engrafted into nonhuman primate hearts remains to be determined.

Despite these issues, the pioneering studies of Blin et al. are very encouraging. Their imaginative work using a nonhuman primate model of myocardial infarction to test the capacity of SSEA-1+ CPCs to engraft and differentiate into mature cardiac myocytes represents an important milestone. Future studies on the long-term survival, functional integration, physiological compatibility of engrafted cells, and beneficial effects on cardiac function will provide new insights into the potential use of SSEA-1+ CPCs for cardiovascular regenerative medicine. Most importantly, the ability to isolate nonhuman primate CPCs using a cell surface marker brings us one step closer to the ultimate dream of cell-based therapies for some of the most devastating forms of heart disease.

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Prevention trumps treatment of antibody-mediated transplant rejection
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Belying the spectacular success of solid organ transplantation and improvements in immunosuppressive therapy is the reality that long-term graft survival rates remain relatively unchanged, in large part due to chronic and insidious alloantibody-mediated graft injury. Half of heart transplant recipients develop chronic rejection within 10 years — a daunting statistic, particularly for young patients expecting to achieve longevity by enduring the rigors of a transplant. The current immunosuppressive pharmacopoeia is relatively ineffective in preventing late alloantibody-associated chronic rejection. In this issue of the JCI, Kelishadi et al. report that preemptive deletion of B cells prior to heart transplantation in cynomolgus monkeys, in addition to conventional posttransplant immunosuppressive therapy with cyclosporine, markedly attenuated not only acute graft rejection but also alloantibody elaboration and chronic graft rejection. The success of this preemptive strike implies a central role for B cells in graft rejection, and this approach may help to delay or prevent chronic rejection after solid organ transplantation.

Acute and chronic rejection

Newly transplanted organs are susceptible within a week to acute rejection, mediated dominantly by T cells, but are usually effectively protected from this form of inflammation and injury by currently used immunosuppressive agents such as calcineurin inhibitors, antiproliferative agents, mTOR inhibitors, and prophylactic therapy with T cell–specific antibodies. When acute rejection occurs, as it does in 5%–25% of solid organ recipients within the first year, it can typically be successfully treated with steroid therapy or, if needed, T cell–specific antibodies. However, the current immunosuppressive pharmacopoeia is relatively ineffective in preventing or treating rejection mediated by B cells and the antibodies they produce. Antibody-mediated allograft injury, which occurs in 50% of heart transplant patients within 10 years, typically manifests more than a year after transplantation, more insidiously than T cell–mediated injury, and in a process characterized by complement deposition and microvascular obliteration that leads to tissue ischemia and eventually fibrosis with loss of graft function. Chronic graft rejection refers to this antibody-mediated process.

While factors contributing to chronic injury of organ transplants are multiple and include ischemia/reperfusion injury, preexisting donor disease, drug toxicities, and recurrence of original disease, the...
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subtle development in the graft recipient of antibodies specific for the foreign donor tissue (alloantibodies) in the months and years following organ transplantation has been shown to be an accurate predictor of graft failure (1, 2). The magnitude of this problem is compounded by the practical difficulties in designing feasible clinical trials to evaluate methods for preventing alloantibody development and by the paucity of proven strategies to prevent alloantibody development in large animal models or humans. Nevertheless, data suggest that if preexisting alloantibody levels can be reduced, the risk of graft loss is lower (3).

B cell depletion as treatment for established antibody-mediated rejection

In the medical literature, organ transplant patients experiencing antibody-mediated rejection have been treated with rituximab (a CD20-specific monoclonal antibody that depletes the B cell population) or by targeting of their plasma cells (antibody-secreting differentiated B cells), and in most cases these patients possessed preexisting alloantibody or suffered from early antibody-mediated rejection (4, 5). As expected, it is difficult to reverse the damage done by alloantibody in the setting of an established B cell immune response, and the efficacy of targeting B cells with rituximab under these posttransplant circumstances has been difficult to clearly establish. The combination of B cell depletion with profound T cell immunosuppression may also be complicated by loss of protective immunity (6). In other words, infection or malignancy may ensue, especially when both T cell– and B cell–depleting antibodies are administered simultaneously or sequentially. Therefore, an alternative strategy, that being prevention as opposed to treatment of the B cell alloimmune response, even if resorting to B cell depletion, may be attractive.

Figure 1

B cell– and antibody-related biologics in transplantation. (i) CD20-specific mAb (i.e., rituximab) (anti-CD20), as reported in the current issue of the JCI by Kelishadi et al (7), binds and selectively depletes CD20+ B cells, thereby reducing alloantibody levels. Third generation CD20-specific mAbs are under development (e.g., ocrelizumab, ofatumumab). (ii) Inhibitors such as belimumab neutralize BAFF, while inhibitors such as atacicept (TACI-Ig) inhibit both BAFF and APRIL. (iii) Proteasome inhibitors (e.g., bortezomib) reversibly bind to the proteasome and disrupt various cell signaling pathways including the NF-κB pathway. (iv) Complement inhibitors, such as eculizumab (an antibody specific for complement component 5 [C5]), bind the complement protein C5, leading to cessation of complement-mediated cell lysis via the membrane attack complex (MAC). Since activation of the complement system is initiated by binding of 2 alloantibody molecules to a multivalent antigen followed by formation of the C1 complex, C1 inhibitor (C1-INH) prevents initiation of the serial complement cascade by inhibiting proteolytic cleavage of later complement components (specifically C2 and C4) and formation of C3 convertase. (v) Abatacept and belatacept (LEA29Y) are CTLA4-Ig molecules that bind the B7 costimulation molecule and block T cell costimulation of B cell activation and thereby production of alloantibodies. (vi) CD40-specific mAb (anti-CD40) binds the CD40 costimulation molecule. Blocking CD40L/CD40 interactions with CD40-specific antibody prevents T cell help to B cell activation, and consequently alloantibody production is inhibited.
Preemptive B cell depletion
In their study in this issue of the JCI, Kelishadi et al. (7) show that preemptive treatment of cynomolgus monkeys transplanted with an allogeneic heart with rituximab on the day of the transplant substantially eliminated the injury attributable to B cells. In particular, infiltration of the graft by B cells was markedly reduced, as were intragraft levels of B cell–activating factor (BAFF; also known as B lymphocyte survival factor [BlyS]) and the B cell costimulatory molecules CD80 and CD86. In addition, the downstream effects of B cell activation were attenuated; for example, levels of alloantibody in the blood were reduced and less complement deposition in the graft was observed. Perhaps most importantly, these mechanistic changes were reflected by a substantial improvement in the microvascular integrity of the transplanted hearts (i.e., there was less chronic allograft vasculopathy) and by improved cardiac function, with four of four hearts beating well by 90 days compared with only three of seven in cynomolgus monkeys treated with cyclosporine treatment alone, suggesting a role for B cells in acute rejection as well as chronic rejection.

Implications for human transplant patients
Therapeutic targeting of CD20 in transplantation may be appealing because of CD20’s stable expression primarily on B cells in the peripheral blood and its absence from plasma cells, pro-B cells, and hematopoietic stem cells, thus permitting maintenance of serum IgG levels and posttreatment recovery by spared pro-B and stem cells (8). For the same reasons, therapeutic targeting of CD20 may not be as effective in treating recipients known to have donor-specific alloantibody prior to transplantation, since memory B cells and plasma cells capable of producing antibody specific for the donor organ would already be primed. Since many T cell–mediated immune responses include a B cell component, the impact of B cell depletion may extend beyond suppression of measurable antibody (9), as is suggested by the observation in the current study that acute rejection was reduced from a 57% incidence in cynomolgus monkeys treated with cyclosporine alone to zero by addition of rituximab to the treatment regimen (7).

Nonhuman primate (NHP) models, such as the one used by Kelishadi et al. (7), are far closer, genetically, to the human condition than any rodent model might be, and thus the current report is expected to predict better than any rodent model of transplantation how humans might respond to B cell depletion. Nevertheless, it is worth noting that even observations in NHPs in the field of organ transplantation have sometimes been difficult to translate directly into the clinic (10, 11). By analogy, human heart transplant patients usually receive three or four simultaneous immunosuppressive agents to prevent T cell–mediated rejection, whereas the cynomolgus monkeys in the study by Kelishadi et al. received high-dose cyclosporine as their sole immunosuppressive agent (7). The applicability of the findings of the current study to human organ transplantation will therefore require rigorous testing in order to determine whether preemptive CD20 monoclonal antibody treatment in the setting of more intense T cell immunosuppression is accompanied by opportunistic infection.

Other B cell strategies for transplantation
Targeting B cell immunity without depleting these cells in order to prevent alloantibody development may also lead to opportunities to prevent allograft injury (Figure 1). Such strategies include targeting complement pathway components (12) and B cell cytokines and/or chemokines such as BAFF and/or a proliferation-inducing ligand (APRIL), which may influence both B and T cell responses (13, 14). Other biologics being considered for development for the targeting of B cell responses in the setting of transplantation are those that affect the costimulatory pathways. Interactions between CD28 on CD4+ T cells and CD80/CD86 on B cells, as well as between CD40 ligand (CD40L; also known as CD154) on activated CD4+ T cells and CD40 on B cells have been shown to participate in providing T cell help to B cells (15). The CD40/CD40L interaction stimulates B cell proliferation and isotype switching in the appropriate cytokine milieu (16, 17). CD28 and CTLA4 expression have also been shown to be involved in germinal center formation (18).

Each of these potential therapies is under active investigation. It will be important to compare the relative safety and efficacy of such strategies with that of profound B cell depletion with rituximab. Additionally, it will be necessary to determine the durability and need for repeated application of B cell therapy in the setting of constant exposure to alloantigen, as is the case with an organ transplant. Nevertheless, the current report by Kelishadi et al. (7) offers clear experimental evidence in a large animal model that B cell targeting in parallel with T cell inhibition can prevent alloantibody development and lead to improved long-term graft histology and better small blood vessel patency. Prevention of chronic rejection would represent a major advance for the field of transplantation, and prevention of alloantibody development is more likely to succeed than are strategies to reverse ongoing antibody-mediated graft injury.

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Pathogen-specific antibodies: codependent no longer

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Antibody-mediated defense against pathogens typically requires complex interactions between antibodies and other constituents of the humoral and cellular immune systems. However, recent evidence indicates that some antibodies alone can inhibit pathogen function in the absence of complement, phagocytes, or NK cells. In this issue of the JCI, McClelland et al. have begun to elucidate the molecular bases by which antibodies alone can impact pathogen growth and metabolism. They show that mAbs specific for the polysaccharide capsule of the human pathogenic fungus Cryptococcus neoformans elicit diverse effects on fungal gene expression, lipid biosynthesis, susceptibility to amphotericin B, cellular metabolism, and protein phosphorylation. These data suggest that pathogens have the capacity to generate broad metabolic responses as a result of surface binding by pathogen-specific antibodies, effects that may hold therapeutic promise.

Evolving concepts of antibody defense

The word immunity derives from the Latin “immunitas” meaning exemption and has come to mean protection from disease. Immunity has been observed over the centuries during plagues in Athens and Byzantium, epidemics of bubonic plague and smallpox, as well as with snake bites and vaccinations (1). The identification of specific pathogens in the late 19th century was associated with the concept that such organisms were ultimately inhibited by depleting their environment of required nutrients, by their own metabolic by-products, or by the inhospitability of infected tissues. Enter host defense. Initial conflicts arose between advocates of a predominately soluble or humoral basis for immunity and those favoring a cellular basis. These disparate viewpoints were ultimately reconciled in large part when antibodies, the key mediators of humoral immunity, were shown to rely on other soluble factors, particularly complement, and cells known as phagocytes to provide protection against and mediate resolution of infection. For its part, the microbe itself often expresses a range of protective defenses. These microbial virulence factors may bind, mask, or subvert the activity of antibodies by binding to their effector Fc constant regions (e.g., via staphylococcal protein A or streptococcal protein G) that otherwise direct pathogens to an Fc receptor–bearing phagocyte. The protective effects of antibodies are classically mediated through their specificity for the pathogen (facilitated via their variable regions) and the ability of their Fc constant region to act as a bridge or scaffold. Other host defense mechanisms (e.g., complement, phagocytes, and NK cells) use this foundation to induce the fatal injuries on the pathogen, on which antibody defense is dependent (Figure 1A).

However, in their study in this issue of the JCI, McClelland et al. advance an intriguing conceptual paradigm that binding of specific antibodies alone can elicit a range of metabolic perturbations in the fungal pathogen Cryptococcus neoformans (2). C. neoformans, widespread in the environment, is well-controlled and rarely symptomatic in healthy individuals, in large part because of antibody-dependent mechanisms. However, the organism causes recalcitrant disease and high mortality in patients with advanced cell-mediated immunodeficiency, such as those individuals with HIV/AIDS who have very low CD4+ T cell counts and patients who have undergone solid organ transplantation (3, 4). McClelland and colleagues show that three antibodies that bind to distinct topological sites on the polysaccharide capsule of C. neoformans elicit varying effects on its gene expression (2). The effects are direct and due to the antibodies in the absence of other soluble or cellular host elements, providing evidence that pathogens can recognize and respond to antibody binding by modulating distinct microbial genetic pathways (Figure 1B). These findings raise the intriguing possibility that the physiology of a pathogen and its susceptibility to clearance may be manipulated by rational antibody design.

Building on the past

Previous studies have revealed that, independent of the presence of complement or phagocytes, antibody-pathogen interactions can disrupt microbial integrity, although the genetic mechanism(s) remained undetermined (5–14). Antibodies

Conflict of interest: Edward N. Janoff’s laboratory has received research funding from Pfizer Pharmaceuticals (previously Wyeth Vaccines) and Vaxdinate Corporation for vaccine studies unrelated to Cryptococcus.

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