Review

The Puzzle of Immunosuppressive Drugs

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Academic Editor: Maurizio Salvadori

Special Issue: Immunosuppression in Kidney Transplantation

Received: August 30, 2020
Accepted: December 17, 2020
Published: January 11, 2021

Abstract

Kidney transplantation has become the preferred treatment option in end-stage chronic renal failure as it provides significant improvements over dialysis in terms of both quality and duration of life. Even after several randomized studies conducted in the last 20 years, the combination of CNI, MMF, and steroids continues to be considered the gold standard for kidney transplantation. However, novel molecules aimed at minimizing renal and cardiovascular toxicity, particularly with CNI sparing, are being identified. The present review assesses various such molecules available currently and briefly discusses the existing combination strategies and novel perspectives for the redesigning of protocols based on our novel therapeutic arsenal.

Keywords
Renal transplantation; calcineurin inhibitor; mTOR inhibitor; CTLA4-Ig; thymoglobulin

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1. Introduction

In 2020, transplantation is undoubtedly considered the best treatment option in end-stage kidney disease as it offers a better quality of life and increased survival compared to dialysis [1, 2]. Various developments in the field of immunosuppression have led to considerable progress in terms of short-term graft survival, particularly due to the decreased frequency of acute rejection and its consequences. However, not much progress has occurred in terms of long-term graft survival, which presents an overall graft failure rate of 20% even after ten years since it was first introduced [3]. Calcineurin inhibitors (CNI) probably provide the best immune protection during the first year, although there are frequent reports describing chronic renal toxicity and adverse metabolic effects caused by CNI. Novel molecules aimed at minimizing treatment toxicity, particularly by reducing the recourse to CNIs, have been developed, although only a few of these have been proven to be suitable alternatives to CNIs as their benefits are frequently counterbalanced by the disappointing results in terms of rejection or adverse effects. In the present review, the various molecules currently available are assessed, and an overview of the existing combination strategies and novel perspectives in the redesigning of treatment protocols are provided based on our already extensive therapeutic arsenal.

2. Available Molecules

2.1 The Alloimmune Response (Figure 1)

![Figure 1 T-cell activation. APC: Antigen-presenting cell; MHC: Major histocompatibility complex; TCR: T-cell receptor; MAPK: Mitogen-activated protein kinase; PKC: Protein kinase C; PI3K: Phosphatidylinositol 3 kinase; NFAT: Nuclear factor of activated T cells; AP-1: Activator protein 1; NF-KB: Nuclear factor kappa B; IL: Interleukin; mTOR: Mammalian target of rapamycin.](image-url)
T cells are considered the chief orchestrators in the alloimmune response and are, therefore, the main target in anti-rejection therapy. The activation of T cells requires three signals, the first of which results from the interaction of T cells with an antigen-presenting cell (APC) that presents an allopeptide to the T-cell receptor via type I or type II MHC.

The second costimulatory signal results from the interaction of CD80/CD86 located on the APC with CD28 present on the T cell. These two signals activate the calcium/calcineurin, RAS MAP kinase, and NFκB pathways, leading to the following three important events: (i) the expression of CD40L, a key element in the stimulation signal; (ii) the production of various cytokines, including the interleukin 2 which is the cornerstone of T-cell proliferation, and (iii) the transcription of the alpha chain of the IL-2 receptor (α-CD25), which improves the affinity of IL2 toward its receptor. The third signal involves IL-2 binding via an autocrine/paracrine pathway, leading to the activation of the PI3K/AKT/mTOR pathway, thereby triggering a lymphocyte-proliferation signal.

The different cytokines released by the T cells are also involved in the activation of other cells that participate in graft rejection. The inhibition of these three signals involved in the activation of T cells is, therefore, expected to favor long-term graft acceptance.

2.2 Therapeutic Arsenal

Various molecules with the capability of controlling the activation and proliferation of T cells have been developed. Some of these molecules eliminate T cells to facilitate the restoration of a T-cell population with a different repertoire, while the other molecules aim to control the existing T cells by inhibiting their cytokine production.

2.2.1 Polyclonal Antilymphocyte Sera

Antilymphocyte sera are produced by the animals (rabbits or horses) immunized with lymphoid cells derived from the human thymus or a human cell lineage (such as Jurkat cells). These antibodies cause profound T-cell and B-cell depletion within a few hours of administration, followed by immune reconstitution shortly after the perfusion ends. However, total immune reconstitution occurs over months following the initial perfusion, with memory T lymphocytes reappearing more rapidly compared to naive lymphocytes. This approach has been demonstrated to prevent acute rejection in highly sensitized patients more efficiently compared to an absence of induction or induction with anti-CD25 antibodies [4]. However, the lack of specificity and the higher levels of immunosuppression increase the risk of infection and neoplasia [5-7]. Furthermore, xenogeneic molecules could be associated with a risk of manifestations of the serum sickness disease, which occurs less frequently since the treatment duration was reduced from three to five days.

2.2.2 Humanized Monoclonal Anti-CD52 Antibody

Alemtuzumab is a humanized IgG1 monoclonal antibody directed against CD52, a glycoprotein present on circulating T cells and B cells, monocytes, macrophages, natural killer cells, and granulocytes. Alemtuzumab was initially used for treating chronic lymphocytic B-cell leukemia, and now, it is also used off-label as an induction agent in renal transplantation. Similar to polyclonal antilymphocyte sera, alemtuzumab also induces an immediate and profound depletion, followed by reconstitution with lymphocytes exhibiting a phenotype shift toward a higher proportion of memory
T cells [8]. However, accumulating evidence suggests that this cell population may trigger alloimmunization [9, 10]. The adverse effects of alemtuzumab include cytopenia and, in rare cases, autoimmunity (hemolytic anemia, thrombopenia, and hyperthyroidism), in addition to a higher incidence of infections or tumors. Alemtuzumab has the advantage over thymoglobulins in being a monoclonal agent with less variability between the batches, although it also has a narrower spectrum of effects as thymoglobulins act on several targets.

2.2.3 Monoclonal Anti-CD25 Antibody (Anti-rIL2 Antibody)

The activation of the IL-2 receptor leads to T-cell proliferation (Figure 1). Basiliximab is a chimeric blocking antibody that targets the α-CD25 chain of the IL-2 receptor (rIL2) and is, therefore, used as an induction agent in transplantation. It presents a higher immunological risk as it does not deplete and induces less-marked immunsuppression. Anti-rIL2 antibodies have been demonstrated to be more efficient in preventing acute rejection in patients who are not highly sensitized [11-13]. However, several lines of evidence suggest that anti-rIL2 antibodies may impair the development of regulatory T cells, which require low-dose IL2 stimulation for their expansion [14-17]. However, no major specific adverse effects are reported for anti-rIL2 antibody treatment. In the large SYMPHONY trial [18], induction with an anti-rIL2 antibody (daclizumab) combined with low doses of cyclosporine A and mycophenolate acid resulted in the same rate of acute rejection as was observed in the patients receiving a standard dose of cyclosporine A without rIL2 induction. Anti-rIL2 induction, therefore, appears to allow CNI exposure during the initial phase of transplantation.

2.2.4 Corticosteroids

Steroids exhibit immunosuppressive effects manifested by inhibiting the production of various vasoactive cytokines, such as IL-2 IL1, IL6, and TNFα via two main pathways- one is by binding to the corticosteroid receptor, which leads to the migration of the complex toward the nucleus, where it induces or represses gene expression, and the other is a direct action, in which they regulate the action of transcription factors, such as AP1 or NF-κB, by stabilizing their cytosolic inhibitors. Despite several studies attempting to eliminate these drugs from the maintenance therapy to minimize their adverse cardiovascular, metabolic, bone, or skin effects, steroids continue to be a key induction agent in cancer treatment [19-22]. However, due to the minimal use of steroids in maintenance therapy, fewer studies are being conducted now with steroid withdrawal, highlighting the implication of these molecules in the complicated mechanisms involved in graft acceptance.

2.2.5 Calcineurin Inhibitors (CNIs)

CNIs are small molecules that interact with the cytosolic proteins referred to as immunophilins (FK-BPs or CyPs). These interactions cause the inhibition of calcineurin, a phosphatase implicated in the dephosphorylation of the transcription factor NFAT, which is a key player in the T-cell activation cascade. However, redundancy in this pathway has been observed, warranting the use of other molecules in combination with the CNIs for the induction of graft acceptance. Moreover, CNIs have a narrow therapeutic window and, therefore, require constant monitoring to improve their efficacy and reduce their toxicity. However, since their discovery in the 1980s, CNIs have been a key treatment element in solid organ transplantation, despite their toxicity, which is independent of
their immunosuppressive properties and includes neurological, nephrological, and metabolic effects. The nephrotoxicity of CNIs is a major problem. CNIs cause direct reversible vasoconstriction that may lead to acute kidney injury. CNIs are also implicated in chronic lesions, such as arteriole hyalinosis and tubulointerstitial fibrosis, which appear to be related to the endothelial reticulum stress in various types of renal cells. CNIs may also cause thrombotic microangiopathy (TMA), probably via direct endothelial cell injury and/or dysfunction. They are also reported to be associated with a higher risk of developing hypertension, dyslipidemia, and de novo diabetes, all of which are associated with cardiovascular risk factors or mortality [23-26].

2.2.6 Antimetabolites

The duration between the 1960s and the 1980s was considered the golden era of azathioprine (AZA), which was demonstrated to extend canine allograft survival [27]. AZA suppresses the proliferation of the activated T and B cells and decreases the number of circulating monocytes by inhibiting the cell cycle of bone marrow promyelocytes. The antiproliferative effect of AZA is mediated by its metabolites, which act as chain terminators in DNA replication and also block the de novo purine base synthesis, a fundamental process in lymphocytes that lack the purine rescue pathway, by forming thioinosinic acid. The main complication of AZA treatment is the toxicity caused to the bone marrow and the liver. Since the 2000s, the use of AZA decreased considerably, and the molecule was replaced by mycophenolate mofetil (MMF), which is a reversible inhibitor of inosine monophosphate dehydrogenase (IMDPH) isofrom 2, an enzyme required for the de novo synthesis of purine bases. At the end of the 20th century, MMF contributed greatly to the improvements in kidney survival. Nevertheless, the digestive and hematological toxicities caused by MMF represent a major barrier to its extensive application. Moreover, MMF also increases the risk of infection, particularly for viral diseases, despite being demonstrated to exhibit antiviral properties in vitro [28]. In addition, its pharmacodynamics is complex as its metabolism is regulated by various transporters, and individual drug exposure cannot be easily evaluated and requires the calculation of the area under the curve (AUC) or mini AUC, which are difficult to reproduce during patient follow-up in real life.

2.2.7 Mammalian Target of Rapamycin Inhibitors (mTOR Inhibitors)

Sirolimus and everolimus block the proliferation signal by inhibiting the mTOR kinase. These drugs were originally developed to replace the CNIs as the cornerstone of immunosuppression. However, their generalized use is limited due to their adverse effects and the higher incidence of acute rejection reported in pivotal studies. For instance, in the SYMPHONY study, the rate of acute rejection in the patients treated with rapamycin plus MMF and induction with an anti-rIL2 antibody was 60% higher than that in the patients treated with low doses of cyclosporine combined with the same associated immunosuppressive therapies, and twice that in the patients treated with low doses of tacrolimus. A recent retrospective analysis suggested that the patients treated with mTOR inhibitors have a higher risk of developing de novo donor-specific antibody production. In addition, the mTOR inhibitors may cause renal lesion via various mechanisms, including proteinuria, as evidenced by their effects on the PI3K/AKT/mTOR pathway in podocytes during renal injury [29], and microangiopathy, possibly due to a decrease in the glomerular secretion of VEGF [30]. Furthermore, their use in combination with full-dose CNIs results in increased renal toxicity.
Moreover, sirolimus may interfere with recovery from acute tubular necrosis [31]. However, several studies have reported these drugs to be effective in reducing tumor development or preventing the recurrence of skin cancer in renal transplant patients, while the meta-analyses revealed that mTOR inhibitors are associated with a lower risk of viral infection in the renal transplant patients [32–34].

### 2.2.8 Belatacept (BCT)

Belatacept is a fusion protein that binds to the CD80/CD86 present on the APCs, thereby inhibiting the costimulatory signal, leading to the activation and proliferation of naive T cells (Figure 2). The main advantage of BCT is that it does not exhibit nephrotoxicity. The phase III BENEFIT study [35, 36] reported significant improvements in GFR in the patients on belatacept, although with the disadvantage of higher rates of cellular rejection compared to those in the patients on CNIs. However, the frequency of de novo donor-specific antibody (DSA) production is reported to be lower in the patients treated with belatacept compared to those on CNI regimens, and this effect is correlated with the inhibition of B-cell maturation. These effects have also been confirmed for long-term usage in a seven-year-long follow-up study. In addition to improving the renal function, belatacept appears to improve both metabolic profile and graft survival [37]. Since belatacept is administered as an intravenous infusion, it is convenient to monitor adherence compared to the orally-administered agents, thereby decreasing the risk of occult nonadherence.

![Costimulation blockade](costimulation.png)

**Figure 2** Costimulation blockade. MHC: Major histocompatibility complex; TCR: T-cell receptor; CTLA-4: Cytotoxic T lymphocyte-associated protein 4; APC: Antigen-presenting cell.

The risk of opportunistic infection (OPI) is not elucidated in the literature, although there are cases of disturbing viral reactivation and fungal infection reported quite frequently. In the initial phase of development, a higher incidence of post-transplantation lymphoproliferative disorders (PTLD) related to the primary infection with EBV was observed in the recipients. Recently, in a
A retrospective cohort, Bertrand et al. [38] reported an OPI frequency of approximately 12%, essentially due to CMV reactivation and pneumocystis pneumonia. In the CMV-seronegative recipients of organs from seropositive donors, belatacept usage was associated with a higher incidence of CMV viremia, a higher rate of first-line treatment failure, a longer time for virus clearance, and cases of severe CMV retinitis [39, 40]. Certain cases of progressive multifocal leukoencephalopathy associated with the JC virus are reported. Our research group has also previously described a rapidly fatal case of PML associated with a refractory state of T-cell anergy, potentially due to belatacept therapy. This anergy state was characterized by a functional defect in the lymphocytes (affecting the cytokine secretion, proliferation, and cytotoxicity), which was associated with a strong expression of the inhibitory receptor PD-1. The treatment with a therapeutic anti-PD1 receptor antibody ex vivo failed to improve the T-cell function, which was consistent with the definition of lymphocyte anergy [41]. Although these data require confirmation in a larger case-control study, they nevertheless suggest that such events should be monitored carefully.

3. Conventional Protocols

3.1 Induction

High doses of corticosteroids considerably reduce the rate of rejection in the early stages of transplantation and, therefore, represent the cornerstone of induction treatment. They are used for reducing inflammation in the initial phase, which is followed by maintenance therapy. In several cases, this induction is combined with polyclonal antilymphocyte serum or basiliximab, leading to further lowering of the rates of rejection and graft loss [42].

In patients with low immunological risk, basiliximab is the most frequently used agent for limiting comorbidities or infectious diseases.

In most transplant centers, thymoglobulin (rATG) is routinely administered to immunized renal transplant recipients [43] as the preferred agent for T-cell depletion [44, 45], while the highly sensitized patients receive depleting agents to reduce both cellular and antibody-mediated rejection. In addition to the greater risk of neoplasia and infection, higher cardiovascular mortality is reported in renal transplant recipients [46]. Ducloux et al. characterized the CD4 T-cell lymphopenia as a potential immunological marker of immunosuppression [47, 48] in renal transplantation, and then demonstrated it to be associated with atherosclerosis and cardiovascular death [49]. According to these findings, it appears reasonable to reserve this treatment, as far as possible, to patients with an extremely high immunological risk, and to prefer basiliximab for more general use, particularly because of the increasing evidence of its safety, even in the high-risk patients. Recently, Phanish et al. [50] reported the results for a retrospective study, in which an immunosuppression regimen comprising basiliximab induction, tacrolimus, MMF, and prednisolone combined with early steroid withdrawal was investigated in low-risk patients and with MMF withdrawal in high-risk patients. The authors reported low acute rejection rates in both the groups (15.1% and 13.9%, respectively) along with high rates of graft function. Although a randomized study is required to further confirm these observations, the results are nonetheless highly promising.

Rituximab was recently evaluated as an induction therapy agent in solid organ transplantation. This monoclonal antibody acts against CD20 and induces rapid and profound depletion of B
lymphocytes. Rituximab has received approval for use in the treatment of lymphoma and leukemia. It is also used off-label for induction in the cases of ABO-incompatible (ABOi) transplantation due to its profound effect on B cells, particularly in the B1 cells. Rituximab has completely replaced splenectomy, decreasing the anti-agglutinin titers and generating interesting results, as graft survival in ABOI kidney transplants is now similar to that in the allogeneic kidney transplantation under the standard criteria, despite the potential increase in the risk of acute humoral transplant rejection [51, 52]. Interestingly, the rate of acute cellular rejection is generally reported to be similar to that in transplantation from the donors meeting the standard criteria, although certain studies have reported lower rates as well [53].

Rituximab is proposed as a treatment for humoral acute rejection due to its action on the B cells, although the results obtained to date, albeit in inadequately powered trials, are not convincing, suggesting that rituximab may not be as effective against the memory B-cell compartment and plasma cells. However, rituximab does act on B-cell precursors and has, therefore, been proposed as an induction therapy agent for reducing the production of donor-specific antibodies. Van den Hoogen et al. conducted a double-blind randomized study in patients treated with a conventional regimen and demonstrated that induction with 375 mg/m² rituximab was associated with a lower rate of acute cellular and humoral rejection compared to the same regimen without induction in the sensitized patients but not in non-sensitized patients, with no increase reported in the risk of infection [54]. Tyden et al. had previously reported a trend toward lower rates of acute rejection in the patients receiving induction with 375 mg/m² rituximab in a randomized study. Interestingly, low doses of rituximab (100 or 200 mg/m²) resulted in low rates of DSA in calcineurin-based regimens, although with a lower incidence of acute cellular rejection in the matched cohorts of patients as well. These findings suggest that rituximab also plays an important role in connection to the antigen-presenting cells and that the pre-transplantation use of rituximab may reduce the initial immunization [54-57]. In a recent case-control study concerning 230 kidney transplants from living ABO-compatible donors, half of the patients were administered a low dose (100 mg) of rituximab, and acute cellular rejection was observed in 11% of these patients, and de novo DSA was observed in 13.9% of these patients, versus 21.7% (p < 0.041) and 26.9% (p < 0.005), respectively, in the patients treated with the conventional regimen. Moreover, the use of low-dose rituximab for induction was associated with a lower rate of CMV infection, suggesting the safety of this strategy. Low-dose rituximab may, therefore, be beneficial in sensitized patients as well as in the patients not receiving the induction therapy. These results require further confirmation in larger prospective studies.

### 3.2 Maintenance

Most of the trials concerning the maintenance treatments focus on a few molecules (CNI, AZA, MMF, mTOR inhibitors, prednisone, and belatacept), used in different combinations, mostly in the form of tritherapy. This combination of different molecules allows the blocking of T-cell activation via different pathways simultaneously and decreasing the doses in each treatment, thereby limiting the toxicity of each molecule individually.

On the basis of several randomized controlled trials, the combination of CNIs, MMF, and steroids has been identified as the gold standard for reducing the rate of graft rejection at one year after the kidney transplantation. The main concerns regarding this treatment relate to the nephrotoxicity of
CNIs. Combinations of the rIL2-blocking antibody with tacrolimus, MMF, and steroids are reported to be superior to the similar combinations, including cyclosporine A in place of tacrolimus, as evidenced by the acute rejection rates of 12.3% vs. 24% at one year (p < 0.001) and the glomerular filtration rates of 65.4 ±27 mL/min vs. 59.4 ±25.1 mL/min (p < 0.001) for tacrolimus-based vs. cyclosporine A-based regimens [18].

An alternative strategy based on the combination of mTOR inhibitor with MMF and a steroid is reported. In the SYMPHONY study, a fourth arm was added to this combination, and the patients were treated with daclizumab, sirolimus, mycophenolate acid, and steroids. In comparison to the groups receiving tacrolimus or cyclosporine A (see above), the rejection rate was significantly higher at one year (37.2%), and the GFR was 56.7 ±26.9 mL/min. In the ORION study [58], the same combination of sirolimus with steroids was reported to present a significantly higher acute rejection rate compared to MMF/tacrolimus (31.3% vs. 8.2%). Moreover, a higher rate of acute rejection (15.2%) was observed in the third group of patients who initially received sirolimus plus tacrolimus, with a gradual withdrawal of CNIs. Two-thirds of the patients included in the study population were withdrawn from the study as they presented a high rate of rejection, and consequently, these strategies involving mTOR inhibitors without CNI had to be abandoned. Furthermore, despite a small improvement in renal function reported by certain studies, a higher frequency of rejection and, more importantly, a higher risk of the de novo development of DSA was observed in the patients [18]. However, the patients treated with mTOR inhibitor-based regimens presented a lower rate of viral infections and CMV- and BKV-associated nephropathy. In addition, as the mTOR inhibitors are known to block the cell cycle in various cell types, they are used for the treatment of renal or lung cancer. The mTOR inhibitors are also reported to decrease the incidence of skin cancer recurrence, suggesting the additional potential benefits of preventing tumor lesions in transplant patients.

Belatacept was the first biotherapy to be used as a maintenance treatment in kidney transplantation. The lack of nephrotoxicity and a better metabolic profile compared to the other treatments present belatacept as an ideal treatment drug. Belatacept has been used in combination with MMF and steroids for the induction therapy based on anti-rIL2 antibodies. The use of belatacept was validated in 2011 by two studies, BENEFIT (for donors with standard criteria) and BENEFIT EXT (for donors with extended criteria), which compared the introduction of this molecule de novo, without CNIs, into a standard triple-therapy that included cyclosporine. Belatacept is demonstrated to improve long-term renal function compared to cyclosporine-based regimens. In the BENEFIT study, belatacept also increased the predicted mean half-life of the kidney grafts, although this regimen was associated with a high rate of PTLD, limiting the use of this molecule in the EBV-positive patients. In addition, higher rates of rejection and more severe rejection were observed, with no impact on graft function, in the patients treated with belatacept, although surprisingly, fewer of these patients developed de novo DSA. Moreover, a lower risk of new-onset diabetes after transplantation (NODAT) was observed. The potential of belatacept in decreasing the development of anti-HLA antibody, together with the improvement observed in renal function, have prompted the researchers to search for appropriate partner molecules for belatacept.

Therefore, despite providing improvements in the renal function or decreases in the incidence of infection, the novel regimens based on mTOR inhibitors or belatacept have, so far, failed to fulfill the hopes of the physicians and patients. All have been developed in head-to-head comparisons, and it is possible that other combinations might be more powerful in allogeneic transplantation.
4. Novel Perspectives

4.1 CNI Minimization via mTOR Inhibitors

Despite the poor results obtained with the combination of cyclosporine A and mTOR inhibitors in the initial study conducted in 1996, in which rapid development of renal fibrosis was observed, the low rate of acute rejection was encouraging as it suggested the synergetic action of these two molecules [59]. Improvements in the understanding of the therapeutic target of everolimus and the reports describing the beneficial effects of everolimus on tumor growth and viral infection prompted researchers to design novel protocols based on the combination of low doses of tacrolimus and everolimus. In the TRANSFORM study [60], the authors selected the lowest possible doses of CNIs, and 2,000 patients with low immunological risk were included in this study comparing a combination of everolimus (trough concentration of 5 to 7 ng/mL) and a CNI at a trough concentration half the usual level, with an arm in which MMF and CNI were administered to achieve the standard trough concentrations. No differences in renal function or rejection rate were observed at two years, although the viral infection rates were lowered by 60% in case of CMV infections and 45% in case of BKV-associated nephropathy [61]. These results have encouraged several research teams to review their protocols and propose similar CNI minimization strategies for the patients at low immunological risk.

4.2 CNI Sparing via Immunotherapy

The development of Belatacept regimens was hampered due to the acute rejection rates of 5%–40% in the absence of CNIs, in contrast to the low rates of de novo DSA. Major efforts were put to eliminate the phenotype of the T cells resistant to belatacept. Several studies reported a correlation between the presence of high rates of CD4 or CD8 T cells and a memory or exhausted phenotype at the time of transplantation [62]. In primates, T cells lose CD28 during maturation, which potentially accounts for their CD28 independence and the lack of sensitivity to belatacept [63]. However, the CD28 T cells occur less frequently in humans, and the expression of CD28 is maintained on memory T cells. These cells appear to be Belatacept-resistant and are activated and continue to proliferate in the presence of belatacept, indicating that the combinations of mTOR inhibitors and belatacept should be used instead. Data from experiments on mice suggest that the concomitant blockade of costimulation and use of mTOR inhibitors promotes the apoptosis of activated alloreactive T cells and immunotolerance [64]. The other options are to deplete the memory compartment using the depleting agents in combination with belatacept or to use molecules capable of controlling memory T cells in combination with belatacept for the initiation of the transplantation. Three different strategies have been envisaged. The first one involves induction therapy with T cell-depleting antibodies, the second one involves an initial combination of belatacept with tacrolimus followed by rapid withdrawal of tacrolimus, and the third one uses a combination of belatacept with the mTOR inhibitors.

4.2.1 Induction with Depleting Agents

Pivotal studies were conducted with belatacept treatment developed with the use of basiliximab, an anti-rIL2 antibody for induction. Several studies have demonstrated that the number of memory
cells on Day 0 correlates with the risk of developing acute rejection [65], which led to the development of an interesting strategy that involved the use of depleting antibodies for induction, along with belatacept, to eliminate the memory T cells. As reported by Ferguson [66] and the BEST study [67], the use of thymoglobulin or alemtuzumab in combination with Belatacept and MMF, along with rapid steroid withdrawal, is associated with a significantly higher rate of acute rejection in the belatacept arms compared to those in the tacrolimus arms. The BELACOR study conducted in 2019 prospectively assessed the potential benefits of Belatacept use for preventing antibody-mediated rejection (AMBR) in patients with low MFI for preformed DSA; the induction therapy was based on thymoglobulins. Interestingly, while AMBR was not more frequent in the Belatacept group, the rate of T cell-mediated rejection (TCMR) was significantly higher in this group compared to the group receiving treatment with CNIs (25.4% vs. 5.64%, respectively), with no difference in terms of graft survival or renal function [68]. Overall, these results suggest that depleting agents are ineffective in the long-term depletion of Belatacept-resistant T lymphocytes even in non-sensitized patients.

4.2.2 Induction with CNI and Belatacept

Most of the acute rejections occur in the first three months post-transplantation in the patients on Belatacept regimens. Therefore, for the first few months, the combination of a calcineurin inhibitor with belatacept, along with basiliximab for induction, was considered. Afterward, the CNI is gradually withdrawn to prevent acute rejection and avoid long-term toxicity [69]. Using this combination, the rate of acute rejection observed was lower than that observed in the BENEFIT regimen and higher than that in the comparator group. The curves of acute rejection-free survival revealed that acute rejection was delayed until the CNI was withdrawn or its dose was reduced, suggesting that CNIs may partly control Belatacept-resistant T cells but cannot eliminate them.

4.2.3 Combination of Belatacept with mTOR Inhibitors

The occurrence of acute rejection in patients receiving belatacept has led to the notion that Belatacept-resistant lymphocytes can proliferate and participate in graft rejection [62, 70, 71]. Fergusson et al. and Kirk et al. separately performed pilot studies evaluating the addition of an mTOR inhibitor to Belatacept treatment to control the rate of Belatacept-resistant T cells [59, 51]. The treatment also included induction with thymoglobulin. Interestingly, in a small group of patients, this strategy resulted in a low rate of acute rejection compared to the control group using a CNI (3% vs. 4%, respectively), and despite the rapid withdrawal of steroids. Several patients discontinued the initial treatment because of the adverse effects, attributable, in most cases, to the use of mTOR inhibitors without steroids. Nevertheless, this pilot study was encouraging, and these results were again confirmed by two recent studies, in which treatment with belatacept plus mTOR inhibitors, along with the induction based on alemtuzumab or thymoglobulin, resulted in extremely low rates of acute rejection, suggesting a synergetic effect of these two molecules. However, these findings should be confirmed by larger, randomized studies.
4.3 Viral Infection in the Context of Kidney Transplantation and Immunosuppressive Treatment Management

Viral infections are common causes of opportunistic infections after transplantation. The risk of a viral infection depends on various parameters, including the pathogen encountered, the immunosuppressive treatment used for preventing graft rejection, and other host factors, such as the cellular antiviral response. The treatment for viral infections generally includes antiviral agents and/or reduction of the immunosuppressive treatment to ensure an antiviral immune response without increasing the risk of rejection [73]. Since the antiviral response is mediated by CD8 T cells, an initial reduction or withdrawal of the antimetabolites and/or calcineurin inhibitors could be considered, although with an increased risk of graft rejection. Switching from the "standard" regimen to another protocol is also proposed as a means of preventing the viral infection. The mTOR pathway is involved in both lymphocyte expansion and viral replication. Recent data from studies in humans have suggested that the mTOR inhibitors exhibit antiviral effects in transplant patients, resulting in a lower risk of CMV, polyomavirus, and HHV8 infection, compared to the treatments combining calcineurin inhibitors, mycophenolate mofetil, and steroids [74]. The prospective ATHENA study, which evaluated a combination of everolimus, an mTOR inhibitor (mTORi), and a low dose of calcineurin inhibitor (tacrolimus or cyclosporine A) in comparison to a combination of tacrolimus and mycophenolic acid, reported lower rates of viral infections, but not of bacterial infections, at one year after transplantation [25.7% for everolimus + tacrolimus, 11.6% for everolimus + cyclosporin A, 40.7% for tacrolimus + mycophenolic acid], while the rates for the CMV infection were 6.2%, 2.5%, and 20.6%, respectively, and those for BKvirus infection were 17.1%, 9.1%, and 22.5%, respectively [75]. The lower rate of viral infection was not associated with a higher rate of acute rejection. However, the renal function was poorer in the group that received everolimus plus a calcineurin inhibitor.

Therefore, three different strategies may be proposed for non-immunized patients. The first is the use of mTORi + tacrolimus + steroid to decrease the rate of viral infection. The second is the use of a standard of care maintenance regimen with tacrolimus, mycophenolic acid, and steroids, and switching to an everolimus-based strategy in the cases of viral infection. In both these strategies, the physician should adapt the treatment according to the presence or absence of viral replication, regardless of whether a specific antiviral response is observed.

The third approach is an innovative one and is designed to increase the risk of viral infection without increasing the risk of rejection. This approach is based on the assessment of the ability of the cellular antiviral response to control viral reactivation [76]. Virus-specific T-cell monitoring is proposed as a means of optimizing the management of virus reactivation in transplant patients. Functional cellular immune responses are reported to be associated with the control of viral replication in the cases of infection with CMV or polyomavirus [77, 78]. Immunovirological monitoring could be used to provide personalized medical management to the patients through an individual assessment of the risk of viral reactivation. For instance, CMV-specific cellular immune monitoring has been demonstrated to predict CMV control after solid organ transplantation. An undetectable CMV-specific cellular immune response is associated with a higher risk of developing uncontrolled CMV reactivation. Therefore, CMV immune monitoring, in addition to clinical and DNA-based monitoring for CMV, could be included in the standard follow-up to improve CMV management [79]. Similarly, BKV-specific cellular immune responses are crucial for the control and
clearance of BKV. BKV-specific T-cell dysfunction increases the risk of uncontrolled BKV infection, while the increases in the levels of BKV-specific CD8 T cells are associated with a better prognosis in BKV-associated nephropathy. Nevertheless, BKV-specific cellular immune monitoring in the management of BKV infection should be explored because such assessments may guide the decreases in the intensity of immunosuppressive treatment as well as the increases after clearance [80].

4.4 Transplantation in the Context of Cancer

Numerous reports have highlighted the higher incidence of certain cancers in solid organ transplant patients, with non-melanoma skin cancers being the most frequently appearing ones. Since the mTOR inhibitor can inhibit the mTOR pathway involved in the cancer cell growth, the use of mTOR inhibitors to reduce the recurrence of non-melanoma skin cancer appears to be attractive when combined with calcineurin inhibitors, compared to calcineurin inhibitor maintenance [81]. Similarly, in the CONVERT study, switching from a calcineurin inhibitor-based regimen to sirolimus-based immunosuppression was observed to be associated with a reduced incidence of malignancy in kidney transplant recipients [81]. Meta-analyses have confirmed this tendency mostly in terms of the occurrence of secondary non-melanoma skin cancer [82, 83]. In these initial reports, mTOR inhibitors were used in place of CNI. However, recent observations of a higher risk of DSA occurrence in the patients treated with mTOR inhibitors along with mycophenolate acid and the interesting findings of the Transform and Athena study [84, 85] have suggested that the combination of an mTOR inhibitor with a low dose of CNI could be an effective strategy for the patients with the first appearance of non-melanoma skin cancer. The long-term outcomes of the Transform and Athena study would provide the essential information to decipher the effect of these approaches on the occurrence of skin cancer. The effect of the combination of an mTOR inhibitor with belatacept is currently unknown.

4.5 Transplantation in the Context of Pregnancy

Since transplantation was first performed, several female transplant recipients have retained the capability of becoming pregnant and having babies. In 2011, over 11,000 deliveries were recorded worldwide for female kidney recipients [86]. It is generally recommended to delay the pregnancy for at least a year after the transplantation due to the risk of graft failure [87]. The predictors of good maternal and fetal outcomes include young age of the mother, stable graft function with no recent episodes of graft rejection, a serum creatinine concentration of < 1.5 mg/dL, a proteinuria level of <500 mg a day, and normal or well-controlled hypertension [86, 88, 89]. Immunosuppressive treatments, antihypertensive medication, and clinical parameters such as blood pressure may affect fetal development and pregnancy outcomes. For instance, angiotensin-converting enzyme and angiotensin 2 receptor inhibitor treatments should be terminated during pregnancy due to the risk of renal agenesis.

The data regarding the use of maintenance therapy during pregnancy or lactation are limited. Generally, calcineurin inhibitor treatments are maintained during pregnancy. Both cyclosporine and tacrolimus are associated with fetal growth retardation and prematurity, although not with malformations, and their withdrawal during lactation is generally not required. On the contrary, mycophenolic acid, being a teratogen, should be replaced with azathioprine at least a few weeks
prior to attempting the conception. In the best-case scenario, the absence of proteinuria or the absence of an increase in proteinuria should be verified, together with the serum creatinine concentration and azathioprine tolerance (absence of leukopenia), prior to conception. Three months before any attempt at conception is a more comfortable time for treatment equilibration. In regard to the other molecules, limited evidence is available concerning the safety of the mammalian target of rapamycin inhibitors and belatacept, although a few pregnancies have been successful in the patients treated with Belatacept or Abatacept for autoimmune disease [90, 91]. mTOR is essential for placenta implantation as well as for the growth of most fetal organs, and its absence is embryo-lethal as mTORC1 and 2 play key roles in embryonic development and growth [92, 93]. Finally, prednisone may be used safely in pregnant women [8]. The preferred regimen is, therefore, a combination of a calcineurin inhibitor with azathioprine and prednisolone. After delivery, azathioprine is frequently replaced with another molecule because of the stronger association of the former with skin carcinomas.

5. Conclusions

Kidney transplantation is becoming an increasingly complex procedure as its outcomes improve. Interestingly, novel combinations of the therapies that were initially developed independently have emerged as a potential alternative to the standard of care for reducing the CNI toxicity, decreasing the occurrence of infections or tumors, and limiting antibody-mediated rejection. In this context, the mTOR inhibitors and belatacept have re-emerged after a difficult departure and could serve as promising candidates in novel molecular combinations. Therefore, more trials are required to confirm the encouraging results presented by them and to respond to the diversity of patients undergoing kidney transplantation.

Author Contributions

All authors contributed to this work as follows: Writing: AC, JO, MD, AD; Editing: AC, JO, MD, AD.

Competing Interests

The authors have no competing interests to declare.

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