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Immunomodulators as an antimicrobial tool
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The spectrum of infectious diseases has shifted in the past 50 years to include those caused by microbes that cause disease predominantly in immunocompromised individuals. This phenomenon has underscored the dependence of microbial virulence on the immune status of the host. The limited efficacy of the available antimicrobial armamentarium in immunocompromised individuals, combined with increasing resistance to these agents, has led to an urgent need for new therapies for infectious diseases. Immunomodulators represent a novel approach to antimicrobial therapy that depends on bolstering host immunity, rather than direct antimicrobial activity. Immunomodulators can be divided into those that are specific to pathogens (pathogen-specific) and those that are not specific to pathogens (non-specific). However, to date only a few immunomodulators have been evaluated for their efficacy as antimicrobial tools.

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Introduction: immunomodulation in the context of the Damage-response framework
Immunomodulators are usually products of the immune system [1\(^\text{**}\)]. As such, it is useful to consider immunomodulation approaches to infectious diseases in the context of microbial pathogenesis. In contrast to microbe-centric views, in which microbial pathogenesis and virulence are considered to reflect singular microbial functions, the Damage-response framework provides a flexible construct that accounts for the contribution of the host, as well as the microbe, to these entities [2]. The Damage-response framework considers host damage to be the common denominator in microbial pathogenesis. Based on this tenet, host damage can be plotted against the host immune response as a U-shaped curve, whereby the maximal host damage resulting from a given host–microbe interaction occurs both when the immune response is weak and when it is overly strong (Figures 1 and 2). The inherent flexibility afforded by this curve lies in its ability to account for the fact that certain microbes only cause disease in certain hosts, a phenomenon that cannot be explained by views of microbial pathogenesis that consider virulence to be a singular microbial trait [2].

A logical corollary of the Damage-response framework is that infectious diseases only occur in susceptible hosts. Host immune mechanisms protect against infectious diseases by preventing or reducing the damage that can result from host–microbe interaction. The relationship between host immunity and microbial pathogenesis is clearly exemplified in immunocompromised hosts, by diseases that are caused by commensal microbes, such as *Candida albicans* and *Staphylococcus epidermidis*, and fungi, such as *Cryptococcus neoformans* and *Pneumocystis jiroveci*, and the success of immune reconstitution in preventing HIV-associated diseases caused by these microbes. The phenomenon of immune reconstitution disease that follows antiretroviral therapy with highly active anti-retroviral therapy, or HAART, in patients with AIDS (acquired immunodeficiency syndrome) illustrates how rebounding immunity can produce disease (discussed in [3]). Because the success of antimicrobial therapy is a function of its ability to ameliorate disease, and disease is a manifestation of host damage, the Damage-response framework provides a useful construct to consider approaches to treating infectious diseases that reduce host damage resulting from the host–microbe interaction.

The crisis in antimicrobial therapy, which has stemmed from antibiotic overuse, misuse and the limited number of new antimicrobial drugs on the near horizon is well documented [4\(^\text{**}\)]. However, another area that limits the utility of antimicrobial drug-based therapy is that antimicrobial agents are frequently ineffective in individuals with impaired immunity, often despite being highly active *in vitro* or in individuals with intact immunity. This underscores the crucial relationship between host immunity and microbial virulence and provides a powerful rationale for approaches to antimicrobial therapy that regulate the immune response to reduce, ameliorate or prevent host damage.

Immunomodulators as antimicrobial tools
Approaches to immunomodulation can be divided into those that are specific to pathogens (pathogen-specific) and those that are not (non-specific). Pathogen-specific
immunomodulators include antibody reagents and vaccines. With the exception of the rabies and varicella-zoster vaccines, currently licensed vaccines are administered to prevent acute infectious diseases rather than for therapy and are not discussed further here. Non-specific immunomodulators include cytokines, antimicrobial peptides, certain antimicrobial drugs and microbes such as probiotics. At present, clinical experience with non-specific immunomodulators as antimicrobial tools has been predominantly limited to cytokines.

**Pathogen-specific immunomodulators: antibody-based agents**

There are powerful historical precedents for the use of antibody-based therapies to treat infectious diseases (reviewed in [5]). The first era of antimicrobial therapy, early in the 20th century, was based on serum therapy with antibody preparations. Hence, the inaugural antimicrobial agents were immunomodulators [6]. First-generation antibody reagents were abandoned because of their toxicity, which was a result of their impurity and derivation from non-human species, and the arrival of antimicrobial drugs that acted directly on the microbe. Nonetheless, there was evidence for synergism between antibiotics and serum therapies [7]. Serum therapy was validated in animal models before being administered to humans [7]; however, the mechanism by which it ameliorated infectious diseases or enhanced the efficacy of antimicrobial drugs was largely unknown. For most of the 20th century, the mechanisms of antibody action that were thought to influence antibody efficacy included their ability to neutralize, promote opsonization and...
phagocytosis or antibody-dependent cell-mediated cytotoxicity (ADCC) and/or to activate complement [8]. However, a significantly more robust menu of potential mechanisms of antibody action, which includes direct antimicrobial action, immunomodulation and generation of oxidative species, has emerged over the past decade [8]. For example, the efficacy of a pneumococcal capsular polysaccharide-specific antibody was associated with modulation of the cellular response to pneumococcus in the lungs of mice with pulmonary infection [9**]. This finding suggested that the well-documented ability of serum therapy to ameliorate fever and other clinical symptoms of pneumococcal pneumonia [10] could have reflected a downregulation of the host inflammatory response (or damage control). Perhaps the observation that pneumococcal pneumonia was only responsive to serum therapy with the homologous capsular polysaccharides-specific antisera in the first three days of symptoms indicated that its capacity to mediate immunomodulation was limited to the early stages of disease. Antibodies to other microbes, including *C. neoformans* and *Histoplasma capsulatum* are able to modulate the cellular immune response to pulmonary infection (see [11*]). Studies of antibody action have shown that the older dichotomous view that antibody immunity was only effective against extracellular pathogens, whereas cellular immunity was responsible for immunity against intracellular pathogens has been deconstructed by evidence that antibody reagents can be effective against classical intracellular pathogens, such as *Mycobacterium tuberculosis*, *C. neoformans*, *H. capsulatum* and scores of viruses [11*].

**Monoclonal antibody-based agents**

Currently, there is only one antibody reagent licensed for use against an infectious disease in the United States — Palivizumab. Licensed in 1998, Palivizumab is a neutralizing, humanized monoclonal antibody (mAb) to protein F on respiratory syncytial virus (RSV) that reduced hospitalization for RSV in premature and other high-risk infants when given as prophylaxis [12]. Because the antiviral activity of Palivizumab was associated with a reduction in inflammatory mediator release in a murine model of RSV [13*], its mechanism of action probably involves immunomodulation. Despite ongoing controversies about the cost and target population of Palivizumab, its success in reducing the risk of RSV in high-risk infants promoted the development of second generation reagents and vaccine candidates [14].

Recently, Mycograb, a human recombinant antibody fragment was shown to significantly improve the response to amphotericin B in patients with invasive candidiasis [15***]. Patients who received Mycograb and amphotericin B showed a higher rate of complete overall response on day 10 of therapy, a significantly better mycological response and less *Candida*-attributable mortality than patients who received amphotericin B and a placebo. Mycograb was safe and well-tolerated; however, episodes of hypertension occurred more frequently in patients who received Mycograb than those who received a placebo. Mycograb is a recombinant antibody fragment lacking an Fc region, and is produced from a human anti-Hsp90 (heat-shock protein 90) cDNA library with an epitope that inhibits fungal Hsp90, NILKVIRKNIVKK [16]. The development of this antibody was driven by the observation that recovery from invasive candidiasis was associated with the appearance of antibodies to a 47 kDa determinant [17] that was found to be a fungal homolog of human Hsp90. Mycograb was tested in patients in comparison to standard therapy. As such, the question of whether or not its efficacy in *vivo* depends on synergy with antifungal drugs is unanswered. Nonetheless, the in *vitro* activity of Mycograb (with amphotericin B and other antifungal agents) against resistant *Candida* and other fungal species [18,19] suggests it could hold promise as a broadly active antifungal agent.

The first mAb used to treat a fungal disease in humans was the mouse mAb 18B7, which binds to the cryptococcal capsular polysaccharide glucuronoxylomannan [20***,21]. Extensive preclinical testing revealed that 18B7 augmented host defense mechanisms against *C. neoformans*, in *vitro* and in *vivo* (reviewed in [22]). In the clinical trial, administration of a single 1 mg kg⁻¹ dose of 18B7 to HIV-infected patients treated for cryptococcal meningitis was well-tolerated and was associated with a reduction in serum glucuronoxylomannan levels [20***]. The tolerability and promising effect of this reagent in HIV-infected patients bolsters the prospect that immunotherapeutic interventions have the potential to augment host immune mechanisms in the treatment of infectious diseases in immunocompromised individuals.

Now, mAbs have been developed against a myriad of microbes responsible for emerging infectious diseases and/or those that cause disease in the setting of immune impairment. A human mAb to *Bacillus anthracis* toxin has recently successfully completed Phase I trials and stands as a potentially useful therapeutic in the event of an anthrax biological attack [23*]. Several human mAbs to the SARS (severe acute respiratory syndrome) coronavirus have been developed that might be useful if the disease reappears [24*]. Remarkably, these mAbs were developed to the point that clinical use was possible in less than five years. Human mAbs were highly effective against experimental shiga-toxin producing *Escherichia coli* in piglets [25]. Studies in experimental models have revealed that the efficacy of certain mAbs depends on intact cellular immunity (see [11,26]). As such, the use of mAbs in immunocompromised patients could depend on whether its efficacy requires the immune function (a subset or element of) that is lacking in the relevant patient(s). mAbs have the advantage of homogeneity and high specific activity. Although there is concern that...
mAbs could have limited usefulness for microbes that demonstrate high antigenic variation and mutability, combinations of mAbs have shown promise in overcoming this limitation [27**].

**Polyclonal antibody-based agents**

Another type of antibody-based therapy for infectious diseases consists of polyclonal immunoglobulin-based agents, including intravenous immunoglobulin (IVIG) and specific immune globulins (sIgs, sometimes called hyperimmune globulin). Treatment and prevention of rabies depends entirely on the combination of two immunomodulators: rabies vaccine and rabies immune globulin [28]. The sIgs are the mainstay of managing exposures to viral agents in susceptible individuals who are not candidates for live vaccines, such as pregnant women and patients with impaired immunity [29]. A polyclonal preparation derived from individuals with high serum antibody titers to staphylococcal fibrinogen-binding proteins, serine aspartate dipeptide repeat G and clumping factor A is under clinical development [30*]. An evaluation of this preparation in very low-weight infants revealed a trend toward fewer staphylococcal and candidal infections [31]. The ability of preformed antibodies to provide immediate defense against infectious diseases in susceptible individuals provides a potent justification for the use of antibodies, for example, in the setting of an act of bioterrorism or of epidemic diseases [32].

The use of IVIG in infectious diseases remains controversial. IVIG has also been shown to be useful for the treatment of Kawasaki Disease, cytomegalovirus pneumonitis in organ transplant recipients [33*] and parvovirus in patients with HIV infection [34]. IVIG is also useful in patients with toxic shock syndrome [35], West Nile virus infection [36] and sepsis [37], and is also invaluable in the management of patients with hypogammaglobulinemia who are at increased risk for infectious diseases, such as enteroviral meningitis [33*].

**Non-specific immunomodulators: cytokines as antimicrobial tools**

Cytokine-based therapies contrast with antibody reagents in that they are not pathogen-specific. The rationale for the use of cytokines as adjunctive immunomodulators for infectious diseases is based on the concept that replacement or augmentation of natural mediators of host defense should enhance the antimicrobial effect of host immune mechanisms and/or antimicrobial agents. Despite the logical basis for this concept, the potential antimicrobial power of these agents has been difficult to harness clinically. Currently, there are only a few examples of the use of adjunctive cytokines against infectious diseases. Notable exceptions are the use of recombinant α-interferons and nucleoside analogs for hepatitis B virus (HBV) and pegylated interferons and ribavirin for hepatitis C virus (HCV) [38]. The efficacy of interferons against HCV has been attributed to the induction of Th1 immunity [39]. A side effect of interferon-based therapies that limits their use in certain patients is depression [40*].

The efficacy of adjunctive interferon-gamma 1b (IFN–γ1b) with amphotericin B was studied in a Phase II, double-blind placebo-controlled trial for AIDS-associated cryptococcal meningitis [41***]. There was a trend towards mycological response and clinical improvement among interferon recipients, with 26% showing improvement, compared to 8% of placebo-controlled subjects. Although this difference did not reach statistical significance, the trend towards a beneficial effect of adjunctive interferon is encouraging, calling for further, larger scale studies and studies to identify those patients in whom adjunctive immunotherapy could be beneficial. The rationale for interferon therapy for cryptococcosis has a strong basis in preclinical studies in mice [42] and a human study showing an association between cerebrospinal fluid levels of IFN–γ and treatment in HIV-infected patients with cryptococcal meningitis [43*]. In light of the established benefit of interferon therapy for the prevention of bacterial diseases in patients with chronic granulomatous disease [44], adjunctive interferon could hold promise as an adjunctive agent for HIV-associated cryptococcal meningitis. However, the absence of surrogate markers that can predict the patients who would benefit from interferon therapy underscores the potential pitfalls in study-design and patient selection for clinical trials. This is particularly problematic for cryptoccocal meningitis, a disease that can occur in the case of weak or reconstituted immunity in patients with HIV infection [3]. The failure to demonstrate the effect of a pro-inflammatory immunomodulator could reflect the induction of an excessive inflammatory response that promotes disease. Figure 1 provides a schematic interpretation of the use of IFN–γ therapy in the context of the Damage-response framework. Depending on the immunological status of the affected patient, adjunctive IFN–γ therapy could be beneficial or detrimental.

Preclinical data, demonstrating the importance of Th1 helper T-cell responses in protection against fungi, in experimental models has led to the proposal that adjunctive cytokines be used with antifungal agents for invasive fungal infections [45]. The rationale for the use of colony stimulating factors (CSFs), derived from granulocytes (G-CSF) or macrophages (GM-CSF), is in part based on their ability to alleviate neutropenia [45]. Data on the clinical use of adjunctive CSFs for fungal diseases is limited to small studies or case reports (see [45] for review). Nonetheless, administration of GM-CSF with antifungal agents in patients with invasive fungal infections was associated with decreased patient mortality or better response rates compared to a placebo [46] or antifungal therapy alone [47]. In view of the small number
of affected patients and medical and ethical considerations in the design of randomized, double-blind placebo-controlled studies, it is possible that safe, well-tolerated candidate immunomodulators that are beneficial in preclinical studies will find their way into the antimicrobial armamentarium through compassionate use protocols and salvage therapy [48**].

**Anti-inflammatory immunomodulators**

The rationale for the use of adjunctive pro-inflammatory cytokines and certain antibodies for treating infectious diseases is to enhance the host response. However, a relatively underappreciated principle of microbial pathogenesis is that the damage resulting from host-microbe interaction can be the result of an overly exuberant host response [2]. Hence, there is a rationale for use of therapeutic interventions that dampen or reduce, as well as those that enhance or augment, the inflammatory response. It has been proposed that the beneficial effect of IVIG against inflammatory diseases involves engagement of the inhibitory Fc receptor, which downregulates the inflammatory response [49]. Probiotics, which are live bacteria derived from the human gastrointestinal tract, have been used as therapy for inflammatory bowel diseases, including antibiotic-associated diarrhea [50]. Probiotics remain outside the established antimicrobial armamentarium and are fraught with the potential for harm in immunocompromised hosts [51]. Nonetheless, it is logical to postulate that replacement and/or enhanced activity of the human microbiota could be beneficial for treatment of infectious diseases that reflect a failure of mucosal surfaces or of the innate mechanisms they bolster to protect against microbe-mediated damage (see [1**]).

Corticosteroids are important anti-inflammatory agents. Despite decades of controversy, corticosteroids have been validated as an important adjunct to antimicrobial therapy for bacterial meningitis [52*], HIV-associated Pneumocystis pneumonia [53] and tuberculosis meningitis [54*]. However, their mechanism of efficacy in these diseases might not be a direct anti-inflammatory effect [55]. Intriguingly, agents which have been proposed to have anti-inflammatory effects are macrolide antibiotics [56*]. The ability of macrolides to suppress the release of inflammatory mediators from phagocytes in vitro has been recognized for some time, but the clinical relevance of this phenomenon remains uncertain. Nonetheless, it has been hypothesized that the benefit of low-dose macrolide therapy in chronic pulmonary diseases, could be as a result of macrolide-induced reductions in levels of inflammatory cytokines, neutrophil recruitment and biofilm formation, which have been observed in animal models and/or in vitro [57]. In contrast to macrolides, amphotericin B [58] and penicillin [59] induce the release of inflammatory mediators in vitro through Toll-like receptor 2 (TLR2)-mediated stimulation. The inflammatory properties of these drugs raise the question of whether their immunomodulatory properties have an independent contribution to their therapeutic efficacy.

**Conclusions: the future of adjunctive immunomodulators as antimicrobial tools**

The future use of adjunctive immunomodulators for infectious diseases requires a better understanding of microbial pathogenesis and the relative need for immune activation versus immune modulation in the context of the immune response of the affected individual. In light of the fact that certain infectious diseases reflect an insufficient response, whereas others reflect an overly exuberant response, different types of interventions are likely to be required, depending on the immune status of the patient (Figure 2). The Damage-response framework can be a useful tool when considering the type of intervention that might be successful, but such predictions require experimental validation to be translated to the bedside and clinic.

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