Clinical Applications of Regulatory T cells in Adoptive Cell Therapies

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Abstract

Interest in adoptive T-cell therapies has been ignited by the recent clinical success of genetically-modified T cells in the cancer immunotherapy space. In addition to immune targeting for malignancies, this approach is now being explored for the establishment of immune tolerance with regulatory T cells (Tregs). Herein, we will summarize the basic science and clinical results emanating from trials directed at inducing durable immune regulation through administration of Tregs. We will discuss some of the current challenges facing the field in terms of maximizing cell purity, stability and expansion capacity, while also achieving feasibility and GMP production. Indeed, recent advances in methodologies for Treg isolation, expansion, and optimal source materials represent important strides toward these considerations. Finally, we will review the emerging genetic and biomaterial-based approaches on the horizon for directing Treg specificity to augment tissue-targeting and regenerative medicine.

Now over 20 years since the discovery of regulatory T cells (Tregs) within the CD4⁺CD25⁺ T cell fraction [1], investigations of adoptive Treg cellular therapies are well underway to restore or induce immunological tolerance in settings of autoimmunity, allogeneic transplantation, allergy, protein replacement therapy, graft versus host disease (GvHD), and more [2–19]. Though once thought to be a single T cell lineage, diverse Treg subsets...
are now identified according to an array of surface proteins, transcription factors, and mechanisms of suppression. Indeed, there may be organ-specific autoimmune disease applications where FOXP3+Helios+ thymic-derived Tregs (tTregs) are optimal for adoptive cell therapies (ACT), whereas other diseases may preferentially benefit from the bystander mechanisms employed by peripherally-induced Tregs (pTreg) (reviewed in [20]), IL-10-producing Tr1 cells (reviewed in [21]) or TGF-β secreting LAP+ Th3 cells [22]; hence, not all Tregs are equal and may need to be tailored to a given pathology.

In vitro suppression assays have been used to demonstrate the capacity for Tregs to attenuate responder cell proliferation through both antigen-specific and bystander mechanisms in a dose-dependent manner [23,24]; therefore, increasing the Treg ratio relative to pathogenic effector cells in vivo, through ACT, may bolster immunoregulation to achieve clinical benefits for immune-mediated diseases resulting from autoimmunity or allore cognition. Animal studies in various disease models have justified this rationale [25–28] and led to the initiation of human clinical trials in patients with type 1 diabetes (T1D) [2–4], Crohn’s disease [5], solid organ transplantation (i.e., liver [6] and kidney [7]), and allogeneic hematopoietic stem cell transfer (allo-HSCT) [13–18]. Herein, we review the lessons learned from these early-phase studies and the recent advances that will enable progress toward safe, efficacious, and cost-efficient Treg therapy for widespread use in the clinic.

**CLINICAL APPLICATIONS OF TREGS**

Tregs display wide-ranging mechanisms of action and have displayed efficacy in ameliorating a large number of inflammatory and autoimmune conditions. However, translating early proof-of-concept animal experiments into clinical practice raises a number of practical considerations, including:

- What are the specific target antigen(s)?
- Is the disease progressive or relapsing and remitting?
- Is there a potential for Tregs to reverse a disease process that is already underway?

In terms of antigen-specificity, polyclonal Tregs may be appropriate for systemic diseases such as systemic lupus erythematosus (SLE) [29], while other conditions with tissue or protein-directed pathologies might benefit from antigenic targeting via TCR or chimeric antigen receptor (CAR)-directed approaches [30,31] (discussed in greater detail below). CAR-modified T cell (CAR-T) immunotherapy has provided remarkable efficacy against various forms of cancer [32–38]. These cell-based therapies are more specifically targeted than traditional chemotherapeutic treatments, and strategies are being developed to manage the most common adverse events (e.g., cytokine release syndrome) [39]. Hence, there is clear incentive to apply these tissue-targeting principles toward ACT involving Tregs to limit potential off-target immunosuppression (Table 1).

**Autoimmunity**

Deficient numbers and/or functional suppression by Tregs have been reported as contributing factors in the pathogenesis of a number of autoimmune diseases (reviewed in

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This concept emanates from the profound systemic autoimmune syndrome observed in subjects presenting with a mutation in the Treg lineage-defining transcription factor FOXP3 resulting in immune dysregulation, polyendocrinopathy, enteropathy, and X-linked (IPEX) syndrome (reviewed in [41]). Similar autoimmune symptoms have been observed in other monogenic disorders, such as autoimmune polyendocrine syndrome type 1 (APS-1) characterized by mutations in the Autoimmune Regulator (AIRE) resulting in defects in thymic negative selection as well as impaired Treg suppression with reduced FOXP3 expression (reviewed in [42]). Genes controlling Treg development and function have also been identified as a source of heterogeneity related to immune regulatory defects in polygenic/multifactorial autoimmune diseases such as SLE, multiple sclerosis (MS), rheumatoid arthritis (RA), inflammatory bowel disease (IBD), T1D, juvenile idiopathic arthritis (JIA), and more [43–53]. Furthermore, Treg ACT has been reported to prevent or reverse autoimmunity in a variety of animal models [54–58]. Hence, human trials are planned or underway to treat a number of autoimmune and auto-inflammatory conditions including ulcerative colitis (UC) [59,60] and T1D [2–4,61] (Table 1).

Early clinical trials of *ex vivo* expanded autologous Treg from peripheral blood of adults or children with T1D have already provided evidence of safety with few or no adverse reactions [2–4]. Importantly, infused autologous adult peripheral blood (APB) Tregs were shown to persist in the circulation of T1D patients for at least one year following transfer and were not associated with increased risk of infection [4]. Although conclusions about efficacy (i.e., the capacity to preserve endogenous β-cell function) cannot be formally demonstrated from these small cohorts, early trials have not resulted in reversal of disease. This may be due to several factors, such as disease staging (i.e., the pre-symptomatic stage of T1D [62,63] prior to clinical diagnosis may represent a more suitable therapeutic window), but may also be related to challenges such as Treg specificity, engraftment, persistence, and/or trafficking, as discussed below.

### Allergy

Analogous to the findings in autoimmunity, Tregs from patients with allergic asthma are reduced in number and exhibit compromised suppressive capacity [64–66]. For this application, Treg ACT is still in the preclinical testing phase. Initial efforts were directed at adoptive transfer of allergen-specific Tregs [8] or *in vivo* induction of pTregs and Th3 cells via nasal delivery of soluble antigen [8,67]. Studies have since demonstrated adoptive transfer of polyclonal Treg or polyclonal *ex vivo* induced Tregs to have similar efficacy in reducing allergic airway inflammation in mice [10]. More recently, Skuljec and colleagues reported that adoptive transfer of CAR-Tregs specific for carcinoembryonic antigen (CEA) expressed in lung epithelial tissues suppressed airway inflammation, mucus production, Th2 cytokine production, allergen-specific IgE production, bronchoalveolar eosinophilia, and airway hyperreactivity more effectively than polyclonal Treg therapy in a murine model of allergic asthma (Table 2) [11]. Efforts to examine models of other allergic conditions, including contact hypersensitivity and food allergy, have focused primarily on *in vivo* pTreg and Th3 cell induction [68–70], though studies of Treg ACT are warranted in these settings given the efficacy observed in models of allergic asthma.
**Protein replacement therapy**

Inhibitory antibody formation occurs in approximately 30% of patients receiving life-saving protein replacement therapy for the blood clotting disorder hemophilia; similarly, the development of neutralizing antibodies occurs in a subset of patients during enzyme replacement therapy for monogenic lysosomal storage disorders (e.g., Pompe disease) and monogenic metabolic disorders (e.g., Fabry disease) [71–74]. Hence, Treg-mediated tolerance induction represents a critical goal to prevent/overcome inhibitor formation for successful long-term disease management (reviewed in [75,76]). In murine models of Pompe disease and hemophilia, oral delivery of plants expressing the disease-relevant antigen (i.e., acid alpha-glucosidase (GAA) or coagulation factor VIII or IX, respectively) induced Tr1, Th3, and pTreg mediated tolerance and prevented antibody inhibitor formation; however, tolerance was lost upon cessation of oral treatment [77–80]. In contrast, exogenous polyclonal Treg therapy induced durable tolerance for two months post-infusion in mice with hemophilia [12]. Recently, two coagulation factor VIII specific human Tregs applications (i.e., TCR transgenic and ASN8 CAR-Tregs) were shown to suppress cognate T and B cell responses in vitro and in a humanized mouse model (Table 2) [31,81]. As Good Manufacturing Practice (GMP) grade production of either polyclonal or antigen-specific exogenous Treg therapies becomes more streamlined, clinical investigation is clearly needed to test Tregs’ ability to prevent or suppress inhibitory antibodies in patients with Pompe disease and hemophilia. Beyond this, we anticipate that the knowledge gained from early studies in hemophilia and Pompe disease may translate effectively for a large number of rare metabolic, lysosomal, and clotting disorders for which protein replacement therapies exist [82]. In fact, these diseases may hold some potential advantages, including the capacity to control the timing of ACT therapy with protein replacement and the potential for more permissive background genetics when compared to individuals carrying autoimmune disease susceptibility alleles.

**Allotransplantation & GvHD**

There is a rapidly growing body of literature regarding the use of Treg ACT for applications in allotransplantation (reviewed in [83]) and GvHD. In a pilot study of ten adult liver transplant recipients, autologous polyclonal Tregs were expanded ex vivo in the presence of irradiated donor lymphocytes plus anti-CD80/86 monoclonal antibodies. Following Treg infusion, seven of the ten patients were completely weaned from immunosuppressive drugs while the remaining three continued taking low dose immunosuppression [6]. Importantly, isolation and expansion of Tregs was possible from kidney transplant recipients taking immunosuppressive drugs, and expanded polyclonal Tregs were stable, persistent, and efficacious in reducing kidney inflammation post-infusion [7]. Moreover, cryopreservation of expanded polyclonal human Tregs did not alter their phenotype or function, suggesting Tregs can be expanded, manipulated (genetically or otherwise), and banked for future use [84]. Finally, CAR-directed Tregs, specific for graft donor HLA class I, were recently shown to prevent cutaneous graft rejection in a model of xenograft transplantation, supporting human trials of these antigen-specific, “designer” Tregs in allotransplantation [85].

GvHD represents a severe and significant risk associated with allo-HSCT that confers lifelong and even terminal complications (reviewed in [86]). While first-in-man studies
of Treg ACT were performed in response to GvHD [13], the first report of Treg ACT prior to allo-HSCT prevented GvHD in the absence of immunosuppressive drugs in 24/26 human leukemia patients who achieved full engraftment [15]. In a subsequent trial, co-transfusion of Tregs with conventional T cells (Tconv) and CD34+ HSCs at the time of allo-HSCT, again without immunosuppressive drugs, prevented GvHD in 85% of patients with only 0.05 cumulative incidence of relapse, suggesting that Tregs do not impede graft-versus-leukemia efficacy and provide appreciable protection against GvHD [18]. In a dose escalation trial, Tregs isolated and expanded from non-autologous umbilical cord blood (UCB) significantly reduced the incidence of both acute and chronic GvHD following double UCB transplantation compared to controls who did not receive UCB Tregs [17]. Altogether, the data support polyclonal Tregs, obtained from autologous or donor peripheral blood or UCB, as a safe and effective cellular drug in settings of GvHD.

**Regeneration**

Tregs, commonly thought of as mediators of immunoregulation, are now being studied for their capacity to induce tissue regeneration and remodeling. Treg therapy may enhance hepatic recovery in fulminant or chronic hepatitis, the latter of which eventually results in liver cirrhosis and liver cancer (recently reviewed in [87]). Indeed, polyclonal Treg ACT prevented liver injury and enhanced hepatic regeneration in mouse models of fulminant hepatitis [88–90]. Endogenous memory Tregs become highly enriched in the liver of human patients with chronic Hepatitis B infection during phases of inflammatory relapse, likely signifying an attempt to control hepatic injury [91]. Hence, it seems plausible that adoptive Treg therapy combined with antiviral medication might improve clinical outcomes and reduce liver damage in patients with viral hepatitis; however, there is an ongoing debate as to whether Tregs might potentiate chronic infection by suppressing antiviral immunity, necessitating further investigation (reviewed in [92,93]).

A specific subset of muscle-resident Tregs expressing amphiregulin and IL-10 has been shown to support muscle regeneration and repair following acute injury [94]. Moreover, Treg depletion exacerbated muscle degeneration in the mdx mouse model of muscular dystrophy while Treg potentiating treatment with low-dose IL-2 reduced muscle inflammation [95]. Muscle biopsies from human muscular dystrophy patients exhibit increased Treg frequencies versus healthy controls, similar to mdx versus control mice, suggesting endogenous repair mechanisms are insufficient to ameliorate the disease [95]. Hence, there is potential for ACT with Tregs modified to overexpress amphiregulin or other regenerative factors to support tissue repair in settings of injury or chronic degenerative disease such as muscular dystrophy.

**PRACTICAL CONSIDERATIONS**

Treg-based therapies hold great promise for inducing tolerance without many of the known toxicities observed with global immunosuppressive drugs (reviewed in [96–99]). In order to make Treg therapies feasible for widespread clinical use, it is essential to optimize the pipeline from isolation and expansion to infusion.

Treg purity of the final cell product will be a major factor determining therapeutic efficacy. Since most Treg therapy production strategies are likely to include expansions, elimination...
of non-Treg contaminants during the isolation step (Figure 1) is the most efficient way to achieve the purest product. In addition, specific Treg lineage and source (i.e., UCB Tregs or naïve peripheral blood Tregs, discussed below) may be considered when designing an enrichment and purification approach. Therefore, lineage and source selection, as well as isolation and enrichment strategies must be optimized to minimize the incidence of unintended immune activation and inflammation induced by the Treg ACT. Herein, we will discuss isolation strategies capable of generating a high yield and purity Treg product.

**Identification of human Treg**

FOXP3 serves as a master regulator of Treg lineage commitment and function. In human T cells, however, it alone does not define a Treg cell given that Tconv also transiently express intermediate levels of FOXP3 following activation [100]. FOXP3 expression in combination with another transcriptional regulator, Helios, denotes a thymic-derived tTreg population noted for phenotypic stability, suppressive potency, and ability to bestow these qualities unto daughter cell generations [101,102]. Within tTreg, the high degree of demethylation of the Treg-specific demethylated region (TSDR), found within the conserved non-coding DNA sequence 2 (CNS2) enhancer in the FOXP3 locus confers lineage stability [101–105]. Additionally, tTregs express homing receptors, including chemokine receptors and adhesion molecules that enable trafficking both systemically and to disease-relevant tissues [106–110]. As compared to pTreg populations, tTregs exhibit a higher degree of lineage stability upon in vivo transfer [101,104]. Hence, tTregs are commonly targeted for ACT, though head-to-head comparisons of Treg subsets are ongoing to determine which subset may be optimal in certain disease settings (reviewed in [111]).

With the identification of FOX-P3\(^{+}\)Helios\(^{+}\) Tregs, numerous advances in our understanding of immune regulation have been achieved. Unfortunately, labeling intracellular proteins (including these transcription factors) renders cells inviable; therefore, Treg isolation must rely on surface markers. In 2007, human Tregs were found to express low surface levels of the alpha chain of the IL-7 receptor, CD127, compared to Tconv [112], thereby enabling identification and isolation of live Treg based on the CD3\(^{+}\)CD4\(^{+}\)CD25\(^{+}\)CD127\(^{-/lo}\) phenotype [61]. Importantly, cells isolated by this phenotype are highly enriched for FOXP3\(^{+}\)Helios\(^{+}\) co-expression [113]. This observation represented a key step enabling future Treg cellular therapies in humans, given the ostensible importance of purity prior to reinfusion.

**Human Treg isolation**

Currently two methods exist for live Treg isolation: magnetic bead-based isolation or fluorescence-activated cell sorting (FACS). The former involves the closed-system isolation of CD4\(^{+}\)CD25\(^{+}\) T cells and was widely used for early trials, given prior experience with CD34\(^{+}\) HSC sorting and ACT. Strategies can be implemented to increase Treg purity from magnetic isolation such as: 1) depletion of CD8\(^{+}\) and CD19\(^{+}\) subsets before Treg isolation [15,16,18] or from the final post-expansion product [60]; 2) depletion of CD127\(^{+}\) population [114]; 3) perform multiple rounds of CD25\(^{+}\)-selection; or 4) limit the amount of anti-CD25 antibody loaded on the bead [115]. Unfortunately, these strategies dramatically reduce the overall yield and fail to eliminate activated and memory CD4\(^{+}\)CD25\(^{+}\) populations from the final product. Generally, this can be overcome by addition of rapamycin and/or all trans
retinoic acid (ATRA) to the Treg expansion media, which inhibit Tconv proliferation and promote conversion of naive T cells to induced Tregs, respectively [116].

FACS allows for precise gating based on marker expression level in contrast to microbead bulk enrichments. Importantly, emerging GMP-compliant, closed-loop sorting systems for FACS are expected to enable optimized isolation of CD4*CD25*CD127−/lo Tregs.

Soon to follow will likely be GMP-compatible reagents enabling the identification of precise cell subsets, potentially based on yet-to-be-defined surface proteins or technologies relying on transcriptional markers of phenotype [117]. For example, it is well recognized that CD4*CD25*CD127−/loCD45RA+ naïve Tregs exhibit greater stability during ex vivo expansion, while CD4*CD25*CD127−/loCD45RA-memory Tregs are more likely to downregulate FOXP3 and upregulate their expression of proinflammatory cytokines [118–120]. We recently demonstrated that the co-inhibitory receptor T-cell immunoglobulin and immunoreceptor tyrosine-based inhibitory motif domain (TIGIT) and Helios were highly co-expressed by Tregs [121]. Conversely, CD226, which directly competes with TIGIT to bind CD155, was most highly expressed by Tconv and FOXP3+Helios− pTregs. Of note, CD226 depleted Tregs exhibited comparable expansion kinetics and enhanced suppressive capacity compared to non-depleted cells. Eventually, protocols will narrow until the most functionally appropriate Treg, perhaps tailored to the disease/condition, can be delivered in the safest manner.

**Expansion approaches**

The number of peripheral blood mononuclear cells (PBMCs) that can be safely obtained from patients via peripheral blood draw or leukapheresis is limited according to body weight. Hence, obtaining clinically relevant numbers of Tregs sufficient to induce immunoregulation likely requires ex vivo expansion. Optimal conditions for Treg expansion involve multiple (i.e., 1-3) rounds of re-stimulation over the course of a 14-28 day culture in the presence of exogenous IL-2 [122–126]. Culture durations beyond this time frame are contraindicated because subsequent rounds of activation can compromise Treg suppressive function and stability [119]. Various stimulation approaches exist (i.e., antigen presenting cells [APCs] loaded with antigen, artificial APCs [aAPCs], allo-APCs, beads), each offering unique merits (Table 1, Figure 1).

The earliest aAPCs (K562 cells) are well described in the literature [127–129]. Briefly, K562 cells have been genetically modified to achieve long-term and stable expression of human CD32 and CD64 (KT32/64), which together bind the Fc region of anti-CD3 (OKT3) antibodies; hence, K562 aAPCs are capable of inducing robust Treg proliferation [128]. Moreover, Tregs stimulated through the CD28 pathway (by KT32/86 cells that constitutively express CD86) in addition to the CD3 pathway and in the presence of rapamycin showed 1000-fold expansion in three weeks with consistent suppression of xenogeneic GvHD in immunodeficient mice [130]. Cell-based aAPC expansion approaches provide a standardized platform for consistent Treg expansion and thus, are attractive for therapeutic applications.

Importantly, anti-CD3 and anti-CD28 coated microbeads offer an off-the-shelf GMP-compliant alternative to APC-based expansion. Just two rounds of activation (on day 0 and day 9) and 300U/ml IL-2 replenishment every two days routinely achieved >1000 fold...
Treg expansions [131,132]. We have also found that different GMP-grade products provide varying activation thresholds for TCR activation and co-stimulation impacting both the expansion efficiency and stability of the final Treg product [132].

**Treg expansion capacity and lineage stability**

The need to generate a sufficiently large, pure Treg population represents a significant challenge in Treg ACT. The optimal Treg dose is unknown and likely differs for each disease application. However, in mice that received a solid organ transplant, tolerance was achieved when approximately 30% of the organ-infiltrating CD4+ T cells were Tregs [133]. Supposing this value might translate to Treg cell therapy in settings of autoimmunity, Tang et al. calculated a theoretical target dose of 53x10^9 Treg to achieve a systemic Treg frequency of 30% among total CD4+ T cells in human patients [134]. Such a high dose is likely unfeasible due to practical/technical limitations and far exceeds the highest doses tested to date (100 × 10^6 Treg/kg [17]) but may support cryopreservation of Treg cells for subsequent repeat dosing. Additionally, efforts to optimize Treg ACT by selecting optimal cell sources and/or modulating their specificity, trafficking, or function, prior to reinfusion are ongoing in order to circumvent this potential limitation.

Efforts to expand either peripheral blood [4,7] or UCB-derived Tregs [61] have yielded cellular products that far surpass clinical release criteria for viability and/or CD8+ T cell contamination [4,7]. However, we recently reported that post-expansion Treg stability (i.e., FOXP3 and Helios co-expression as well as TSDR demethylation), purity (calculated from CD8+ T cell contamination), and Treg naivety (assessed by CD45RA expression) were significantly greater when Tregs were derived from UCB versus adult peripheral blood (APB) [61]. UCB Tregs also exhibited greater proliferative capacity compared to APB Tregs [61]. Moreover, expansion kinetics and post-expansion assessments (i.e., purity, stability, naivety) were comparable for Tregs isolated from fresh or cryopreserved UCB [61]. Hence, UCB may represent an optimal Treg source for ACT, eliminating the need for leukapheresis procedures, with cryopreserved UCB offering the potential for autologous Treg treatment.

**Treg Specificity**

Tregs are classically considered to modulate Tconv responses through both antigen-dependent and bystander suppression mechanisms [24], but they also exert their regulatory effects on the innate immune cells [135,136]. As a result, polyclonal Tregs may be particularly suitable for treatment of autoimmune or autoinflammatory conditions where the target antigen remains unknown or where a variety of autoantigens/tissues are targeted [137,138], but they might also offer a valuable treatment option for organ-specific diseases without the need for extensive ex vivo manipulation [4,7]. In contrast, antigen-specific Tregs, generated through TCR gene transfer, introduction of a CAR or selective expansion (Table 2), may traffic more efficiently to the site of cognate antigen expression and tissue draining lymph nodes [139–141], making it possible to achieve an efficient dose of Tregs in the target tissue/organ with a lower systemic Treg frequency. In a humanized mouse model of allotransplantation, selectively expanded alloantigen-reactive autologous Tregs were more effective at preventing allograft rejection compared to polyclonally expanded autologous Tregs [142]. Similarly, compared to polyclonal Tregs, 10-fold fewer β-cell
antigen-specific Tregs were able to reverse autoimmune diabetes in NOD mice [143]. However, it is important to consider that in human T1D various epitopes are targeted, including native β-cell antigens (i.e., insulin, GAD, IA-2, ZnT8) as well as hybrid insulin peptides (HIPs) and defective ribosomal insulin gene products (DRiPs) [144–147], so polyclonal or oligoclonal Treg preparations may provide greater autoantigen coverage. Moreover, we recently demonstrated that two TCRs reactive against the same peptide epitope with different avidities exhibited significantly different capacity to suppress Tconv [24].

Additional considerations, such as location of antigen expression/mode of antigen presentation, must guide receptor selection for antigen-specific Treg ACT. TCR redirected Tregs offer the benefit of recognizing intracellular antigen presented in the context of HLA, but the requirement for HLA compatibility requires use of autologous cells. In contrast, CAR-Tregs, which express the antigen binding region of an immunoglobulin (Ig) fused to intracellular T cell signaling domains [37,38], do not engage HLA but rather, recognize extracellular antigen. Extensive studies of CAR-T cells in the cancer field have set the stage for CAR and TCR-directed Tregs for ACT (summarized in Table 2). Ultimately, clinical studies are needed to identify the most appropriate Treg application (i.e., polyclonal or antigen-specific and TCRs or CARs) for use in various disease states.

**Engraftment & trafficking**

While the efficacious Treg dose has yet to be determined for the treatment of autoimmunity, a dose escalation study in adults with T1D demonstrated an exceptional safety profile even with high doses (up to 2.6x10⁹ cells) of ex vivo expanded autologous peripheral blood Treg. Importantly, Bluestone et al. reported that Tregs labeled with deuterated glucose were still detectable from circulation 12 months following transfer [4]. This suggests Tregs may display a similar long-term engraftment potential observed in cancer immunotherapy settings [148–151]. Additionally, IL-2 adjunct therapy may enhance Treg engraftment, similar to the effect of Ibrutinib with CAR-T treatment of chronic lymphocytic leukemia [152].

*Ex vivo* expanded human Tregs were shown to downregulate their expression of CCR5 [153], which directs T cell homing to sites of inflammation [154,155]. In contrast, human T cells exposed to the vitamin D analogue TX527 exhibited upregulated expression of the inflammatory homing receptors, CCR5, CXCR6, and CXCR3 [156]. CCR4⁺ human Treg migration into ovarian tumors was dependent upon CCL22 [157]. Furthermore, in a murine model of IBD, adoptive transfer of CCR4 competent Tregs prevented colitis, but CCR4 deficient Tregs were unable to traffic to the mesenteric lymph nodes resulting in disease [158]. Modification of chemokine receptor(s) and integrin expression could further guide Treg trafficking post-infusion. These approaches theoretically offer the additional benefit of reducing the likelihood of off-target suppression, which might impair protective immunity against an infection or cancer.
Regulatory requirements

Regulatory agencies, such as the U.S. Food and Drug Administration (FDA) and the European Medicines Agency [159], provide several guidelines about the cell therapy manufacturing process to ensure that each batch meets stringent quality standards concerned with cell collection, manufacturing area, the instruments and reagents used, and the release criteria of the final product prior to application. In addition to passing sterility screenings conducted throughout isolation and expansion, the final Treg product must satisfy purity criteria (e.g., composition of <5% CD8\(^+\) T cell contamination as well as >60% FOXP3\(^+\) Treg; or >96.7% cell viability, 99.6% CD4\(^+\), 98.9% CD25\(^+\), 0.0% CD127\(^+\), 0% CD8\(^+\), 0.0% CD19\(^+\) and 0.0% CD56/16\(^+\)) [4,60] and potentially, confirmation of functional suppression in vitro [60] prior to infusion. Thought must also be given to the timing of Treg infusion, particularly in children, with relation to vaccination schedules.

For research use, FACS occurs in an open system; therefore, the clinical application requires a modified, closed instrument, in Grade A “clean rooms”, equipped with high efficiency particulate air (HEPA) filters to limit air particulates and avoid microbial contamination. In contrast, cell isolation using CliniMACS (Miltenyi) is performed in a closed system with disposable tubing limiting the potential for contamination inherent in droplet based FACS approaches. As a result, magnetic bead-based isolation currently offers better affordability and a shorter cell sorting process but is limited to basic bimodal sorts. However, as mentioned above, emergent closed-loop FACS technologies are highly anticipated. Indeed, this interface will enable more widespread application of cellular therapies offering more precise and affordable CMP isolation of CD4\(^+\)CD25\(^+\)CD127\(^{-/lo}\) Tregs for a highly pure cell product.

FUTURE DIRECTIONS FOR TREG CELLULAR THERAPY

Moving forward, advances in isolation, expansion, and cellular engineering technologies are expected to enable precise engineering of optimal Treg products for various therapeutic applications (reviewed in [160]). Gene editing techniques, such as endogenous TCR ablation, are expected to reduce potential heterologous chain pairing. Moreover, allogeneic cell sources for TCR-redirected Treg or CAR-Treg therapies could eliminate the need for invasive leukapheresis procedures. Given the inherently different argument for equipoise in chronic conditions such as autoimmunity and allergy versus terminal cancer, efforts are underway to introduce additional safeguards, including suicide genes [37,161,162], to enable depletion of the cells post-infusion in the event of off target suppression or an unforeseen adverse reaction [161,162].

Gene editing could theoretically be used to optimize “designer Tregs” via knockdown of molecules known to inhibit suppressive function, introduction of regulatory elements (e.g., TIGIT [121,163]), expression of growth factors (e.g., amphiregulin [94,95]), and/or site-specific mutagenesis of single nucleotide polymorphism (SNPs) associated with disease risk/pathogenesis specific to the intended application. For example, Tregs from donors carrying a SNP risk variant in protein tyrosine phosphatase, non-receptor type (PTPN22), which confers risk in several autoimmune disorders and outside of HLA and insulin, has the highest odds ratio for T1D (reviewed in [164]), can be gene-edited to the protective
variant prior to ex vivo expansion. Ectopic overexpression of FOXP3 in autologous Tconv cells could transform our ability to treat IPEX syndrome (reviewed in [165]). Additionally, we envision that it may eventually be possible to treat both allograft rejection and autoimmunity in pancreas or islet transplant recipients with T1D via ACT of CAR-Tregs against donor HLA plus polyclonal or TCR redirected Tregs specific for β-cell autoantigens, albeit likely in combination with additional immunomodulatory agents.

Our lab and others are currently pursuing biomaterials applications to optimize Treg survival and function in vivo following infusion. Poly(d-lactide-co-glycolide) (PLGA) is degraded in the body via bulk erosion and hydrolysis into glycolic acid and lactic acid, conferring no harmful side effects [166]; consequently, PLGA is used in numerous devices/therapeutics approved by the FDA, such as biodegradable surgical sutures and drug delivery products, which makes it extremely attractive for development of products quickly translatable to clinical use. PLGA can be used to produce micro- or nanoparticles (NPs), which have been widely investigated for delivering immunotherapeutics [167,168]. By altering the lactide/glycolide ratio, PLGA NPs can be designed to provide tunable sustained release of encapsulated immunomodulatory molecules [169]. Moreover, NPs can be engineered to simultaneously deliver therapeutic factors and adjuvant drugs, each with tailored release kinetics while maintaining the tissue-targeting capacity of a CAR or TCR. Indeed, controlled release of immunomodulatory agents by PLGA particulate systems are being explored as immunotherapeutic tools for treatment of cancer, autoimmunity and transplant-related complications [170–172]. An attractive strategy is to focus drug and growth factor action on adoptively-transferred, therapeutic Tregs through the conjugation of agent-loaded NPs directly onto the cell surface via the incorporation of reactive groups into the PLGA formulation [173]. This flexible approach allows for concurrent in vivo delivery of therapeutic agents in both an autocrine and paracrine manner. As result, it may be possible to simultaneously conjugate Tregs to PLGA NPs loaded with Treg growth factors (e.g., IL-2) and to PLGA NPs loaded with therapeutic agents designed to impact the extracellular milieu upon Treg migration to sites of inflammation.

Clearly, a vast number of possible targets for cellular manipulation exist, supporting a synthetic biology approach to therapeutic T cell engineering [174]. By establishing a standardized series of ex vivo modifications for each target tissue/disease application, scientists and clinicians will be able to ensure infusion of a consistent therapeutic product, essential for both safety and efficacy. The feasibility of using Tregs as a future ACT to restrain inflammation is supported by numerous studies utilizing either allogeneic or autologous effector T cells to augment immunity in cancer patients (reviewed in [175–177]). The therapeutic potential of these cells has been further augmented by genetic engineering, conferring upon T cells the ability to target a specific tumor-associated antigen while simultaneously increasing their lifespan and reducing potential toxicity [32–38]. As a result, ACT-based immunotherapies have been extraordinarily successful in treating cancer and infectious diseases, and the lessons learned from those efforts will afford an advantage toward the design and implementation of novel Treg therapies.
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Figure 1. *Ex vivo* human Treg expansion protocols reported in the literature.

Isolation of Tregs from UCB or APB is accomplished by either MACS or FACS. FACS is often used to sort out an enriched population of CD4+CD25+CD127lo/-CD45RA+ cells directly from the starting sample, but MACS can be employed prior to sorting to remove contaminating CD8+ and CD19+ cell types [84,114,125]. Likewise, MACS can be utilized to pre-enrich for CD25+ cells prior to FACS to obtain a highly enriched population of CD4+CD25+CD127lo/-CD45RA+ cells. Regardless of the isolation protocol, these highly enriched cells are then cultured in the presence of cytokines to support Treg proliferation and inhibitory compounds such as rapamycin to maintain Treg purity. Expansion of the Treg population is dependent on the introduction of stimulatory agents in the form of either anti-CD3/CD28 monoclonal antibody coated beads, allogenic DCs, or artificial APCs. Expansion of Treg populations requires up to 28 days of culture with multiple re-stimulations and frequent cytokine supplementation of the growth media. Expanded cells are validated for fold expansion, stable expression of Treg markers, purity, and stability before being used in therapeutic applications or experimental studies designed to enhance Treg specificity and potency.

APB: Adult peripheral blood; APC: Antigen presenting cell; DC: Dendritic cell; FACS: Fluorescence activated cell sorting; GVHD: Graft versus host disease; MACS: Magnetic activated cell sorting; NK: Natural killer; PBMC: Peripheral blood mononuclear cell; UCB: Umbilical cord blood.
TABLE 1

GMP polyclonal Treg isolation and/or expansion strategies from peripheral blood mononuclear cells or umbilical cord blood reported in the literature.

| Treg Source | Isolation | Depleted Subsets | Depletion pre- or post-isolation | Activation | Stimulation (days) | Expansion time (days) | IL-2 (IU/ml) | Rapamycin | Fold-expansion | Final cell number | Purity (%) | Target Dose | Disease | Ref. |
|-------------|-----------|-----------------|---------------------------------|------------|-------------------|----------------------|---------------|------------|----------------|-----------------|------------|-------------|---------|------|
| PBMC        | ClinMACS  | CD8+            | Pre                             | Dynabeads  | 0                 | 10                   | 25            | –          | 31±16         | –               | 57         | –           | Autoimmunity, transplantation | [114] |
| PBMC        | ClinMACS  | CD8+            | Pre                             | alogeneic: PBMC | 0                 | 10                   | 25            | –          | 28±14         | –               | 54         | –           | Autoimmunity, transplantation | [114] |
| PBMC        | ClinMACS  | CD8+, CD19+     | Pre                             | Dynabeads  | 0                 | 10                   | 25            | –          | 34±17         | –               | –          | –           | Autoimmunity, transplantation | [114] |
| PBMC        | ClinMACS  | CD8+, CD19+     | Pre                             | alogeneic: PBMC | 0                 | 10                   | 25            | –          | 34±11         | –               | –          | –           | Autoimmunity, transplantation | [114] |
| PBMC        | FACS      | –               | –                               | Dynabeads, aAPCs | 0, 9               | 14                   | 300           | –          | 480           | –               | 92         | 0.5 x 10^6/kg | –       | [141] |
| PBMC        | ClinMACS  | CD8+, CD19+     | Pre                             | aAPCs (KT64/86) | 0, ~10, ~20, ~30, ~40 | ~45                   | 300           | 109nM      | 5x10^6         | –               | 60         | –           | GvHD    | [125] |
| PBMC        | AutoMACS  | CD8+, CD19+     | –                               | aAPCs (KT64/86) | 0, ~13, ~26, ~42, ~56 | 70                   | 300           | 109nM      | 31x10^6        | –               | 63         | –           | GvHD    | [125] |
| PBMC        | ClinMACS  | CD8+            | Pre                             | MACS ExpAct | 0, 12, 24          | 36                   | 500           | 100nM      | 1430–2080      | 680x10^6–2.680x10^6 | 89–91     | 4.5 x 10^6/kg | LT       | [84] |
| UCB         | FACS      | –               | –                               | MACS ExpAct | 0, 9, 18           | 27                   | 600           | –          | 15,000        | 3x10^10          | >90        | 30 x 10^6/kg | T1D     | [61] |
| UCB         | FACS      | –               | –                               | aAPC & Dynabeads | 0, 9               | 14                   | 600           | –          | 1,200         | 4x10^9          | >90        | 30 x 10^6/kg | T1D     | [61] |
| UCB         | FACS      | –               | –                               | Dynabeads  | 9                  | 16                   | 300–600       | –          | 2,100         | 1.1x10^10        | >90        | 30 x 10^6/kg | T1D     | [61] |
| PBMC        | ClinMACS  | –               | –                               | MACS ExpAct | 0                  | 21                   | 500           | 100ng/ml   | 22–778        | –               | 90         | –           | UC      | [59] |
| PBMC        | ClinMACS  | CD8+, CD19+     | Post                            | MACS ExpAct | 0, 14              | 21                   | 1000          | 100ng/mL   | 59.4–117       | 528x10^6–1,440x10^6 | 94        | –           | Various autoimmune and inflammatory disorders | [60] |

Clinical trials of Treg adoptive cell therapy
| Treg Source | Isolation | Depleted Subsets | Depletion pre- or post-isolation | Activation | Stimulation (days) | Expansion time (days) | IL-2 (IU/ml) | Rapamycin | Fold-expansion | Final cell number | Purity (%) | Target Dose | Disease | Ref. |
|-------------|-----------|-----------------|-------------------------------|-----------|-------------------|----------------------|--------------|-----------|---------------|------------------|------------|-------------|---------|------|
| PBMC (Buffy Coat) | MACS/ FACS | CD8+, CD19+ | – | Dynabeads | 21 | 1000 | – | – | – | – | >95 | 3x60x10⁶/kg | aGvHD | [13] |
| UCB | ClinMACS | – | – | Dynabeads | 0 | 18±1 | 300 | – | 13–1796 | 74 x 10⁶–12.6 x 10⁷ | 62–97 | 0.1–3 x 10⁹ | aGvHD | [14] |
| dPBMC | ClinMACS | CD8+, CD19+ | Pre | – | – | – | – | – | – | – | 93 | 2–4 x 10⁹/kg | GvHD | [15] |
| PBMC | Ficoll, limiting dilution | – | – | S2 cells, expressing CD80, CD58, antiCD3, IL-2/4, OVA | 0 | 7+ | Feeder cell derived | – | – | 1 x 10⁹ | 55 | 10⁶–10⁹ | CD | [5,178] |
| PBMC | ClinMACS | CD8+, CD19+ | Pre | – | – | – | – | – | – | – | 81 | 2.5±1 x 10⁹/kg | GvHD | [18] |
| dPBMC | ClinMACS | CD8+ | Pre | Dynabeads, MACS ExpAct | 0, 7/8 | 7–12 | 300–1000 | 100ng/ml | 2.5–17.7 | 1.6 x 10⁸–23 x 10⁷ | 78–92 | 1.19–5.47 x 10⁹/kg | cGvHD | [16] |
| PBMC | FACS | – | – | Dynabeads | 0, 9 | 14 | 300 | – | 30–1367 | 0.19–13 x 10⁹ | 76–97 | 5–2940 x 10⁶ | T1D | [4] |
| UCB | ClinMACS | – | – | aAPCs (KT6/86) | 0, 12±1 | 18±1 | 300 | – | 1352–27183 | 1.88 x 10⁸–4.26 x 10⁹ | 83–95 | 3–100 x 10⁹/kg | GvHD | [17] |
| dPBMC | ClinMACS FACS | – | – | ExpAct | 0, 7 | 13 | 1000 | – | 177 | 1.18 x 10⁸–2.85 x 10¹⁰ | 90 | – | Autoimmunity, transplantation | [179] |
| PBMC | FACS | – | – | Dynabeads | 0, 9 | 14 | 300 | – | 300 | ≥1 x 10⁹ | 93–97 | 319–363.8 x 10⁶ | KT | [7] |

Treg sources were autologous PBMC, donor PBMC, autologous UCB, or donor UCB. Cell isolation was performed by ClinMACS or FACS with or without magnetic depletion of CD19+ and/or CD8+ cells pre- or post-expansion. Method and regimen for Treg activation (schedule, IL-2 dose, Rapamycin dose), subsequent fold expansion, final cell number, and purity are listed along with the target Treg dose and the proposed or tested disease application, including GvHD (aGvHD, cGvHD), LT, UC, T1D, KT, and CD.

aGvHD: Acute graft versus host disease; aPBMC: Autologous peripheral blood mononuclear cells; aUCB: autologous umbilical cord blood; CD: Crohn’s Disease; ClinMACS: Magnetic bead-based cell separation; cGvHD: Chronic graft versus host disease; dPBMC: donor peripheral blood mononuclear cells; dUCB: Donor umbilical cord blood; FACS: Fluorescence activated cell sorting; GvHD: Graft versus host disease; KT: Kidney transplant; LT: Liver transplant; PBMC: Peripheral blood mononuclear cells; T1D: Type 1 diabetes; UC: Ulcerative colitis; UCB: Umbilical cord blood.

– indicates not applicable, not performed, or not reported.
### TABLE 2

Studies of CAR-Tregs and TCR modified/enriched Tregs from mice and humans.

| Treg Modification | Method of Receptor Modification/Enrichment | Receptor Specificity | Disease/model | Ref. |
|-------------------|-------------------------------------------|----------------------|---------------|------|
| TCR               | Incubation w/ host allogenic splenocytes   | Alloantigen          | Graft-versus-host disease | [180] |
| TCR               | Transgenic mouse model                     | Ovalbumin            | Colitis       | [181] |
| TCR               | Transgenic mouse model                     | Beta cell antigen (BDC2.5) & pancreatic islet antigen (GAD286) | Type 1 diabetes | [25] |
| TCR               | Transgenic mouse model                     | Beta cell antigen (BDC2.5) | Type 1 diabetes | [143, 182] |
| TCR               | Transgenic mouse model/antigen specific beads (p31-I-Ag^7) | Beta cell antigen (BDC2.5), pancreatic islet antigen (GAD286), p31-I-Ag^7 | Type 1 diabetes | [183] |
| TCR               | Incubation w/ host allogenic antigen-presenting cells | Alloantigen | Graft-versus-host disease | [184] |
| TCR               | Incubation w/ peptide-pulsed antigen-presenting cells | H2 Kb allelopeptide | Transplantation | [185] |
| TCR               | Retroviral transduction                    | H-2Kd haplotype      | Transplantation | [186] |
| TCR               | Donor strain-derived antigen-presenting cells | Directly and indirectly presented alloantigen | Transplantation | [187] |
| TCR               | Transgenic mouse model                     | Myelin basic protein | Multiple sclerosis | [188] |
| TCR               | Retroviral TCR gene transfer               | Ovalbumin (OTII-TCR) | Rheumatoid Arthritis | [189] |
| TCR               | Lentiviral TCR gene transfer               | Melanoma antigen tyrosinase | Adoptive cell therapy | * |
| TCR               | Mixed lymphocyte reaction                  | HLA-DRb1 Alloantigen | Transplantation | * [190] |
| TCR               | Native to isolated Treg population         | Ovalbumin            | Crohn's disease | * |
| TCR               | Mixed lymphocyte reaction (CD40L-stimulated B cells) | Diverse TCR | Transplantation | * [142] |
| TCR               | Retroviral transduction                    | Factor VIII (FVIII)  | Hemophilia     | * [81] |
| TCR               | Lentiviral TCR gene transfer               | Pancreatic islet antigen | Type 1 diabetes | * [30] |
| TCR               | Lentiviral TCR gene transfer               | GAD55-567            | Type 1 diabetes | * [24] |
| CAR               | Transgenic mouse model                     | Antigen MBP89-101    | Autoimmunity (EAE) | [191, 192] |
| CAR               | Transgenic mouse model                     | 2,4,6-trinitrophenol | Colitis        | [193] |
| CAR               | Lentiviral transduction                    | Myelin oligodendrocyte glycoprotein | Multiple Sclerosis | [194] |
| CAR               | Retroviral transduction                    | Carcinobryonic antigen | Colitis        | [195] |
| CAR               | Lentiviral transduction                    | HLA-A2               | Graft rejection | * [196] |
| CAR               | Retroviral transduction                    | HLA-A2               | Graft rejection | * [197] |
| Treg Modification | Method of Receptor Modification/Enrichment | Receptor Specificity | Disease/model | Ref. |
|-------------------|------------------------------------------|---------------------|---------------|-----|
| CAR               | Lentiviral transduction                   | HLA-A2              | Graft rejection* | [85] |
| CAR               | Retroviral transduction                   | Factor VIII (FVIII) | Hemophilia*    | [31] |
| CAR               | Transgenic mouse model                    | Carcinoembryonic antigen | Asthma | [11] |
| mAb-CAR           | Transient genomic DNA plasmid transgene   | Modular (monoclonal antibody-directed) | Alloreactivity | [198] |

* Human study.