Cardiac allograft immune activation: current perspectives

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Abstract: Heart transplant remains the most durable option for end-stage heart disease. Cardiac allograft immune activation and heart transplant rejection remain among the main complications limiting graft and recipient survival. Mediators of the immune system can cause different forms of rejection post-heart transplant. Types of heart transplant rejection include hyperacute rejection, cellular rejection, antibody-mediated rejection, and chronic rejection. In this review, we will summarize the innate and adaptive immune responses which influence the post-heart transplant recipient. Different forms of rejection and their clinical presentation, detection, and immune monitoring will be discussed. Treatment of heart transplant rejection will be examined. We will discuss potential treatment strategies for preventing rejection post-transplant in immunologically high-risk patients with antibody sensitization.

Keywords: heart transplant, innate immunity, adaptive immunity, rejection, immunosuppression

Introduction
The burden of heart failure continues to grow, with an excess of 5 million patients in the United States with this clinical syndrome. End-stage, or stage D, heart failure affects approximately 5% of this population, in which 1-year mortality can be as high as 75% with medical therapy alone. Options for stage D heart failure are limited, but include guideline-directed medical therapy, palliative care (including chronic inotropic therapy), mechanical circulatory support, heart transplant, and experimental therapies, such as stem cell trials. Heart transplantation remains the most durable treatment for end-stage heart disease that is not amenable to other treatment modalities, such as anti-ischemic or antiarrhythmic therapies. In a recent report from the International Society for Heart and Lung Transplantation (ISHLT), the median survival after heart transplant is 11 years. According to this report, however, less than 2,500 heart transplant surgeries are completed each year in North America. Thus, donor availability for heart transplant is far below the need from potential heart transplant recipients. For this reason, left ventricular assist devices and cardiac replacement therapies, such as the Total Artificial Heart (SynCardia Systems Inc., Tucson, AZ, USA), can be used as a bridge to heart transplant. These devices are being utilized with increasing frequency prior to heart transplant as approximately 33% of heart transplant recipients had a mechanical device as a bridge to transplant. Left ventricular assist devices and Total Artificial Heart therapies offer promise to patients who need mechanical support prior to heart transplant. Complications after placement of mechanical circulatory support include infection and blood transfusion. These factors, among others, can lead
Mechanisms of immune responses

The innate and adaptive immune systems work in conjunction to ward off pathologic infection. In dysregulation, these systems can also cause autoimmune disease in non-immunosuppressed patients or cause rejection in solid organ transplant recipients. While there is significant interaction between the innate and adaptive immune systems, to date, modulation of the adaptive immune system has been the focus of chronic immunosuppression post-transplant. The role of the innate immune system post-transplant is a topic of current interest in translational research.

Components of the innate immune system include leukocytes (neutrophils, eosinophils, and mastocytes), macrophages, dendritic cells, and natural killer (NK) cells. Cells of the innate immune system interact against a defined set of molecules including pathogen-associated molecular patterns in the context of infection and damage-associated molecular patterns in the process of tissue injury. Toll-like receptors (TLRs) are one type of pattern-recognition receptor (PRR) involved in initiation and amplification of innate immune responses. Cells of the innate immune system, including dendritic cells, can recognize TLRs. Different TLRs are expressed on different cell types (endothelial or epithelial cells) or in intracellular compartments. TLRs assist in early detection of microbes, but can also recognize markers of cell damage, including heat shock proteins and nucleic acids released by necrotic cells. TLR activation is amplified through signaling, primarily through myeloid differentiation factor 88 (MyD88). Recognition of TLR by dendritic cells aids in the maturation of the dendritic cell, allowing dendritic cell interactions with effector T lymphocytes. In this manner, stimulation of the innate immune system can, in turn, activate the adaptive immune system. Cytokines and chemokines are released in response to activation of the innate immune system via TLR signaling. Chemokine release can, in turn, recruit mediators of the adaptive immune response, including leukocytes, to sites of tissue injury. At the time of heart transplant, reperfusion injury and tissue injury can activate such components of the innate immune system to stimulate cytokine release to activate the complement cascade leading to further cell injury and cell death. The complement cascade has been implicated in acute and chronic rejection processes. The C5a and C3a components of the complement cascade may assist in T-cell co-stimulation. NK cells impact the innate immune response by recognizing foreign cells in the absence of antigen-specific interactions. In this manner, NK cells can directly kill donor cells. Antigens from deceased donor cells can be processed by antigen-presenting cells (APCs), leading to indirect stimulation of the adaptive immune system by the NK cell.

Primary mediators in the adaptive immune system include T lymphocytes and B lymphocytes. T lymphocytes develop in the thymus through processes of positive and negative selection. T-cells can be subdivided into cytotoxic T-cells (CD8+ T-cells) and CD4+ T-helper cells. The CD4+ T-helper cells can function as effector or regulatory T-cells. Differential cytokine milieu can influence differing CD4 T-cell phenotypes. Interferon (IFN)-gamma and interleukin (IL)-12 promote Th1 cells that are important in host defense mechanisms against intracellular pathogens. IL-4 drives Th2 CD4 T-cells that combat extracellular parasites. IL-6 and transforming growth factor (TGF)-beta lead to Th17 cells that combat yeast, fungi, and extracellular bacteria. Th17 cells can also influence autoimmune diseases including multiple sclerosis, psoriasis, rheumatoid arthritis, and Crohn’s disease. Tfh cells, or T follicular helper cells, aid germinal center B-cells in plasma cell differentiation. B lymphocytes develop in the spleen and lymph nodes. B-cells are the primary antibody-producing cells of the adaptive immune system. Both T- and B-cells can proliferate in response to activation and form a memory response. With repeat activation, the response is augmented and amplified. T-cells interact with B-cells and other APCs (including macrophage and dendritic cell lines of the innate immune system) via the MHC class II cell surface receptor on the APC and the T-cell receptor (TCR) of the T-cell. MHC I antigens are constitutively expressed on a majority of cells and help delineate self versus non-self (non-self or foreign protein/s that would elicit an immune response). TCR engagement of class I or class II MHC leads to T-cell activation. Co-stimulation via B7-CD28 and CD40-CD40L interactions between APCs and T-cells are also required for T-cell activation. These activated T-cells are a main mediator of the adaptive immune response and contribute to heart transplant rejection. In fact, most standard chronic immunosuppression post-heart transplant is directed at suppressing T-cell activation via suppressing T-cell signaling or T-cell proliferation.

Activated effector T- and B-cells are pathologic to the transplanted graft. Regulatory T-cells may serve to downregulate an activated immune system and antagonize the process
of allograft immune activation. There are numerous types of regulatory cells. Regulatory T-cells include CD4+ T-cells that express regulatory transcription factor forkhead box (Fox) p3 (Tregs), CD8+Foxp3+ Tregs, CD4–CD8– T-cells, and NK T-cells.13 Tregs in humans can express Foxp3, Fas receptor (CD95), and cytotoxic T-lymphocyte antigen 4 (CTLA4). These Tregs are of considerable interest as they can exert potent immunoregulatory effects, including APC killing, consumption of IL-2, and downregulation of the immune system. Degranulation of mastocytes can lead to loss of Tregs and can contribute to heart transplant rejection.14

Heart transplant rejection

Post-heart transplant, mediators of innate and adaptive immunity influence the possible development of rejection. The risk of rejection is highest in the first year post-heart transplant.15 Risk of rejection in the first year post-transplant has decreased from 33% of recipients in 2004 to 25% of recipients in 2010. Moreover, rejection requiring treatment has decreased from 25% in 2004 to 14% in 2010.3 Hyperacute rejection, acute cellular rejection (ACR), acute antibody-mediated (humoral) rejection, and chronic rejection can affect the functional status and longevity of the heart transplant recipient. The main goal of immunosuppression is to prevent heart transplant rejection. Further understanding of the mediators of rejection will aid in prevention and treatment of these disease processes.

Hyperacute rejection can occur immediately post-heart transplant and is manifest in critical cardiogenic shock and profound organ hypoperfusion. Diagnosis is made usually on clinical suspicion, and mechanical support with extracorporeal membrane oxygenation is often required along with aggressive antirejection therapies. This process is difficult to distinguish from severe primary graft dysfunction. Hyperacute rejection is thought to be an immunological process caused by preformed recipient antibodies to the graft. It has been reported in ABO blood type incompatibility as well.16 Hyperacute rejection is currently a rare clinical scenario due to the development of the prospective cytotoxic crossmatch17,18 and, more recently, use of the virtual crossmatch.19,20 The prospective cytotoxic crossmatch, or complement-dependent, cytotoxicity-based assay, combines recipient serum (potential source of donor-specific antibodies [DSA]) with donor lymphocytes. Cytotoxicity with addition of exogenous complement represents a positive crossmatch and significant DSA against donor class I and/or class II HLA. The prospective crossmatch is still used in select highly sensitized recipients, but, more frequently, the virtual crossmatch is utilized, as the prospective crossmatch requires local expertise and is time-consuming. The virtual crossmatch utilizes information from solid-phase assays that allow detection of the specificity and binding strength of an antibody.21 With use of these assays, the immunologic MHC class I and II status of the recipient and donor can be virtually matched for compatibility. Prior to transplant, the recipient’s probability of potential donor compatibility can be calculated against a panel of reactive antibodies.22 Calculated panel reactive antibody (cPRA) data are based on allele frequencies from HLA phenotypes of deceased kidney donors. While there are differing definitions of antibody sensitization pre-transplant, a patient with a cPRA >10% may be considered sensitized. Highly sensitized individuals have cPRA >50% and may require desensitization treatment prior to transplant.23 A novel assay to detect the potential functional significance of antibodies has been described. This C1q assay detects a subset of IgG antibodies that are capable of fixing complement. The C1q assay was useful in prediction of C1q+ DSA contributing to antibody mediated rejection (AMR) episodes and graft loss following kidney transplant.24

ACR is primarily a T-cell-mediated process with graft infiltration by leukocytes and macrophages. ACR requiring treatment may present insidiously. It may also manifest with new-onset symptomatic heart failure, with rhythm disturbances, or with cardiogenic shock. The most common rhythm disorder seen in acute rejection is atrial fibrillation. Bradycardia in the setting of heart transplant rejection is an ominous finding. The diagnosis of cellular rejection is definitively made by endomyocardial biopsy.25 Sampling error may lead to false negative biopsy results. Endomyocardial biopsy, however, remains the gold standard in the diagnosis of heart transplant rejection. Grading of cellular rejection is based on standardized guidelines (see Table 1).26 In this grading scale, grade 2 R and 3 R cellular rejections generally require treatment. Decrement in left ventricular systolic function by echocardiogram is highly suggestive of rejection as well. Subtherapeutic immunosuppressant trough levels predispose recipients to cellular rejection. Inadequate immunosuppression may be due to medication noncompliance, nonadherence to medications due to side effect or cost, poor medication absorption, infection, or drug interactions. Risk factors for ACR include a higher number of HLA mismatches, age of recipient, female sex, race, and use of induction therapy.27 Young patients are at higher risk of rejection, and African-American recipients may have a genetic predisposition to require higher doses of effective immunosuppressant therapy, possibly due to polymorphisms in cytochrome P450 enzymes.
Histologic findings are present and immunopathologic findings are negative—Negative for pathologic AMR

Diffuse infiltrate with multifocal myocyte damage ± edema, ± hemorrhage ± vasculitis

Notes: "R" denotes revised grade to avoid confusion with 1990 scheme. Reprint from the Journal of Heart and Lung Transplantation, 24(11), Stewart S. Winters GL, Fishbein MC, et al, Revision of the 1990 working formulation for the standardization of nomenclature in the diagnosis of heart rejection, 1710–1720, Copyright © 2005, with permission from the International Society for Heart and Lung Transplantation.

Abbreviation: ISHLT, International Society for Heart and Lung Transplantation.

Table 1: ISHLT standardized cardiac biopsy grading: acute cellular rejection

<table>
<thead>
<tr>
<th>Grade</th>
<th>Definition</th>
<th>Substrates</th>
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<tbody>
<tr>
<td>Grade 0 R‡</td>
<td>No rejection</td>
<td>Histologic and immunopathologic studies are both negative</td>
</tr>
<tr>
<td>Grade 1 R, mild</td>
<td>Interstitial and/or perivascular infiltrate with up to 1 focus of myocyte damage</td>
<td>Histologic findings are present and immunopathologic findings are negative</td>
</tr>
<tr>
<td>Grade 2 R, moderate</td>
<td>Two or more foci of infiltrate with associated myocyte damage</td>
<td>Histologic findings are negative and immunopathologic findings are positive (CD68+ and/or C4d+)</td>
</tr>
<tr>
<td>Grade 3 R, severe</td>
<td>Diffuse infiltrate with multifocal myocyte damage ± edema, ± hemorrhage ± vasculitis</td>
<td>Histologic and immunopathologic findings are both present</td>
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Notes: Histologic changes include interstitial capillary injury and activated mononuclear cells, which are characterized by endothelial cell swelling and intravascular macrophage accumulation. Severe AMR is characterized by hemorrhage, neutrophilic or mixed inflammatory cell infiltrates, intravascular thrombus, and myocyte necrosis. Reprint from the Journal of Heart and Lung Transplantation, 32(12), Berry GC, Burke MM, Andersen C et al, The 2013 International Society for Heart and Lung Transplantation Working Formulation for the standardization of nomenclature in the pathologic diagnosis of antibody-mediated rejection in heart transplantation, 1147–1162, Copyright © 2013, with permission from the International Society for Heart and Lung Transplantation.

Abbreviations: AMR, antibody-mediated rejection; CD68+, cluster of differentiation 68; C4d+, complement factor 4 deposition; H+, histopathologic; I+, immunopathologic; ISHLT, International Society for Heart and Lung Transplantation; pAMR, pathologic AMR.

Table 2: The 2013 ISHLT working formulation for pathologic diagnosis of cardiac antibody-mediated rejection

<table>
<thead>
<tr>
<th>Grade</th>
<th>Definition</th>
<th>Substrates</th>
</tr>
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<tbody>
<tr>
<td>pAMR0</td>
<td>Negative for pathologic AMR</td>
<td>Histologic and immunopathologic studies are both negative</td>
</tr>
<tr>
<td>pAMR1 (H+)</td>
<td>Histopathologic AMR alone</td>
<td>Histologic findings are present and immunopathologic findings are negative</td>
</tr>
<tr>
<td>pAMR1 (I+)</td>
<td>Immunopathologic AMR alone</td>
<td>Histologic findings are negative and immunopathologic findings are positive (CD68+ and/or C4d+)</td>
</tr>
<tr>
<td>pAMR2</td>
<td>Pathologic AMR</td>
<td>Histologic and immunopathologic findings are both present</td>
</tr>
<tr>
<td>pAMR3</td>
<td>Severe pathologic AMR</td>
<td>Intertitial hemorrhage, capillary fragmentation, mixed inflammatory infiltrates, endothelial cell pyknosis, and/or karyorrhexis, and marked edema and immunopathologic findings are present. These cases may be associated with profound hemodynamic dysfunction and poor clinical outcomes</td>
</tr>
</tbody>
</table>

Notes: Histologic changes include interstitial capillary injury and activated mononuclear cells, which are characterized by endothelial cell swelling and intravascular macrophage accumulation. Severe AMR is characterized by hemorrhage, neutrophilic or mixed inflammatory cell infiltrates, intravascular thrombus, and myocyte necrosis. Reprint from the Journal of Heart and Lung Transplantation, 32(12), Berry GC, Burke MM, Andersen C et al, The 2013 International Society for Heart and Lung Transplantation Working Formulation for the standardization of nomenclature in the pathologic diagnosis of antibody-mediated rejection in heart transplantation, 1147–1162, Copyright © 2013, with permission from the International Society for Heart and Lung Transplantation.

Abbreviations: AMR, antibody-mediated rejection; CD68+, cluster of differentiation 68; C4d+, complement factor 4 deposition; H+, histopathologic; I+, immunopathologic; ISHLT, International Society for Heart and Lung Transplantation; pAMR, pathologic AMR.
are unlikely to present with angina. Routine angiography should be considered to evaluate for presence and severity of CAV. Conventional coronary angiography remains the gold standard with respect to assessment of coronary vasculature. Alternatives to the invasive coronary angiogram include exercise-based or pharmacologic stress testing, cardiac positron emission tomography, and coronary computed tomography angiography. Additional modalities, such as optical coherence tomography, may augment conventional coronary angiographic and IVUS data in the evaluation of CAV.

**Detection of rejection and immune monitoring**

The gold standard for diagnosis of ACR or AMR remains the assessment of the endomyocardial biopsy. Cardiac magnetic resonance imaging has shown promise in diagnosis of rejection. This modality may be particularly useful in cases of biopsy-negative rejection. As clinical and echocardiographic manifestations may present late, many transplant centers will have a protocol-based schedule for endomyocardial biopsy with right heart catheterization to routinely evaluate for ACR, AMR, and cardiac hemodynamics. Biopsies are most frequent in the first month post-transplant and taper gradually in time. When rejection is present and treated, our institutional practice involves follow-up biopsy within 2 weeks to ensure that the findings on biopsy improve or normalize after treatment. Routine assessment post-biopsy with echocardiography will give further information regarding cardiac function and structure. Data are limited regarding accurate diagnosis of heart transplant rejection by noninvasive echocardiographic-based methods.

To reduce patient discomfort and anxiety due to the invasive nature of endomyocardial biopsy, there are alternative tests that can be performed to assess for heart transplant rejection. The most commonly used noninvasive blood test used to assess the possibility of ACR in low-risk individuals is the AlloMap® (CareDx, Inc., Brisbane, CA, USA). This test uses gene expression profiling and was first studied between 6 months and 5 years post-transplant in patients at low risk for rejection. The test has a reported excellent negative predictive value, but the positive predictive value is low. Abnormal AlloMap studies prompt a safety endomyocardial biopsy to further assess for cellular rejection. Use of the AlloMap in the first year post-transplant has also been examined. There are a number of limitations of the AlloMap. Its use has been studied in a low-risk patient population in single-organ (heart) transplant. It is important to note that the AlloMap does not assess AMR. Hemodynamics from a right heart catheterization, which would routinely follow an endomyocardial biopsy, are not obtained. Finally, an abnormal AlloMap would prompt a safety endomyocardial biopsy. Despite its limitations, in appropriately selecting patients, the AlloMap test does have clinical utility in the assessment of cellular rejection.

Another noninvasive assay with clinical utility to monitor levels of immunosuppression is the Cylex test (ImmuKnow®, Cylex Inc., Columbia, MD, USA). The Cylex test assesses adenosine triphosphate production from phytohemagglutinin (PHA)-stimulated T-cells. Low Cylex values can suggest
over-immunosuppression and risk of infection. Elevated values of the Cylex score in theory may reflect under-immunosuppression and risk of rejection. Elevated Cylex values have been reported to correlate with increased plaque progression by IVUS even at 2 months post-transplant, which suggests that elevated Cylex scores could potentially signal risk of AMR, given the association of AMR with CAV.

**Chronic maintenance immunosuppression and treatment of rejection**

Chronic maintenance immunosuppressant therapy post-heart transplant generally involves therapies directed to suppress the T-cell. These therapies include a combination of calcineurin inhibition (CNI) with tacrolimus (Prograf®; Astellas Pharma Inc., Tokyo, Japan) or cyclosporine (Gengraf®; Abbvie Inc., North Chicago, IL, USA) to inhibit signaling downstream of the TCR; purine antagonism with mycophenolate mofetil or azathioprine to attenuate T-cell proliferation; and pan-immunosuppressant steroid therapy. Ideally, patients can be weaned off prednisone in the first year post-heart transplant should they maintain normal cardiac function, be free of rejection, and not have development of DSA. CNI monotherapy has been used to attempt to minimize the adverse sequelae of immunosuppression. With monotherapy-based immunosuppression, target trough levels may be targeted in a higher range, which may lead to differences in the side effect profiles observed. For example, in CNI monotherapy, higher target trough levels could, in theory, lead to renal insufficiency. In combination immunosuppressive therapy, should a patient have an episode of rejection, one could consider transition from purine antagonist to proliferation signal inhibition (PSI) with sirolimus (Rapamune®; Wyeth Pharmaceuticals Inc., Philadelphia, PA, USA) or everolimus (Zortress®; Novartis International AG, Basel, Switzerland). This transition may reduce the risk of rejection given the results of a clinical trial comparing three arms of immunosuppressant therapies. A recent trial reported success with early PSI use and CNI withdrawal within 3 months post-heart transplant. The risk/benefit profile of purine antagonist versus PSI therapy must be weighed on a case-by-case basis, as some clinical trials for immunosuppression with PSI have shown possible deleterious effects on renal function and risk of infection.

Treatment of rejection depends on patient symptoms and the severity of the rejection episode. With treatment for rejection, there is always a balance between augmented immunosuppression with potential for improvement in graft function and the risk of infection. Treatment for AMR in asymptomatic patients with preserved cardiac function is controversial, as its benefit has not been proven. ACR can be treated with oral or intravenous (IV) steroid bolus and taper and CNI target levels can be increased. For patients experiencing ACR and AMR with reduced cardiac function or with clinical symptoms, pulse-dose IV steroids, rabbit anti-thymocyte globulin (ATG), and/or IV immunoglobulin (IVIG) can be utilized. ATG is a cytolytic antibody that binds the CD3 moiety associated with the TCR, leading to T-cell depletion. The mechanism of IVIG is not fully understood, but is likely pleomorphic. IVIG can affect immune activation by neutralization of pathologic antibodies via anti-idiotypic interactions, competition for fragment, crystallizable portion of an antibody (Fc) binding sites, and inhibition of complement activity. Patients that present with heart failure or cardiogenic shock may have hyperacute rejection immediately post-transplant, delayed hyperacute rejection within 1 week post-transplant, or, later, a combination of ACR and AMR. These patients are at high risk for significant morbidity and death and are treated with IV pulse steroids as a pan-immunosuppressant, plasmapheresis to remove pathologic antibodies, cytolytic therapy with ATG to deplete T-cells, and IVIG. There are limited data supporting the use of B-cell-targeted therapies in acute AMR. Treatment of acute rejection with bortezomib (Velcade®; Millenium Pharmaceuticals, Inc., Cambridge, MA, USA) in conjunction with plasmapheresis is another option that has not been studied in detail in heart transplant recipients, but has shown benefit in renal transplant patients with ACR and AMR. Bortezomib is a proteasome inhibitor that selectively depletes antibody-producing plasma cells. Patients that are critically ill due to rejection require close hemodynamic monitoring and may require directed inotropic therapy, mechanical support with an intra-aortic balloon pump, or potentially, extracorporeal membrane oxygenation support. With aggressive therapy, there is hope that the immune activation process will be downregulated and that cardiac function will normalize (Table 4).

After treatment for rejection, patients who develop DSA may benefit from further treatment with IVIG and rituximab (Rituxan®; Genentech Inc., South San Francisco, CA, USA), a humanized mouse monoclonal antibody that depletes CD20+ B-cells. One such approach would use IVIG at 1 g/kg daily for 2 days, followed by rituximab therapy commenced at 1 week (1 g or 375 mg/m² if the patient is <50 kg). Repeat IVIG dosing can be commenced at 1 month. These therapies need to be temporally separated, as IVIG may bind to and attenuate the efficacy of rituximab. Serial flow cytometric HLA should be assessed 2 weeks after treatment. After
treatment for acute rejection, for patients that have recurrent or persistent rejection on endomyocardial biopsy, additional intense immunosuppressant treatment may be warranted. Further treatments of recurrent or persistent rejection include total lymphoid radiation and photopheresis. Total lymphoid radiation is not often used due to concerns for future development of malignancy. Photopheresis has been used for immunomodulation. The process involves treatment of a minority (~5%) of peripheral lymphocytes with 8-methoxypsoralen and ultraviolet light, inducing T-cell apoptosis. This process of T-cell depletion may have an impact on Tregs. Photopheresis, in contrast to other modalities of antirejection treatment, may not increase infectious risk. Patients with rejection are not candidates for repeat heart transplant within 6 months of an episode of rejection as the immunologic milieu of the patient may acutely reject an organ upon retransplantation. Rarely, durable mechanical assist devices are used as a bridge to recovery or retransplantation.

**Treatment of immunologically high-risk sensitized patients prior to and after heart transplant**

While immunosuppressant therapies are required post-heart transplant to prevent or treat cardiac allograft activation, the use of immunomodulating therapies prior to heart transplant is under investigation. There is evidence that supports desensitization strategies prior to renal transplant with IVIG and rituximab.\(^1\)\(^2\) Prior to heart transplant, patients with elevated cPRA may benefit from desensitization therapies to potentially reduce the risk of post-transplant rejection and expand the number of compatible donor candidates available for heart transplant. As described above for patients who develop DSA post-transplant, IVIG and rituximab can be used in a desensitization protocol prior to heart transplant.

For patients that do not reduce their antibody burden with IVIG and rituximab therapy, a pilot study showed utility of plasmapheresis and bortezomib in reduction of cPRA.\(^3\) Use of plasmapheresis and bortezomib in this context followed the protocol used for treatment of acute rejection post-kidney transplant. Infectious risk may be higher in patients who undergo desensitization treatment. It is uncertain if the benefit of therapy outweighs the risk of infection due to desensitization strategies.

Immediately post-heart transplant, induction therapy can be used for sensitized patients to potentially reduce their risk for rejection. A number of different modalities have been used in an induction strategy. Common induction agents include T-cell-depleting agents, including ATG, and IL-2 receptor antagonists, including basiliximab (Simulect®; Novartis International AG). Routine induction therapy due to physician choice occurs in approximately 50% of heart transplants. The use of routine induction therapy at the time of heart transplant is controversial, as there is no clear survival benefit with its use.

Another strategy under current investigation, which may serve as an alternative to induction therapy, involves use of the terminal complement inhibitor eculizumab (Soliris®; Alexion Pharmaceuticals, Cheshire, CT, USA) at the time of heart transplant and protocol-based administration in the first 2 months post-transplant. Eculizumab is a humanized anti-C5 antibody which impairs C5 cleavage to C5a and C5b. This inhibition prevents formation of the membrane attack complex (MAC) and C5a-induced chemotaxis of inflammatory cells. Cd4 deposition, C3a activity, and other upstream complement cascade steps are not affected. Eculizumab significantly reduced, but did not abolish, early AMR in renal transplant recipients with known DSA against their living related donor.\(^4\) This suggests the possibility of complement-independent mechanisms of

<table>
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<th>Table 4 Treatment modalities for cellular and/or humoral rejection</th>
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<td><strong>Type of rejection</strong></td>
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<tr>
<td>Cellular</td>
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<td>Antibody-mediated</td>
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Abbreviations: ATG, rabbit anti-thymocyte globulin; CNI, calcineurin inhibitors; DSA, donor-specific antibodies; ECMO, extracorporeal membrane oxygenation; EF, ejection fraction; IABP, intra-aortic balloon pump; iv, intravenous; MMF, mycophenolate mofetil; PSi, proliferation signal inhibition.
acute AMR. There did not appear to be significant infectious risk with use of this agent. Since the complement cascade acts as an intermediary between the innate and the adaptive immune systems, there is a chance that terminal complement inhibition will modulate the immune system of patients post-transplant, reduce rates of AMR, and possibly impact CAV.

Further understanding of the mechanisms of AMR and CAV will potentially lead to therapeutic targets to prevent the development or slow the process of CAV. Antigens of various endothelial cells, including angiotensin II type 1 receptor, anti-MHC class I chain-related A (MICA), anti-MHC class I chain-related B (MICB), vimentin, and adhesion or trafficking receptors may play a role in CAV development. Inflammatory modulators that affect cytokine signaling may have a mechanistic role in the development of CAV.\(^{57,68}\)

**Conclusion**

Cardiac allograft immune activation and heart transplant rejection remain major contributors toward morbidity and mortality after heart transplant. Antibody sensitization presents additional challenges in the care of heart transplant recipients. Further understanding of the interactions between the innate and adaptive immune systems is key to effective immunomodulation and prevention of cardiac allograft activation and heart transplant rejection. Targeted therapies could lead to improved patient care with improved graft and patient outcomes and reduced treatment side effects. Complement deposition is part of the process of AMR. The utility of terminal complement inhibition is under current examination and represents one example of modulation of the innate and adaptive immune systems.

**Disclosure**

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