision, sensory loss, bladder dysfunction, neuropathic pain, and in the most severe cases respiratory failure [1]. The prevalence of NMOSD in Caucasians is relatively low at 0.5–10 per 100,000, but severe and frequently rapid impairment observed in patients can lead to an unfavourable prognosis, including death.

The mechanism of NMOSD is associated with the presence in the serum of patients’ IgG1 antibodies against aquaporin-4 (AQP4 Ab), the major water channel present within the CNS specifically on astrocyte endfeets at the blood-brain-barrier (BBB) [2]. This antibody can be found in more than 80% of patients. The discovery of AQP4 Ab has contributed significantly to our understanding of the pathology of NMOSD, also allowing for a much more precise diagnosis. AQP4 Ab appears to have a pathogenetic role in the mechanism of NMOSD [3]. Recombinant AQP4 Ab after passive transfer in rats has induced NMOSD-specific immunopathology [4]. Serum AQP4 Ab titers have been shown to correlate with clinical attacks and with the extent of spinal cord lesions on MRI [5, 6]. In addition, serum AQP4 Ab titers have been shown to drop after immunosuppressive treatment, and to remain low during remissions [7]. This antibody, produced by T-helper dependent B cells from the peripheral immune compartment, crosses the blood-brain-barrier (BBB) and interacts with AQP4 on astrocyte endfeets leading to astrocyte oedema and dysfunction. Subsequent inflammation involves complement activation, increased BBB permeability, and a massive influx of neutrophils and eosinophils [8]. Thus, inflammatory lesions in NMOSD are clearly different from CNS inflammation in multiple sclerosis (MS), where T and B cells constitute the majority of invading
cells. These findings, demonstrating significant mechanistic differences between MS and NMOSD, have allowed for a definitive separation of these two clinical entities.

More recently, another antigen was found to be associated with AQP4 seronegative NMOSD. Antibodies against myelin oligodendrocyte glycoprotein (MOG) were detected in 4–11% of seronegative patients [9]. Unlike AQP4 Ab, anti-MOG Ab did not induce astrocitary pathology. It was found that anti-MOG Ab led to demyelination with limited immune cell infiltration [10]. Anti-MOG Ab almost never co-exists with AQP4 Ab and occurs much more frequently in children than in adult patients.

Despite the discovery of AQP4 Ab and anti-MOG Ab, there are still patients who meet the clinical diagnostic criteria of NMOSD, but in whose sera the two Abs cannot be detected. The prerequisite of a NMOSD diagnosis requires the presence of optic nerve and spinal cord symptoms [11]. The current 2015 International Consensus Diagnostic Criteria for NMOSD are stratified according to the presence of AQP4 Ab. For seropositive patients, they require the presence of at least one of the core manifestations, whereas for seronegative patients two manifestations including optic neuritis, myelitis or area postrema syndrome are required (Tab. 1). Recently, preliminary results have suggested the role of antibodies against glial fibrillary acidic protein (GFAP) in seronegative NMOSD [12].

**Current treatment of NMOSD**

Until recently, there was no approved treatment for NMOSD, and patients were restricted to off-label therapies bringing uncertain benefits. Based on its clinical course, NMOSD therapy can be divided into relapse treatment and preventive treatment.

**Relapse treatment**

For relapse, patients are usually treated with pulsed steroid therapy. Methylprednisolone at a dose of 1g is given intravenously (IV) for 3–5 days but therapy can be extended in a very severe relapse [13]. Relapses that respond poorly to methylprednisolone can be treated with plasma exchange (PLEX), usually 5–7 procedures every other day. PLEX can also be used as a first line therapy for NMOSD relapse [14]. Since the progression of disability in NMOSD is mainly driven by relapses, it is critically important that NMOSD relapse is treated as early as possible and with the most effective procedure.

Shortly after steroid pulsed therapy or PLEX, patients usually are started with immunosuppressive therapies. To protect patients from disease progression before the immunosuppressive drugs start to work, prednisone or prednisolone can be given orally for up to six months. Relapses in NMOSD dependent on anti-MOG Ab usually respond better to steroid treatment than in AQP4 dependent disease [15]. Limited data indicates that intravenous immunoglobulins (IvIg) might show a benefit in NMOSD relapse treatment. In one study, the effectiveness of IvIg in the treatment of acute relapses in NMOSD was assessed in a retrospective review of 10 patients unresponsive to treatment with steroids with or without PLEX [16]. Improvement was noted in five of 11 (45.5%) relapses, and the remaining relapses had no further worsening. The study concluded that IvIg may have a role in treating acute NMOSD relapses.

**Immunosuppressive treatment**

Based on the autoimmune mechanism of NMOSD involving the production of auto-antibodies against AQP4 and MOG, several immunosuppressive agents have been tested in this disease over the past 30 years. Unfortunately, on most occasions the studies only involved a small number of patients and were designed as open label trials. None of these drugs have been formally approved for NMOSD treatment.

Traditionally, Azathioprine (AZT) was widely used as a first line NMOSD treatment. AZT was recommended as

<table>
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<th>Table 1. IPND 2015 diagnostic criteria for NMOSD</th>
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<tr>
<td><strong>NMOSD with AQP4-Ab</strong></td>
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<tr>
<td>At least one core clinical characteristic plus positive test for AQP4-IgG using best available detection method*</td>
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<tr>
<td>Exclusion of alternative diagnoses</td>
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<tr>
<td><strong>NMOSD without AQP4-Ab</strong></td>
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<tr>
<td>At least two core clinical characteristics occurring as a result of one or more clinical attacks and meeting all of the following requirements:</td>
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<tr>
<td>1. At least one core clinical characteristic must be optic neuritis, acute myelitis with LETM, or area postrema syndrome.</td>
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<td>2. Dissemination in space (two or more different core clinical characteristics).</td>
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<tr>
<td>3. Additional MRI requirements, as applicable.</td>
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<tr>
<td>4. Negative tests for AQP4-IgG using best available detection method* or testing unavailable.</td>
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<tr>
<td>5. Exclusion of alternative diagnoses.</td>
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<td><strong>Core clinical characteristics</strong>: Optic neuritis; acute myelitis; area postrema syndrome (hiccups, nausea and vomiting); acute brainstem syndrome; symptomatic narcolepsy or acute diencephalic clinical syndrome with NMOSD-typical diencephalic MRI lesions; symptomatic cerebral syndrome with NMOSD-typical brain lesions.</td>
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*AQP4-IgG serology: cell-based assay is strongly recommended.
a first line preventive treatment by the EFNS panel on guidelines on diagnosis and management of NMO [17]. Several small studies have tested the efficacy of AZT alone, or in combination with steroids [18]. Some of these studies showed an approximately 70% reduction in the NMOSD relapse rate and reduced disability after several years of treatment. In a large retrospective review of the medical records of 103 AQP-4 antibody-positive NMOSD patients, 89% (n = 92) had reduced their median annualised relapse rates from 1.5 (IQR 0.6–4.0) to 0 (IQR 0–0.27, p < 0.00005) during treatment. Sixty-one per cent (n = 63) remained relapse-free at a median follow-up of 18 months. Neurological function improved or stabilised in 78%. However, the discontinuation rate was 47%, reflecting the poor tolerability of this drug [19].

The other problem with AZT treatment in NMOSD is the delayed mechanism of action of this drug. AZT on average requires 3–6 months to demonstrate its effect on the immune system. Mitoxantrone [20], cyclophosphamide [21] and some other immunosuppressive treatments including methotrexate have demonstrated beneficial effects in NMOSD only in case reports and only in a limited number of patients. In addition, safety profiles of these drugs reduce the enthusiasm for their use in NMOSD. Little more data is available regarding Mycophenolate Mofetil (MMF). In a prospective study including 67 NMOSD seropositive and seronegative patients, MMF reduced the annual relapse rate by 49% and stabilised EDSS in 80% of patients within a 24 month observation period. There was no difference between seropositive and seronegative patients [22]. Several retrospective studies have assessed the efficacy of MMF in AQP4- and MOG- seropositive and double negative patients. In one of these studies [23], the median post-MMF annualised relapse rate was significantly lower than the pre-MMF annualised relapse rate (0.0 vs 1.5; p < 0.001). EDSS scores also significantly decreased after MMF treatment (3.0 vs 2.5; p = 0.005). Thirty-five patients (60%) were relapse-free with a median treatment duration of 20 months, and EDSS scores were stabilised or improved in 53 patients (91%). In a similar way to AZT, MMF requires an extended period of time to demonstrate its effect on the immune system, and a substantial number of patients discontinued treatment due to side effects.

**Intravenous immunoglobulins**

In recent years, intravenous immunoglobulins (IvIg), which have been proven to be effective in some other antibody-mediated autoimmune conditions including inflammatory demyelinating polyneuropathies [24], have attracted significant attention in terms of NMOSD treatment. IvIg have been shown to induce diminished activity on membrane-damaging components of the complement system, B cell activating factor (BAFF), and several other immune mechanisms including interference with antigen recognition, downregulation of cytokine secretion, adhesion molecules expression, and suppression of T-cell activation relevant to NMOSD [25]. Although popular views and practical experience are strong advocates for the use of IvIg in NMOSD, there is relatively little evidence to support these conclusions.

The first study to look into the role of IvIg treatment in NMOSD included eight patients [26]. Five experienced relapsing optic neuritis with or without myelitis, and the remaining three had relapsing longitudinal extensive transverse myelitis (LETM). After a total of 83 infusions (4–21 per patient) and a mean follow-up duration of 19.3 months (6–39 months), it was observed that the mean relapse rate had decreased from 1.8 in the 12 months pre-IvIg therapy to 0.06 during follow-up (p = 0.0001), while the mean EDSS score had declined from 3.3 ± 1.3 to 2.6 ± 1.5 (p = 0.04). In another study [27], the use of IvIg was evaluated in preventing relapses in patients with NMOSD. Six NMOSD patients who were treated with an IvIg induction dose followed by infusions every 2–3 months were retrospectively analysed. ARR and EDSS pre- and post-IvIg were recorded. The median number of relapses and the median ARR were significantly reduced (8.0 to 1.0 and 0.75 to 0.15; p < 0.05) during IvIg treatment. EDSS remained the same during four years of treatment.

**Cell depletion therapy**

The discovery of a pathological role of autoantibodies against AQP4 and MOG in NMOSD has prompted the application of B cell depletion therapy which might help to eliminate antibody-producing cells and improve treatment of this disease. Rituximab (RTX), a chimeric monoclonal antibody against CD20, a pan-B cell surface marker, has been tested in NMOSD in several small or medium-sized open labelled studies [28, 29]. RTX very efficiently depleted B cells over a duration of 6–9 months. Importantly, the cells from early stages of B cell lineage development and plasma cells were preserved, securing immune homeostasis during RTX treatment.

Most of the RTX studies have shown its profound beneficial effect on ARR and on stabilisation and reduction of disability measured with EDSS. Some of these studies have even shown complete suppression of relapses over a 2–3 year period. A recently published meta-analysis analysed 26 studies, in which differences in the ARR ratio and EDSS score before and after RTX therapy were used as the main efficacy measures [30]. This meta-analysis involved 577 patients. Antibodies against aquaporin-4 were present in 435 (75.39%) patients. The findings suggested diminished mean ARR ratio after rituximab therapy by 1.56 (95% CI, -1.82 to -1.29). No significant correlation was detected between the outcome of ARR ratio change and the following variables: age at onset, duration of disease, follow-up time, dose of infusion or AQP4-IgG serostatus. The findings of this meta-analysis disclosed also a reduction in the mean EDSS score by -1.16 (95% CI, -1.36 to -0.96) during RTX treatment. A total of 330 out of 528 patients (62.9%) achieved relapse-free status. RTX showed acceptable tolerance, and there were no serious safety issues in NMOSD patients treated with RTX. All of these findings have led to increased off label use of RTX in NMOSD in recent years.
Novel emerging treatments of NMOSD

Progress in the understanding of NMOSD pathologic mechanisms has led to unprecedented attempts to intensify research into the development of new therapies in this disease in recent years. New therapeutic strategies have emerged which involve targeting novel molecules believed to be operating in the pathomechanism of NMOSD. These new strategies include the prevention of complement activation, interfering with IL6 receptor (IL6-R) signalling, and the depletion of AQP-4 and MOG antibodies producing cells (Fig. 1). More importantly, these new strategies were tested in studies designed as double blinded, randomised and controlled phase 2 and 3 trials. Within the last few months, very encouraging results of these studies have become available and have generated renewed hope of combating this devastating disease. It is expected that the results of these recent studies will allow for the formal approval of all of these therapies for NMOSD in the near future.

Eculizumab

Eculizumab is a humanised monoclonal antibody which inhibits the terminal complement protein C5 and prevents its cleavage into C5a and C5b fragments [31]. Preclinical data indicates that AQP4 Ab triggers the complement cascade [32], which leads to inflammation and the formation of a membrane attack complex. The membrane attack complex is implicated in astrocyte destruction and neuronal injury. C5a expresses proinflammatory activity, and C5b induces the formation of a membrane attack complex. Experimental studies have shown that complement inhibitor efficiently suppresses NMOSD development [33]. Eculizumab has been approved in the USA and Europe for paroxysmal nocturnal haemoglobinuria, atypical haemolytic uraemic syndrome, and myasthenia gravis.

In recently published results of a phase 3, randomised, double-blind, placebo-controlled, time-to-event trial (PREVENT) it was shown that eculizumab significantly suppressed disease activity in 143 AQP4-Ab-positive patients with a moderate to severe course of the disease [34]. Patients were randomised 2:1 to either treatment with eculizumab or a placebo. The drug was administered intravenously at a dose of 900 mg weekly for the first four doses; subsequently patients received a maintenance regimen of 1,200 mg every two weeks until relapse or until the end of the trial. Immunosuppressive treatment used prior to study entry was allowed (with the exception of rituximab) and the group of immunosuppressive concomitant treatment was 108 patients out of the total group of 143. The primary efficacy endpoint was the first adjudicated relapse. The primary endpoint occurred much less often in the eculizumab group (3 of 96 patients – 3%) than in the placebo group (20 of 47 – 43%) (HR, 0.06; 95% CI, 0.02 to 0.20; p < 0.001). At 48 weeks, 97.9% of patients receiving eculizumab were relapse-free, compared to 63.2% of patients receiving the placebo. Symptoms of most of the relapses were related to myelitis. Eculizumab was associated with a lower adjudicated annualised relapse rate after adjustment than the placebo, which represented the first secondary end-point, 0.02 and 0.35, respectively (p < 0.001). No conclusions could be made regarding the remaining secondary endpoints because the difference between groups for the next endpoint in the hierarchy, which was a change in the EDSS score, was not significant. The lack of between-group differences in disability progression resulted from the trial design precluding follow-up beyond six weeks after a single relapse. This implied that there were no benefits of the drug on disability progression during the short period of the trial. Patients from the eculizumab group had higher rates of upper respiratory tract infection and headache than did patients in
the placebo group: 31 versus 19 events of upper respiratory tract infection per 100 patient-years, and 55 versus 38 headache events per 100 patient-years. One patient in the eculizumab group who was receiving concomitant azathioprine died from pulmonary empyema. Eculizumab increases the risk of meningococcal and encapsulated bacterial infection [35], and all patients received meningococcal vaccination prior to study entry. No cases of meningococcal infection were reported during the trial.

Eculizumab (Soliris) was approved by FDA for NMOSD treatment following an expedited six-month priority review in June 2019. EMA approved Eculizumab for treatment of AQP4-Ab-positive patients with relapsing course of the disease in August 2019.

**Tocilizumab**

Tocilizumab was the first anti-IL6-R mAb tested in the treatment of NMOSD. The rationale for using tocilizumab in NMO was related to the potential role of IL-6 in the mechanism of NMOSD [36]. This drug has been already used for the treatment of severe rheumatic arthritis patients.

Tocilizumab has been tested in NMOSD in several small open label studies and case reports [37, 38]. These small series have demonstrated a beneficial effect in NMOSD. In one study [39], eight female patients with highly active AQP4-Ab-seropositive NMOSD (n = 6) and NMOSD patients (n = 2) whose disease had been resistant to previous medications including B-cell depletion, were switched to tocilizumab (6–8 mg/kg of body weight per dose). The patients were followed up for 30.9 months after switching to tocilizumab. Two of the eight patients received add-on therapy consisting of monthly corticosteroid pulses or azathioprine. During tocilizumab treatment, the median annualised relapse rate significantly decreased from 4.0 in the year before tocilizumab therapy to 0.4 (p = 0.008), and the median EDSS score significantly decreased, from 7.3 to 5.5 (p = 0.03). Active magnetic resonance imaging lesions were seen in 6/8 patients at tocilizumab initiation and in 1/8 patients at the last magnetic resonance imaging. Three patients remained relapse-free during tocilizumab treatment. The AQ4P-Ab titers (p = 0.02) and pain levels (p = 0.02) dropped significantly during tocilizumab treatment. Adverse effects included moderate cholesterol elevation in 6/8 patients, infections in 4/8, deep venous thrombosis in 1/8, and neutropenia in 1/8.

Tocilizumab development for NMOSD treatment has been delayed since another anti-IL6-R mAb, satralizumab, was designed to improve pharmacokinetics of IL6-R binding and provide better efficacy (see below). However, at the recent ECTRIMS meeting in Stockholm, the results were presented of an investigator-initiated study (Tango) in China [40]. This was a randomised, open-label, parallel-group study comparing tocilizumab with azathioprine. Patients (n = 118) were randomly assigned 1:1 to receive 8 mg/kg intravenous tocilizumab monthly or 2–3 mg/kg oral azathioprine daily. Treatment was administered in conjunction with a gradual discontinuation of the previous treatments, followed by monotherapy for 12 months; 85% of patients were seropositive for AQP-4. The primary endpoint was the time to first relapse. After a mean observation period of 48 weeks, the percentages of relapse-free patients were 91.5% in the tocilizumab group and 67.8% in the azathioprine group (HR = 0.32, 95% CI 0.14–0.70, p = 0.004). Sustained reduction in disability was more likely among patients treated with tocilizumab than in patients with azathioprine (HR = 0.34, 95% CI 0.13–0.90, p = 0.03). Serum levels of anti-AQP4-Ab were reduced significantly, by 42% with tocilizumab compared to 15% with azathioprine (p = 0.03). In the azathioprine group, there was a higher frequency of lymphopenia (46% vs 7%) and anaemia (27% vs 41%) than in the tocilizumab group. During the Tango study, two patients died (one in each arm): in tocilizumab because of a severe recurrence of NMOSD, and in azathioprine because of meningococcal sepsis. In both groups, there was an increased number of patients with elevated transaminases (31% and 46%) respectively for tocilizumab and azathioprine.

**Satralizumab**

Satralizumab is a humanised IgG2 subtype recombinant anti-IL6-R monoclonal antibody [41]. IL-6 is a pro-inflammatory pleiotropic cytokine produced by a large number of cell types, including T and B lymphocytes, monocytes and fibroblasts [42]. IL-6 plays a role in several immunopathological processes such as T-cell activation, induction of immunoglobulin secretion, and enhancing macrophage activity, and it has been implicated in the mechanism of NMOSD [43]. IL-6 specifically contributes to the development of autoimmunity by promoting the generation of Th17 cell lineage [44]. Th17 cells are recognised as the primary T cell sub-population contributing to the development of autoimmune conditions. IL-6 has been found to be significantly elevated in the serum and cerebrospinal fluid of patients with NMOSD; it induces AQP4-Ab production by plasmablasts, and thus represents a novel therapeutic target for NMOSD.

The molecule of satralizumab was designed to improve pharmacokinetics of anti-IL6-R Ab by applying so-called ‘antibody recycling technology’ [45]. This technology leads to increased dissociation of anti-IL6-R Ab from IL6-R within the acidic environment of the endosome, while maintaining its binding affinity to IL6-R in plasma. Thus, in the endosome acidic environment, IL6-R Ab after degradation of IL6-R is dissociated from this complex and can again bind another IL6-R in the plasma, increasing its efficacy of IL6-R elimination. Satralizumab efficacy in NMOSD was evaluated in 83 patients in a randomised, double-blind, phase 3 study (SAkuSky) [46]. AQ4P-positive patients represented 66.3% and AQ4P-negative patients 33.7%. In this study, satralizumab was compared to a placebo as an add-on to baseline treatment with a stable dose of immunosuppressive and/or corticosteroids.
Subjects were randomised to satralizumab (120 mg s.c.) or placebo administered at weeks 0, 2, 4, and Q4W thereafter. The primary endpoint was time to first protocol-defined relapse (PDR), adjudicated by a clinical endpoint committee. Pre-specified subgroup analyses included assessing the response to treatment by AQP4-Ab serostatus, baseline treatment, and region. Satralizumab showed a 79% risk reduction of PDR compared to placebo in the NMOSD AQP4-Ab positive subgroup (HR, 0.21; 95% CI, 0.06–0.75). At weeks 48 and 96, the percentages of relapse-free patients were 91.5% (95% CI, 69.6%–97.8%) and 91.5% (95% CI, 69.6%–97.8%) with satralizumab and 59.9% (95% CI, 36.3%–77.3%) and 53.3% (95% CI, 29.3%–72.4%) with placebo, respectively. However, for the NMOSD AQP4-Ab negative patients, satralizumab showed a risk reduction of PDR that was significantly lower compared to the AQP4 Ab positive group, 34% compared to the placebo group (HR, 0.66; 95% CI, 0.20–2.23), and the percentages of relapse-free patients at weeks 48 and 96 were 84.4% (95% CI, 50.4%–95.9%) and 56.3% (95% CI, 24.2%–79.2%) with satralizumab, and 75.5% (95% CI, 41.6%–91.4%) and 67.1% (95% CI, 34.2%–86.2%) with placebo, respectively.

More recently, results have become available for satralizumab monotherapy in NMOSD. In the SAkuraStar study, the efficacy and safety of satralizumab was compared to placebo for relapse prevention in patients with NMOSD [47]. In this phase 3, double-blind, placebo-controlled study, 95 patients were randomised 2:1 to satralizumab (120 mg s.c.) or placebo, administered at weeks 0, 2, 4 and every four weeks thereafter. Unlike the SAkuraSky study, concomitant immunosuppressant medications were prohibited. Patients with AQP-4 antibodies represented 65.1% in the satralizumab group and 71.9% in the placebo group. All patients had had ≥1 documented relapse, including first attack, in the year prior to screening. The primary endpoint was time to first protocol-defined relapse (PDR) adjudicated by a clinical endpoint committee. Satralizumab monotherapy significantly reduced the risk of PDR by 55% compared to placebo (HR 0.45; 95% CI 0.23–0.89; p = 0.018).

Inebilizumab

Inebilizumab is a humanised mAb of IgG1 subtype directed against the extracellular B cell marker CD19 leading to depletion of a broad range of B cells, including autoantibody-secreting plasmablasts and CD19-expressing plasma cells [48]. Inebilizumab induces a cytotoxic T-lymphocyte response and a strong antibody dependent cell cytotoxicity (ADCC) against B cells.

Inebilizumab was evaluated in a phase 3, double-blind, randomised, placebo-controlled trial (N-MOmentum) in 231 NMOSD patients, both AQP4 Ab-positive (91%) and AQP4 Ab-negative (9%) [49]. Enrollment of AQP4 Ab-negative patients required approval of an eligibility committee that confirmed the entry criteria. Participants were randomised 3:1 to either treatment with inebilizumab, 600 mg iv in two doses of 300 mg each two weeks apart, or a placebo, with no further doses occurring after day 15. Concurrent treatment with other immune suppressants was prohibited.

This means that inebilizumab was the first monotherapy tested in NMOSD free from the confounding influence of other background immunosuppressive treatments. B cells were depleted within approximately four weeks of treatment initiation, and this depletion was sustained throughout the randomised, controlled period of the study. The patients were followed for 28 weeks, after which time the blinded control period was stopped early for efficacy. The primary outcome measure was time to first adjudicated relapse. Following the blinded period, patients were given the option of entering an open-label extension period, in which they received 300 mg of inebilizumab every six months. Inebilizumab met the primary efficacy endpoint with a 77% reduction in the risk of developing an NMOSD relapse when compared to placebo in AQP4-Ab seropositive patients after 28 weeks of treatment (HR: 0.227; p < 0.0001).

A similar effect on relapse risk (73% reduction) was observed in the total inebilizumab-treated patient population, inclusive of AQP4-Ab seronegative patients, (HR: 0.272; p < 0.0001). At 28 weeks, at the end of the randomised-controlled period, 89% of AQP4-Ab seropositive patients treated with inebilizumab were relapse-free, versus 58% in the placebo group. Inebilizumab had also met most of the secondary endpoints. Reduction of disability worsening measured with EDSS in the inebilizumab-treated patients was significantly lower, 15.5%, than in the placebo group where it was 33.9%, (p = 0.0049). In inebilizumab-treated patients, the reduction in NMOSD-related hospitalisations was significantly lower, 5.7% of patients, versus placebo 14.3% (p = 0.01). In this study, MRI was also used as a secondary endpoint and showed a reduction in the frequency of cumulative total active MRI lesions in inebilizumab-treated patients (45.4% patients) versus placebo (57.1%) (p = 0.0034). Visual acuity, another secondary endpoint, did not demonstrate a statistically significant difference.
between inebilizumab-treated patients and placebo. Inebilizumab demonstrated favourable safety and tolerability profiles, with an adverse event rate similar to that of the placebo. The rate of infusion-related reactions was low in both arms. The rates of serious and/or ≥ Grade 3 severity adverse events were similar in the inebilizumab (10.3%) and placebo (14.3%) groups. Two deaths occurred in the open-label period; one related to a severe NMO relapse, and the other related to a brain event of unclear aetiology without a definite diagnosis.

Based on data from the pivotal N-MOMentum study, the FDA has granted Breakthrough Therapy Designation (BTD) for the development of inebilizumab for the treatment of NMO.

Breakthrough Therapy Designation is designed to expedite the development and regulatory review of medicines intended to treat a serious condition that have shown encouraging early clinical results which may demonstrate substantial improvement on a clinically significant endpoint over available medicines.

**Future perspectives**

The pathogenic role of AQP4-Ab in NMO is associated with the production of AQP4-Ab by peripheral plasma cells, which can enter the CNS and bind to AQP4 on perivascular astrocytes. This binding initiates activation of the terminal complement complex and the induction of inflammatory lesion formation. Impaired blood-brain barrier function allows for a massive influx of neutrophils and eosinophils into the CNS of NMO patients. Neutrophil counts are elevated in CSF in about 60% of NMO patients during relapse, and about 20% during remission [50]. Eosinophils are also present in the CSF of NMO patients. Degranulation of these cells and the release of several toxic proteins and enzymes provide a direct mechanism of damage to astrocytes, followed by oligodendrocyte injury and neuronal death [51].

Future NMO therapies will need to address the inhibition of AQP4-Ab and prevention of neutrophils and eosinophils activation leading to CNS infiltration. 

*In vitro* and *in vivo* studies have demonstrated that AQP4-Ab deglycosylation or cleavage reduce the complement-dependent cytotoxicity and the antibody-dependent cell-mediated cytotoxicity, leading to decreased astrocyte damage and reduced development of inflammation [52]. Similarly, IgG-degrading enzyme produced by *Streptococcus pyogenes* efficiently cleaved AQP4-Ab in mice *in vivo* and greatly reduced lesion formation in an experimental mice model of NMO [53]. The bacteria-derived endoglycosidases may target AQP4-Ab and reduce its pathogenicity by inhibition of the AQP4-Ab binding to AQP4 and preventing NMO pathology. Another strategy to inhibit interaction between AQP4-Ab with AQP-4 protein involved Aquaporumab which is a synthetic IgG that competes with AQP4-Ab for AQP 4 binding [54]. In contrast to pathogenic AQP4-Ab, the mutated Fcγ portion of Aquaporumab does not activate the antibody-dependent complement and cell-dependent mediated cytotoxicity. Its competitive inhibition of AQP4-Ab binding depends also on the greater affinity of aquaporumab to the AQP4 protein, compared to that of pathological AQP4-Ab. Its efficacy to compete with AQP4-Ab binding has already been proven in a preclinical study [55].

The dominant presence of neutrophils in inflammatory infiltrates of the CNS, and the proven role of these cells in NMO lesions formation, points at their inhibition as a new strategy of NMO treatment.

This concept was supported by findings that intracerebral injection of AQP-4 Ab in neutropenic mice induced less inflammation and demyelination than in mice with normal neutrophil counts [56]. The potential utility of neutrophil protease inhibitors might have particular significance in this regard [57]. It has already been shown that Sivelestat, an inhibitor of neutrophil elastase, demonstrated a beneficial effect in animal models of NMO, as evidenced by reduced NMO lesion formation [56]. Sivelestat was applied intraperitoneally or intracerebrally either alone or in combination with cathepsin G inhibitor. In addition to its inhibition of proteolytic activity, Sivelestat also reduced the production of inflammatory cytokines and suppressed neutrophil-induced capillary permeability and leukocyte kinetics in other conditions [58].

Eosinophil infiltration is another prominent feature of NMO lesions, and eosinophils have been found to be elevated in the CSF of NMO patients [59]. Accordingly, hypoeosinophilic mice showed diminished potential for NMO development. Eosinophil inhibition, either by anti-IL-5 or gene depletion, led to reduced lesion severity in experimental models of NMO.

These findings confirm the involvement of eosinophils in NMO's pathogenesis, and suggests the therapeutic utility of eosinophil-targeted drugs [60]. The inhibition of eosinophil degranulation has demonstrated promising results in animal models of NMO. The histamine H1 receptors antagonists have been shown to influence eosinophil activity. Cetirizine, a selective antagonist of the H1 receptor, was administered orally before and during AQP4-Ab intracerebral injection and significantly reduced eosinophil infiltrates and lesion formation in mice. These results prompted the testing of cetirizine in a pilot, open-label, add-on trial to standard therapy for 16 NMO patients. Cetirizine was administered at a dose of 10 mg daily. After one year of treatment, ARR was reduced fourfold in these patients [61]. Thus, antihistaminic drugs affecting eosinophilic function might be beneficial as an add-on therapy in NMO treatment.

Another future strategy for NMO treatment might be associated with an attempt to enhance apoptotic death of plasma cells. This strategy might reduce the number of cells producing pathogenic AQP4-Ab and demonstrate benefit for NMO patients.
Bortezomib is a selective inhibitor of the 26S proteasome subunit leading to enhanced cell death. Bortezomib has been tested in an open-label study including five NMOSD AQP4-Ab-positive patients. All patients were refractory to previous therapies, and two of them were resistant to RTX. Four of these five patients remained stable or improved within 12 months of the study. Patients treated with bortezomib had a lower plasma cell count and diminished levels of serum AQP4-Ab [62].

Conclusions

Recent years have witnessed unprecedented progress in the understanding and treatment of NMOSD (Tab. 2). 2019 has been called the Year of NMOSD. The discovery of the pathogenic role of AQP4-Ab and anti-MOG Ab has allowed the targeting of the basic immune mechanism of this disease.

The exciting findings of the recent randomised and controlled trials have provided a realistic hope that the era of unproved therapies in NMOSD will come to an end.

The findings from studies with novel drugs targeting complement activation, interfering with IL6-R activation and depleting antibody-producing plasma cells, should soon bring about a new and effective treatment of this devastating disease. These results are very welcome for NMOSD patients who have been relegated to off-label therapies with uncertain benefits for many years. The approval of new therapies for NMOSD will undoubtedly prove to be another breakthrough in modern neurology.

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