Controlling graft-versus-host disease (GVHD) remains a major unmet need in stem cell transplantation, and new, targeted therapies are being actively developed. CD28-CD80/86 costimulation blockade represents a promising strategy, but targeting CD80/CD86 with CTLA4-Ig may be associated with undesired blockade of coinhibitory pathways. In contrast, targeted blockade of CD28 exclusively inhibits T cell costimulation and may more potently prevent GVHD. Here, we investigated FR104, an antagonistic CD28-specific pegylated-Fab’, in the nonhuman primate (NHP) GVHD model and completed a multiparameter interrogation comparing it with CTLA4-Ig, with and without sirolimus, including clinical, histopathologic, flow cytometric, and transcriptomic analyses. We document that FR104 monoprophylaxis and combined prophylaxis with FR104/sirolimus led to enhanced control of effector T cell proliferation and activation compared with the use of CTLA4-Ig or CTLA4-Ig/sirolimus. Importantly, FR104/sirolimus did not lead to a beneficial impact on Treg reconstitution or homeostasis, consistent with control of conventional T cell activation and IL-2 production needed to support Tregs. While FR104/sirolimus had a salutary effect on GVHD-free survival, overall survival was not improved, due to death in the absence of GVHD in several FR104/sirolimus recipients in the setting of sepsis and a paralyzed INF-γ response. These results therefore suggest that effectively deploying CD28 in the clinic will require close scrutiny of both the benefits and risks of extensively abrogating conventional T cell activation after transplant.
CD28 blockade controls T cell activation to prevent graft-versus-host disease in primates

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Controlling graft-versus-host disease (GVHD) remains a major unmet need in stem cell transplantation, and new, targeted therapies are being actively developed. CD28-CD80/86 costimulation blockade represents a promising strategy, but targeting CD80/CD86 with CTLA4-Ig may be associated with undesired blockade of coinhibitory pathways. In contrast, targeted blockade of CD28 exclusively inhibits T cell costimulation and may more potently prevent GVHD. Here, we investigated FR104, an antagonistic CD28-specific pegylated-Fab’, in the nonhuman primate (NHP) GVHD model and completed a multiparameter interrogation comparing it with CTLA4-Ig, with and without sirolimus, including clinical, histopathologic, flow cytometric, and transcriptomic analyses. We document that FR104 monophrophylaxis and combined prophylaxis with FR104/sirolimus led to enhanced control of effector T cell proliferation and activation compared with the use of CTLA4-Ig or CTLA4-Ig/sirolimus. Importantly, FR104/sirolimus did not lead to a beneficial impact on Treg reconstitution or homeostasis, consistent with control of conventional T cell activation and IL-2 production needed to support Tregs. While FR104/sirolimus had a salutary effect on GVHD-free survival, overall survival was not improved, due to death in the absence of GVHD in several FR104/sirolimus recipients in the setting of sepsis and a paralyzed INF-γ response. These results therefore suggest that effectively deploying CD28 in the clinic will require close scrutiny of both the benefits and risks of extensively abrogating conventional T cell activation after transplant.

Introduction

Graft-versus-host disease (GVHD) represents a major cause of mortality after hematopoietic stem cell transplant (HCT) (1–4), and despite substantial effort, the control of GVHD remains one of the key unmet needs in the field. Moreover, prevailing calcineurin-based therapies are associated with significant off-target toxicities (5, 6) and are not associated with immune tolerance induction (7, 8). This has intensified investigation into calcineurin-free regimens, employing strategies that have been shown in preclinical models to be protolerogenic. Of the new approaches, those targeting T cell costimulation are being increasingly investigated in auto- and alloimmunity (9, 10) and are beginning to be explored to prevent and treat acute GVHD (aGVHD) (11–13). One of the most potent T cell costimulation pathways occurs through CD28:CD80/86 ligation (14). Blockade of the CD80/86 component of CD28:CD80/86 costimulation with CTLA4-Ig (using either of the 2 FDA-approved formulations, abatacept or belatacept) represents the first successful clinical translation of costimulation blockade (9, 11, 12, 15–20). However, targeting CD80/86 has not been entirely successful in downregulating auto- and alloimmunity (10, 21–23) and may be associated with untoward immune-activating effects through blockade of CD80/86-dependent coinhibitory pathways (24–35). Given these challenges with CTLA4-Ig, there has been growing interest in more selectively targeting CD28 (36–41), and these approaches have shown efficacy in selected preclinical models (42–49). Although the molecular mechanisms by which CD28 blockade controls T cell alloactivation have been proposed (50), the comprehensive interrogation of the relative efficacy of targeting CD28 versus CD80/86 and associated safety signatures (both alone and when combined with other immunomodulators) have been undetermined, and these aspects, which have not been comprehensively interrogated, represent key questions in the field, given that reagents targeting both CD80/86 and CD28 are now clinically available.

To thoroughly address these mechanistic issues, we used the nonhuman primate (NHP) GVHD model to evaluate a clinically available therapeutic anti-CD28 Fab’, FR104 (42, 43, 45, 46, 48), in order to determine its impact on T cell function, reconstitution, and GVHD, both as a monophrophylaxis and combined with the
monitored either as monoprophylaxis or in combination with sirolimus for its ability to prevent GVHD in the setting of MHC-haploidentical T cell–replete HCT (Figure 1A). Figure 1B shows the resulting FR104 pharmacokinetics (PKs) in the current study. During the period of antibody dosing, the average FR104 trough was 28.6 μg/ml ± 2.3 μg/ml, a level previously shown to successfully block CD28 in healthy controls (HCs) (46). After antibody discontinuation at day 54, FR104 levels steadily declined, with a terminal half-life of 5.6 ± 1.0 days, resulting in undetectable FR104 levels after day 66 after transplant (Figure 1B) (45, 46). To directly assess CD28 receptor occupancy with FR104 in the setting of HCT, we measured the crossblocking of CD28 expression flow cytometrically using a mouse anti-CD28 antibody (clone CD28.2) prior to each FR104 dose (Figure 1C). This analysis confirmed more than 97.5% and more than 99% CD28 blockade on CD4+ and CD8+ T cells.

Results

CD28 blockade: pharmacokinetics, pharmacodynamics, and donor engraftment. In this study, CD28 blockade with FR104 was evaluated either as monoprophylaxis or in combination with sirolimus for its ability to prevent GVHD in the setting of MHC-haploidentical T cell–replete HCT (Figure 1A). Figure 1B shows the resulting FR104 pharmacokinetics (PKs) in the current study. During the period of antibody dosing, the average FR104 trough was 28.6 μg/ml ± 2.3 μg/ml, a level previously shown to successfully block CD28 in healthy controls (HCs) (46). After antibody discontinuation at day 54, FR104 levels steadily declined, with a terminal half-life of 5.6 ± 1.0 days, resulting in undetectable FR104 levels after day 66 after transplant (Figure 1B) (45, 46). To directly assess CD28 receptor occupancy with FR104 in the setting of HCT, we measured the crossblocking of CD28 expression flow cytometrically using a mouse anti-CD28 antibody (clone CD28.2) prior to each FR104 dose (Figure 1C). This analysis confirmed more than 97.5% and more than 99% CD28 blockade on CD4+ and CD8+ T cells.

mTOR inhibitor sirolimus, and to compare this agent with CTLA4-Ig–based GVHD prevention strategies (51). These experiments provide strong evidence for superior control of T cell proliferation and activation when CD28 is directly targeted, as compared with when CD80/86 is targeted. However, while the combination of anti-CD28 and sirolimus led to superior GVHD-free survival, it did not enhance Treg homeostasis, nor did it improve overall recipient survival as compared with CTLA4-Ig/sirolimus. These studies underscore the complex nature of targeted blockade of cosignaling pathways and suggest that head-to-head comparisons of these agents are indicated in clinical trials for GVHD prevention and treatment.

Results

CD28 blockade: pharmacokinetics, pharmacodynamics, and donor engraftment. In this study, CD28 blockade with FR104 was evaluated either as monoprophylaxis or in combination with sirolimus for its ability to prevent GVHD in the setting of MHC-haploidentical T cell–replete HCT (Figure 1A). Figure 1B shows the resulting FR104 pharmacokinetics (PKs) in the current study. During the period of antibody dosing, the average FR104 trough was 28.6 μg/ml ± 2.3 μg/ml, a level previously shown to successfully block CD28 in healthy controls (HCs) (46). After antibody discontinuation at day 54, FR104 levels steadily declined, with a terminal half-life of 5.6 ± 1.0 days, resulting in undetectable FR104 levels after day 66 after transplant (Figure 1B) (45, 46). To directly assess CD28 receptor occupancy with FR104 in the setting of HCT, we measured the crossblocking of CD28 expression flow cytometrically using a mouse anti-CD28 antibody (clone CD28.2) prior to each FR104 dose (Figure 1C). This analysis confirmed more than 97.5% and more than 99% CD28 blockade on CD4+ and CD8+ T cells.
On day 7 after transplant, each of the other monoprophylaxis cohorts displayed moderate-to-severe clinical aGVHD, while animals receiving FR104 monoprophylaxis were still significantly controlling disease (Figure 2C). However, as shown in Figure 2, A, D, and E, GVHD did ultimately develop in the FR104 monoprophylaxis cohort. The delay in the onset of aGVHD in the FR104 monoprophylaxis cohort was associated with a survival advantage compared with both the No Rx and CTLA4-Ig cohorts (MST for the FR104 monoprophylaxis cohort = 21 days, \( P < 0.01 \) versus No Rx, \( P = 0.013 \) versus CTLA4-Ig, \( P = 0.95 \) versus sirolimus; Figure 2B).

Transcriptomic analysis of T cells purified at day 14 from the peripheral blood of recipients in the FR104 monoprophylaxis cohort revealed more complete blockade of CD28 expression in the bone marrow, gastrointestinal (GI) tract, spleen, and lymph node (LN), with more variable blockade in the liver (Figure 1, D and E). However, as expected from the PK analysis, after day 66 after transplant, we observed desaturation of CD28 occupancy in both blood and tissue T cells (Figure 1E).

**Figure 2. Clinical aGVHD scoring and aGVHD-free survival.** (A and B) Longitudinal clinical aGVHD scoring (A) and GVHD-free survival curves (B) of untreated (No Rx, \( n = 11 \); red), CTLA4-Ig (\( n = 4 \); pink), sirolimus (\( n = 6 \); orange), FR104 (\( n = 3 \); blue), FR104/sirolimus (\( n = 9 \); navy), and CTLA4-Ig/sirolimus (\( n = 7 \); purple) cohorts. Scoring was based on our previously described NHP aGVHD clinical scoring system (12). GI aGVHD scores from FR104/sirolimus recipients with documented enteric infections were censored. Data are shown as mean ± SEM. For GVHD-free survival analysis, FR104/sirolimus recipients with documented graft rejection were excluded from this analysis. The Kaplan-Meier product-limit method was used to calculate survival. (C) Clinical aGVHD scoring in untreated (No Rx, \( n = 11 \)), CTLA4-Ig (\( n = 4 \)), sirolimus (\( n = 6 \)), and FR104 (\( n = 3 \)) cohorts on day 7. GI aGVHD scores from FR104/sirolimus recipients with documented enteric infections were censored. Data are shown as mean ± SEM. Statistical analysis was performed using 1-way ANOVA with Holm-Šidák multiple-comparison post test. (D) Clinical aGVHD scoring in the following cohorts, each at terminal analysis: untreated (No Rx, \( n = 11 \)), CTLA4-Ig (\( n = 4 \)), sirolimus (\( n = 6 \)), FR104 (\( n = 3 \)), FR104/sirolimus recipients undergoing terminal analysis before day 66 (\( n = 6 \)), FR104/sirolimus recipients undergoing terminal analysis after day 66 (\( n = 3 \)), CTLA4-Ig/sirolimus (\( n = 7 \)), and autologous controls (\( n = 3 \)). Data are shown as mean ± SEM. Statistical analysis was performed using 1-way ANOVA with Holm-Šidák multiple comparison post test. (E) Terminal aGVHD histopathologic scores. The scores shown represent the total score for the skin, liver, and GI tract (83). GI aGVHD scores from FR104/sirolimus recipients with documented enteric infections were censored. Data are shown as mean ± SEM. Statistical analysis was performed using 1-way ANOVA with Holm-Šidák multiple-comparison post test. *\( P < 0.05 \); **\( P < 0.01 \); ***\( P < 0.001 \).
cohorts. These included gene sets indicating relative preservation of naive T cells in the FR104 versus No Rx comparison (Figure 3A) and underrepresentation of gene sets associated with cell proliferation, T cell antigen-dependent activation, and effector differentiation in the FR104 cohort (Figure 3B). In contrast, when compared with HCs, the day 14 FR104 monoprophylaxis T cells were clearly not quiescent and demonstrated relative enrichment for these same gene sets (Figure 3C). Thus, while relatively controlled compared with fulminant hyperacute GVHD, immune escape during FR104 monoprophylaxis was nonetheless occurring.

The monoprophylaxis cohorts also permitted a direct comparison of the transcriptome of T cells in the setting of CD28 blockade versus blockade of CD80/86, and as documented in Figure 3, Supplemental Table 2, there were numerous gene sets

cohort (Figure 3 and Supplemental Tables 1 and 2; supplemental material available online with this article; https://doi.org/10.1172/JCI98793DS1) provided insights into the immune pathways responsible for the bioactivity of FR104 in delaying GVHD as well as the ultimate immune escape that occurred despite anti-CD28 monoprophylaxis. Differential expression analysis highlighted the ultimate similarities in the severity of GVHD in both the FR104 and No Rx cohorts in that there were no differentially expressed (DE) genes between these 2 cohorts using a differential expression threshold of 1.4-fold and a P value of less than 0.05, corrected for multiple testing (Supplemental Table 1). However, despite the lack of individual transcript differences, gene set enrichment analysis (GSEA) (54), revealed relative over- and underrepresentation of multiple gene sets in the FR104 monoprophylaxis versus No Rx cohorts. These included gene sets indicating relative preservation of naive T cells in the FR104 versus No Rx comparison (Figure 3A) and underrepresentation of gene sets associated with cell proliferation, T cell antigen-dependent activation, and effector differentiation in the FR104 cohort (Figure 3B). In contrast, when compared with HCs, the day 14 FR104 monoprophylaxis T cells were clearly not quiescent and demonstrated relative enrichment for these same gene sets (Figure 3C). Thus, while relatively controlled compared with fulminant hyperacute GVHD, immune escape during FR104 monoprophylaxis was nonetheless occurring.

The monoprophylaxis cohorts also permitted a direct comparison of the transcriptome of T cells in the setting of CD28 blockade versus blockade of CD80/86, and as documented in Figure 3, D–F, and Supplemental Table 2, there were numerous gene sets...
that were significantly different in T cells purified from these recipients. Importantly, this analysis uncovered evidence for the absence of CTLA4 signaling in the CTLA4-Ig cohort compared with the FR104 cohort (Figure 3D). In addition, GSEA identified significant enrichment for resting/naive T cell gene signatures in the FR104 cohort compared with the CTLA4-Ig cohort (Figure 3E). In contrast, T cells isolated from recipients in the CTLA4-Ig monophrophylaxis cohort demonstrated enrichment for gene sets involved in (a) cell cycle/proliferation, (b) effector T cells, (c) antigen response, and (d) cytotoxicity (Figure 3F).

**Long-term control of GVHD with FR104/sirolimus combination prophylaxis.** While both sirolimus and FR104 as monoprophylaxis only modestly delayed the onset and lethality of aGVHD (Figure 2, A–D), the combination of these agents synergistically controlled disease. Two FR104/sirolimus cohorts were examined. In cohort 1 (short-term, recipients R.26, R.27, R.28, and R.29; Table 1), animals underwent a planned terminal analysis at days 33–35 after transplant, with 3 of 4 recipients reaching the predetermined experimental endpoint. The single recipient that did not reach the experimental endpoint: R.222 died at day 21 of a hemorrhagic stroke prior to platelet engraftment (platelet count 14 at terminal analysis); R.213 died at day 66 unexpectedly after anesthesia, in the setting of intercurrent giardia infection; R.250 died at day 72 of culture-negative sepsis. None of these animals had signs of clinical GVHD at the time of their deaths.

As shown in Figure 2B, despite the early deaths, the clinical protection from aGVHD resulted in prolongation of GVHD-free survival in the FR104/sirolimus cohort compared with multiple other cohorts, including the No Rx, sirolimus, and CTLA4-Ig monophrophylaxis cohorts as well as the CTLA4-Ig/sirolimus cohort (51): \( P = 0.0001 \) vs. No Rx (MST = 8 days); \( P = 0.0001 \) vs. CTLA4-Ig (MST = 10 days); \( P = 0.0005 \) vs. sirolimus (MST = 17 days); \( P < 0.0014 \) vs. FR104 (MST = 21 days); and \( P = 0.004 \) vs. CTLA4-Ig/sirolimus (MST = 34 days). As shown in Figure 2, D and E, histopathologic GVHD scores for animals in the FR104/sirolimus cohort were divided into 2 groups: (a) those necropsied during ongoing FR104 exposure (defined as animals necropsied prior to day 66 after transplant, with this cutoff based on the PK analysis depicted in Figure 1B) and (b) those that were necropsied after FR104 washout (Figure 2, D and E). The histopathologic GVHD scores of animals necropsied during ongoing FR104 exposure aligned with clinical aGVHD scores, with significantly less GVHD-associated pathology in comparison with the No Rx cohort (2.2 ± 0.5 vs. 6.9 ± 0.6; \( P = 0.003 \)), which was similar to what was found in autologous transplant controls. Of note, while animals necropsied after FR104 washout did not show significant clinical signs of GVHD at terminal analysis (Figure 2D), histopathologic analysis did indicate subclinical pathology in these animals in the skin, liver, and colon (Figure 2E and Supplemental Table 3), with a total histopathologic score of 4.2 ± 0.8 (\( P = 0.94 \) compared with No

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**Table 1. FR104 and FR104/sirolimus cohorts: transplant and outcomes summary**

<table>
<thead>
<tr>
<th>Animal ID</th>
<th>Day of neutrophil engraftment</th>
<th>Transplant-associated events</th>
<th>Day of event</th>
<th>WBC count (ANC count) at terminal analysis</th>
<th>WB chimerism at terminal analysis (%)</th>
<th>Day of death/euthanasia</th>
<th>Reason for euthanasia</th>
<th>Inclusion in transcriptomic analysis</th>
</tr>
</thead>
<tbody>
<tr>
<td>R.22</td>
<td>9</td>
<td>None</td>
<td>NA</td>
<td>9.3(5.8) × 10^6/ml</td>
<td>99</td>
<td>21</td>
<td>Skin GVHD</td>
<td>Yes</td>
</tr>
<tr>
<td>R.23</td>
<td>19</td>
<td>None</td>
<td>NA</td>
<td>1.5(1.2) × 10^6/ml</td>
<td>100</td>
<td>27</td>
<td>Liver GVHD</td>
<td>Yes</td>
</tr>
<tr>
<td>R.24</td>
<td>12</td>
<td>None</td>
<td>NA</td>
<td>2.3(1.3) × 10^6/ml</td>
<td>100</td>
<td>16</td>
<td>Skin GVHD</td>
<td>Yes</td>
</tr>
<tr>
<td>R.26</td>
<td>14</td>
<td>GI Infection with E. coli</td>
<td>30</td>
<td>2.9(1.4) × 10^6/ml</td>
<td>100</td>
<td>33</td>
<td>Timed terminal analysis</td>
<td>Yes</td>
</tr>
<tr>
<td>R.27</td>
<td>12</td>
<td>Giardiasis</td>
<td>10</td>
<td>4.5(3.4) × 10^6/ml</td>
<td>100</td>
<td>34</td>
<td>Timed terminal analysis</td>
<td>Yes</td>
</tr>
<tr>
<td>R.28</td>
<td>15</td>
<td>None</td>
<td>NA</td>
<td>3.7(0.4) × 10^6/ml</td>
<td>99</td>
<td>34</td>
<td>Timed terminal analysis</td>
<td>Yes</td>
</tr>
<tr>
<td>R.29</td>
<td>22</td>
<td>Wound infection</td>
<td>10</td>
<td>1.3(1.0) × 10^6/ml</td>
<td>88</td>
<td>22</td>
<td>Wound infection</td>
<td>Yes</td>
</tr>
<tr>
<td>R.213</td>
<td>15</td>
<td>Giardiasis/unexpected death</td>
<td>44/66</td>
<td>1.98(1.15) × 10^6/ml</td>
<td>58</td>
<td>66</td>
<td>Unexpected death</td>
<td>Yes</td>
</tr>
<tr>
<td>R.222</td>
<td>11</td>
<td>Hemorrhagic stroke</td>
<td>21</td>
<td>9.7(6.7) × 10^6/ml</td>
<td>100</td>
<td>21</td>
<td>Hemorrhagic stroke</td>
<td>No</td>
</tr>
<tr>
<td>R.249</td>
<td>18</td>
<td>None</td>
<td>NA</td>
<td>6.4(3.3) × 10^6/ml</td>
<td>53</td>
<td>105</td>
<td>Timed terminal analysis</td>
<td>Yes</td>
</tr>
<tr>
<td>R.250</td>
<td>20</td>
<td>Fatal sepsis; loss of myeloid chimerism</td>
<td>6S</td>
<td>18.0(17.3) × 10^6/ml</td>
<td>15</td>
<td>72</td>
<td>Sepsis</td>
<td>No</td>
</tr>
<tr>
<td>R.251</td>
<td>10</td>
<td>Secondary graft failure</td>
<td>61</td>
<td>0.8(0.2) × 10^6/ml</td>
<td>78</td>
<td>100</td>
<td>Timed terminal analysis</td>
<td>No</td>
</tr>
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</table>
Flow cytometric and transcriptomic evidence for synergistic control of T cell effector maturation with FR104/sirolimus. The prevention of clinical aGVHD during ongoing FR104 exposure in the FR104/sirolimus cohort correlated closely with control of T cell proliferation, cytotoxicity, and effector function. Thus, as we have previously described (12, 53), and in agreement with the transcriptomic data discussed above, the No Rx, sirolimus, and CTLA4-Ig cohorts developed increasing expression of Ki-67 (Ki67\textsuperscript{lo}) that denotes proliferation and overexpression of granzyme B (GZMB\textsuperscript{lo}) after transplant (Figure 4A). In contrast, FR104 monophosphorylation partially controlled Ki67 expression (but not GZMB overexpression; Figure 4A), while combination prophylaxis with FR104/sirolimus resulted in synergistic control of both proliferation and granzyme B overexpression in the blood and tissues (Figure 4, A and B). This control of proliferation resulted in restraint of effector maturation and improved maintenance of the naive T cell phenotype in the FR104/sirolimus cohort, most prominently in CD4\textsuperscript{+} T cells (Figure 4C), which persisted in the FR104/sirolimus cohort for the length of FR104 exposure. However, when FR104 was discontinued, and concomitant with desaturation of CD28 occupancy on donor T cells (Figure 1, C and E), loss of naive CD4\textsuperscript{+} T cells did occur (Figure 4, C and D). This loss of naive cells occurred earlier for CD8\textsuperscript{+} T cells and did not correlate with FR104 exposure (Figure 4, C and D).

The impact of FR104/sirolimus on T cell proliferation and effector maturation was also observed transcriptomically. Thus, GSEA performed on T cells purified on day 14 after transplant (or at the time of necropsy if this occurred prior to day 14) confirmed underrepresentation of gene sets involved in the control of cell proliferation and in the control of antigen response and overrepresentation of gene sets associated with the naive T cell phenotype in the FR104/sirolimus cohort compared with both FR104 and sirolimus cohorts (Figure 4E). The complete list of gene sets over- and underrepresented in the day 14 FR104/sirolimus cohort is shown in Supplemental Table 2.

Previous reports have suggested that the control of T cell signaling through CD28 blockade may be mediated by coinhibitory 2B4 signaling (44). However, and consistent with the lack of effector maturation in the FR104-prophylaxed cohorts (55) (Figure 4, A-E), we did not observe increased expression of individual coinhibitory receptors (including PD-1, LAG3, 2B4, CTLA4 and TIM-3; Figure 4F) in the FR104/sirolimus cohort and, indeed, observed decreased expression of these molecules in the FR104/sirolimus versus No Rx cohorts.

Transcriptional consequences of FR104/sirolimus versus CTLA4-Ig/sirolimus for GVHD prophylaxis. T cell transcriptome analysis allowed us to further interrogate the mechanisms underlying the control of GVHD with FR104/sirolimus and the relative impact of FR104/sirolimus on T cell signaling compared with CTLA4-Ig/sirolimus. We first probed our data set for pathways that defined the synergistic impact of combining CD28 blockade with mTOR inhibition. To do this, we quantified the number of genes DE in the FR104, sirolimus, and FR104/sirolimus transcriptomes compared with those expressed in the No Rx transcriptome (Figure 5A). Gene arrays for this analysis were performed on T cells purified from recipients at day 14 after transplant (or at the time of necropsy, if this occurred prior to day 14). To avoid potential confounding associated with the secondary graft failure/loss of donor chimerism that occurred in 2 animals in the FR104/sirolimus cohort (recipients R.250 and R.251; Table 1 and discussed in detail below), we excluded the transcriptomes from these recipients from this analysis. As demonstrated in Figure 5A, T cells purified from the FR104/sirolimus cohort demonstrated a distinct gene expression landscape in comparison with the No Rx, sirolimus, and FR104 cohorts. Thus, as shown in this figure and Supplemental Tables 1 and 4, using an absolute fold change cutoff of greater than 1.4, there were no DE genes in the FR104 monophosphorylation versus No Rx comparison and only 32 DE genes in the sirolimus monophosphorylation versus No Rx comparison. In striking contradiction, there were 2,162 DE genes in the FR104/sirolimus versus No Rx comparison. Importantly, of these 2,162 transcripts, 2,139 (99%) were uniquely affected by combination prophylaxis, suggesting deep synergy between the 2 agents. Next, we performed a similar DE gene analysis comparing the CTLA4-Ig/sirolimus, sirolimus, and CTLA4-Ig transcriptomes to No Rx (as before, gene arrays were performed on day 14 or at the time of necropsy if it occurred earlier than day 14). As shown in Figure 5A, again using an absolute fold change cutoff of greater than 1.4, there were no
Figure 5. FR104/sirolimus synergistically modulates T cell effector and proliferative transcriptional signals dysregulated during aGVHD. (A) Top panel: number of genes DE in the FR104 (n = 3; at day 14), sirolimus (n = 4; at terminal analysis), and FR104/sirolimus (n = 6; at day 14) cohorts compared with the No Rx cohort at terminal analysis. Each bar represents the number of DE genes that are unique for each comparison. Bottom panel: number of genes DE in the CTLA4-Ig (n = 3; at day 14), sirolimus (n = 4; at terminal analysis), and CTLA4-Ig/sirolimus (n = 6; at day 14) cohorts compared with the No Rx cohort at terminal analysis. Each bar represents the number of DE genes that are unique for each comparison. (B) Venn diagram showing the number of genes uniquely DE in either the FR104/sirolimus or the CTLA4-Ig/sirolimus cohorts compared with the No Rx cohort and the degree of overlap between these 2 DE gene lists. Red text depicts the number of overrepresented transcripts; blue text depicts the number of underrepresented transcripts. (C) Functional characterization of pathways enriched in both the overrepresented and underrepresented genes, defined as shown in B: underrepresented genes unique for FR104/sirolimus vs. No Rx comparison (left); overrepresented genes unique for FR104/sirolimus vs. No Rx comparison (middle); and underrepresented genes shared between FR104/sirolimus vs. NoRx and CTLA4-Ig/sirolimus vs. NoRx comparisons (right). A complete list of the individual pathways identified is found in Supplemental Table 5. Pathway identification used a Benjamini-Hochberg-corrected P value of less than 0.05. (D) Representative GSEA underrepresented plots of cell cycle– and immune response–related gene sets and an overrepresented naive T cell–related gene set in the FR104/sirolimus (n = 6) transcriptome in comparison with the CTLA4-Ig/sirolimus transcriptome (n = 6; transcriptomes derived from T cells isolated at day 28 from both cohorts) with FDR of q < 0.05.
DE genes in the CTLA4-Ig monophylaxis versus No Rx comparison and the same 32 DE genes in the sirolimus monophylaxis versus No Rx comparison. There were far fewer DE genes in the CTLA4-Ig/sirolimus versus No Rx comparison compared with the FR104/sirolimus versus No Rx comparison: 51 DE genes in the CTLA4-Ig/sirolimus versus No Rx comparison, with 45 genes (88%) being uniquely affected by combination CTLA4-Ig/sirolimus prophylaxis versus No Rx (Figure 5A).

To develop a deeper understanding of the pathways uniquely modified by sirolimus combined with either CD28 or CD80/86 blockade, we determined the unique transcripts that were DE in the FR104/sirolimus versus No Rx comparison compared with the CTLA4-Ig/sirolimus versus No Rx comparison. We identified 2,098 genes that were uniquely DE in the FR104/sirolimus cohort (but not in the CTLA4-Ig/sirolimus cohort) when compared with No Rx (659 transcripts were overrepresented and 1,439 were underrepresented; Figure 5B). There were many fewer transcripts uniquely DE in the CTLA4-Ig/sirolimus compared with No Rx (4 total transcripts, with 2 overrepresented and 2 underrepresented in the CTLA4-Ig/sirolimus versus No Rx comparison; Figure 5B, Supplemental Table 4), with no pathways identified among these 4 transcripts. While the vast majority of DE transcripts compared with No Rx were unique between the FR104/sirolimus and CTLA4-Ig/sirolimus comparisons, we did identify 41 transcripts (18 overrepresented and 23 underrepresented) that were DE in both FR104/sirolimus and CTLA4-Ig/sirolimus cohorts (Figure 5B, discussed below).

Pathway analysis using DAVID (56) allowed functional annotation of the cellular processes represented by these DE genes, which are depicted in Figure 5C and enumerated in Supplemental Table 5. With respect to pathways uniquely perturbed in the FR104/sirolimus cohort, we identified 5 overrepresented pathways and 55 underrepresented pathways. Of note, of the 5 overrepresented pathways uniquely identified in the FR104/sirolimus versus No Rx comparison, 2 (TGF-β signaling pathway and signaling by BMP) belonged to TGF-β/Smad-related signaling with an additional pathway (signaling pathways regulating pluripotency of stem cells) that also contained many mediators of TGF-β signaling (Figure 5C and Supplemental Table 5). The other overrepresented pathways involved cytokine signaling (cytokine-cytokine receptor interaction and FoxO-signaling pathway). In contrast with the highly focused upregulation of TGF-β signaling, we identified many more pathways uniquely downregulated in the FR104/sirolimus versus No Rx comparison. As shown in Figure 5C, these pathways belonged to the following major functional categories: (a) cell cycle (36.4% of pathways); (b) DNA repair (23.6%); (c) gene expression (transcription) (9.1%); (d) replication and repair (5.5%); (e) signal transduction (signaling by Rho GTPase; 5.5%); (f) immune system (3.6%); and (g) cell growth, senescence, and death (5.4%).

This analysis also permitted us to determine which pathways were affected by both FR104/sirolimus and CTLA4-Ig/sirolimus prophylaxis regimens compared with No Rx in or der to determine the mechanisms potentially shared by these 2 GVHD-prevention strategies (Figure 5, B and C). This analysis revealed no upregulated and 6 downregulated pathways that were shared by both FR104/sirolimus and CTLA4-Ig/sirolimus versus No Rx. The shared downregulated pathways were predominated by those controlling the cell cycle (4 of 6 pathways; Figure 5C and Supplemental Table 5), underscoring the shared impact of both regimens on T cell proliferation and expansion.

While both CTLA4-Ig/sirolimus and FR104/sirolimus downregulated cell-cycle-related transcripts compared with No Rx (Figure 5, A–C), GSEA comparing these 2 cohorts identified better control of GVHD documented with FR104/sirolimus compared with CTLA4-Ig/sirolimus (Figure 5D and Supplemental Table 2), with concomitant enrichment in naive T cells in the FR104/sirolimus versus CTLA4-Ig/sirolimus comparison (Figure 5D and Supplemental Table 2). These transcriptional results provide important mechanistic associations with the improved control of GVHD documented with FR104/sirolimus versus CTLA4-Ig/sirolimus (Figure 2).

To most rigorously define the transcriptional hallmarks of FR104/sirolimus versus CTLA4-Ig/sirolimus, we applied unsupervised weighted gene correlation network analysis (WGCNA) (57, 58) to construct a gene coexpression network from our data set, using the transcriptomes from the FR104/sirolimus (n = 6)
and CTLA4-Ig/sirolimus \((n = 6)\) cohorts (microarray data collected at day 28) in comparison with the No Rx cohort (microarray data collected at terminal analysis, which occurred prior to day 28 for all No Rx animals; \(n = 11\)) and with HCs \((n = 62)\). After applying WGCNA, we then performed hierarchical clustering of the matrix, thereby identifying sets of genes that met a predetermined threshold of coexpression, which resulted in the identification of 9 discrete self-assembling modules (Figure 6, A and B, and Supplemental Table 6). The module eigengenes \((59)\) were then clustered with the 4 experimental cohorts (No Rx, HC, CTLA4-Ig/sirolimus, and FR104/sirolimus) in order to identify metamodules (Figure 6B). This analysis determined that the No Rx and CTLA4-Ig/sirolimus cohorts were strongly clustered in a metamodule with the blue eigengene (Figure 6B) and that this blue eigengene demonstrated no correlation with either the FR104/sirolimus or the HC cohorts (Figure 6B), suggesting that it could reveal genes and pathways.

**Figure 7. Visualization of the blue gene module.** (A) Visualization of gene coexpression network connections between the most connected genes in the blue module using Cytoscape. Shown are nodes with network connections whose topological overlap is above a threshold of 0.1. Edges with network connections above the threshold of 0.25 are shown. Mean expression fold change values of the FR104/sirolimus cohort versus HC for each gene are visualized using a false-color scale. Pathways were then identified using the DAVID database, using a cutoff derived from the Benjamini-Hochberg statistic, \(P < 0.05\). In addition, 7 representative submodules containing genes from the top statistically ranking pathways (enumerated with the Reactome Database) are shown. (B) Functional distribution of all pathways enriched in the blue module. A complete list of the individual pathways identified is found in Supplemental Table 7. Pathway identification used a Benjamini-Hochberg–corrected \(P\) value of < 0.05.
that linked the breakthrough aGVHD that occurred with CTLA4-Ig/sirolimus with the severe GVHD observed in the No Rx cohort. In contrast, the brown eigengene clustered with the FR104/sirolimus cohort and with HCs, but not with the No Rx and CTLA4-Ig/sirolimus cohorts (Figure 6B), suggesting that it may reveal genes associated with the T cell quiescence induced by FR104/sirolimus. The transcripts and gene correlations identified in the blue and brown eigengenes were visualized using Cytoscape (60) (Figure 7A and Figure 8), and the pathways encapsulated by the blue module are enumerated in Supplemental Table 7. The resulting pathway analysis using DAVID (interrogating the Reactome, KEGG, and Biocarta databases) revealed that the blue module was enriched in multiple pathways of T cell activation and proliferation (Figure 7, A and B). These prominently included pathways associated with cell cycle and DNA repair, but also included pathways associated with the regulation of gene transcription, cellular senescence, sig-
The relative number (percentage of total CD4 T cells; top panel), the absolute number (middle panel), and the Treg/100 Tconv ratio, normalized to the corresponding pretransplant values (bottom panel) were tracked longitudinally by flow cytometry in the No Rx (n = 7), sirolimus (n = 6), FR104 (n = 3), FR104/sirolimus (n = 9), and CTLA4-ig/sirolimus (n = 7) cohorts. Tregs were defined as CD3+CD4-CD25-CD127hiFoxP3+; Tconv cells were defined as CD3+CD8+ and CD3+CD4+CD25 CD127lo by flow cytometric analysis. Data are shown as mean ± SEM. The solid red threshold line represents the Treg/100 Tconv ratio in the No Rx cohort at terminal analysis (62), with dotted lines above and below the threshold line representing the SEM interval. (B) The relative number (percentage of total CD4 T cells) of Tregs in the peripheral (axillary and inguinal) LNs and spleen in HC animals and recipients from KY1005/sirolimus undergoing terminal analysis before or after day 66. (C) The normalized Treg/100 Tconv ratio (left panel) and the percentage of CD28+CD4+ T cells (right panel) before (white circles) and after (black circles) discontinuation of FR104 in FR104/sirolimus cohort recipients who survived more than 66 days after transplant (R.249, R.250, and R.251). Each line represents a single experiment.

Figure 9. The impact of FR104/sirolimus on Treg homeostasis after transplant. (A) The relative number (percentage of total CD4 T cells; top panel), and the absolute number (middle panel), and the Treg/100 Tconv ratio, normalized to the corresponding pretransplant values (bottom panel) were tracked longitudinally by flow cytometry in the No Rx (n = 7), sirolimus (n = 6), FR104 (n = 3), FR104/sirolimus (n = 9), and CTLA4-ig/sirolimus (n = 7) cohorts. Tregs were defined as CD3+CD4-CD25-CD127hiFoxP3+; Tconv cells were defined as CD3+CD8+ and CD3+CD4+CD25 CD127lo by flow cytometric analysis. Data are shown as mean ± SEM. The solid red threshold line represents the Treg/100 Tconv ratio in the No Rx cohort at terminal analysis (62), with dotted lines above and below the threshold line representing the SEM interval. (B) The relative number (percentage of total CD4 T cells) of Tregs in the peripheral (axillary and inguinal) LNs and spleen in HC animals and recipients from KY1005/sirolimus undergoing terminal analysis before or after day 66. (C) The normalized Treg/100 Tconv ratio (left panel) and the percentage of CD28+CD4+ T cells (right panel) before (white circles) and after (black circles) discontinuation of FR104 in FR104/sirolimus cohort recipients who survived more than 66 days after transplant (R.249, R.250, and R.251). Each line represents a single experiment.

CD28 blockade negatively affects Treg homeostasis after HCT. While the transcriptional analysis described above provided noticeable evidence for the impact of FR104/sirolimus on T cell proliferation and activation, it is notable that we did not identify pathways associated with Tregs in this analysis. This may be due to the fact that CD4+FoxP3+ Tregs express CD28 (confirmed in this study; see Supplemental Figure 2A) and that they rely on CD28 signaling for their survival and suppressive function (61). We have previously shown that the development of aGVHD in NHP is associated with a significant decrease in the Treg/Tconv ratio (Figure 9A, Supplemental Figure 2B, and ref. 62), and in the current study, we observed that, consistent with the development of GVHD in the FR104 monoprophylaxis cohort, these recipients were unable to maintain either their pretransplant numbers of Tregs or the pretransplant Treg/Tconv ratio. Furthermore, the decay in these parameters coincided with the development of clinically significant GVHD (Figure 9A). Importantly, while combination prophylaxis with FR104/sirolimus initially preserved absolute Treg numbers as well as the Treg/Tconv ratio, this effect was not durable for the length of the experiment, with peripheral blood Treg counts dropping by approximately day 25 after transplant (Figure 9A) and with a decrease in the proportion of Tregs compared with the pretransplant level also observed in the LNs and spleens of FR104/sirolimus recipients (Figure 9B). The inhibitory effect on Tregs may have been due to FR104. Thus, in 2 of 3 recipients in the FR104/sirolimus cohort that survived for more than 66 days after transplant, we observed a rebound of the Treg/Tconv ratio after discontinuation of FR104, concomitant with the desaturation of T cell CD28 expression (Figure 9C), as well as a modest increase in the absolute number of Tregs after discontinuation of FR104 (Figure 9A). Importantly, we did not observe any negative impact of FR104 on the in vitro suppressive capacity of Tregs, as measured by the sustained ability of Tregs that were pretreated with FR104 to suppress Tconv alloproliferation in a mixed lymphocyte reaction (MLR) assay (Supplemental Figure 2, C and D).

Transplant-related mortality in the FR104/sirolimus cohort. As noted above, while combination prophylaxis with FR104/sirolimus was successful in preventing clinical GVHD, the regimen was not uniformly successful in supporting posttransplant survival,
In order to interrogate the possible causes of the increased risk of sepsis and graft loss in the FR104/sirolimus cohort, 2 experiments were performed. In the first, we determined whether other immune cells also expressed CD28 in NHP and thus could have been affected by FR104 blockade. While each of 4 distinct anti-CD28 clones labeled T cells, none of the antibodies bound to the surface of B cells, NK cells, dendritic cells, monocytes, or granulocytes (Supplemental Figure 3A). Intracellular labeling techniques demonstrated similar results, with only 1 of 4 clones (15E8) labeling non–T cell leukocytes at low levels after permeabilization. This labeling was likely nonspecific to CD28, however, given that it was only minimally affected by preincubation with FR104 (Supplemental Figure 3, A–C). In addition to the expression analyses described above, functional assays were also performed, which demonstrated that short-term preincubation with FR104 did not affect phagocytosis or the respiratory burst of granulocytes (Supplemental Figure 3, D and E). Together, these experiments suggest that the impact of FR104 on protective immunity was unlikely to be due to the expression of CD28 on non–T cells.

The second experiment was designed to determine other possible mechanisms of the increased risk of sepsis in transplant recipients. We observed the following posttransplant complications in both the short-term and long-term FR104/sirolimus cohorts (Table 1): (a) 1 episode of hemorrhagic stroke associated with thrombocytopenia (R.222) and (b) 2 episodes of infection-related death (R.29 and R.250). R.29 developed an injection-site infection that led to necropsy on day 22, and R.250 died of culture-negative sepsis on day 71 in the setting of a high WBC count (18.0 × 10⁶/ml), absolute neutrophil count (ANC) (17.3 × 10⁶/ml), C-reactive protein (CRP) count (44.5 mg/l). In addition to these deaths, 1 additional animal (R.213) died unexpectedly during sedation. At the time of death, R.213 had a confirmed GI infection with *Giardia*. In addition to the infection-related morbidity and mortality, 2 animals in the 9-animal FR104/sirolimus cohort had signs of loss of engraftment (Figure 10, A and B). Thus, R.250 demonstrated a loss of myeloid chimerism (Figure 10A) and R.251 demonstrated secondary graft failure (Figure 10, A and B). This resulted in a dichotomous impact on survival after transplant in the FR104/sirolimus cohort. While GVHD-free survival was improved compared with the CTLA4-Ig/sirolimus cohort (Figure 2B) and R.251 demonstrated secondary graft failure (Figure 10, A and B), this resulted in a dichotomous impact on survival after transplant in the FR104/sirolimus cohort. While GVHD-free survival was improved compared with that in the CTLA4-Ig/sirolimus cohort (Figure 2B), combination FR104/sirolimus prophylaxis did not improve overall survival compared with CTLA4-Ig/sirolimus prophylaxis (Figure 10C).

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The second experiment was designed to determine other possible mechanisms of the increased risk of sepsis in transplant
recipients exposed to FR104/sirolimus by determining the ability of these animals to mount an appropriate cytokine response during infection (Figure 10D and Supplemental Figure 4A). While cytokines classically associated with innate immune responses, including IL-1RA, IL-6, and IL-12 (63–65), were appropriately elevated in the serum during infection in the FR104/sirolimus cohort, it was notable that an infection-associated response by IFN-γ, which represents the major T cell cytokine produced during sepsis (66–68) (and which was elevated in a transplant recipient from a separate costimulatory blockade protocol who also developed sepsis after allogeneic HCT; Supplemental Figure 4B), did not occur. This result is consistent with the transcriptomic analysis shown in Supplemental Figure 4C, which documented substantial control of IFN-γ production/lack of IFN-γ signaling in the FR104/sirolimus cohort compared with the CTLA4-Ig/sirolimus cohort and suggests that inhibition of an appropriate IFN-γ response may have increased the risk of infection in the FR104/sirolimus cohort.

Discussion
Here, we provide evidence for broad synergy between CD28 blockade and mTOR inhibition in preventing GVHD and, using a systems biology approach, have determined the molecular mechanisms associated with disease control. Importantly, this study has also allowed us to measure the comparative impact of targeting CD28 versus CD80/86 in controlling T cell activation and preventing GVHD, both in single-agent experiments and in combination with mTOR inhibition.

One of the most striking findings in the current study was the depth of synergy between mTOR inhibition and CD28 blockade in controlling aGVHD. The impact was impressive, with 2,098 genes identified that were exclusively normalized in the FR104/sirolimus cohort compared with the No Rx cohort. The other critical result of this study was the evidence for more complete control of T cell activation, proliferation, and effector maturation with CD28 blockade versus CD80/86 blockade. Two other key observations made in this study deserve special discussion, each of which has important clinical relevance. The first is the impact that CD28 blockade had on Treg reconstitution and homeostasis in the NHP GVHD model. Given the contribution that CTLA4 signaling makes to Treg function, it has long been thought that targeting CD80/86 (and thus potentially blocking signaling from both CD28 and CTLA4) would not be ideal in terms of Treg homeostasis. However, with regard to the impact that CD28 (and its blockade) has on Treg development, function, and homeostasis, the literature supports complex mechanistic relationships, with evidence for both salutary and inhibitory effects of CD28 deficiency on Tregs (61, 69–73). In the experiments presented here, we provide evidence for the inhibition of Treg homeostasis in the FR104/sirolimus cohort. These results strengthen the inference that control of allostreactivity in the FR104/sirolimus cohort occurred predominantly through the inhibition of conventional T cell activation rather than the promotion of Treg predominance.

The second critical observation concerns possible drawbacks associated with the substantial control of conventional T cell activation and functional maturation that occurred in the FR104/sirolimus cohort. Thus, we observed a number of transplant-related toxicities in this cohort that resulted in a lack of a beneficial impact on overall survival when compared with CTLA4-Ig/sirolimus, despite improved GVHD-free survival. While the number of recipients analyzed was relatively small (n = 9 total, with 5 followed long term), we did observe fatal sepsis in 2 animals, loss of donor chimerism in 1 animal, and secondary graft loss in 1 animal. These complications occurred in the setting of intact reconstitution of CD4 cells with autologous transplant controls, but slower reconstitution of CD8 cells (Supplemental Figure 1). Death from these clinical complications was not observed in any of the 7 animals that we have previously reported in the CTLA4-Ig/sirolimus cohort (51). While sepsis and loss of chimerism are expected complications associated with transplant, it is also possible that they were directly affected by the impact of combined FR104/sirolimus on T cell activation. Thus, it is well established that T cells function within an allograft to promote donor engraftment and protection against infection and that both failure to engraft and infectious complications accompany T cell depletion (74–77). The significant control of T cell activation with FR104/sirolimus may blunt these T cell functions, and indeed, previous results in murine models suggested that deficiency of CD28 increased the risk of nonengraftment after allo-HCT (39) and that intact CD28 signaling in T cells was required to mount an effective protective immune response against bacterial infections (78–80). Moreover, and further supporting the hypothesis that FR104/sirolimus-mediated control of T cell activation may have increased the risk of infection and sepsis in this cohort, we found that FR104/sirolimus recipients were unable to mount an appropriate (66–68) IFN-γ response to sepsis (Figure 10D). While the ability to discern a positive impact on overall survival may have been limited by a lack of statistical power with the current sample size, given the combined risk of infection and engraftment failure that we encountered in the FR104/sirolimus transplants, it was not feasible to add enough additional animals to provide a more definitive evaluation of this end point. It is important to note, however, that the NHP model may have accentuated the risks of both infection-related death and graft rejection compared with what would be seen in the clinic. Thus, the ability to provide successful prophylaxis and treatment of infection is more challenging in NHP than in patients, which likely contributes to the increased risk of life-threatening sepsis in this animal model. Furthermore, the NHP transplant model employed used only total body irradiation (TBI) for pretransplant conditioning; thus, the risk of rejection could be mitigated in patients by augmenting the pretransplant regimen with additional conditioning agents. Given the prominent control of T cell activation and GVHD that we observed, and if clinical risks were mitigated as described above, translation of this regimen would be appropriate and potentially of considerable value to transplant patients at high risk of GVHD.

The striking control of T cell activation when FR104 was combined with sirolimus suggests that this combinatorial strategy may be an important approach for clinical translation (81). However, the complications that were also observed in the FR104/sirolimus cohort suggest that effectively deploying CD28 blockade in the clinic may be complex. Our results suggest that there may be considerable trade-offs associated with the comprehensive down modulation of conventional T cell activation that occurred with FR104/sirolimus, which may impede functional engraftment if
used in the early peritransplant period. Nonetheless, the potency of the control of T cell activation with FR104/sirolimus suggests that it may be of use in settings characterized by refractory T cell pathophysiology, for example, in treating refractory acute or chronic GVHD. Although the current NHP model does not speak directly to either of these clinical entities, the transcriptomic evidence for potent downmodulation of effector T cell activation when CD28 blockade was combined with mTOR inhibition suggests that this combination may have efficacy in these hard-to-treat scenarios. Given the unique balance of risks and benefits that may be inherent to anti-CD28–based regimens, careful clinical investigation, with robust immunologic analysis linked to clinical outcomes, is warranted.

Methods

Experimental animals and transplant protocol. Transplants were performed as previously described (12, 53), with details of conditioning regimens, immunomodulation regimens, and stem cell infusions described in Figure 1A, Table 1, and Supplemental Methods.

Flow cytometric and immunofluorescence analysis. For phenotypic analysis and T cell sorting, multicolor flow cytometry was performed on all transplant recipients, as previously described (12). A complete description of flow cytometry methods is found in Supplemental Methods.

Microarray and data analysis. CD3 CD20+ lymphocytes were purified flow cytometrically using a FACS Jazz Cell Sorter (BD Biosciences). Following T cell purification, RNA was stabilized in T cell lysates with RLT buffer (QIAGEN) and RNA was purified using the RNeasy Column Kit (QIAGEN). RNA was quantified using a Nanodrop Spectrophotometer (Thermo Fisher Scientific) and purity confirmed with an RNA 6000 Nano Kit (Agilent). The purified RNA was sent to either the Vanderbilt Technologies for Advanced Genomics Core (Nashville, Tennessee, USA) or to the Oregon Health Sciences University Gene Profiling Shared Resource (Portland, Oregon, USA), where RNA quantity and quality were verified. This was followed by cDNA/cRNA synthesis and target hybridization to the GeneChip Rhesus Macaque Genome Array (Affymetrix). The resultant fluorescent signals were processed and normalized using the robust multichip averaging (RMA) method (82). The microarray was performed in 7 batches, with batches containing samples from both HCs and transplanted recipients. Details of microarray statistical analysis and full details of the computational analysis of the transcriptome are found in the Supplemental Methods.

The NHP microarray data discussed in this publication were deposited with the Gene Expression Omnibus (GEO) (GSE116070).

Statistics. Survival statistics were calculated using Kaplan-Meier analysis. Other variables were expressed as mean ± SEM, and P values less than 0.05 were considered statistically significant. Statistical analysis was performed using Prism version 6 for Mac OS X (GraphPad Software). Details of the statistical analysis used for the transcriptome studies are described in Supplemental Methods.

Study approval. This study was conducted in strict accordance with USDA regulations and the recommendations in the Guide for the Care and Use of Laboratory Animals (National Academies Press, 2011). It was approved by the Animal Care and Use Committees (IACUC) at Emory University and at the University of Washington.

Author contributions

BKW designed and performed experiments and wrote the paper. VT designed and performed experiments and wrote the paper. SNF analyzed data. DJH performed experiments. KB performed experiments. AY performed experiments. MB performed experiments. NP performed experiments and analyzed data. HBZ performed experiments. AT performed experiments. LC performed experiments. CM performed experiments and analyzed data. GB analyzed data and wrote the paper. JPS performed experiments and analyzed data. A Garcia performed experiments. TD performed experiments. BRB analyzed data and wrote the paper. BV analyzed data and wrote the paper. LSK conceived the study, analyzed data, and wrote the paper.

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