TNF-α-induced upregulation of p100 is somehow blocked under inflammatory conditions, such as in that in RA synovial joints, or do pathologic conditions induce signaling molecule(s) that suppress the inhibitory effect of p100? Second, while p100 acts as an IκB-like molecule that binds to RelB, p100 also binds to the RelA/p50 dimer (11, 18). Given that Rela−/− or Relb−/− single-knockout mice do not show an altered bone phenotype, p100 may suppress TNF-α-induced osteoclast formation by retaining both Rela/p50 complex and RelB in the cytoplasm. Thus, it will be interesting to investigate the bone phenotype of Rela−/−Relb−/− mice. Third, the mechanism of how TNF-α suppresses the degradation of TRAF3 has not been clarified. It will also be intriguing to test whether TNF-α−induced osteoclast formation is enhanced in Traf3−/− mice. Given that Traf3−/− mice die soon after birth (14), transfer of Traf3−/− bone marrow cells to WT mice would be a feasible way to investigate the role of TRAF3 in TNF-α−induced osteoclast formation in vivo. Finally, what is the role of bone-forming osteoblasts in TNF-α−induced pathologic bone resorption, since TNF-α is known to induce RANKL expression in osteoblasts? These intriguing questions remain for future study.

In conclusion, Yao et al. (13) have convincingly demonstrated that NF-κB2p100 plays a negative role in suppressing TNF-α−induced osteoclast formation under pathologic conditions using various animal models. Thus, blockade of the processing of p100 might be a novel strategy to treat various bone diseases such as RA, in which TNF-α−induced osteoclast formation plays a crucial role in the progression of diseases.

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Pathogenic antibodies are active participants in spinal cord injury

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The role of B cells and autoimmunity as contributing factors to poor neurological outcomes following spinal cord injury (SCI) is poorly understood. The study by Ankeny et al., in this issue of the JCI, identifies a new immunopathological mechanism arising after SCI in mice (see the related article beginning on page 2990). The study shows that B cells produce pathogenic antibodies that impair lesion repair, resulting in worse neurological outcome. This new understanding of SCI disease pathogenesis, if confirmed in humans, reveals potential avenues for the development of novel neuroprotective immunotherapies.

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Primary trauma to the CNS initiates a series of interrelated responses, including edema, excitotoxicity, and inflammation, that lead to secondary injury, resulting in further expansion of the initial lesion and additional loss of neurological function. The treatment of neuroinflammation in the context of both traumatic brain injury and spinal cord injury (SCI) still lacks a standard, universally accepted therapy that leads to improved neurological outcomes. There exists a clear, unmet medical need for an effective antiinflammatory treatment for the acute and chronic stages of traumatic brain injury and SCI arising in general civilian as well as injured military personnel populations. Currently, an important area of SCI research focuses on...
Understanding the dual nature of post-SCI neuroinflammation. It can lead to increased damage to the neural tissue, yet it is an essential part of the wound healing process (1, 2). It is becoming increasingly clear that injury to the CNS results in alterations to the systemic immune system that in turn affect the competing pathogenic and wound healing processes evolving at the site of injury (3–5). The study reported by Ankeny et al. in this issue of the JCI (6) offers a new perspective on a poorly understood aspect of CNS inflammation in response to traumatic injury in mice, namely the impact of B cells and autoimmunity on neurological outcomes. This new understanding of SCI disease pathogenesis, if confirmed in humans, reveals avenues for the development of novel neuroprotective immunotherapies.

**B cells in the normal and diseased CNS**

B lymphocytes are a key component of the adaptive immune system. They arise from bone marrow hematopoietic stem cells that differentiate in an antigen-independent manner to the immature B cell stage. In the presence of antigen, further differentiation leads to mature and activated B cells. Activated B cells convert into short-lived, antibody-secreting plasma cells or into memory B cells and then into long-lived, antibody-secreting plasma cells (7, 8). Activated B cells and long-lived plasma cells migrate not only to the bone marrow and secondary lymphoid organs but to the CNS via normal homeostatic processes (7). This B cell recruitment mechanism is upregulated during CNS autoimmune diseases, such as MS (7, 9). There are several B cell–specific factors and receptor interactions that are critical to B cell function and are potential therapeutic targets. B cell–activating factor (BAFF), lymphotoxin-β, and a proliferation-inducing ligand (APRIL) have roles important to B cell survival, differentiation, germinal center formation, and antibody synthesis (7, 8). These factors are secreted by macrophages and, within the CNS, by astrocytes (7, 8). Thus, B cells have an established mechanism that allows them to traffic to and be supported in the CNS. The normally assumed role of B cells is to produce antibodies, but it is now clear that B cells can serve as potent regulatory and antigen-presenting cells (8, 10). It is well known that under normal circumstances human cerebrospinal fluid (CSF) harbors low levels of antibody, produced by long-lived plasma cells, some of which are autoreactive (11). The role of B cells in various CNS autoimmune conditions is also well known. However, until recently the role of B cells was deemed secondary to that of T cells in disease pathogenesis. There is now clear evidence that B cells and associated autoantibodies can play an important primary role in CNS autoimmune disease (8, 12).

**SCI leads to pathogenic autoantibody production**

The results presented in this issue by Ankeny et al. (6) clearly demonstrate that, in a mouse model of SCI, trauma of moderate severity at thoracic level 9 (T9) leads to a surprisingly robust B cell response that produces pathogenic antibodies. This important conclusion is supported by experiments demonstrating that spontaneous neurological recovery after injury was greatly improved in B cell–knockout mice compared with WT mice. Following SCI, coordinated stepping involving all four limbs was achieved in 88% of B cell–knockout mice but in only 35% of WT mice at the...
end of the nine-week observation period. Consistent with this improved functional recovery, the neuropathology observed in the B cell–knockout mice was also markedly less pronounced compared with WT animals. This suggests that, in WT mice that received an SCI, B cells play a role in the evolving inflammatory response that impedes neurological recovery. Importantly, passive transfer (injection) of purified pathogenic antibody into the spinal cord of WT mice under sterile conditions induced a similar type of neurotoxicity to that observed in mice with SCI. This confirmed that neurotoxic product of the SCI-induced B cell activation was likely pathogenic antibodies.

This article (6) raises the question as to why B cells produce pathogenic antibodies when the SCI is in the lower half of the spinal cord (T9–T10). Yet, these same authors previously reported that when the SCI occurs at a higher level (T4–T5) profound immune suppression occurs, including that of B cell function (13). A likely explanation may lie in the fact that a high SCI disrupts the cholinergic antiinflammatory pathway by removing the sympathetic contribution (via the splenic nerves) due to injury to the intermediolateral column sympathetic fibers. This pathway plays a key role in regulating systemic inflammation (4). Further investigation is required to understand the underlying pathological mechanisms created by the loss of sympathetic control that results in immune suppression and whether this is a permanent feature of SCI located above T4/T5. Alternatively, the appearance of autoimmune neuropathogenic antibodies may only be delayed.

The presence of pathogenic antibodies in the spinal lesion is in part derived from systemic sources via a compromised blood–spinal cord barrier after injury. However, evidence was presented by Ankeny et al. (6) that B cell “follicle-like” structures were present in the lesion area, suggesting that local antibody production may be critical to the pathogenic outcome. Both activated B cells and plasma cells are present in these structures. The presence of B cell follicle-like structures in the diseased CNS is not unique to SCI but has been described in other CNS autoimmune diseases, including MS (9). The local SCI inflammatory environment likely would support the development of B cell follicle-like structures in two ways. Infiltrating monocyte/macrophages as well as the large number of proliferating astrocytes responding to the SCI likely could supply sufficient BAFF and APRIL to support B cell survival, proliferation, differentiation as well as follicle development (Figure 1). Secondly, SCI-induced neuroinflammation dysregulates an important immune regulatory mechanism known as the tryptophan–kynurenine pathway (14–17). The critical component of this tryptophan catabolic pathway is the rate-limiting enzyme, indoleamine 2,3-dioxygenase (IDO), which is primarily induced by IFN-γ but can be superinduced in the presence of TNF-α (14, 15). Both cytokines are present following SCI. IDO is a key regulator of tolerance and a precursor of autoimmunity (18, 19). IDO is expressed by activated microglia and infiltrating leukocytes (14). T cells, but not B cells, are particularly sensitive to the effects of IDO. The tryptophan metabolites generated by IDO, such as quinolinic acid, are well known neurotoxins and also render T cells, but not B cells, highly susceptible to apoptosis and other aspects of IDO-mediated regulation (4, 14, 17, 19). This alters the local cytokine environment to one that favors B cell autoimmunity and antibody production (17, 19) (Figure 1). This within the SCI lesion, the local environment is conducive to the support of B cells and antibody production.

**Potential new immunotherapies for SCI**

The results reported by Ankeny et al. (6) suggest opportunities for the development of new immunotherapies for SCI. Therapies that target the B cell or block the effects of pathogenic antibodies have demonstrated considerable promise in early phase I/II clinical trials (7, 10, 20). However, there are no FDA-approved, licensed anti–B cell products yet available for CNS autoimmune indications. Plasmapheresis is one approach that might improve neurological outcomes in SCI patients, by removing pathogenic autoantibodies at crucial times in the early stages of recovery, leading to a potential neuroprotective benefit. Whether plasmapheresis would lead to sustained improved neurological outcomes for SCI patients needs to be determined. Intravenous immunoglobulin (IVIG) administration has the potential for significant neuroprotection. IVIG is known to block, via an anti-idiotypic mechanism, the binding of pathogenic antibodies to their targets, and IVIG also blocks the complement system (20) (Figure 1). The results of Ankeny et al. (6) support the testing of IVIG for SCI treatment, as both autoantibody binding to spinal tissue and complement were demonstrated to be involved in SCI-associated autoantibody-induced pathology. IVIG therapy is generally well tolerated with few adverse side effects (20).

B cell depletion as a neuroprotective therapy is another viable option for SCI (Figure 1). Clinical trial results testing the beneficial effects of B cell depletion in various CNS autoimmune conditions support the approach, but with caution (7, 10, 12). Anti-CD20 antibody–mediated depletion of B cells is fast, effective, long-lasting, and largely free of adverse events in short-term applications (7, 10). Anti-CD20 does not block anamnestic antibody-mediated immune responses and is not immunosuppressive in that context (8). Long-term application of anti-CD20 therapy is still a concern (8) especially since SCI individuals are already immunosuppressed (3). For SCI, the real issue is whether anti-CD20 antibody–mediated B cell depletion will lead to improved outcomes. CD20 is expressed on early, mature, and activated B cells but not on antibody-secreting plasma cells (8). Consequently, long-term B cell depletion only has small to moderate affects on reducing plasma antibody levels and is not effective in eliminating B cells in follicles (8, 12). Pre-existing plasma cells that secret potentially pathogenic antibodies will not be eliminated by anti-CD20 therapy. It may be affective in preventing the appearance of de novo–created short-lived plasma cells by elimination of their precursors (7, 8). If antigen presentation or other B cell functions are critical to the induction of the SCI-induced pathogenic autoantibodies, then anti-CD20 therapy will likely be effective. These are important issues that need to be resolved. Alternatively, there are monoclonal antibody therapies being developed to block the actions of BAFF, lymphotoxin-β, and APRIL (8). Providing sufficient levels of neutralizing antibody could be delivered to target areas, preventing the development of B cell follicles in the SCI lesion area and in secondary lymphoid tissues may reduce the levels of autoantibody, such that the pathogenic consequences are minimal.

It remains to be determined whether pathogenic autoantibodies are a primary cause of human secondary SCI. However, there is now sufficient evidence to support the development of strategies, in appropriate animal models, to test whether potential immunotherapies restricting B cell responses following SCI warrant translation to the treatment of human SCI.
The aging process affects all aspects of the immune system, particularly the T cells. The immune system in older individuals is often characterized by lower T cell numbers, lower naive/memory T cell ratios, and lower T cell diversity. Most measures of inflammation increase with age. Why this happens, and why there is so much person-to-person variability in these changes, is not known. In this issue of the JCI, Sauce and colleagues show that removal of the thymus during infancy results in premature onset of many of these age-associated changes to the immune system (see the related article beginning on page 3070). The effect of thymectomy was particularly notable in those individuals who acquired CMV infection. Data from this study, as well as data from other observational settings, suggest that reduced thymic function and persistent viral infections combine to accelerate a decline in immunologic function.

The aging process affects all organs, including the immune system. Immunologic aging, generally referred to as “immunosenescence,” clearly affects T cell function, but changes also occur in B cells, antigen-presenting cells, NK cells, and perhaps even stem cells. With respect to T cells, the major age-associated changes include a decline in the total number of cells, a shift from a naive to a memory/effector T cell population, and a decline in CD8+ T cell receptor repertoire diversity. Generalized inflammation also increases with age, perhaps due to a loss of immunoregulatory function. These age-associated alterations result in immune compromise with potential susceptibilities to infection, autoimmunity, and neoplasia. Nearly all of the typical age-associated complications, including heart disease, cancer, infection, dementia, and frailty, have been epidemiologically linked (but not necessarily causally linked) to the process of immunosenescence.

Effect of thymectomy and CMV on immunosenescence

Although the pathogenesis of immunosenescence has been well studied in experimental models, very little mechanistic work has been performed in humans. In this issue of the JCI, Sauce and colleagues studied a group of young adults who as infants underwent complete removal of their thymus during the surgical correction of a congenital heart defect (1). Compared with age-matched controls, the thymectomized adults had a number of classic age-associated immunologic abnormalities, including lower total T cell counts and a preferential loss of naive T cells. These observations were not surprising, because the thymus is an estab-