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B Cells with Regulatory Function in Animal Models of Autoimmune and Non-Autoimmune Diseases

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Abstract

Although the identification of B cell subsets with negative regulatory functions and the definition of their mechanisms of action are recent events, the important negative regulatory roles of B cells in immune responses are now broadly recognized. There is an emerging appreciation for the pivotal role played by B cells in several areas of human diseases including autoimmune diseases and non-autoimmune diseases such as parasite infections and cancer. The recent research advancement of regulatory B cells in human disease coincides with the vastly accelerated pace of research on the bridging of innate and adaptive immune system. Current study and our continued research may provide better understanding of the mechanisms that promote regulatory B10 cell function to counteract exaggerated immune activation in autoimmune as well as non-autoimmune conditions. This review is focused on the current knowledge of BREG functions studied in animal models of autoimmune and non-autoimmune diseases.

Keywords

Bregs; Animal Models; IL-10; Autoimmune Disease; Immune Regulation

1. Introduction

Historically, B cells have been characterized as positive regulators of humoral immune responses and are distinguished by their ability to terminally differentiate into antibody (Ab)-secreting plasma cells [1] [2] or serve as antigen (Ag)-presenting cells (APCs), for optimal Ag-specific CD4⁺ T-cell expansion, memory formation, cytokine production [3]–[5] and positively regulate CD8⁺ T-cell responses by expression of co-stimulatory molecules [6]–[8].

Evidence for B-cell negative regulatory function has accumulated over the past 30 years. The hypothesis that suppressor or regulatory B (Bregs) cells orchestrate the immune system was originally proposed in the 1970s and maintains that the suppressive function of B cells

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was mainly restricted to their ability to produce "inhibitory" antibodies [9]. These initial findings were later followed by a flurry of seminal papers supporting a "suppressive framework" for B cells and a link with T-cell tolerance [10]–[13]. Later studies showed that a B-cell-restricted IL-10 deficiency had a similar exacerbating effect on EAE [14] [15] and rheumatoid arthritis [16], suggesting that activated B cells exerted regulatory activity that resolved the inflammation. From this, this term "regulatory B cell" was coined.

More recently, a relatively rare negative regulatory B-cell subset was identified that was predominantly contained within a phenotypically unique CD1d^{hi}CD5⁺CD19^{hi} subset in the spleens of naive wild-type mice [17]. This regulatory B-cell subset is Ag-specific and significantly influences T-cell activation and inflammatory responses through IL-10 production [17] [18]. Given that multiple regulatory B cell subsets are likely to exist, as now recognized for T cells, it has been specifically labeled this IL-10 competent CD1d^{hi}CD5⁺CD19^{hi} regulatory subset as B10 cells because they are responsible for most IL-10 production by B cells and they appear to only produce IL-10 [19].

Recently, many studies have used animal models of human diseases to demonstrate that B cells have regulatory functions *in vivo*, and most of them use B-cell-deficient μ MT mice or the adoptive transfer of bulk B cells. In addition, multiple groups have identified IL-10-producing regulatory B cells of varying phenotypes in different disease models following diverse stimulation and culture protocols [14]–[16] [20]–[27].

2. Bregs in Autoimmune Diseases

2.1. Experimental Autoimmune Encephalomyelitis

B cells play a dual role in experimental autoimmune encephalomyelitis (EAE), which is the animal model of the human autoimmune disease multiple sclerosis (MS). The first one is contributing to the pathogenesis of EAE through the production of anti-myelin antibodies that contribute to demyelination. The other one is playing an essential role in the spontaneous recovery from EAE.

In 1996, Wolf found that B cells were not required for the onset of EAE, revealed that μ MT mice failed to spontaneously recover from EAE [28]. This was the first indication in animal autoimmune models that B cells play a regulatory role in down-regulating inflammation. While in the later study, C57BL/6µMT mice were either not able to recover from EAE when an EAE induction protocol that allowed for recovery from EAE was used, which indicating that B cell production of IL-10 and expression of CD40 were requirements for their regulatory activity [14].

In regards to TLR signaling TLR2/4 ligands are present in the CFA adjuvant not only used to induce EAE but also induce the production of IL-10 by B cells [29]. Thereby, IL-10-producing regulatory B cells, most likely B10 cells, are important for controlling EAE severity and resolution. More specifically, Myd88 expression by B cells was required for recovery from EAE in MOG-peptide EAE [29].

The Tedder laboratory studied that both genetic disruption of CD19 and the transfer of $CD19^{-/-}$ B cells from MOG peptide-primed mice exacerbated EAE which demonstrated that

signaling was important for regulatory B cell functions [30]. They analysed $CD19^{-/-}$ and human CD19 transgenic (hCD19tg) mice, with the later harboring hyperactive B cells, indicating that $CD19^{-/-}$ mice exhibit enhanced T cell-mediated inflammation (contact hypersensitivity (CHS)), and inflammation in hCD19tg mice was reduced [17].

The B cell regulatory effects were recently shown not to be IL-10 dependent [31]. On the contrary, the Tedder laboratory showed that the adoptive transfer of CD1d^{hi}CD5⁺B cells from CD19^{-/-} mice or from MOG-sensitized animals prior to EAE induction by MOG peptide attenuated EAE disease severity. Meanwhile, B cells from IL-10^{-/-} mice did not reduce disease severity, indicating an IL-10-dependent mechanism [1]. B cell depletion therapy in early MS clinical trials with anti-CD20 showed remarkable efficacy in preventing disease progression [32] [33]. However, much still needs to be learned regarding how each B cell function can harnessed to either prevent or induce the recovery from MS.

2.2. Type 1 Diabetes

Studies on B10 cells and mouse models of diabetes are limited to the nonobese diabetic (NOD) mouse, a spontaneous model of type 1 diabetes in which autoimmune destruction of the insulin-producing pancreatic β cells is primarily T cell mediated [34].

Although B cells play the pathogenic role in T1D initiation [35], B cells activated *in vitro* can maintain tolerance and transfer protection from T1D in NOD mice, both delay the onset and reduces the incidence of T1D. Protection from T1D is IL-10 dependent since the transfusion of activated NOD-IL- $10^{-/-}$ B cells does not confer protection from T1D or the severe insulitis observed in NOD recipients [36] [37]. In another study, LPS-activated B cells were transferred into prediabetic NOD mice and found that Fas ligand and secreted transforming growth factor- β were upregulated, which were considered to contribute to inhibit autoimmunity [37].

Although the animal studies in TID have shed some light on the limitation of the rarity of circulating B10 cells, the possibility of therapeutic transfusion of autologous, IL-10-producing, BCR-activated B cells or B10 cells in order to protect human subjects at risk for T1D remains elusive.

2.3. Arthritis

CIA is a model for human rheumatoid arthritis that develops in susceptible mouse strains immunized with heterologous type II collagen emulsified in complete Freund's adjuvant [38] [39], which shares in common with rheumatoid arthritis having an association with a limited number of MHC-II haplotypes that determine disease susceptibility [40] [41].

B cells are important for initiating inflammation and arthritis [42]. By contrast, IL-10producing B-cell sub-sets regulate inflammation during CIA. Activation of arthritogenic splenocytes with Ag and agonistic anti-CD40 mAb induces a B cell population that produces high levels of IL-10 and low levels of IFN γ [16]. Specifically, multiple studies have tested whether the adoptive transfer of activated B cells could inhibit CIA. Mauri's lab injected CD40 mAb and collagen-activated B cells from the spleens of arthritogenic mice into recipient mice, observed that arthritis incidence (>50% reduction), disease severity (>90%),

and Th1 cell differentiation are inhibited. Moreover, adoptive transfer of B cells also partially inhibits arthritis incidence and severity, even after disease initiation. However, the adoptive transfer of IL-10^{-/-} B cells does not prevent arthritis in this model system [16]. Evans has tested the adoptive transfer of B cells into mice immunized with bovine collagen (type II collagen) inhibits TH1 responses, prevents arthritis development, and is effective in ameliorating established disease, while the adoptive transfer of CD21^{hi}CD23⁺IgM⁺ B cells from DBA/1 mice in the remission phase could prevents CIA and reduces disease severity through IL-10 secretion [22]; Gu also found a substantial reduction in the number of T_{H17} cells [43]. Other studies administered apoptotic thymocytes to mice up to 1 month before the clinical onset of CIA is also protective for severe joint inflammation and bone destruction [23].

Collectively, activated spleen B cells responded directly to apoptotic cell treatment, increasing secretion of IL-10, which is important for inducing T-cell-derived IL-10. Moreover, the passive transfer of B cells from apoptotic cell-treated mice provided significant protection from arthritis.

2.4. Systemic Lupus Erythematosus

Studies in the NZB/W spontaneous lupus model therefore suggest that B10 cells have protective and potentially therapeutic effects. In wild type NZB/W mice, the CD1d^{hi}CD5⁺B220⁺ B cell subset, which is enriched in B10 cells, is increased 2.5-fold during the disease course, whereas CD19^{-/-} NZB/W mice lack this CD1d^{hi}CD5⁺ regulatory B cell subset [44]. Mature B cell depletion initiated in NZB/W F1 mice, hastens disease onset, which parallels depletion of B10 cells, suggesting that B cell-negative regulatory effects are important in NZB/W mice [45]. Moreover, the potential therapeutic effect of B10 cells in lupus is highlighted by the prolonged survival of CD19^{-/-} NZB/W recipients following the adoptive transfer of splenic CD1d^{hi}CD5⁺ B cells from wildtype NZB/W mice [44]. However, in the MRL.Fas(lpr) mouse lupus model, B cell-derived IL-10 does not regulate spontaneous autoimmunity [46]. The study suggests fundamental differences in the pathogenesis and immune dysregulation in the NZB/W lupus model compared with the MRL.Fas(lpr) model.

2.5. Inflammatory Bowel Disease

Early studies showed that B cells and their autoantibody products suppress colitis in T cell receptor alpha chain-deficient mice that spontaneously develop chronic colitis, while B cells are not required for disease initiation [47].

Mizoguchi's group has subsequently demonstrated that B cells are the regulatory mediators and were the first to identify a B-cell subset with up regulated CD1d expression that is induced in the gut-associated lymphoid tissues of mice with intestinal inflammation, and they also suggested IL-10-producing B-cell subsets with varying phenotypes and origins regulate intestinal inflammation during inflammatory bowel disease [15] [48].

Dextran sulfate sodium-induced intestinal injury is more severe in $CD19^{-/-}$ mice where B10 cells are absent than in wild type mice [49], suggesting these inflammatory responses are negatively regulated by $CD1d^{hi}CD5^+$ B cells producing IL-10. Moreover the other study

observed the adoptive transfer of mesenteric lymph node B cells also suppresses inflammatory bowel disease through a mechanism that correlates with an increase in regulatory T-cell subsets [50]. Thus, cytokine-producing B cells can regulate immunemediated gut inflammation. B10 cells therefore emerge during chronic inflammation in mouse models of inflammatory bowel disease, where they suppress the progression of inflammatory responses and ameliorate disease manifestations.

3. Bregs in Parasitic Infection

The first clue for an anti-inflammatory role of Bregs in infections with parasites came from a study demonstrating a regulatory role of B cells during *Schistosoma mansoni* worm plus egg infection. Infection of μ MT mice led to an enlargement of hepatic granuloma and a decreased lifespan [51] [52]. Recently, it has been demonstrated that B cells induced by *Schistosoma mansoni* worms is in an IL-10-dependent manner, which are responsible for protecting mice against fatal, experimentally induced anaphylaxis [53]. Breg cell function during different stages of natural *S. mansoni* infections and showed the existence of active regulatory mechanisms during chronic, but not acute infection [54]. A similar liver pathology was observed in schistosome-infected, Fcg-chain receptor knockout mice [52], suggesting that B-cell regulation is mediated either by the production of antibodies neutralizing egg-derived inflammatory molecules or by triggering the production of anti-inflammatory mediators from FcR⁺ cells.

The concept that helminth-induced B cells can protect against allergic inflammation has been extended to other helminth infections that are natural for mice: in H polygyrus-infected mice, adoptive transfer of mesenteric lymph node B cells suppressed both DerP1-specific airway inflammation and EAE [55]. Interestingly, Breg cell development can also be seen during the infections caused by Leishmania major [56]; IL-10-producing B cells were critical for the development of unprotective TH2 responses and susceptibility to infection. In addition, murine cytomegalovirus has been shown to induce IL-10-producing Breg cells, resulting in decreased virus-specific CD81 T-cell responses and plasma cell expansion [57]. Other studies investigated the role that B cells play in infection with the nematode *Brugia pahangi*. Adoptively transferred peritoneal B cells, isolated from wild-type mice that had been immunized with *B. pahangi*, have been shown to protect athymic recipient mice against infection by *B. pahangi* infection [58].

In another study, the depletion of B cells from splenocytes of infected mice resulted in a reduced level of antigen-specific CD4⁺ T-cell proliferation paralleled by a reduced level of CD80 and CD86. Similar results were obtained if IL-10 was neutralized at the time of infection, suggesting that B cells producing IL-10 might modulate immune response in filarial-infected mice, via the suppression of CD80 and CD86 expression on Bregs [59]. Taken together, these studies show that helminths can induce Breg cells that can protect against allergic diseases via the release of IL-10 and that this process is particularly active during the chronic stage of infection. Furthermore, it can be concluded that the suppressive ability of Breg cells is not restricted to TH1 immune responses associated with autoimmunity, and the effect that the humoral immune response and FcR interactions might play a major role in controlling.

4. Bregs in Cancer

Given that Bregs have been shown to suppress autoimmunity via the inhibition of autoreactive T cells, it might be anticipated that Bregs could also downregulate the protective cytotoxic T lymphocyte responses directed against tumor cells. Terabe found that enhanced antitumor immunity can be seen in B-cell-deficient mice and is associated with an increased activity of T and NK cells, both of which are important for the promotion of natural tumor surveillance [60]. Increased CD4⁺ and CD8⁺ T-cell responses to TS/A tumors are observed in µMT mice [61]. In B16 melanoma, mature B-cell depletion using CD20 mAb dramatically exacerbates tumor progression and metastasis, arguing that B cells primarily support antitumor immune responses in this model [1]. Thus, these results suggest that B cells can also negatively regulate tumor immunity.

Scott and colleagues have shown that the interaction between $CD40^{L}$ expressed on tumors and CD40 on B cells induces IL-10 production by B cells, indicating that the release of IL-10 is probably responsible for the diminished IFN γ production by CD8⁺ T and NK cells and the decreased CD8⁺ T-cell memory development. However, there is some evidence to suggest that IL-10 can suppress angiogenesis and, thus, encourage tumor regression [62] [63]. Furthermore, the ubiquitous expression of CD40 *in vivo* will probably lead to a more complex cascade of anti- and pro-inflammatory cytokines, which might overcome the inhibitory effect of Bregs. But it remains to be formally proven whether or not administration of anti-CD40 can generate Bregs *in vivo*. Another study demonstrated an excessively zealous Breg population may promote tumor cell growth, as one of the mechanisms used by tumor cells to escape from the immune response consists in activation of Bregs that produce TGF-beta [64].

A role for B cells in the development of tumor immunity has been assessed using μ MT mice given Friend murine leukemia virus gag-expressing mouse EL-4, D5 melanoma, or MCA304 sarcoma cells. Inoue and colleagues have tested wild-type mice were unable to control tumor progression, whereas EL-4 gag and D5 tumors (but not MCA304) were eliminated in μ MT mice, which developed tumor-specific cytotoxic T lymphocytes after tumor challenge. Similar study suggested the growth of EL4 thymoma, MC38 colon carcinoma, and B16 melanoma was prevented or slowed in μ MT mice in contrast to control mice [62] [65].

By contrast, B-cell depletion using CD20 mAb in a syngeneic lymphoma model dramatically enhances tumor clearance through B10-cell elimination. Thus, as in autoimmunity, B10 cells are likely to be involved in regulating antitumor immunity. However, this regulation will be significantly influenced by the immunogenicity of the tumor and the nature of the antitumor immune response [66].

5. Summary

Despite the extensive efforts on the characterization of BREG subtypes and their mechanism of action in different animal models, questions still exist to unravel the mechanisms underlying Bregs biology and function. The precise phenotype and characteristic markers of Bregs are still the subject of debate. It remains unclear whether Bregs require self-reactive

BCRs for function. Are Breg cells a developmentally distinct B-cell subset? Do Breg cells display a specific transcriptional signature such as FoxP3 for regulatory T cells? Understanding these questions may open novel avenues for the treatment of inflammatory diseases such as allergy and autoimmunity.

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