Continued dysregulation of the B cell lineage promotes multiple sclerosis activity despite disease modifying therapies

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Abstract
A clear understanding of the origin and role of the different subtypes of the B cell lineage involved in the activity or remission of multiple sclerosis (MS) is important for the treatment and follow-up of patients living with this disease. B cells, however, are dynamic and can play an anti-inflammatory or pro-inflammatory role, depending on their milieu. Depletion of B cells has been effective in controlling the progression of MS, but it can have adverse side effects. A better understanding of the role of the B cell subtypes, through the use of surface biomarkers of cellular activity with special attention to the function of memory and other regulatory B cells (Bregs), will be necessary in order to offer specific treatments without inducing undesirable effects.

Keywords
Multiple sclerosis, antibody secreting cell, memory B cell, naïve B cell, B regulatory cell

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Introduction
Multiple sclerosis (MS) is a chronic, neuroinflammatory disease of autoimmune origin, which causes demyelination and neurodegeneration of the central nervous system (CNS). It is the leading cause of disability among young adults with neurological diseases. Current MS diagnostic methodologies are based on criteria that include clinical presentation, determination of oligoclonal bands (OCB) and other biomarkers such as the IgG index and IgG synthesis in 24 hours in the cerebrospinal fluid (CSF), as well as presence of inflammatory and/or demyelinating lesions in the magnetic resonance imaging (MRI) analysis. Currently, the follow-up and response to treatment of patients with MS is based on the concept of “no evidence of disease activity” (NEDA) based on clinical presentation, imaging, and disability progression, without taking into account any other biomarkers of disease. Over the last three decades, the prognosis of disease has dramatically improved due to the availability of multiple disease modifying therapies (DMT).

It has been established that both T and B cells play a role in the pathogenesis of MS. The MS-promoting role of B cells can be carried out through the secretion of antibodies such as OCB, presentation of antigens, activation of T cells and/or production of cytokines. Perturbation of T cell homeostasis due to a reduction in cells of thymic origin, and reduction in the diversity of T cell repertoires, among others, has been documented in MS. Antibody secretion is the most studied function in the pathogenesis of MS and, in recent years, exploration of the role of cytokines in the regulation of immunity, for therapeutic purposes, has begun. In addition, the use of anti-CD20 therapies for MS, which do not affect plasmablasts (PB) and plasma cells (PC), has led to a better understanding of the role of B cells in the pathogenesis of the disease. Although DMT have the ability to slow down the progression of the disease, they are not a cure. In the present article, we show how the dysregulation of the B cell lineage could be strongly linked to the activity and progression of the disease and how selective therapy, guided by cell surface markers, could become key in controlling it.

B cell lineage
The family of B cell subsets results from an evolutionary process of embryonic cells expressing different surface markers (especially CD19, CD20, and CD38) through their lifespan, in different organs, until they reach the state of antibody secreting cells (ASC), thus culminating the evolution process with the presence of effector B cells. The change of stage from membrane-linked antibody cell to ASC represents the terminal differentiation toward B cells that do not proliferate (Figure 1). The B cell lineage begins in fetal life from pluripotent hematopoietic stem cells (SC), located in the fetal liver and in the postnatal bone marrow (BM). Henceforth, they evolve into multipotent myeloid/lymphoid progenitors (MPP), which continue their evolution towards the common lymphoid progenitors (CLP). CLP from the BM evolve to a pro-B state in which they express the CD19 marker and then transform into pre-B cells; those, in addition to expressing CD19, begin to express CD20, and later progress to immature B cells that express IgM as a surface marker. While transiting from the BM to the secondary lymphoid organs (SLO), the B cells express the B cell receptor (BCR) surface markers IgM and IgD, thus evolving into transitional B cells; this step requires a checkpoint that entails clonal deletion and receptor editing before entering the SLO (spleen, lymphoid node, tonsils, and mucosa-associated lymphoid tissue [MALT]) where they become mature naïve B cells. The mature naïve B cells, at this point, can have three possible destinations: a) they enter the marginal zone of the spleen where they may become short-lived plasma cells (SLPC) that produce IgM and rapidly enter apoptosis (since these are B cells involved in rapid and transitory defense); b) move to the intestine and the pulmonary epithelium (B1 cells); c) migrate to splenic follicles and lymphoid nodules, becoming follicular B cells. Naïve B cells that carry the BCR IgD go through early class switch recombination (CSR) in the extrafollicular zone, with support from T cells, then enter the germinal center (GC) where they undergo somatic hypermutation (SHM), after which they express BCR IgG. Subsequently, the resulting memory B cells, PB and PC will have the ability to secret high-affinity antibodies for decades, or for the lifespan of an individual, and most of them will be able to migrate to the bone marrow to establish as long-lived plasma cells (LLPC). Dysregulation of the GC has been associated with autoimmune disease. Evidence suggests that the origin of B cell autoreactivity occurs in the GC due to dysfunction of thymus-derived follicular T helper cells and follicular regulatory T cells. PB may develop from...
any type of activated B cell (including naïve, marginal zone, follicular, and memory B cells), but it is not clear if PB that originated from these cells (except for memory B cells) are competent to mature to LLPC.6 PB will carry the CD19+CD20-CD27++CD38++IgG+/- markers, and will express the chemokine receptor CXCR4, which will help them get attracted to the chemokine CXCL12 in the BM niches. As an alternative, PB expressing the receptor CXCR3 will become LLPC in the spleen and lymph nodes (assisted by the chemokine CXCL12) or in inflamed tissue (assisted by the chemokines CXCL9-CXCL12), and subsequently undergo apoptosis upon resolution of inflammation.7

**Physiopathology of multiple sclerosis**

Memory B cells access the CNS through the disrupted blood brain barrier (BBB). They are identified in perivascular spaces, demyelinating lesions in the brain and spinal cord, and disperse in the meninges where they can form aggregates known as tertiary lymphoid organs (TLO). These TLO emulate GC function, supporting the formation and persistence of cortical lesions.16–19 In addition, they are a local source of class-switch IgG that contribute to the immune process and are subsequently determined as OCB in the CSF of patients with MS.20 In the meninges, the inflammatory infiltrates are composed of CD3+ T cells, CD20+ B lymphocytes and PC.16 In the white matter lesions, the inflammatory infiltrates are localized in the perivascular spaces containing T and B lymphocytes and PC.16 In the diffuse infiltrates and normal-appearing white matter, CD8+ T lymphocytes predominate almost exclusively.16 The presence of PB in CSF has been reported.21,22 Despite the knowledge accumulated to this date, the complete understanding of the evolution of the B cell lineage is still in progress. Activated lymphocytes are able to access the CNS, both in health and disease, through the BBB, the blood meningeal barrier, and the blood-CSF barrier.12 In normal conditions the amount of B cells that access the CNS is very low.23 These cells primarily exit the CNS via lymphatic drainage through nasal blood vessels, or via meningeal lymphoid vessels to the lymphoid cervical nodules.24

Naïve and memory B cells are crucial within the B cell lineage and they have been shown to negatively correlate in their function: increased memory B cells and decreased naïve B cells have been linked with a worsening of the disease, and vice-versa. In fact, when the presence of memory B cells induce the auto-proliferation of CD4 T cells, which tend to home in the brain, naïve B cells are decreased.3 Memory B cells can be heterogeneous, i.e., originate from different cells or express different phenotypes, including class-switched (CD19+CD27+IgM-IgD-) and class-unswitched (CD19+CD27+IgM+IgD-).25 Inhibition of memory B cells prevents relapsing MS.25 In a study in naïve patients with

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**Figure 1. The B cell lineage cycle.** The B cell lineage begins in fetal life from pluripotent hematopoietic stem cells (SC) in the fetal liver and in the postnatal bone marrow developing B cell receptors and migrating to different locations, including peripheral blood, and the secondary lymphoid organs (SLO) where they will acquire, in the germinal center (GC), the ability to recognize antigens and produce highly specific antibodies. In the pathogenesis of multiple sclerosis (MS), these cells may cross the BBB and may be found in the CSF, perivascular spaces (PVS), white matter (WM) demyelinating lesions and in the tertiary lymphoid organs (TLO). Abbreviations: SC: stem cell; SLO: secondary lymphoid organ; CSR: class switch recombination; SHM: somatic hypermutation; GC: germinal center; TLO: tertiary lymphoid organ; PVS: perivascular space; WM: white matter.
Regulatory B cells
Within the B cell lineage, the regulatory B cells subset (Bregs) stands out, since there is still no agreement on its origin and classification. Bregs are not a specific subtype of B cells, but represent a regulatory functional state resulting from inflammation. Although it has been assumed that interleukin 10 (IL10) is the hallmark of Bregs, other factors such as IL35, transforming growth factor (TGF) β, and direct cell-to-cell contact are also mechanisms of Bregs function. Immature B cells, mature B cells, PB and PC are believed to function as Bregs. The Bregs can express the following markers: IL10, CD27, CD5, CD25, CD86, CD24 and CD28. Based on the production of IL10, three important subtypes of regulatory Bregs have been identified, including the transitional (CD19⁺CD24hiCD38hi), naïve (CD19⁺CD24⁺CD38⁻), and memory (CD19⁺CD24hiCD38⁻) subtypes, among which the transitional cells are the main producers of IL10. Transitional B cells are capable of suppressing differentiation of naïve T cells into Th1 and Th17, which are dependent on the co-stimulatory molecules CD80 and CD86. Survival of Bregs is linked to the B cell activating factor (BAFF) and to a proliferation inducing ligand (APRIL). Under normal conditions, the population of Bregs is kept low in order to maintain immune homeostasis. In newborns, 50% of umbilical cord blood B cells correspond to the transitional B cell subtype whereas, in adults, it represents only 4% of the cell population in peripheral blood. It is estimated that, in human peripheral blood, the Bregs subtype represents only 1-2% of all B cells. In an experimental allergic encephalitis (EAE) animal model, it was documented that IL10 contributed to the reduction of the inflammatory response mediated by microglia and astrocytes in the CNS. Bregs transfer reversed the increase in Th1 and Th17 cells in an arthritis model lacking IL10. In mice with low expression of the IL35 subunit p35, or EBi3 in B cells, an inability to recover from EAE was detected, with evidence of activation of macrophages and inflammatory T cells, and an increased activity of B cells such as APC. IL10 regulates the differentiation of the lineage from IL10-secreting B cells to PB that secrete IgG or IgM. In a post-mortem analysis of MS patients with high levels of meningeal inflammation and cortical demyelination, Magliozzi et al., reported an increase in IL10 expression, among other proinflammatory cytokines and molecules related to B cell activity and lymphogenesis in meninges and CSF. Additionally, an increase in IL10 was found in the CSF of MS patients with high cortical involvement at the time of diagnosis. Early development of MS in individuals with the clinically isolated syndrome (CIS), or radiologically isolated syndrome (RIS), seems to correlate with a reduced production of IL10 by B cells. In a cohort of MS patients followed for 10 years, Farian et al. found that patients with positive OCB at diagnosis advanced, more frequently and earlier in the course of the disease, to a progressive phase. This can be explained by the presence of a greater intrathecal inflammatory component that causes greater cortical involvement. Besides, the authors reported an over-expression of inflammatory molecules, including IL10. PB and PC inside the MS lesions presented a high IL10 expression.

Double negative and CD21⁺LOW cells
Another subset of CD19⁺ B cells that could be relevant to MS pathogenesis are the double negative (DN) B cells (IgD⁻CD27⁻) and the CD21⁺LOW cells, which have been associated with aging and autoreactivity. These cells are believed to develop outside the GC, are independent from T-cells, and display a pro-inflammatory cytokine profile. These cells have been found in healthy subjects, and have also been found at higher levels in the CSF of MS patients younger than 60 years when they were compared to age-matched healthy donors (DN B cells 19.5% against 3.03%, and CD21⁺LOW 21.95% against 6.06%, respectively). Most DN B cells display an IgG⁺ phenotype while CD21⁺LOW B cells originate from a heterogeneous population that includes CD27⁻ naïve, CD27⁺ memory, and IgG⁺ and IgM⁺ B cells. Both DN and CD21⁺LOW B cell frequencies were higher in the CSF compared to blood levels for these patients. Fraussen et al. have suggested that the DN B cells may have multiple origins, considering IgG⁺ cells better linked to the class-switched memory B cells, while IgM⁺ cells share more similarity with the naïve and the non-class-switched IgD⁺CD27⁺ memory B cells.

B cells in the CSF compartment in MS
Inflammation of the CNS is reflected in the presence of B cells in the CSF. Cepok et al. evaluated the B cell subtype in CSF in MS patients, finding that the majority of detectable cells were memory B cells (CD19⁺CD27⁺), whereas a minority were naïve B cells (CD19⁺CD27⁻); those were different from naïve B cells that predominated in peripheral blood. In addition, they detected PB (CD19⁺CD27⁺CD138⁻) subtypes representing between 30-50% of cells in CSF, and were present in the course of the disease, without correlation with the level of PB present in peripheral blood, while short lived PB (CD19⁺CD27⁺CD138⁺HLADR⁺) and PC (CD19⁺CD27⁺CD138⁺HLADR⁻) were absent from CSF. In contrast, Corcione et al. reported the predominance of both memory B cells and PC in the CSF of MS
patients without treatment. In patients with RRMS and primary progressive MS (PPMS) with positive B cells for G1m1 (IgG1 heavy chain gene), Lossius et al. detected IgG1 ASC with a phenotype compatible with highly proliferating PB (CD19dimCD27hiCD38hi) and with high expression of CD138+, HLA-DR+ and Ki67+ in CSF. In pediatric MS, an increase in memory B cells in CSF, with a predominance of non-switched memory B cells and PB, was found, while in adults with MS, class-switched memory B cells and PC predominated in CSF during relapses of MS. Using a deep repertoire sequencing of IgG heavy chain variable genes (IgG-Vh) in paired CSF and peripheral blood from patients with MS, Von Budingen et al. found that there was a cluster of clonally related B cells involved in a bidirectional cell exchange across the BBB, with some of them being present primarily in the CNS while others were present in the periphery or in both compartments. Additionally, using the same protocol, they found evidence of clonally related B cell receptors in a patient’s blood and CSF, after seven years of therapy with rituximab, indicating a prolonged presence in this compartment during the disease span due to recirculating memory B cells or LLPC. Greenfield et al. found that clonally related B cells were present as class-switched IgG and CD27+ in the CSF of patients with MS, leading to the conclusion that, despite having been on DMT, there were complex patterns of persistence of clonal B cells in CSF and blood. A significant depletion of CD20+ B cells has been detected in the blood, with a partial and transient depletion in the CSF and the CNS perivascular spaces, after therapy with rituximab. Table 1 presents a summary of the different surface markers that characterize the B cell lineage through its lifespan.

### Table 1. B-cell subsets surface markers and corresponding compartments in MS. Up-to-date reported B-cell subtypes with reference to the B-cell lineage in MS. Sub-types from other inflammatory conditions are also mentioned. Abbreviations: SLO: secondary lymphoid organ; GC: germinal center; BM: bone marrow; ASC: antibody secreting cells; LLPC: long lived plasma cells. ‘Compartment’ makes reference to the organ where the biomarkers were found.

| B-cell subtype                  | Surface biomarkers                  | Compartment | Reference |
|--------------------------------|-------------------------------------|-------------|-----------|
| Stem cell                      | CD34, HLA-DR                        | BM          | 12        |
| Pro-B cell                     | CD19+CD34+IgM                       | BM          | 20        |
| Pre-B cell                     | HLA-DR, CD19, CD20, Pre-BCR         | BM          | 12        |
| Immature B cell                | CD19+CD34+IgM                       | BM          | 20        |
| Transitional B cells           | CD19+CD27+CD38hiCD24hi             | Blood       | 27        |
|                               | HLA-DR, CD19, CD20, IgM, IgD, CD38  | Blood       | 12        |
|                               | CD19+IgD+CD27+CD38+                | Spleen/blood| 20        |
|                               | CD19+CD38+CD24+                     | Spleen/blood| 20        |
|                               | CD24highCD38high                    | Blood, SLO  | 85        |
|                               | CD38+CD10+IgD+                      | Blood       | 88        |
| Naive B cell                   | HLA-DR, CD19, CD20, IgM, IgD        | SLO/Blood   | 12        |
|                               | CD19+CD27                           | CSF         | 21        |
|                               | CD19+CD27+IgD                       | Blood       | 27        |
|                               | CD19+CD27+IgD+                      | Spleen/blood| 20        |
|                               | CD19+CD27+IgD+                      | CSF         | 97        |
|                               | CD27+IgD+                           | Blood       | 88        |
| Non-class-switched memory B cell| CD19+IgD+CD27+                      | Spleen/blood| 20        |
| Pre-class-switched memory B cell| CD27+IgD+                           | Blood       | 88        |
| Memory B cell                  | CD19+CD27+IgD                       | Blood       | 27        |
|                               | HLA-DR, CD19, CD20, CD27            | Blood, SLO  | 12        |
|                               | CD19+CD27                           | CSF         | 21        |
|                               | CD19+CD27+CD80+CD86+               | CSF         | 97        |
| Centroblast                    | CD19+CD38+CD77+Ki67+Bcl-2+          | CSF         | 97        |
| Centrocyte                     | CD19+CD38+CD77                      | CSF         | 97        |
Mapping of cell markers is essential to evaluate treatment response

Advances in immunotherapy have made it possible to limit the presence and expansion of B cells, thus reducing relapses and the progression of disability. However, the determination of which cells from the lineage could be responsible for clinical deterioration, or improvement, is still to be investigated. In the treatment of other autoimmune diseases, such as pemphigus, the mapping of cell markers has been used to evaluate the response to treatment, finding alterations in the function of CD19+CD24hiCD38hi Bregs cells, which are present in significantly higher numbers in patients in an active state compared to patients in the remitting state of pemphigus.46 Late antibody-mediated rejection continues to be a problem for patients undergoing kidney transplantation and, for many years, it was believed that tolerance and rejection of transplantation were mediated by T cells.47 However, it was recently shown that a population of Bregs may be playing a deleterious role in transplant immunity, and be responsible for the production of alloantibodies.48 Recently, B cells (CD19+CD24hiCD38hi) dysfunction has been reported in peripheral blood, with decreased production of IL10 in patients with RRMS compared to healthy subjects.49 Later antibody-mediated rejection continues to be a problem for patients undergoing kidney transplantation and, for many years, it was believed that tolerance and rejection of transplantation were mediated by T cells.47 However, it was recently shown that a population of Bregs may be playing a deleterious role in transplant immunity, and be responsible for the production of alloantibodies.48

Available therapies affecting the B cell lineage

a. Interferon B (IFNβ) acts in the periphery, inducing apoptosis of CD27+ memory B cells through a mechanism that requires FAS receptor/transmembrane activator and calcium-modulating cyclophilin ligand interactor (TACI) signaling, leading to a specific depletion of these memory B cells (which carry the ability to induce the production of autoantibodies).28

Table 1. Continued

| B-cell subtype                          | Surface biomarkers                                                                 | Compartment | Reference |
|----------------------------------------|------------------------------------------------------------------------------------|-------------|-----------|
| Post-class switched memory B cell      | CD27*IgD*                                                                          | Blood       | 88        |
| Class-switched Memory B cell           | CD19*IgD-CD27**                                                                   | Spleen/blood| 20        |
| Short lived plasmablast               | CD19*CD27**CD138*CD38*HLA-DR**                                                    | CSF         | 21        |
| Plasmablasts                           | HLA-DR, CD19, CD20 low, CD27, CD38                                                | SLO/blood   | 12        |
|                                        | CD19*CD27*CD38*                                                                   | Blood       | 27        |
|                                        | CD38*CD27*IgD*                                                                    | Blood       | 88        |
|                                        | CD19*CD27*CD38*                                                                   | Spleen/blood| 20        |
|                                        | CD19*CD138**                                                                      | Spleen/blood| 20        |
|                                        | CD19*CD27**CD138*CD38**                                                           | CSF         | 21        |
|                                        | CD19dimCD24hiCD38*CD138HLA-DR KI67                                               | CSF, blood  | 40        |
| Plasma cells                           | CD19*CD27*CD138*CD38                                                              | CSF         | 21        |
|                                        | CD19 low, CD27, CD38, CD138                                                        | SLO/blood   | 12        |
|                                        | CD38*CD138*                                                                        | Spleen/blood| 20        |
|                                        | CD19 CD138*                                                                        | CSF         | 97        |
|                                        | CD19*CD138*                                                                        | Blood       | 27        |
| Bregs cell                             | IL10, CD27, CD5, CD25, CD86, CD24, CD38                                            | Blood       | 27        |
|                                        | CD24highCD27*                                                                     | Blood, SLO  | 85        |
|                                        | CD19*CD24**CD38**                                                                  | Spleen/blood| 20        |
|                                        | CD19*CD5*CD14**                                                                    | Spleen/blood| 20        |
| Double Negative B cell                 | CD27IgD*                                                                           | Blood       | 88        |
| LLPC                                   | CD19IgD CD27CD138CD38hi                                                           | BM          | 103       |
|                                        | CD19 CD138CD38hi                                                                   | BM          | 103       |
| ASC in CSF                             | CD19*CD27*CD38*IgG                                                                  | CSF         | 40        |
| B cells in CSF                         | CD19*CD138                                                                        | CSF         | 21        |
|                                        | CD19dimCD24hiCD138ASC                                                             | CSF         | 40        |
to harbor EBV) and an increase in the CD27+ cell subtype that contains naïve B cells secreting IL10.52 Furthermore, it was observed that memory B cell depletion was accompanied by a reduction in EBV markers.53 IFNβ leads to the inhibition of leukocyte proliferation and antigen presentation.53 It also changes the cytokine profile towards an anti-inflammatory profile in both peripheral blood and the CNS, and reduces T cell migration by inhibiting the activity of the T cell matrix proteinase.53 IFNβ increases naïve B cells and decreases memory B cells in peripheral blood.54 Ersöz et al. showed that IFNβ induces the production of high amounts of IL10 compared to therapy with azathioprine in patients with RRMS.55 A meta-analysis study in patients with MS who received IFNβ showed that there was a lower proportion of Th17 cells in the peripheral CD4+ T cell pool and a reduction of IL17 and IL23 levels in serum.56

b. **Fingolimod** is a sphingosine-1-phosphate (S1P) modulator that binds to the S1P receptor on lymphocytes, and retains naïve B cells and central memory B cells in lymphoid nodes.57 B cell subsets in the periphery are susceptible to being modified by fingolimod, thus leading to a reduction of memory B cells and an increase in the number of transitional B cells and Bregs in the periphery,58 with an associated increase in the production of IL10.59 Treatment with fingolimod has also been associated with a reduction in the lymphocyte count in peripheral blood and with an increase in the percentage of naïve B cells.59 Fingolimod also causes an increase in DN B cells.54 Fingolimod does not affect the exchange of B cells through the BBB, but it affects the intrathecal clonal expansion, thus inhibiting the activity of the GC.50

c. **Dimethyl fumarate (DMF)** causes long-term lymphopenia through its effect on two genes: it induces NFR2 (antioxidant effect) and inhibits NFκB which, in turn, induces a change from the Th1 to the Th2 subtype.57 DMF increases the ratio of naïve B cells to memory B cells and increases the number of transitional and IL10 producing B cells.57 DMF increases the percentage of naïve B cells, with a relative reduction in memory B cells and DN B cells.54

d. **Teriflunomide** is a drug that inhibits the dihydroorotate dehydrogenase, thus interfering with the biosynthesis of pyrimidines and leading to a reduced cell proliferation. It can significantly reduce Bregs (CD24+CD38high), mature B cells (CD24+CD38low) and, to a lesser extent, memory B cells (CD24+CD38−) in the peripheral blood of patients with RRMS.54 Yilmaz et al. reported a reduction of PC in the peripheral blood of patients with RRMS, who were treated with teriflunomide.52

e. **Natalizumab** blocks the entry of T cells (mainly CD4) into the CNS by neutralizing VLD4 or α4β1 integrins.57 Natalizumab in peripheral blood lowers PB and increases memory B cells.54 Kemmerer et al. reported an insignificant increase in the number of B cells and memory B cells in patients treated with natalizumab.54 In addition, PB were reduced due to a mechanism of natalizumab that alters their traffic through the BBB. Natalizumab decreases the exchange of peripheral and intrathecal B cells, but does not modify their intrathecal clonal expansion and can induce a reduction of OCB production in some cases.50,53 Traub et al. found that natalizumab promotes the activation and proinflammatory differentiation of peripheral B cells in MS.54

f. Although **glatiramer acetate (GA)** is a compound affecting T cells, no effect in the maturation and differentiation of B cells has been detected.54 GA interferes with antigen presentation and promotes switching from the pro-inflammatory Th1 state to an anti-inflammatory Th2 state, on top of inducing CD8+ T reg cell production.57 The pro-inflammatory pattern, mediated by the secretion of IL6 by peripheral B cells, has been shown to abate and switch to a pattern mediated by IL10-secreting Bregs in MS patients treated with GA.56 However, other studies on the efficacy of GA on B cells in patients with RRMS have reported a reduction in the total numbers of B cells, PB and memory B cells in peripheral blood.54,55,56 By reducing the expression of intracellular adhesion molecule (ICAM-3), GA contributes to reducing the migration of B cells to the CNS.20

g. **Rituximab** blocks the CD20 receptor, thus removing pathogenic B cells.57 Although rituximab depletes naïve and memory B cells in the circulation and is not as effective in depleting B cells in tissues, effector and regulatory cells are balanced during cell repopulation after therapy.57 Palanichamy et al. found that rituximab induced depletion of memory B cells in blood for up to 12 months.57 A depletion of T cells by more than 50% and B cells by 95%, in CSF, has also been reported after treatment with rituximab in patients with RRMS.58 Using the surface B cell marker CD21 in patients with secondary progressive MS, who received IV rituximab on days 0 and 15, and intrathecal rituximab on day 0, six weeks and twelve months later, Komori et al. found a significative reduction in CD21 expression in the serum of patients, suggesting a
complete and lasting depletion of B cells, as opposed to an insignificant change in CSF corresponding to an incomplete and transitory depletion of B cells in the CNS compartment. In patients with neuromyelitis optica (NMO) seropositive for aquaporin-4, treatment with rituximab was followed by no relapse while their memory B cells were below 0.05% in peripheral blood. Hausler et al., working on a model of EAE induced by myelin oligodendrocyte glycoprotein (MOG) and another model in naïve mice observed, after treatment with the murine surrogate of rituximab, a persistence of mature B cells in the spleen; an early reconstitution of B cells in the bone marrow and in the spleen before being released into the periphery; and a presence of reactive B cells against myelin when the model included activation of B cells. Altogether, these findings suggest that pathogenic B cells were able to persist despite an anti CD20 treatment. In addition, they reported a fast depletion of B cells in the peripheral blood, which upon discontinuation of treatment, began to repopulate, proving that cells have different sensitivities to therapy with anti CD20.

h. Ocrelizumab is a humanized monoclonal antibody version of rituximab capable of causing more severe CD19+ cell depletion than rituximab in patients with rheumatoid arthritis. Ocrelizumab is also associated with a very long therapeutic effect, up to 22 months, after the last dose as demonstrated by the RRMS clinical trials OPERA I and OPERA II. Recent studies have shown that patients who received treatment with rituximab, or ocrelizumab, for RRMS and NMO for several years, developed hypogammaglobulinemia or a defective recovery of B cells, which could be asymptomatic or could present with bacterial infections or recurrent viral diseases. The duration of hypogammaglobulinemia fluctuated between one month and eleven years. Marcinno et al. recommended that, in patients who receive anti CD20 therapy, the serum levels of IgA, IgG, and IgM should be determined before initiation of treatment, and repeated yearly with special attention to patients who present a drop in IgG and IgM early in the course of therapy, and who should receive protection against tetanus.

i. Alemtuzumab depletes the CD52 marker in B and T cells with very long periods of CD4 T cell depletion. Alemtuzumab is able to deplete 70 to 95% of CD4 T cells in active relapsing MS. During the reconstitution of B cells after treatment with alemtuzumab, there is a predominance of immature transitional cells, which is followed by a predominance of mature naïve B cells, accompanied by an increase in BAFF, while the reappearance of memory B cells is slow. Mohn et al. reported a change in the distribution of B cells toward a B cell-naïve phenotype in MS patients treated with alemtuzumab, observing negativization of OCB in two patients. The adverse effect of alemtuzumab, including autoimmune disease of the thyroid gland, kidney, platelets and lungs, are well known and correlate with the early recovery of the B cell population with persistence of CD4 T cell depletion, especially during the first year of therapy.

j. Atacicept binds to BAFF and to APRIL, blocking the maturation, differentiation and survival of B lymphocytes. Atacicept depletes transitional and naïve B cells, PB and PC, and IL10-producing B regs. Atacicept causes B cell depletion without affecting progenitor cells (pre- and pro-B cells) and memory B cells. Treatment of MS patients with atacicept, unexpectedly, induced more relapses in the ATAMS trial.

k. Cladribine is a chlorinated deoxyadenosine analog, partially resistant to adenosine deaminase. The role of cladribine as an immune reconstitution therapy (IRT) has been proven by its prolonged depleting effect on CD4+ T and B lymphocytes in the periphery. Cladribine has the ability to reduce class-switched and unswitched memory B cells to a level comparable to that seen in therapy with alemtuzumab.

l. Inebilizumab is an anti-CD19+ B cell drug with the ability to deplete the B cell lineage from pro-B cell to PC stage, which was recently reported to induce rapid depletion of B cells and PC in MS patients in a phase I study. A new generation of anti-CD20 therapies capable of depleting B cells in the resident organ is under development, including obinutuzumab, although it has not been tested in the treatment of MS yet.

m. Human immunoglobulin G (IVIg) acts on steady-state B cells, inhibiting the homeostatic proliferation of B cells accompanied by an induction of cell aggregation.

n. Autologous haematopoietic stem cell transplantation (AHSCT) is another alternative treatment that achieves a therapeutic effect by depleting all lymphocytic cell population involved in MS; however, its efficacy depends on the type of cell ablation used, since, as described by Hausler et al. while reporting an animal model of EAE, the reconstitution of B cells after anti-CD20 therapy stems from the B cell population that has survived in the bone marrow and spleen. An analysis of peripheral blood lymphocyte reconstitution after AHSCT, with high-dose immunosuppressive therapy in patients with RRMS followed for two...
years, disclosed a greater progressive expansion of the population of naïve B cells in the first and second year post-transplant. At one month, patients with systemic sclerosis who underwent AHSCT had a transient increase in transitional B cells and PB with an increase in the percentage of naïve B cells up to 14 months; their cytokine profile also changed in the long term, increasing IL10 secretion. The B cell compartment also showed decreased percentages of pre- and post-switch memory, as well as DN B cells. In a study conducted by Niederhäusern et al, memory B-cell subpopulations were found to recover slowly and to remain below normal levels with a reduced repertoire diversity one year after AHSCT. A successful shifting of B-cell populations, from a predominantly transitional to a naïve immune phenotype, was seen as cells began to recover within 3 months and increased above normal levels 12 months after AHSCT. Memory B-cells that had survived DMT were mainly class switched and persisted in the early posttransplant stage suggesting that not only plasma cells (PC) but also memory subsets had survived conditioning. The patients in the study received AHSCT with drugs of the BEAM regimen (BCNU, melphalan, and to a limited extent also etoposide and cytarabine) that can penetrate the CNS compartment, but it remained unknown whether tissue-resident B cells were affected by the conditioning regimen of the transplant.

Bruton tyrosine kinase (BTK) inhibitors appear promising as potential therapeutic agents, since evobrutinib has previously been shown to prevent the activation of B cells and improves the clinical course in EAE.

Discussion
The origin of MS still remains enigmatic, although different animal models of EAE have been developed, emulating a peripheral attack compromising the CNS, or an intrinsic CNS pathology process with effect in the peripheral blood. Sabatino et al. have suggested that the paradigm of autoimmune reaction occurring within the CNS may coexist with the outside-in paradigm. Either way, B and T cells are interdependent in the pathogenesis of MS. Inside the brain, the TLO found in the meninges are the driving force of the autoimmune pathogenic process. It has been proposed that previous EBV infection, vitamin D deficiency, and/or a genetic substrate may be the initiators or determinants of the disease process in MS. The B cell lineage plays a crucial role in the pathogenesis of MS and remains active during the course of the disease, in the periphery and CNS, and an aggressive depletion with current therapies can only control the clinical activity and slow down the progression toward disability. Deciphering the intricate variety of phenotypes and the role of the different B cell subsets in MS would be paramount for a complete understanding of this disease.

The acknowledgement of the role of B cell subsets in the presentation of several inflammatory diseases has stemmed from observations in autoimmune conditions such as rheumatoid arthritis, end-stage renal disease secondary to nephritis, bullous pemphigus, and granulomatosis with polyangiitis. Patients with end-stage kidney disease, who have an increase in transitional B cells and Bregs in the blood before transplant, and who present a significant reduction in post-transplant Bregs, are more likely to suffer acute and chronic rejection. Although Bregs appear as anti-inflammatory cells, there is evidence that they may play a pro-inflammatory role in certain pathologies. In another study with bullous pemphigus, Liu et al. confirmed that identifying the role of each cell subtype in the pathophysiology of the disease is crucial. In the same study, a dysfunction of Bregs exhibiting a pro-inflammatory phenotype was observed to contribute to the production of autoantibodies.

In MS, it has been documented that memory B cells can lead to an exacerbation of RRMS through the activation of T cells in the periphery. Furthermore, the fact that memory B cell numbers are decreased under the action of various DMT, and the fact that they were not found to be eliminated by atacicept, confirms their pathogenic role. Most of the immunomodulatory therapies currently available generally induce a reduction in memory B cells and an increase in naïve B cells in peripheral blood, which translates into clinical improvement. In contrast, natalizumab blocks the passage of B cells, mainly memory B cells, through the BBB, increasing their number in peripheral blood. The effect of B cell intrathecally depleting agents is not fully understood.

In relation to Bregs, Matsushita et al. observed that depletion of B cells, before the induction of an EAE model, exacerbated the severity of the pathology, due to the depletion of the Bregs population and its suppressive capacity; in contrast, depletion during the acute phase decreased symptoms by affecting the effector cells, which prevented the activation of CD4+ T cells. Identifying the subtypes of B cells which may be responsible for the inflammatory process in MS, in the periphery and in the CNS, is essential to achieve a selective and timely intervention in order to modulate or neutralize their function and to avoid disease progression, without interfering with the functions of immune surveillance and decreasing the anti-inflammatory response of Bregs. The ability of some cells of the B lineage to transform into Bregs, counteracting inflammation through the production of IL10, is remarkable and warrants to be considered for the development of better therapeutic strategies. Several studies conducted on patients who received kidney transplantation...
and patients with other autoimmune diseases, have shown that treatment with anti-CD20 is effective in the restoration of the balance between effector B cell and Bregs, and that the repopulation of B cells might predict a clinical relapse.98

Current consensus dictates that early initiation of therapy in patients with MS leads to a better prognosis. However, a common dilemma in the MS clinics entails deciding when patients with CIS should start treatment. It is usually considered that CIS patients with high risk factors such as presence of OCB, uptake lesions on MRI, and marked severity of the clinical episode are most likely to evolve to clinically definitive MS or RRMS. Another dilemma is observed in patients with CIS who have been started on DMT, based on risk factors, but who, after four or five years of follow up, do not display evidence of disease activity yet.99 Mapping B cell subtypes in peripheral blood and CSF could be considered as an additional tool to determine alterations in the B cell lineage, which could be suggestive of disease activity in these subjects. Novel diagnostic and prognostic biomarkers in MS, including determination of the kappa free light chain (KFLC) and KFLC index in CSF, have shown to be promising tools for diagnosis, especially taking into consideration that, in clinical practice, KFLC index might replace OCB as a first line biomarker of disease and it has been shown to predict the evolution from CIS to RRMS.100 Another biomarker, chemokine (C-X-C motif) ligand 13 (CXCL13) may be used in the diagnosis of MS in clinical practice for assessment of drug treatment response and disease progression.95,100

We still believe that the meningeal TLO works as an operation center with the ability to magnify an auto-immune response by maintaining antibody diversity, B cell differentiation isotype switching, oligoclonal expansion and local production of autoreactive PC.101 However, recent studies with intrathecal rituximab have shown an inadequate effect in progressive MS and failed to show an early effect, with persistence of markers of inflammation in CSF and leptomeningeal enhancement, in PPMS.69,102 Factors involved in a decrement of CNS efficacy of intrathecal rituximab include a decreased complement-dependent cytotoxicity (due to a low complement concentration in the CSF), a decreased antibody-dependent cytotoxicity (due to a lower proportion of CD56dimNK cells) and a poor bioavailability of rituximab for the B cells embedded in the CNS due to the dynamics of the CSF flow from the lumbar cistern to the arachnoid granulations.69

The main goal of this review entailed the summary of the B cell lineage diversity, following the transformation that cells undergo in each specialization stage allowing them to fulfill distinct roles in their attack on the CNS. Simultaneously, it raises the need to give a directed treatment that could improve drug delivery in the CNS, and a more ingenious monitoring of individual responses to therapies, in order to personalize treatment protocols. Finally, the complexity of the function of LLPC and the extraordinary role that they play in the B cell lineage require further investigation, as well as a deeper review of the contemporary medical literature.103

Conclusion
It is becoming evident that a better identification of the role of different B cell subsets, in the periphery and CNS during the lifespan of MS, will be of paramount significance for the understanding of the pathogenesis of the disease. Specifically, a careful evaluation of the expression of surface markers of transitional, naïve, memory B cells and Bregs, in blood and/or CSF, could contribute to a prompt identification of patients who are not responding to therapy and who may be susceptible to undergo relapses and disease progression. A better understanding of the role of these cell subsets would be useful for engineering intelligent cell therapies that, hopefully, may permit a better control of the disease in the future. This approach would encourage us to rethink the current therapeutic strategy in order to improve the prognosis and quality of life of patients with MS.

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Maria Teresa Cencioni
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I have read the revised manuscript and I am satisfied with the revision. The authors have revised the manuscript addressing my comments and I agree with the indexing.

Competing Interests: No competing interests were disclosed.

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.

Version 2

Reviewer Report 11 July 2023

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The immune system plays an essential role in the pathogenesis of multiple sclerosis. Recent publications reporting the outcomes from clinical trials and DMT have contributed to improving our understanding of the role of B and T cells in MS even though the mechanism underlying the
disease is still unknown. Although the relevance of the topic, I have some concerns about the originality of the review as is similar in contents to other reviews published in other journals.

Minor revision:

1. It is not clear in which type of MS relapsing-remitting or progressive this type of investigation can improve the disease treatments. The generation of ectopic follicles is associated with a secondary progressive disease where the inflammation compartmentalizes in the leptomeninges. In this stage, inflammation fuels the neurodegeneration processes and treatment-depleting B cells can reduce inflammation but not block the progression of the disease. I would suggest introducing a paragraph regarding the pathology of MS and clarifying at which stage of the disease this investigation can help in selecting a personalised treatment in patients with MS.

2. As the therapies affect the B cell lineage, I would suggest adding also studies comparing the effectiveness of different DMTs and the modulation of the B cell compartment, if it has been investigated. If not, it could be suggested in the discussion as further investigations.

3. In addition, it would be good to also discuss the effectiveness of each treatment in terms of disease activity and progression of disease in MS patients.

4. (The second sentence in the introduction) Could you please explain which other biomarkers in the cerebrospinal fluid have been used for MS diagnosis? Which of them are specific to B cells and in which stage of MS disease? Please add references.

5. Autologous haematopoietic stem cell transplantation: please add the publication.

Von Niederhäusern V et al.  B-cell reconstitution after autologous haematopoietic stem cell transplantation in multiple sclerosis. Neurol Neuroimmunol Neuroinflamm. 2022.

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Is the topic of the review discussed comprehensively in the context of the current literature?
Yes

Are all factual statements correct and adequately supported by citations?
Yes

Is the review written in accessible language?
Yes

Are the conclusions drawn appropriate in the context of the current research literature?
Yes
Competing Interests: No competing interests were disclosed.

Reviewer Expertise: Immunology and neurology

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard, however I have significant reservations, as outlined above.

Author Response 27 Jul 2023

Carlos Mora

1. We thank Dr. Cencioni, manuscript reviewer, for her valuable comments to the second version of this article. Although our review, at first glance, might give the impression that it establishes a parallel with other reviews on the topic of the role of the B-cell lineage in multiple sclerosis, published in other journals, we believe that this review brings specific recognition to the role of the regulatory B cells including naive, transitional and memory B cells, as well as the role of double negative B cells and CD21 low cells, in the pathogenesis of multiple sclerosis. A better understanding of the role of these cells will be necessary to establish biomarkers for diagnosis, follow-up and prognosis in relapsing-remitting multiple sclerosis (RRMS) and in the chronic progressive forms of the disease.

2. We agree with the reviewer on the fact that treatment–depleting B cells can reduce inflammation, but they cannot block the progression of the disease. However, in the section titled ‘Available therapies affecting the B-cell lineage’ we also described the effects that at least six of the well-known disease modifying therapies, approved for the treatment of relapsing-remitting multiple sclerosis, have on the B-cell subtypes, on top of the effects that the aggressive B-cell depleting drugs (such as rituximab, ocrelizumab, alemtuzumab, inebilizumab and cladribine) have on the B cell lineage subtypes. Certainly, more studies are needed to understand the B-cell dysregulation in MS, so that we can decipher the mechanisms to control the deleterious effect that B cells exert on the CNS once they establish themselves at the ectopic follicles in the tertiary lymphoid organs.

3. We modified the first paragraph of the section titled ‘B-cell lineage’ by separating its final segment and creating a new paragraph titled ‘Physiopathology of multiple sclerosis’ since we believe the content of this segment is more akin to be displayed under that sub-title.

4. Following the request of the reviewer, and in order to provide clarification to the content of the second sentence in the introduction paragraph, we modified its content as follows: ‘Current MS diagnostic methodologies are based on criteria that include clinical presentation, determination of oligoclonal bands (OCB) and other biomarkers such as the IgG index and IgG synthesis in 24 hours in the cerebrospinal fluid (CSF), as well as presence of inflammatory and/or demyelinating lesions in the magnetic resonance imaging (MRI) analysis’. 
5. In the ‘Discussion’ section, we are adding a new reference in relation to the presence of biomarkers of disease in the spinal fluid [Zhang et al, reference 100]. Accordingly, the following sentences were added to the text: ‘Novel diagnostic and prognostic biomarkers in MS, including determination of the kappa free light chain (KFLC) and KFLC index in CSF, have shown to be promising tools for diagnosis, especially taking into consideration that, in clinical practice, KFLC index might replace OCB as a first line biomarker of disease and it has been shown to predict the evolution from CIS to RRMS [Zhang et al, reference 100]. Another biomarker, chemokine (C-X-C motif) ligand 13 (CXCL13) may be used in the diagnosis of MS in clinical practice for assessment of drug treatment response and disease progression [references 99-100].’

6. With reference to the effect of autologous hematopoietic stem cell transplantation (AHSCT) on B-cell reconstitution in patients with MS, we expanded our comments and included the reference recommended by the reviewer [von Niederhäusern et al, reference 89] as follows: ‘In a study conducted by Niederhäusern et al, memory B-cell subpopulations were found to recover slowly and to remain below normal levels with a reduced repertoire diversity one year after AHSCT. A successful shifting of B-cell populations, from a predominantly transitional to a naive immune phenotype, was seen as cells began to recover within 3 months and increased above normal levels 12 months after AHSCT. Memory B-cells that had survived DMT were mainly class switched and persisted in the early posttransplant stage suggesting that not only plasma cells (PC) but also memory subsets had survived conditioning. The patients in the study received AHSCT with drugs of the BEAM regimen (BCNU, melphalan, and to a limited extent also etoposide and cytarabine) that can penetrate the CNS compartment, but it remained unknown whether tissue-resident B cells were affected by the conditioning regimen of the transplant [reference 89].’

7. Table 1: The subtitle for Table 1 was modified so that the table is presented as ‘B cell subsets surface markers and corresponding compartments in MS.’ We believe this addition will encourage the readers to match reported B-cell subtypes with the mentioned biomarkers and the compartments where they were found, with the corresponding reference.

**Competing Interests:** We have no competing interests to disclose.
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We agree with the responses given by authors. We regret that the diversity of combined surface markers provided in Table 1 was not commented (we are still not sure that each combination was a different subtype). Although we are not convinced by the conclusion stating that identification of B-cell subtypes will help to predict drug/relapse/progression, the article is well written and provides a large review concerning B cell roles in MS.

Is the topic of the review discussed comprehensively in the context of the current literature?
Yes

Are all factual statements correct and adequately supported by citations?
Yes

Is the review written in accessible language?
Yes

Are the conclusions drawn appropriate in the context of the current research literature?
Yes

Competing Interests: No competing interests were disclosed.

Reviewer Expertise: MS, NMOSD

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.

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This review article deals with the variety and roles of B cells subtypes in MS pathology. Authors tried to engulf diverse data concerning B cell lineage, in general and throughout compartments and MS stages. They finally examined how B cell lineages are affected by disease-modifying
treatments (DMT).

Although this review provides a general overview of the literature concerning B cell lineage in MS, important points were not expanded. Tertiary lymphoid organs (TLO), which drive cortical pathology and prognosis are shortly described. Mutual exchange of cells across the BBB, local heterogeneity of B cell clusters in the brain, location of affinity maturation are too shortly described.

Our main criticism is nested in figure 2: how consistent and reproducible are data obtained concerning B cell lineages according to the variety of definitions used to qualify each state of maturation?

*Figure 1: plasmablasts and plasmocytes are only depicted in CSF. Homing of these cells was also described in brain and TLO (i.e. ref 16). Moreover it is easy to evaluate that rare floating ASC (cf ref 21) cannot account of the amount of intrathecal IgG synthesis (they account in a range of ~1%). CSR may also occur in brain TLO. Lastly, bidirectional exchange of B cells, which is major process (e.g. ref 42), was not emphasized in the figure.

*Sentence: ‘They are identified in perivascular spaces, demyelinating lesions in the brain and spinal cord, and disperse in the meninges where they can form aggregates known as tertiary lymphoid organs (TLO)’. TLO are real and complete lymphoid organs, lacking a conjunctive external capsule and occurring in a non-genetically driven location. Memory B cells are only part of TLO, which are mostly driven by CD3−CD4+CD45+ lymphoid tissue inducer (LTi) cells then stromal cells.

*Sentence: ‘A significant depletion of CD20+ B cells has been detected in the blood, CSF and perivascular spaces in the CNS after therapy with rituximab and ocrelizumab’. Indeed, B-cells are totally depleted from blood after antiCD20 infusion, whereas CSF compartment remains partly and transiently depleted. This is the point: antiCD20 fails to deplete intrathecal compartment from CD20+ cells, possibly due to the lack of effectors (low complement concentration in CSF, rare NK cells). Intrathecal infusion, although increasing bioavailability in CSF, does not add any efficacy.

*Table 2. We did not understood the importance of Table 2, except giving example that stringent definition of cell classes is not as stringent as it could be among authors and papers. This table may suggest that surface markers of B cells in MS could be specific. We do not understand the column ‘compartment’: does it mean that the lineage was found in this compartment? If so, it is worth to also indicate which lineage was NOT found in each compartment.

*In Conclusion. Sentence: ‘Specifically, a careful evaluation of the expression of surface markers of transitional, naïve, memory B cells and Bregs, in blood and/or CSF, could contribute to a prompt identification of patients who are not responding to therapy and who may be susceptible to undergo relapses and disease progression.’ Although a real improvement of CSF FACS availability remains possible, we do not believe that the study of B cells will drive therapeutic opportunities or help to monitor the disease. CSF is especially difficult to obtain and is probably not a target for scheduled biological tests. Moreover, as the authors demonstrated throughout the text, the precise role of each B cell subtype is far from being understood in MS.

Is the topic of the review discussed comprehensively in the context of the current
Author Response 01 Aug 2022

Carlos Mora

We thank Dr. Bonnan (article reviewer) for his thorough comments to the first version of our manuscript. We have reviewed and modified Figure 1 to remark that the natural exchange of B-cells across the BBB is bidirectional, that class switch recombination (CSR) may also occur in the brain TLO, and that plasmablasts and plasma cells figure out in the brain (including the TLO), CSF and peripheral blood. We also labelled the T-cells present in the white matter lesion and in the secondary lymphoid organ (SLO).

In the section titled ‘B cells in the CSF compartment in MS’ the sentence remarked by the reviewer has been modified as follows: ‘A significant depletion of CD20+ B cells has been detected in the blood, with a partial and transient depletion in the CSF and the CNS perivascular spaces, after therapy with rituximab.’

In the penultimate paragraph of the ‘Discussion’ section, we have added two new references to the manuscript, which give more detailed information about the relevance of the tertiary lymph organ (TLO) [Londoño AC and Mora CA. Role of CXCL13 in the formation of the meningeal tertiary lymphoid organ in multiple sclerosis. F1000Research 2018, 7:514 https://doi.org/10.12688/f1000research.14556.3] now reference 99) and about the controversial poor response to therapy with rituximab in the CSF compartment with persistence of B cells (Bonnan M, Ferrari S, Courtade H, Money P, Desblache P, Barroso B, Debeugny S. No Early Effect of Intrathecal Rituximab in Progressive Multiple Sclerosis (EFFRITE Clinical Trial). Mult. Scler. Int. 2021, 2021, 8813498 -now reference 100). This paragraph reads as follows: ‘We still believe that the meningeal TLO works as an operation center with the ability to magnify an auto-immune response by maintaining antibody diversity, B cell differentiation..."
isotype switching, oligoclonal expansion and local production of autoreactive PC. However, recent studies with intrathecal rituximab have shown an inadequate effect in progressive MS and failed to show an early effect, with persistence of markers of inflammation in CSF and leptomeningeal enhancement, in PPMS. Factors involved in a decrement of CNS efficacy of intrathecal rituximab include a decreased complement-dependent cytotoxicity (due to a low complement concentration in the CSF), a decreased antibody-dependent cytotoxicity (due to a lower proportion of CD56dim NK cells) and a poor bioavailability of rituximab for the B cells embedded in the CNS due to the dynamics of the CSF flow from the lumbar cistern to the arachnoid granulations.

The first sentence of the ‘Conclusion’ section has been modified as follows: ‘It is becoming evident that a better identification of the role of different B cell subsets, in the periphery and CNS during the lifespan of MS, will be of paramount significance for the understanding of the pathogenesis of the disease.’

With reference to the comments to Table 1, titled ‘B cell subsets surface markers’, the reviewer is right in his assertion that the motivation for the presentation of this table was the recognition of the significant diversity in the terminology and nomenclature used for the identification of the B cell subtypes in different organs (we used the term ‘compartment’ in the table) described by at least nine different articles cited in our review (references 12, 20, 21, 27, 40, 85, 88, 96 and 101). According to the data presented in the table, we also believe that surface markers of B-cells in MS could be specific. Minor changes were introduced to the table including the meaning of the ‘compartment’ heading and a clarification in the ‘compartment’ column for the ‘centroblast’ and ‘centrocyte’ cell markers (the abbreviation ‘GC’ was replaced by ‘CSF’ in both [reference 96]).

**Competing Interests:** No competing interests.