Deconvolution of B cell receptor repertoire in multiple sclerosis patients revealed a delay in tBreg maturation

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Background: B lymphocytes play a pivotal regulatory role in the development of the immune response. It was previously shown that deficiency in B regulatory cells (Bregs) or a decrease in their anti-inflammatory activity can lead to immunological dysfunctions. However, the exact mechanisms of Bregs development and functioning are only partially resolved. For instance, only a little is known about the structure of their B cell receptor (BCR) repertoires in autoimmune disorders, including multiple sclerosis (MS), a severe neuroinflammatory disease with a yet unknown etiology. Here, we elucidate specific properties of B regulatory cells in MS.

Methods: We performed a prospective study of the transitional Breg (tBreg) subpopulations with the CD19+CD24hiCD38hi phenotype from MS patients and healthy donors by (i) measuring their content during two diverging courses of relapsing-remitting MS: benign multiple sclerosis (BMS) and highly active multiple sclerosis (HAMS); (ii) analyzing BCR repertoires of circulating B cells by high-throughput sequencing; and (iii) measuring the percentage of CD27+ cells in tBregs.

Results: The tBregs from HAMS patients carry the heavy chain with a lower amount of hypermutations than tBregs from healthy donors. The percentage of...
transitional CD24\textsuperscript{high}CD38\textsuperscript{high} B cells is elevated, whereas the frequency of differentiated CD27\textsuperscript{+} cells in this transitional B cell subset was decreased in the MS patients as compared with healthy donors.

**Conclusions:** Impaired maturation of regulatory B cells is associated with MS progression.

**KEYWORDS**
B regulatory cells, BCR, CD19+CD24highCD38high, multiple sclerosis, MS, transitional Breg, TrB, activated memory-like transitional cells

**Introduction**

Multiple sclerosis is a highly heterogeneous severe autoimmune neurodegenerative disorder with an evident inflammation component (1). Despite considerable advances in this field, the mechanism triggering it remains elusive, hindering the development of effective therapeutics (2–8). MS progression is mostly associated with the promotion of the T cell response (9, 10). Yet the contribution of B cells to various autoimmune disorders, including MS, should not be underestimated (11–16). Apart from antibody production and antigen presentation, B cells play a crucial role in regulating the immune response through their antibody-independent effector functions (17, 18).

In 1974, professor James Turk and coauthors suggested that B cells could inhibit inflammation during delayed hypersensitivity reactions (19). Further on, regulatory properties of B cells were confirmed in experimental autoimmune encephalomyelitis (EAE), an animal model of MS (20). The subpopulation of B cells performing regulatory functions were termed Bregs. In mice, the Breg population constitutes up to 5% of total B cells in the spleen and lymph nodes, and their number significantly increases during inflammation development (21–23). In humans, Bregs account for less than 5% of blood B cells (24). Abnormalities in the Breg counts or their function were observed in patients with autoimmune diseases (25–29) and allergies (30).

Different phenotypes of Bregs are described so far. Transitional Bregs (tBreg cells) CD19\textsuperscript{+}CD24\textsuperscript{high}CD38\textsuperscript{high} (24, 26, 29, 31) and memory Bregs CD19\textsuperscript{+}CD24\textsuperscript{high}CD27\textsuperscript{+} are the most studied regulatory B cell subpopulations that modulate the immune response in humans (32, 33). Meanwhile, several other Breg phenotypes are reported: CD19\textsuperscript{+}CD25\textsuperscript{+}CD71\textsuperscript{+}CD73\textsuperscript{−} (34), CD19\textsuperscript{+}CD27\textsuperscript{int}CD38\textsuperscript{+}IgM\textsuperscript{+}IgD\textsuperscript{−} (35), CD19\textsuperscript{+}CD24\textsuperscript{high}CD27\textsuperscript{+}CD3\textsuperscript{+}IgD\textsuperscript{+}IgM\textsuperscript{−}CD1c\textsuperscript{+} (36), CD19\textsuperscript{+}CD5\textsuperscript{+}Foxp3\textsuperscript{+} (37), and CD19\textsuperscript{+}CD38\textsuperscript{+}CD11\textsuperscript{a}IgM\textsuperscript{+}CD147\textsuperscript{−} (38). Transitional CD19\textsuperscript{+}CD24\textsuperscript{high}CD38\textsuperscript{high} Bregs can be found in the peripheral blood of healthy adults representing a minor subset (approximately 4%) of all circulating B cells (29). This tBreg subset was previously shown to produce IL-10, regulate CD4\textsuperscript{+} T cell proliferation/differentiation toward T helper effector cells (40), and contribute to the cytokine imbalance during autoimmune diseases (41).

Since the molecular underpinnings of MS onset and progression were revealed, T cell–mediated immunity was believed to play the leading role in it. However, it is evident now that B cells are crucial for MS pathogenesis as well (42, 43). Autoreactive B cells in MS may produce catalytic antibodies, hydrolyzing myelin basic protein (13, 44, 45), and cause humoral cross-reactivity between myelin and viral antigens (46–50). Still, the existing studies on Bregs functioning in MS are controversial and far from conclusive. The percentages of IL-10–producing Bregs in MS patients are shown to be lower than in healthy controls (28, 51). Other works report either unaltered (52, 53) or even increased (25, 54) Breg numbers during MS progression. A recent study shows no association between the reduced peripheral blood Bregs levels and the Expanded Disability Status Scale (EDSS) score in MS (51).

These inconsistencies most likely arise from several Breg populations coexisting (55, 56). Indeed, these regulatory subpopulations can comprise B cells at different stages of development; therefore, the level of certain BCR hypermutations can be lower at the earlier stages of maturation of these B cells. The specificity of Bregs’ BCRs and pathways that mediate their maturation are still poorly elucidated. It is still not known whether the specificity of Bregs BCRs undergoes alteration in autoimmune disorders.

This study aims to identify the possible abnormalities in the structure of BCRs in the tBregs isolated from peripheral blood of MS patients as compared with HD. Thus, we suggest that analyzing BCR repertoires of tBregs may reveal the disease-related alterations occurring at the early stage of B cell development.
Materials and methods

Patients and healthy donors

Peripheral blood was obtained from the Neuroinfection Department of the Research Center of Neurology, Moscow, Russia. Venous blood was collected in EDTA Vacutainers (BD) from 19 MS patients (nine with BMS (57) and 10 with HAMS) (58) and 16 HD (Table 1). The age of the MS patients ranged between 23 and 70 years old. Their EDSS scores ranged between 1.5 and 8.5. The EDSS values (scored on a scale of 0 to 10) were calculated based on the Kurtzke EDSS scale (59). BMS was diagnosed if the EDSS score was less than 4 for at least 10 years after the disease onset in the absence of

| MS phenotype | Age, years | Gender | EDSS | Treatment | Disease duration, years | BCR repertoire analysis | CD27+ phenotypic analysis |
|--------------|------------|--------|------|-----------|-------------------------|------------------------|-------------------------|
| BMS          | 56 female  | 2.5    | No treatment | 11 | + | – |
| BMS          | 61 female  | 3      | No treatment | 26 | + | – |
| BMS          | 43 female  | 1.5    | No treatment | 12 | + | – |
| BMS          | 36 male    | 2.5    | No treatment | 14 | + | – |
| BMS          | 46 female  | 2      | No treatment | 27 | – | + |
| BMS          | 43 female  | 2.5    | No treatment | 27 | – | + |
| BMS          | 43 male    | 4.0    | No treatment | 18 | – | + |
| BMS          | 58 female  | 3.5    | No treatment | 30 | – | + |
| BMS          | 70 female  | 4.0    | No treatment | 30 | – | + |
| HAMS         | 33 male    | 6      | IFNβ1b (2006-2011; 2014-2017) | 12 | + | – |
| HAMS         | 23 male    | 5      | No treatment | 3 | + | – |
| HAMS         | 37 female  | 5      | IFNβ1b (2014-2016); GA (2016-2017) | 5 | + | – |
| HAMS         | 29 female  | 8      | GA (2012-2014); IVIG (2014); IFNβ1b (2015-2016) | 12 | + | – |
| HAMS         | 39 female  | 8.5    | No treatment | 8 | + | – |
| HAMS         | 22 female  | 4.5    | No treatment | 1  | – | + |
| HAMS         | 46 male    | 6      | IFNβ1b (2019) | 2  | – | + |
| HAMS         | 44 male    | 8.5    | IFNβ1b (2014-2015) | 13 | – | + |
| HAMS         | 24 male    | 4.0    | No treatment | 2 | – | + |
| HAMS         | 44 female  | 4.5    | No treatment | 2 | – | + |
| Healthy      | 24 female  | N/A    | N/A | N/A | + | – |
| Healthy      | 40 female  | N/A    | N/A | N/A | + | – |
| Healthy      | 36 male    | N/A    | N/A | N/A | + | – |
| Healthy      | 27 female  | N/A    | N/A | N/A | + | – |
| Healthy      | 42 female  | N/A    | N/A | N/A | + | – |
| Healthy      | 25 female  | N/A    | N/A | N/A | + | – |
| Healthy      | 39 male    | N/A    | N/A | N/A | – | + |
| Healthy      | 42 female  | N/A    | N/A | N/A | – | + |
| Healthy      | 35 female  | N/A    | N/A | N/A | – | + |
| Healthy      | 51 female  | N/A    | N/A | N/A | – | + |
| Healthy      | 24 female  | N/A    | N/A | N/A | – | + |
| Healthy      | 34 male    | N/A    | N/A | N/A | – | – |
| Healthy      | 24 female  | N/A    | N/A | N/A | – | – |
| Healthy      | 35 male    | N/A    | N/A | N/A | – | + |
| Healthy      | 68 female  | N/A    | N/A | N/A | – | + |
| Healthy      | 47 male    | N/A    | N/A | N/A | – | + |

IFNβ1b, interferon-β-1b; GA, glatiramer acetate; IVIG, intravenous immunoglobulin; HAMS, highly active MS; BMS, benign MS; HD, healthy donors; N/A, not applicable. “+” indicates that the corresponding analysis has been carried out. “-” indicates that the corresponding analysis has not been performed.
treatment; HAMS relapsing-remitting MS was diagnosed based on an EDSS score of 4.0 within five years after the disease onset, poor response to disease-modifying treatments, and two or more relapses with incomplete recovery during one year. None of the patients received glucocorticoid treatment or immunomodulatory treatment for at least six months prior to blood collection. Data on the course of the disease, its duration, and history of administration of disease-modifying treatments are presented in Table 1. HDs (24–68 years old) had neither autoimmune nor oncology diseases nor recent infections. The study was approved by the Local Ethics Committee of the Research Center of Neurology and was conducted in full compliance with the WMA Declaration of Helsinki, ICH GCP, and appropriate local legislation. All patients provided written informed consent for enrollment, followed by a discussion of the study with the investigators.

**FACS sorting of tBreg and total B cell subsets**

Blood samples were diluted two times in PBS with 2 mM EDTA and layered onto Ficoll–Paque Plus (GE Healthcare) and then centrifuged at 900 g for 40 minutes at room temperature. PBMC isolated from MS patients were incubated with ACK lysis buffer for complete removal of red blood cells. Cells were washed with PBS, incubated with α-CD19-PE-Cy7, α-CD24-PE, α-CD38-APC, α-CD27-FITC, and α-CD45-APC-Cy7 antibodies (Biolegend, USA) or α-CD19-PE-Cy7, α-CD24-PE, α-CD38-APC, α-CD45-APC-Cy7, and sytox green dead cell stain (ThermoFisher Scientific) for 60 minutes at +4°C in the dark. Human Fc-blocker (Miltenyi Biotec) was added to all samples before cell staining. B cell subsets were identified by the following markers on their surface: transitional Bregs (CD19+CD24highCD38high), T1 transitional cells (CD19+CD24++CD38++), T2 transitional cells (CD19+CD24++CD38++), memory Bregs (CD19+CD24highCD27+), and memory (CD19+CD24hiCD38loCD27+) or naive (CD19+CD24hiCD38loCD27-) peripheral B cells (60). Following incubation, the cells were washed with PBS and resuspended in PBS. To distinguish T1 and T2 cells by CD24 and CD38 expression, we used gates for flow cytometry analysis, which provided a T2/T1 ratio of approximately 3:1 in HDs (61). Leukocyte and total lymphocyte counts per mm³ were determined using a hematology analyzer (Nihon Kohden MEK-7222, Nihon Kohden, Japan). All samples were analyzed for B cell and tBreg frequencies by flow cytometry. For nine MS patients (four with BMS and five with HAMS) and six healthy donors, a live CD19+ pool of total B cells or CD19+CD24highCD38high tBreg was sorted into two distinct populations (Table 1). The cells were sorted directly into 1.5-mL microcentrifuge tubes containing Qiazol lysis reagent (Qiagen, Germany). Sorting was carried out using a BD FACSAria III, and the data were analyzed using FlowJo software 9.7.5 (TreeStar, Ashland, OR, USA).

**Library preparation for immunoglobulin sequencing (RT-PCR)**

RNA extraction was performed using the RNeasy Mini Kit (Qiagen, Germany) according to the manufacturer’s protocol. Reverse transcription (RT) was performed in 20 µL reaction volume using MMLV RT according to the manufacturer’s protocol (Evrogen, Russian Federation). Multiplex PCR with a modified set of the previously described VH- and VL-specific primers was used for cDNA amplification (62). The primers included 15 human VH-specific forward primers and four human JH-specific reverse primers for the IGH chain, 13 human Vk-forward primers and two Jκ-reverse primers for Vk genes, and 16 human Vλ-forward primers and three Jκ-reverse primers for Vλ genes (Table S1). Each VH, Vk, and Vλ primer pair was added to a separate 50 µL reaction mix with an appropriate equimolar mixture of the four JH, two Jκ, or three Jλ reverse primers. Then, 0.5 ng cDNA was used in each PCR reaction using the Hot Start Taq Master Mix Kit (Evrogen, Russian Federation). The conditions of PCR were as follows: 1 step (94°C—3 min), 1 cycle (94°C—25 s, 62°C—25 s, 72°C—25 s), 2 cycles (94°C—25 s, 60°C—25 s, 72°C—25 s), 2 cycles (94°C—25 s, 58°C—25 s, 72°C—25 s), 3 cycles (94°C—25 s, 56°C—25 s, 72°C—25 s), 3 cycles (94°C—25 s, 54°C—25 s, 72°C—25 s), 30 cycles (94°C—25 s, 52°C—25 s, 72°C—25 s), and 1 step (72°C—4 min). The products of 15 PCR reactions for VH genes, 13 PCR reactions for Vk genes, and 16 PCR reactions for Vλ genes were combined for each chain and concentrated to 50–80 µL using Amicon 30 kDa (Merck, Millipore). The PCR products (~400 bp length) of VH, Vk, and Vλ were loaded on 1.5% agarose gels and purified with the Gel Extraction Kit (Monarch, NEB).

**Deep sequencing of VH and VL genes from individual patients**

Next, 1 pg of the PCR product was ligated with adapters using the NEBNext Ultra DNA Library Prep Kit for Illumina with the NEBNext Multiplex Oligos set (NEB). Libraries were sequenced on Miseq using a 2x300 bp paired-ends sequencing kit (Illumina) in the SB RAS Genomics Core Facility (ICBFM SB RAS, Novosibirsk, Russia).

**Sequencing data processing and repertoire analysis**

MiXCR (63) software was used to extract BCR clonotypes from raw sequencing data. Raw reads were aligned to the standard reference set of V, D (for heavy chain), and J gene-segment sequences. Successfully aligned reads were used for
clonotype sequence assembly with the following parameters: OassemblingFeatures={CDR1Begin : CDR3End} -OmaxBadPointsPercent=0. To normalize the repertoire analysis depth, equal numbers (13,000 for IGVH or IGVK and 7000 for IGVL) of read pairs covering the full target sequence (CDR1+FR2+CDR2+FR3+CDR3) were randomly sampled from each data set. A BCR clonotype is referred to here as a unique nucleotide sequence covering BCR from the beginning of CDR1 to the end of CDR3. Repertoire sequencing results are summarized in Table S2.

Repertoire data analysis was performed using the R programming language (R Core Team (2017) R: the language and environment for statistical computing (R Foundation for Statistical Computing, Vienna, Austria. URL https://www.R-project.org/). Unproductive IGVH/IGVK/IGVL sequences were filtered out before analysis. To characterize the germline identity for each clonotype, a nucleotide sequence covering the CDR1-FR3 part was used for calculating the percentage of identity with the corresponding reference V-segment sequence. The VDJ tools software (64) with the CalcCdrAaStats subroutine encoded by the J-segment (65). Repertoire data analysis was performed using the R programming language (R Core Team (2017) R: the language and environment for statistical computing (R Foundation for Statistical Computing, Vienna, Austria. URL https://www.R-project.org/). Unproductive IGVH/IGVK/IGVL sequences were filtered out before analysis. To characterize the germline identity for each clonotype, a nucleotide sequence covering the CDR1-FR3 part was used for calculating the percentage of identity with the corresponding reference V-segment sequence. The VDJ tools software (64) with the CalcCdrAaStats subroutine encoded by the J-segment (65).

**IL-10 secretion assay**

B cells from isolated PBMC were enriched using magnetic Dynabeads (negative selection; Invitrogen, Thermo Fisher) following the manufacturer’s instructions with >95% purity. Enriched B cells were maintained in the complete glutamine-enriched RPMI-1640 medium supplemented with 10% fetal bovine serum (FBS) and 10 mM HEPES at a concentration of 0.5×10^6 cells/mL in six-well culture plates at 37°C with 5% CO2. IL-10 production was induced by the incubation with 10 µg/mL CpG-B ODN 2006 for 20 hours. CpG-stimulated B cells were restimulated by adding up to 50 ng/mL PMA (Sigma-Aldrich) and 0.5 µg/mL ionomycin (Sigma-Aldrich) for four hours.

Stimulated B cells were washed twice with PBS and stained for assessing the cell viability with Zombie Violet Fixable Viability Kit (Biolegend, USA) according to the manufacturer’s instructions. Next, B cells were washed with cold (MACS) buffer containing PBS supplemented with 0.5% BSA and 2 mM EDTA; 10^6 of B cells were then resuspended in 90 µl of cold RPMI-1640 medium supplemented with 10% FBS and incubated with 10 µl IL-10 Catch Reagent (IL-10 secretion assay, Miltenyi Biotec) for five minutes on ice. Subsequently, 1 ml of warm (37°C) RPMI-1640 medium supplemented with 10% FBS was added, and cells were kept for 45 minutes at 37°C under slow continuous rotation. B cells were washed twice with a cold (MACS) buffer, resuspended in 80 µl cold (MACS) buffer with the addition of 10 µl IL-10 detection antibody (APC) (IL-10 secretion assay, Miltenyi Biotec) and 10 µl of the antibody mix (α-CD19-PE-Cy7, α-CD24-PE, and α-CD38-AlexaFluor700). After 20 minutes of incubation on ice, B cells were washed with cold buffer and resuspended in PBS. IL-10⁺ and IL-10⁻ B cells were analyzed using FACS Aria III (BD Biosciences).

**Statistical analysis**

The data were analyzed using Prism 9 software. The significance of differences was assessed using the two-tailed Student’s t-test, Mann Whitney U-test, or ANOVA. p-values <.05 were considered to be significant.

**Results**

The elevated level of transitional CD19⁺CD24high⁺CD38high Bregs in peripheral blood correlates with MS severity

We analyzed patients with two diverging courses of relapsing-remitting MS (RRMS): (i) BMS (57), characterized by an infrequent relapse and low levels of disability over long periods of time, and (ii) HAMS (58) with elevated levels of inflammatory activity and rapid progression that facilitate a shift to secondary progressive MS with severely disrupted control of the immune response. The peripheral blood samples were obtained from 19 MS patients and 16 HDs (Table 1). To gain further insight into the nature of Breg development and maturation, we analyzed the CD19⁺CD24high⁺CD38high subpopulation, which is the most studied phenotype of tBreg (26, 66). Mononuclear cells were stained against CD45, CD19, CD24, and CD38 markers. A gating strategy is shown in Figure 1A. We found that the cell number and frequency of CD19⁺CD24high⁺CD38high B cells were significantly increased in the MS patients (4.5 ± 2.4%) as compared with the HDs (2.6% ± 1.8%). This upregulation was most pronounced in the HAMS patients (5.5% ± 2.7%) (Figures 1B, C). Wherein we observed no differences in the absolute counts for B cells between HDs and MS patients with various courses of MS (Table 2).

Previously, Cherukuri and colleagues showed that the activity of regulatory B cells varies depending on the composition of the tBreg subpopulation (67). A reduced T2/T1 ratio is associated with elevated IL-10 production and the most efficient T cell suppression (61). We studied the ratio between the transitional T1 and T2 subsets (the gating strategy is shown on Figure 4) distinguished by CD24 and CD38 expression (Table 2). The absolute T1 count was significantly elevated in the MS patients (p = .037) in...
comparison with healthy donors. The absolute T2 counts in peripheral blood were also significantly increased in the MS patients as compared with HDs ($p = .0007$), especially during HAMS ($p = .0002$). Whereas the mean value of the T2/T1 ratio is elevated during MS progression, no statistically significant difference in comparison to healthy donors was observed. To carefully distinguish between T1 and T2 cells, we analyzed CD24 and CD38 expression as well as the expression of IgD, being an important marker of B cell developmental stage and maturation (68, 69). We observed no significant difference in the T2(CD24<sup>high</sup>CD38<sup>high</sup>IgD<sup>low/−</sup>) / T1(CD24<sup>high</sup>CD38<sup>high</sup>IgD<sup>low/−</sup>) ratio between the two groups of six MS and four HDs, respectively (Figure S3).

To characterize the BCR repertoire of tBregs cells, we sorted total CD19<sup>+</sup> B cells and CD19<sup>+</sup>CD24<sup>high</sup>CD38<sup>high</sup> subpopulations and performed high-throughput sequencing of BCR cDNA libraries for variable heavy (VH) and light (VL) chains in these cell subsets. On average, we obtained ~40,000 and ~37,000 IGVH- and IGVK-containing reads for tBregs and total B cells, respectively. We obtained ~25,000 and ~17,000 IGVL-containing reads for tBregs and total B cells, respectively. We achieved the minimum depth of two IG sequence-containing reads per cell in most samples. Heavy and light chain clonotypes were assembled by the MiXCR and MiXCR with sequencing error correction (63). To reduce a potential bias, we used the equal repertoire analysis depth for all individuals (see Material and Methods section). For the CD19<sup>+</sup>CD24<sup>high</sup>CD38<sup>high</sup> B cell subpopulation, we obtained and included in our analysis ~4500 functional clonotypes for the heavy chain, ~3500 for the kappa, and ~550 for lambda chains for individuals in the MS and HD cohorts. For total B cells (the CD19<sup>+</sup> subpopulation), we obtained ~4000 functional clonotypes for heavy chain, ~3300 for kappa, and ~830 for lambda chains from the raw sequencing data and included this in our analysis for the individuals from the MS and HD cohorts (Table S2).

**TABLE 2** Total numbers and frequency of T1/T2 subpopulations of transitional B cells in peripheral blood from the MS patients and healthy donors.

| Clinical group                   | HD         | MS         | BMS        | HAMS       |
|----------------------------------|------------|------------|------------|------------|
| Absolute counts B cells per mm<sup>3</sup> | 455 ± 52   | 508 ± 74   | 508 ± 98   | 508 ± 112  |
| Absolute counts tBreg cells per mm<sup>3</sup> | 8.4 ± 1.0* | 24.0 ± 5.2* | 17.7 ± 3.0* | 29.1 ± 9.0* |
| T2/T1 ratio                      | 2.8 ± 0.2  | 3.9 ± 0.7  | 3.2 ± 0.3  | 4.4 ± 1.1  |
| Absolute counts T1 cells per mm<sup>3</sup> | 2.3 ± 0.3* | 6.6 ± 2.1* | 3.5 ± 0.8  | 8.1 ± 3.1  |
| Absolute counts T2 cells per mm<sup>3</sup> | 6.1 ± 0.7* | 16.3 ± 3.6* | 9.9 ± 2.2  | 19.4 ± 5.1* |

All values are expressed as mean values ± SEM. Significantly different values evaluated by a Mann Whitney test between the healthy donors and MS patients are indicated with asterisks.
Transitional CD19+CD24^{high}CD38^{high} Bregs of MS patients are characterized by a lower number of hypermutations compared with the healthy individuals. The tBregs are suggested to be the exogenous antigen-naïve cell population, therefore exhibiting fewer somatic hypermutations than the total pool of CD19+ B cells from peripheral blood (Figure 2A). V_H and V_K genes of CD19^{+}CD24^{high}CD38^{high} tBregs from MS patients are generally less mutated as compared with the HDs. There was a statistically significant difference between the HDs and HAMS patients yet not between the donors and BMS patients (Figure 2A). These data are in line with an increased level of tBregs in the peripheral blood from the HAMS patients. We observed approximately the same number of the IGH clonotypes in tBregs and total CD19^{+} B cells in the MS patients and healthy donors (Figure 2B). Analysis of repertoire diversity of the light chains showed that in the HDs the ratio in number of the IGK and IGL clonotypes in the total CD19^{+} pool was higher than in tBregs, whereas it remained unchanged in MS patients (Figure 2B). The lowest repertoire diversity was observed for the lambda light chain. The ratio of tBreg/total B cell clonotypes between MS patients and HDs was different only for the lambda chain (Figure 2C).

Because in the MS patients we observed fewer somatic hypermutations in the tBreg clonotypes, one could suggest a higher number of CDR3 in this cell subset shared among MS patients. However, we detected only a few tBreg IGH clonotypes common for at least two different donors. Furthermore, we did not observe any significant differences in the number of shared amino acid CDR3 sequences between MS and HD (data not shown).

**Characteristics of the CDR3 region of tBreg clonotypes do not vary between MS and healthy states**

CDR3 is the most variable region of immunoglobulin molecules. It can be used for identifying clonal lineages and characterizing their functional repertoire (70). First, we examined CDR3 characteristics by comparing the amino acid CDR3 length of the heavy, kappa, and lambda chains among HDs, HAMS, and BMS disease states in tBregs and total peripheral B cells. In line with our expectations, we detected a significant difference in amino acid length between the heavy chain (18.3 ± 0.7 a.a.) and both light chain types (11.2 ± 0.1 a.a. for the kappa chain and 11.5 ± 0.3 a.a. for the lambda chain) due to the presence of the D-segment in the IGH.

For the heavy chain, the CDR3 length significantly varied between the tBreg and the total B cell populations (Figure 3A) in contrast to the previously reported data (71). Nonetheless, there was no statistically significant difference in the CDR3 length...
between the MS patients and healthy individuals (Figure 3B). We did not detect any significant differences in the length of CDR3 of the kappa and lambda light chains between the tBreg subset and total B cells (Figures 3C, D, S1).

Furthermore, we compared the physicochemical properties of the CDR3 amino acids for the clonotypes from tBregs and total peripheral B cells. We observed no significant differences in the charge and hydrophobicity or amino acid usage in the CDR3 regions between the B cell subsets of the MS patients and HDs (data not shown).

The CD19+CD24highCD38high subpopulation in MS patients is characterized by less mature phenotype

It is previously reported that the number of mature memory CD27+ peripheral B cells (72) in MS patients tends to decrease as compared with healthy individuals (73). Importantly, the ratio of different B cell populations and especially of the memory B cells varies with age (74). To avoid any age-related bias, the samples from sex- and age-matched MS patients (44 ± 14 y.o.) and HDs (43 ± 13 y.o.) were enrolled in this study (Table 1; Figure 4C). To examine the phenotypic stage of tBreg maturation, we analyzed the percentage of CD27+ cells in this subpopulation (Figure 4A). We found that the CD27+ cell content in the CD19+CD24highCD38high tBregs cells was significantly reduced in the MS patients (1.0% ± 0.5%) as compared with healthy individuals (2.2% ± 1.4%). We observed no correlation between age and the frequency of CD27+ cells in the CD19+CD24highCD38high subpopulation (Figure S2). Our findings correlate well with previously reported data suggesting that Bregs (CD19+IL10+) mostly had the CD27- phenotype in HD and MS patients in remission (28), whereas the percentage as well as the absolute counts of memory Bregs (CD24highCD27+) and naïve and memory B cells were similar in the HDs and MS patients (Figure 4B; Table 3).

CD19+CD24highCD38high B cells are characterized by an elevated production of IL-10 in MS patients and HDs

Because the regulatory properties of B cells are not limited exclusively to the CD19+CD24highCD38high subpopulation (55, 56), we analyzed the frequencies of the IL-10-positive cells among the total pool of B lymphocytes after a short CpG stimulation. Such rapid stimulation (less than 24 hours) allows estimating IL-10 production only in Bregs and not in the B cells predisposed for the IL-10 expression. There were no statistically significant differences in the IL-10-producing B cell subsets between MS and HD, whereas more IL-10-positive B cells were observed in the CD19+CD24highCD38high subpopulation than in total B cells for both MS and HD (Figure 5A).
Conversely, IL-10–positive B cells in MS and HD were enriched with CD19+CD24highCD38high cells (Figure 5B).

**Discussion**

The existence of several alternative methods of Breg differentiation might lead to different subsets of B cells with regulatory functions coexisting in an inflammatory milieu (75). The Breg population can originate from either multiple or a limited set of independent pre-Bregs progenitors. Furthermore, it results in either a diverse or restricted clonal repertoire, the latter being ensured by the clonally related Bregs. Moreover, almost any B cell can become a regulatory B cell at a certain stage of its development upon exposure to permissive environmental stimuli (35, 55). In the present study, we focus on the repertoire of tBregs with the CD19+CD24highCD38high phenotype.

The mechanism of immune regulation mediated by B cells was first proposed by S. Fillatreau and colleagues. They first identified a subpopulation of B cells (B10 cells) that produced IL-

**TABLE 3** Total numbers of B cell subpopulations in peripheral blood of the MS patients and healthy donors (gating strategy is shown in Figure 4).

| Clinical group       | HD    | MS    | BMS   | HAMS  |
|----------------------|-------|-------|-------|-------|
| Memory B cells       | 51 ± 9| 41 ± 6| 34 ± 5| 46 ± 9|
| Naïve B cells        | 192 ± 40| 216 ± 47| 172 ± 42| 246 ± 75|
| Memory Breg cells    | 89 ± 17| 77 ± 22| 44 ± 6| 100 ± 34|

All values are given as absolute counts per mm3 and presented as mean ± SEM.
10, alleviating the clinical manifestations of EAE (76). Bregs can also affect the differentiation of T cells to T regulatory cells (Tregs) (24) and inhibit differentiation of the effector T cells via the IL-10–driven suppression of dendritic cells (35). Bregs are shown to suppress inflammation by producing transforming growth factor-β (TGF-β) (77), IL-35 (i35 Breg) (78), IgM, IgG4, the co-inhibitory receptor TIGIT (T cell immunoreceptor with Ig and ITIM domains) (36), and BTLA (B and T lymphocyte attenuator or CD272) (79). Nonetheless, IL-10 production is still believed to be the key mechanism for Bregs to control the immune response in healthy individuals as well as during immune-related disorders (26, 27, 80–82) and organ transplantation (83).

Breg maturation may be induced via BCR and/or TLR signaling (84). Using EAE, the experimental mouse model of autoimmune diseases, a significant impact of TLR-signaling on B cells was revealed (85). TLR-4 and TLR-9 induce a significant increase in IL-10 secretion by B cells in a MyD88-dependent manner. Their activation via TLR and CD40 ligands is also described for infectious diseases of viral (HIV (31), HBV (86)), bacterial (87, 88), and parasitic origins (89). Exposure of PBMC from both healthy donors and patients with rheumatoid arthritis to CD40L and CpG led to an increased number of IL-10–producing Bregs (90). A significantly lower level of IL-10 production by B cells stimulated in the presence of CD40L was found in the groups with relapsing-remitting (29) and secondary-progressive MS as compared with HDs. A similar effect was observed in the CpG-stimulated B cells (82). The fact that CD40 ligation on B cells plays an important role in this process promoting cell survival (91) is in line with the inductive model of Breg formation. The latter implies that B cells may become regulatory and exhibit suppressive capacity in response to specific environmental stimuli.

Another mechanism to induce Bregs is to activate them through BCR signaling (76). Matsumoto et al. demonstrate the importance of correct antigen recognition by Breg BCR (92). The antigen-mediated calcium influx in Bregs is shown to depend on STIM-1 (stromal interaction molecule) and STIM-2. A deficiency in these molecules impairs the ability of Bregs to produce IL-10, disrupts T cell activation, and eventually prevents alleviating EAE in mice. The number of Bregs was reduced in CD19-deficient mice with disrupted BCR-signaling, whereas CD19 overexpression resulted in an increased number of B10 cells (93–95). The antigen-specific interaction between CD4+ T cells and B10 cells is crucial for generating B10 effector cells, whereas antigen-specific T cell response is downregulated by B10-mediated IL-10 production (96). Therefore, the B10 cell effector function primarily depends on antigen specificity; however, it might also be mediated by suppressing antigen presentation by dendritic cells and macrophages (97).

We observed that CD19+CD24highCD38high tBregs have a significantly longer CDR3 in the heavy chain as compared with the total pool of peripheral B cells. Antigen-experienced B cells were previously shown to have shorter CDR3 than naïve or immature B cells (98, 99), thus indicating that, in tBregs, longer CDR3 can point at their immature state. This can also be related to the differences in V/J-usage between tBregs and total B cells (100, 101). Finally, in tBregs, longer CDR3 may reflect the higher poly-/self-reactive potential of their BCRs (102).
Another piece of evidence underlying the importance of BCR during Breg activation is that human B10 cells are often defined as CD27+. This allows for classifying them as memory cells, which is in line with their in vivo antigen experience (25). Of note, in the RRMS patients at the relapse stage, the ratio of naïve and memory Bregs is decreased, resulting in an elevated memory/naïve ratio (28). The presence of the CD27+ subpopulation in early immature transitional B cells could be explained by the recently proposed class-switching of antibodies in early human B cell development (71). Therefore, even transitional cells can undergo early maturation or, according to the alternative hypothesis, the CD19+CD24hiCD38hiCD27+ subset might belong in the IgM memory population (103, 104). Here, we show that the CD19+CD24hiCD38hi tBregs from MS patients contain a statistically lower number of CD27-positive cells as compared with HDs.

Our findings allow us to conclude that the elevated absolute number and frequency of CD19+CD24hiCD38hi tBregs observed in MS patients is characterized by a greater germline identity as compared with HD. At the same time, the absolute cell counts of the recently immigrated from the bone marrow T1 and more mature T2 cells both increased, thus more or less maintaining the overall T1/T2 ratio as in a healthy state. We propose at least three scenarios explaining these findings: (i) deficient maturation of transitional B cells (TrB); (ii) delayed TrB maturation, and (iii) elevated TrB counts that compensate for deficient maturation (Figure 6). The latter seems to be the most likely scenario because the T1/T2 ratio and, especially, percentage and absolute counts of TrB are increased in MS patients, whereas the numbers of memory and naïve B cells remain unaltered. Nonetheless, current evidence is not conclusive yet.

The tBregs maturation failure could arise from either an intrinsic defect in pre-Bregs or insufficient antigen-induced maturation. The BCR sequences of B10 cells were recently shown to be closer to germline and harbor only rare mutations (105). The authors state that, similarly to the splenic B10 cells, the peritoneal cavity B10 cells expressed clonally diverse BCRs that were predominantly germline encoded. Despite the germline proximity, B10 cells are shown to produce IgG as well as IgM (106). Thus, the so-termed “low differentiated” BCR may provide low-affinity antigen-BCR stimulation during chronic disease and development of the B10 precursor to B10, whereas a strong stimulation may switch the Breg precursor to another differentiation pathway (25, 94, 105). In the present work, we revealed no differences in IL-10 production between MS and HD via functional analysis. Therefore, increased tBreg counts along with the absence of increased IL-10 production can serve as indirect evidence in favor of their disrupted functioning and altered inflammatory profile during MS development. We suggest that, in future studies, tBregs should be analyzed for IL-10 production as well.

![Figure 6](image-url)  
*Abnormalities in transitional B cell maturation in MS. An elevated frequency of CD19+CD24hiCD38hi transitional B cells (TrB) observed in MS patients is characterized by greater germline identity compared with healthy donors. There are least three possible coexisting or independent scenarios explaining these findings: (i) deficient maturation of TrB, (ii) delayed TrB maturation, and (iii) an increased number of TrB compensating deficient maturation.*
as co-expression of other cytokines as IL-10⁺ B cells are previously shown to co-express pro-inflammatory cytokines such as IL-6 and tumor necrosis factor alpha (TNFα) (55).

Here, we put forward that the observed stagnation in the maturation of the transitional Bregs potentially may be the first step in the fatal sequence of events leading to the systemic failure of humoral regulative immunity. Being accompanied by one or multiple factors, such as HLA haplotype, cytokine background, abnormal T cell negative selection, enhanced blood-brain barrier permeability, or lack of vitamin D, it may trigger the uncontrolled breakdown of the immunological tolerance toward myelin antigens. On the other hand, the elevated absolute count of tBregs and T1/T2 B cell subset ratio may be a sign of the already activated compensatory mechanisms, which may be monitored on the border of a clinically isolated syndrome and clinically defined multiple sclerosis. Further studies should elucidate if modulating the immunological checkpoints regulating B cell development may be regarded as opening up the avenue for putative therapeutical applications in MS treatment.

Data availability statement

The datasets presented in this study can be found in online repositories. The names of the repository/repositories and accession number(s) can be found below: https://www.ebi.ac.uk/arrayexpress/, E-MTAB-10859.

Ethics statement

The studies involving human participants were reviewed and approved by Local Ethic Committee of the Research Center of Neurology, Moscow, Russia. The patients/participants provided their written informed consent to participate in this study.

Author contributions

YL, ABJr and AG designed the research and wrote the paper. MNZ, MI and TS performed blood sample acquisition, patient data management and MS diagnosis. YL, LO, MK, AT, MYZ and DS performed research. YL, IZ, LO, AM, ABJr and AG analyzed data. YR, IZ, DC, NB, VM, AF and SI made intellectual contributions to data analysis. All authors contributed to the article and approved the submitted version.

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Conflict of interest

The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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Supplementary material

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fimmu.2022.803229/full#supplementary-material

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