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CD4⁺ CD25⁺ Foxp3⁺ T Regulatory Cells with Limited TCR Diversity in Control of Autoimmunity

Dennis Adeegbe, Takaji Matsutani, Jing Yang, Norman H. Altman, and Thomas R. Malek

The importance of high TCR diversity of T regulatory (Treg) cells for self-tolerance is poorly understood. To address this issue, TCR diversity was measured for Treg cells after transfer into IL-2Rβ⁻/⁻ mice, which develop lethal autoimmunity because of failed production of Treg cells. In this study, we show that high TCR diversity of pretransferred Treg cells led to selection of therapeutic Treg cells with lower TCR diversity that prevented autoimmunity. Pretransferred Treg cells with lower diversity led to selection of Treg cells through substantial peripheral reshaping with even more restricted TCR diversity that also suppressed autoimmune symptoms. Thus, in a setting of severe breakdown of immune tolerance because of failed production of Treg cells, control of autoimmunity is achieved by only a fraction of the Treg TCR repertoire, but the risk for disease increased. These data support a model in which high Treg TCR diversity is a mechanism to ensure establishing and maintaining self-tolerance. The Journal of Immunology, 2010, 184: 56–66.

The factors influencing the thymic and peripheral selection of the T regulatory (Treg) cell TCR repertoire are of much interest as they likely represent a major control point in establishing and maintaining self-tolerance. Considerable data indicate that development of Treg cells requires recognition of self-peptide/MHC at an affinity threshold that usually falls between that required for positive versus negative selection (1–6). Most information concerning the Treg cell TCR repertoire has come from analysis of the entire pool of Treg cells in the thymus and periphery from autoimmune-free mice and humans (4, 6–11). In the mouse studies, TCRβ transgenic (Tg) mice were used so that one dominant rearranged TCR β-chain was expressed by all T cells. Analysis was then focused on the diversity of endogenous TCRs, which defined the TCR specificity. These experiments indicate that TCRs of Treg cells are highly diverse with a repertoire as important question is whether this high TCR diversity is mandatory for peripheral tissue-specific self-Ags in shaping the TCR repertoire. Thus, the TCR specificity on developing thymocytes and subsequent selection events do not solely specify the decision to be Treg versus T conventional cells. In addition, the TCR repertoire of thymic and peripheral Treg cells more closely resembled each other than the TCR repertoire of conventional T cells. Correspondingly, most peripheral Treg cells originate from their development and commitment to the Treg lineage within the thymus rather than from conversion of conventional peripheral T cells into induced Treg cells.

The TCR repertoire expressed by thymic Treg cells is reshaped as reflected by a reduction in the frequency of dominant thymic Treg cell TCR specificities in peripheral Treg cells (4, 6, 8–10). The mechanisms responsible for flattening the peripheral Treg cell TCR repertoire are poorly understood, but may reflect homeostatic regulation and influences of self-Ags and environmental Ags on peripheral Treg cells (10, 12). Furthermore, the TCR repertoire of peripheral Treg cells varied considerably when examined based on anatomical location of draining lymph nodes (LNs) (13, 14), consistent with a role for peripheral tissue-specific self-Ags in shaping the TCR repertoire.

Given the high diversity of the Treg cell TCR repertoire, an important question is whether this high TCR diversity is mandatory to suppress potential autoreactive T cells in the periphery that escape thymic negative selection. The current study was designed to address this issue for a population of polyclonal autoreactive T cells by using IL-2Rβ⁻/⁻ mice as a model. Because of the failed production of an effective population of Treg cells, IL-2Rβ⁻/⁻ mice develop rapid lethal systemic autoimmunity that resembles the disease associated with Foxp3⁻/⁻ cells. Cell transfer studies indicate that this autoimmune syndrome is primarily due to autoreactive CD4⁺ T cells (15). The adoptive transfer of either syngeneic or even fully allogeneic wild type (WT) Treg cells into neonatal IL-2Rβ⁻/⁻ mice fully prevents this autoimmune disease such that the recipient mice live a normal life span (16–18). Importantly, these donor Treg cells, including the allogeneic Treg cells, stably engraft, expand, and persist lifelong through extensive homeostatic proliferation to comprise essentially the entire pool of CD4⁺CD25⁺ Foxp3⁺ Treg cells within these autoimmune-free IL-2Rβ⁻/⁻ mice. In this study, we evaluated TCR repertoire diversity of such donor Treg cells when they were obtained from WT mice or mice that expressed a single TCR β-chain.

Materials and Methods

Mice

C57BL/6, BALB/c, C57BL6 TCRα⁻/⁻, C57L/J-Tg(Tcrrb)93Vbo/J TCRβ Tg mice (designated TCRβ Tg) (19) were obtained from The Jackson Laboratory (Bar Harbor, ME). The TCRβ Tg mice were backcrossed to...
and vitrogen). The full-length Vα pMI sequencing primer. Clones were randomly chosen for sequencing. 3730 amplification conditions described previously. For Vb1 min with 5 min for the last extension. The sense primers for V were typically primers specific for V1 h. PCR was performed using one-twentieth of cDNA, 250 nM sense DTT (Invitrogen) and 20U RNasin (Promega, Madison, WI) at 50˚C for triphosphates (10 nmol each). After cooling, cDNA was synthesized with FAM or 5′ V8.2 in VTR cells were adoptively transferred by i.v. injection into the superficial ony using autoimmune-free breeding pairs as previously described (18). CD25 FITC-anti-TCRV manufacturer’s protocol. FACS analysis was performed as previously de-
pression assay, which was performed as previously described (18). In some experiment, the donor Treg cells FACS Ab (FJK-16s; eBioscience, San Diego, CA) according to the manufacturer’s protocol. FACS analysis was performed as previously de-
scribed, using a LSR1 (BD Biosciences) and CellQuest software (BD Biosciences) (17). There were 40,000–50,000 events typically collected per sample.

CDR3 size and sequence analysis

Primers for TCR Vα and Vβ (20) spectratyping and the determination of CDR3 lengths (21) have been previously described. Primers used in this study are shown in Supplemental Table 1. RNA from purified T cells was isolated using TRIzol reagent (Invitrogen, Carlsbad, CA) according to manufacturer’s instructions. Briefly, total RNA was incubated for 5 min at 65°C with oligo-d(T)16 primer (100 pmol) and 2′-deoxynucleoside 5′- triphosphates (10 mmol each). After cooling, cDNA was synthesized with 200U SuperScript III (Invitrogen) in first-strand buffer containing 5 mM DTT (Invitrogen) and 20U RNasin (Promega, Madison, WI) at 50°C for 1 h. PCR was performed using one-twentieth of cDNA, 250 nM sense primers specific for Vβ 4, 7, 8.2, 8.3, 9, 11, 12, and 14, and Cβ antisense primer. Instead of Cβ primer, MCB2 primer was used for amplification of Vβ8.2 in Vβ8.2 Tg mice. After a DNA denaturation step (1 min at 95°C), PCR conditions were 40 cycles at 95°C for 1 min, 60°C 1 min, and 72°C for 1 min with 5 min for the last extension. The sense primers for Vα (VA1-1, VA2-1, VA3-1, VA8-1, and VA14-1) and MCA2 primer were used for PCR as described previously, except the annealing temperature was 55°C.

For Vβ and Vα spectratyping, 0.5 μL of the PCR reaction described previously was subjected to a second round of PCR for 30 cycles using the amplification conditions described previously. For Vβ spectratyping, 5′-FAM-labeled 5′-PET-fluorescence-labeled Jα1.1 antisense primer (20) and the Vβ sense primer described previously specific for each of the Vβ segment were used. The 5′-fluorescence-labeled MCB3 antisense primer was used for analysis of Vβ8.2 in TCRB Tg mice. For Vα spectratyping, 5′-fluorescence-labeled MCA3 antisense primer and the specific Vα primer, described previously, were used. After amplification, PCR products were diluted 1:20 in 90% distilled water and 0.5 μL of the diluents was loaded to ABI Prism 3730x1 DNA analyzers (Applied Biosystems, Foster City, CA). Fragment analysis sample files were analyzed with Peak Scanner software (Applied Biosystems).

Partial or full-length TCR Vα2 genes were amplified from the RT-PCR products using the VA2-1 and MCA2 primers or the Vα2-Xho primer and Vα2-Eco primer, respectively. Partial Vα2 PCR products were cloned into pCRII-TOPO vector using TOPO TA Cloning Kit (Invitrogen). The full-length Vα2 PCR products were digested with EcoRI and XhoI and cloned into pML retrolvar vector (22) and sequenced using a pML sequencing primer. Clones were randomly chosen for sequencing, CDR3 sequences were analyzed by basic local alignment search tool and IMGT-V-QUEST.

Data analysis

To quantify skewing of the TCR repertoire, the method of Gorochov et al. (23) was used and is represented as diversity (D) scores. In brief, fluo-
rescent intensity (peak height) associated with each peak in an individual spectratyping profile was automatically measured and transferred into Excel spreadsheets. The percent representation of an individual peak height was calculated in relationship to the sum of all peaks heights in that profile. First, these calculations were performed and averaged for multiple related profiles to establish a reference profile. For the reference were the CDR3 length profiles from CD4+CD25+ conventional T cells from eight individual C57BL/6 mice that exhibited a Gaussian distribution characteristic of a highly diverse TCR repertoire. For the experimental samples, the absolute difference in these percentages at each peak was determined in comparison with the reference, summed, and then divided by two. This value represents the extent that the repertoire varied from the reference for an individual Vβ or Vα profile. The D score represents the mean of these values for each of the eight Vβ or four to five Vα segments.

Statistical analysis of CDR3 sequences were performed by using Estima-
torS 8.0.0 software (R.K. Codwell, http://viceryx.eeb.ucconn.edu/EstimateS). Default-shared species settings were used to calculate abundance-based coverage estimator (ACE) values to estimate the number of unique Vα2 se-
quences for each experimental group and the Morisita-Horn sample similarity index to quantify and estimate the similarity between the Vα2 sequences between any two groups. For the Morisita-Horn index, a value of zero indicates complete dissimilarity, whereas one indicates identity. A one-way ANOVA using Tukey’s multiple comparison test was used to assess differ-
ences between the test groups and control (p < 0.05) for D scores and ACE values. Unpaired one-tailed Student t test was used to assess health scores versus D scores or ACE values. The p values < 0.05 were considered sta-
istically significant and are designated by an asterisk (*) in the graphs.

Results

TCR repertoire of IL-2Rα−/− mice

Autoimmunity and lymphoproliferative disease is evident in very young (2 wk old) IL-2Rα−/− mice and systemic autoimmunity progresses rapidly, such that most C57BL/6 and BALB/c IL-2Rα−/− mice die between 8–12 wk and 4–6 wk of age, re-
spectively (18). These mice, especially early during disease, often contain LN with 10-fold increased cellularity, including auto-
reactive CD4+ T cells. The TCR diversity of these unregulated CD4+ T lymphocytes was broadly sampled by measuring CDR3 nucleotide lengths for several TCRβ and TCRα genes and was compared with CD4+ T cells from age-matched control WT mice (Fig. 1A). Similar to IL-2−/− and IL-2Rs−/− mice (24), in most cases, a polyclonal distribution of CDR3 lengths was noted for each Vβ and Vα TCR subgroups for 3–5 wk old IL-2Rα−/− mice (Fig. 1B), although some CDR3 length skewing was noted (e.g., Vβ8.3 for mouse no. 1).

To quantify these differences, D scores were calculated, which represent the mean of the variance of Vα or Vβ profiles from an experimental sample when compared with a highly diverse Gaussian profile from a reference sample (23), which represented the average peak height for each Vα or Vβ profile from CD4+ CD25+ conventional T cells obtained from eight adult WT C57BL/6 mice. In this report, D scores for individual spec-
tratyping profiles ranged from 3–5 for a naive Gaussian fully di-
verse unperturbed TCR repertoire to 77–78 for a monoclonal repertoire represented by expression of a single TCR β-chain.

The D-scores for each Vα and Vβ genes were higher for IL-
2Rα−/− CD4+ T cells (Fig. 1B). When the individual Vα and Vβ D scores were averaged, values of 10.6 and 19.6, respectively, were noted, which were significantly higher (p < 0.02) than the average D scores of 4.9 and 4.8, respectively, for WT CD4+ T cells, which closely approximated the reference samples. Very similar low D scores were noted for conventional WT CD4+ T cells...
T cells after depletion of their Treg cells (see the values for control B6 in Supplemental Fig. 3). The higher scores for IL-2Rβ<sup>b−/−</sup> CD4<sup>+</sup> T cells likely represent some clonal expansion by autoreactive T cells. Nevertheless, these data indicate that T cell lymphoproliferation that accompanies autoimmunity associated with IL-2Rβ deficiency is not dominated by an oligoclonal CD4<sup>+</sup> T cell response.

**Health monitoring of IL-2Rβ<sup>b−/−</sup> mice**

Previous data demonstrated that the adoptive transfer of 2 × 10<sup>5</sup> MHC-matched or fully mismatched donor Treg cells consistently prevented lethal autoimmune disease and the donor Treg cells persisted lifelong in IL-2Rβ<sup>b−/−</sup> recipient mice (16, 18). In this report our objective was to assess the TCR repertoire of such donor Treg cells in control of autoimmunity. Therefore, the autoimmune status of all IL-2Rβ<sup>b−/−</sup> recipients was always assessed in parallel with TCR diversity. To establish objective criteria, we devised a health scoring system by evaluating substantial past and concurrent data from this study from normal and IL-2Rβ<sup>b−/−</sup> C57BL/6 mice with regard to immunological changes that are associated with immune system dysregulation because of absent IL-2Rβ function (Fig. 2A). Early immune changes include lymphoproliferation (assessed by LN cellularity and/or histopathological evidence of lymphocytic hyperplasia) and increased activated CD4<sup>+</sup> T cells (assessed by CD69 expression), followed by autoimmunity [assessed by low hematocrit because of hemolytic anemia (15, 18) and/or inflammatory infiltrates in nonlymphoid tissues], leading to wasting and death. IL-2Rβ<sup>b−/−</sup> mice contain only a few immature Foxp3<sup>+</sup> Treg cells in the peripheral immune compartment. For adoptively transferred mice, assessment of engraftment levels of donor Treg cells represents an additional factor related to control of autoimmunity. A cutoff for normal values was set at 1 SD above or below the mean values from WT mice, as appropriate, except for the hematocrit where a more strict criteria was used for a definitive assignment of hemolytic anemia. These values are represented by the horizontal line in each graph within Fig. 2. A threshold of 1 SD was chosen to establish autoimmune trends that varied from the mean rather than a more stringent threshold that would assign a high probability of an abnormal value, as these higher values generally associate with severe disease.

Health scores were assigned as +1 each for abnormally (1) high LN cellularity and/or histopathological evidence of lymphocytic hyperplasia in the spleen or LNs, (2) increased CD69 expression by CD4<sup>+</sup> T cells, or (3) low levels of donor Treg cells. A +4 is assigned when clear evidence of autoimmunity is observed, namely, hematocrits <40 and/or moderate or greater levels of lymphoplasmacytic infiltrates in nonlymphoid tissues as assigned by a veterinary pathologist. Typical targets tissues include the lung, liver, colon, and salivary gland (15, 18, 25).

**TCR repertoire of donor Treg cells from autoimmune-free IL-2Rβ<sup>b−/−</sup> mice**

An important advantage of the IL-2Rβ<sup>b−/−</sup> model is that the donor Treg cells provide a fixed population of cells to assess their TCR repertoire as it relates to suppression of autoimmunity of polyclonal autoreactive T cells that are continually emerging from the thymus. To directly and broadly investigate TCR diversity within the donor Treg cells, the CDR3 size distribution pattern of various Vβ and Vα TCR subgroups was assessed for donor syngeneic and allogeneic Treg cells from individual recipient mice by spectratyping. Comparing the TCR repertoire of engrafted syngeneic and allogeneic donor Treg cells provided a means to assess whether clonal diversity of self-reactive Treg cells was similar or exceeded cross-reactive alloreactive Treg cells.
Initially, CD4\(^+\) T cells were titrated to determine the point where spectratyping profiles (Supplemental Fig. 1) began to vary from fully Gaussian. After calculating D scores, this was \(1 \times 10^5\) cells for TCR\(\beta\) and \(4 \times 10^4\) cells for TCR\(\alpha\) (Fig. 3A). Spectratyping analysis of all experimental samples was performed using cell numbers that were always greater than these limits and usually \(>2 \times 10^5\) T cells were used as this number of donor Treg cells was typically obtained from individual mice. Thus, deviation from highly diverse Gaussian spectratyping profiles cannot be attributed to insufficient sample size.

As expected, Treg cells directly isolated from C57BL/6 mice, which represent the input cells used for adoptive transfers, exhibited Gaussian TCR\(\alpha\) and TCR\(\beta\) CDR3 size distributions (Fig. 3B, top) and low D scores (Fig. 3C) with an average value of 5.7 and 9.7, respectively, characteristic of a highly diverse TCR repertoire. However, TCR\(\beta\) spectratyping profiles for donor C57BL/6 Treg cells isolated 10–16 wk posttransfer from syngeneic (Fig. 3B, middle) or allogeneic autoimmune-free IL-2R\(\beta\)\(^{-/-}\) recipients (Fig. 3B, bottom) were not Gaussian for many of the V\(\beta\) subgroups, which was more striking for donor MHC-mismatched C57BL/6 Treg cells from autoimmune-free BALB/c IL-2R\(\beta\)\(^{-/-}\) mice. Similar spectratype profiles were obtained for BALB/c Treg cells after adoptive transfer into syngeneic BALB/c or allogeneic C57BL/6 IL-2R\(\beta\)\(^{-/-}\) recipients (Supplemental Fig. 2). Although skewed CDR3 size distributions were noted for many V\(\beta\) subgroups, individual mice exhibited unique patterns of skewing within a particular V\(\beta\) subgroup, consistent with distinctive Treg cell TCR repertoires associated with each recipient. Such skewing of the spectratype profiles was less dramatic for TCR\(\alpha\).

D-score analysis of the spectratyping profiles from all 16 individual donor Treg cell samples for each V\(\alpha\) and V\(\beta\) subgroup revealed three obvious trends (Fig. 3C). First, D scores from donor-engrafted Treg cells were typically higher than Treg cells from normal C57BL/6 mice. Second, D scores for donor Treg cells were generally higher for TCR\(\beta\) than TCR\(\alpha\). Third, D scores from donor allogeneic Treg cells were usually higher than donor syngeneic Treg cells. When considering donor C57BL/6 Treg cells from syngeneic and allogeneic IL-2R\(\beta\)\(^{-/-}\) recipients (Fig. 3C), the averaged D scores were 24.8 and 36.0 for TCR\(\beta\). These values were significantly higher (\(p < 0.05\)) when compared with input C57BL/6 Treg cells. Averaged D scores for TCR\(\alpha\) were 13.1 and 18.2 for syngeneic and allogeneic donor C57BL/6 Treg cells, respectively. Nevertheless, these D scores (\(p < 0.05\)) were greater than noted for the input Treg cells, consistent with some selection of the TCR\(\alpha\) repertoire. Very similar and statistically significant differences were noted for the average D scores of BALB/c Treg cells from syngeneic and allogeneic IL-2R\(\beta\)\(^{-/-}\) recipients (Fig. 3C). These results indicate that fewer alloreactive Treg cells are selected after adoptive transfer and suggest that Ag recognition plays an important role for the Treg cells that persist in the IL-2R\(\beta\)\(^{-/-}\) recipients. Collectively, these findings indicate that a Treg cell population with measurable limitations on their TCR repertoire remains effective in preventing autoimmunity that is potentially imitated by a more diverse autoreactive TCR repertoire.

Spectratyping was also performed on recipient conventional CD4\(^+\) T cells from the autoimmune-free Treg cell “cured” IL-2R\(\beta\)\(^{-/-}\) mice (Supplemental Fig. 3). When considering recipients T cells that received C57BL/6 Treg cells, quantitative analysis revealed averaged D scores for TCR\(\alpha\) and TCR\(\beta\) of 8.5 and 14.7, respectively, in a syngeneic setting (C57BL/6 IL-2R\(\beta\)\(^{-/-}\) recipients) and 7.9 and 16.2, respectively, in an allogeneic setting (BALB/c IL-2R\(\beta\)\(^{-/-}\) recipients). These values are lower than found in untreated IL-2R\(\beta\)\(^{-/-}\) mice, but significantly higher (\(p < 0.05\)) for TCR\(\beta\), but not TCR\(\alpha\), when compared with conventional CD4\(^+\) T cells from normal mice. Very similar averaged D scores and trends were noted for recipient T cells that received BALB/c Treg cells. These results demonstrate equivalent degree of normalization of TCR repertoires in these IL-2R\(\beta\)\(^{-/-}\) recipients by syngeneic and allogeneic Treg cells. This finding is consistent with the ability of syngeneic and allogeneic Treg cells to readily suppress autoimmunity.

Control of autoimmunity by Treg cells with a single TCR\(\beta\) TCR\(\beta\) Tg mice provide a source of Treg cells with substantial limits on their TCR diversity because of allelic exclusion at the TCR\(\beta\) locus as it results in T cells that express a single TCR\(\beta\)-chain. Direct analysis of cells from these mice indicates that their decreased TCR diversity resulted in selection of a lower proportion of Treg cells within the peripheral immune compartment when compared with littermate control mice (Fig. 4A).
FACS analysis confirmed that purified Treg cells from TCRβ Tg+ mice on the H-2b genetic background dominantly expressed a single TCRβ-chain on essentially all CD4+CD25+ Foxp3+ Treg cells (Fig. 4B). When TCRβ Tg+ and Tg2 littermate control Treg cells were assessed in vitro, both types of Treg cells equivalently suppressed T cell proliferation, indicating that Treg cells selected through a single TCRβ-chain did not exhibit any intrinsic loss of suppressive activity (Fig. 4C). Treg cells with a single TCRβ-chain are under similar homeostatic regulation in the periphery as Tg2 C57BL/6 Treg cells as assessed by Ki67, a molecule expressed from mid G1 through G2M phases of the cell cycle, and Bcl-2 staining of Foxp3+ T cells (Fig. 4D).

To directly explore the efficacy of Treg cells with a limited TCR repertoire to prevent autoimmunity, we assessed the ability of graded numbers of WT and TCRβ Tg+ Treg cells to engraft and control autoimmunity in IL-2Rβ2/2 mice. To follow engraftment, we took advantage of past work that demonstrated that essentially all Treg cells in the adoptively transferred IL-2Rβ2/2 mice are of donor origin (16–18). Indeed, a representative example of such an autoimmune-free recipient shows that ~98% of the Foxp3+ T cells were CD45.1+ congenic-marked donor Treg cells that were all CD25high (Supplemental Fig. 4). The few host Foxp3+ T cells were immature Treg cells based on lower expression of Foxp3 and lack of CD25. These features, coupled with the near ubiquitous expression of the Vβ8.2 of the TCRβ Tg+ Treg cells, allowed identification and isolation of donor TCRβ Tg+ Treg cells by selection of CD4+CD25+ T cells.

With respect to engraftment, donor MHC-matched TCRβ Tg+ Treg cells were readily detected in the periphery of C57BL/6 IL-2Rβ2/2 mice that received 2 × 105 Treg cells at a level similar to found for control Tg2 donor Treg cells (Fig. 4E). Furthermore, engraftment of TCRβ Tg+ Treg cells 1-wk posttransfer was comparable (data not shown) to that previously found for WT Treg cells (17), indicating that restricting Treg cell TCR diversity did not affect early steps controlling engraftment and expansion of the donor Treg cells. For the recipients of TCRβ Tg+ Treg cells, ~90% of the their CD4+CD25+ T cells were Vβ8.2 and Foxp3+, whereas Vβ8.2 TCRs were expressed on a small minority of the engrafted WT donor CD4+CD25+ Foxp3+ T cells (Fig. 4E). Engraftment by TCRβ Tg+ Treg cells was similar in most IL-2Rβ2/2
recipients that received $2 \times 10^5$ donor cells (Fig. 4F) and at a proportion of the CD4+ T cells typically seen when WT Treg cells are transferred. However, TCRβ Tg+ Treg cells engraftment was dose-dependent with lower proportional engraftment at lower number of input Treg cells (Fig. 4F).

With respect to health status, there was an obvious correlation between number of input donor TCRβ Tg+ Treg cells, the subsequent proportional engraftment, and abnormal measurements of individual parameters of health status (Fig. 4F). This is particularly evident when examining CD69 expression on CD4+ T cells for recipients that received $0.5-2 \times 10^5$ donor Treg cells (Fig. 4F). Of note for the 200,000 12-wk input TCRβ Tg+ group, the three mice with the highest %CD69+ were the same three mice with the lowest %Treg cell engraftment (Fig. 4F). Although very few of these mice had abnormal readings for hemolytic anemia, extensive histopathology was performed on most of these recipients as another measure of autoimmunity and used in health scoring.

A dose-dependent relationship was noted where at $0.5 \times 10^5$ donor input Treg cells, most recipients of WT or TCRβ Tg+ Treg cells exhibited autoimmunity, whereas both types of Treg cells controlled autoimmunity at $1-2 \times 10^5$ donor input cells (Fig. 4G). The parallel loss of suppression of autoimmunity by $0.5 \times 10^5$ WT or TCRβ Tg+ input Treg cells indicates that factors other than TCR diversity contribute to prevent autoimmunity and this result may be due to an initial lower ratio of Treg cells to autoreactive T cells, such that the autoreactive T cells prevailed. This conclusion is supported by other work that showed multorgan inflammatory disease occurred in settings where lymphopenic mice are reconstituted with T cell populations containing limiting number of Treg cells (26). For IL-2Rβ2/2 recipients of $2 \times 10^5$ Treg cells, all 18 recipients of WT Treg cells (Figs. 2B, 4G) and 11 of 16 recipients of TCRβ Tg+ Treg cells (Fig. 4G) were autoimmune-free (health score #1). Thus, in a model where autoimmune disease penetrance is 100% for untreated mice, a typical curative number of donor Treg cells with a single TCRβ-chain, which inherently constrains the TCR repertoire, often prevents autoimmunity. Nevertheless, a tendency of higher health scores was assigned to recipients that received $1-2 \times 10^5$ TCRβ Tg+ Treg cells. This finding suggests...
that the constraints of a single TCR β-chain may lead to “holes” in the Treg cell TCR repertoire that act as an initiating factor for autoimmunity. The strong association of MHC or HLA polymorphisms and autoimmune diseases may in part reflect such a contraction of the Treg cell TCR repertoire.

The TCRα repertoire of donor Treg cells expressing a single TCR β-chain

The donor TCRβ Treg cells were isolated from most of the recipient IL-2Rβ−/− mice to evaluate the diversity of their TCR α-chains. Spectratyping was performed for Treg cells from recipients that received 50,000 and 200,000 Treg cells to broadly assess the extent TCR diversity was narrowed under conditions of suboptimal versus optimal number of transferred cells and as a function of disease status. Representative spectratype profiles are shown for the input TCRβ Tg+ Treg cells (Fig. 5A, top), selected mice that were autoimmune-free (Fig. 5A, middle), or exhibited autoimmunity (Fig. 5A, bottom). The input T cells expressed a generally polyclonal endogenous TCRα repertoire [(19) and Fig. 5A, top], with an averaged D score of 18.4 (Fig. 5B). Furthermore, direct analysis of conventional CD4+ T cells from these TCRβ Tg mice revealed an averaged D score of 22.8 (Supplemental Fig. 5). Both these values were higher than found for WT C57BL/6 Treg mice that were autoimmune-free (Fig. 5A, bottom), or exhibited autoimmunity (Fig. 5B). Both the input and engrafted Treg cells showed a single monoclonal peak and a D score of 77–78 for Vβ8.2 Importantly, quantitative analysis of these profiles from all individual mice revealed a consistent trend of higher D scores for each of five Vα subgroups that in aggregate assessed ~30% of the Vα genes (Fig. 5B). The averaged D scores for the engrafted Treg cells (29.8–36.6) were higher and significantly different (p < 0.05) from the input Treg cells, although not from each other. This finding suggests that the Treg TCR diversity is independent of the number of transferred Treg cells and health status of the recipient. Indeed, there was no significant difference in the mean averaged D scores when they were plotted as a function of health scores (Fig. 5C).

To further evaluate the restriction of the TCRα repertoire and specificity, CDR3s from Vα2 TCRs were sequenced from three pools, each derived from five to six mice, of pretransferred input donor Treg cells and 18 individual recipients at 8–16 wk post-transfer (Fig. 6A). For each experiment, according to the shown donor/recipient relationship, 100–200 sequences were obtained for each pool of input Treg cells, whereas usually 30–50 sequences were obtained from recipient-derived Treg cells such that when pooled each group also consisted from 100–200 sequences. The diversity of TCRs in each individual sample or after pooling the

**FIGURE 5.** Spectratype analysis for TCRα diversity of input and donor TCRβ Tg+ Treg cells from individual IL-2Rβ−/− recipient mice. At the indicated time posttransfer, TCRβ Tg+ Treg cells were purified from individual C57BL/6 IL-2Rβ−/− recipients. A, Representative Vα spectratype analysis for CDR3 was performed for the indicated Vα subfamily and Ca gene segment for input and donor-derived Treg cells. Spectratype analysis for CDR3 size distribution of Vβ8.2 is shown to illustrate the expression of the TCRβ Tg by these populations of purified Treg cells. The D score is shown to the right of each spectratype profile. B, D scores for Vα spectratype distribution profiles for all mice (n = 4–6 mice/group). Data for the averaged D scores were compared by one-way ANOVA. C, The averaged D score for the recipients that received TCRβ Tg+ Treg cells were plotted against the health scores as assigned in Fig. 4F. Data were compared by unpaired one-tailed t test.
sequences within a group were calculated using the ACE of species richness (Fig. 6B). This analysis revealed statistically significant ($p < 0.05$) higher ACE values, indicative of greater diversity, for the Vα2 TCRs from the pretransferred donor Treg cells when compared with those for the postransferred cells from individual recipients. These latter values were not significantly different from each other. A plot of ACE values and health scores of the corresponding individual recipient mice revealed a trend toward lower ACE values and higher health scores, but this was not statistically significant. (Fig. 6C), which agrees with the spectratyping results. Collectively, these data indicate that each recipient contains TCRs with more limited diversity than the input donor cells and that factors other than TCR diversity are also important to determine whether autoimmunity occurs. One such factor may be the specificities of the Treg TCRs present in each recipient.

The lower ACE values for TCR diversity of Treg cells that engrafted and persisted within each recipient was independent of input cell number for the range tested. However, after the Treg cell TCR sequences from each individual recipient within a group were pooled, the ACE values consistently increased (Fig. 6D). This finding suggests that overall fraction of Treg cell specificities that contribute to the control of autoimmunity is greater than operative within a single IL-2Rα−/− recipient.

Morisita-Horn similarity index values were also compared for these groups of TCR data sets pre- and posttransfer (Fig. 6D). This analysis revealed that TCR sequences for the input pretransferred Treg cells were more similar to each other than to 16 of 18 posttransferred Treg TCR sequences, either when analyzed individually or pooled. Two exceptions were recipients five and six, which received the same distinct group of donor Treg cells. Higher similarity to the input Treg cells was noted, but both recipients exhibited a high health score of 4, indicative of autoimmunity. The lower overall similarity for recipients that received 200,000 donor Treg cells may reflect in part that TCR sequences were not obtained from the pretransferred cells for these two groups. Furthermore, low Morisita-Horn similarity index values were noted for the individual Treg cell TCRs posttransfer when compared with each other, even between individuals that received the same inoculums of donor Treg cells (Supplemental Fig. 6).

The relationship between the Treg TCR specificities were compared for any CDR3 for an individual pre- and posttransfer sample that was detected at a frequency of ≥4%. An overlap was found in sequences for the pre- and posttransferred donor Treg cells (Fig. 7A) and sometimes the most highly frequent sequence was associated with the postransferred Treg cells. The two most prevalent sequences in the input pretransferred Treg cells were not always found in the postransferred cells (Fig. 7B). Moreover, a large majority of these prevalent sequences (74 of 96) were detected only from the postransferred donor Treg cells. Some of these sequences were shared between donor Treg cells from distinct recipients, but most were not. Many of the most frequent postransferred donor Treg TCR CDR3s were unique to particular
FIGURE 7. Frequency of Vα2 CDR3 sequences associated with pre- and posttransferred TCRβ Tg⁺ Treg cells. A. All unique sequences in the Vα2 data set expressed at a frequency ≥4% in any pretransferred sample (input) or any individual recipient posttransferred (output) were identified. The prevalence of these sequences were compared with each other and represented as a heat map. B. The frequency distribution of the two most prevalent sequences in the pretransferred Treg cells in relationship to matched output posttransferred Treg cells. C. Frequency of the three most prevalent sequences within individual samples of pre- and posttransferred Treg cells.
recipients. For a sequence at a frequency of 5% in the posttransferred Treg cells, binomial probability calculations indicate that there is >99% confidence level that this sequence would be found in the pretransferred cells when accumulating 115–188 CDR3s. Considering this calculation and the most prevalent sequence associated with posttransferred Treg cells from individual recipients was found at frequency usually ≥15% (Fig. 7C), many of these dominant specificities must have been derived from a minor constituent of the pretransferred Treg cells. In comparison with the input Treg cells, more highly frequent Vo2 sequences were associated with most of the posttransferred Treg cells that were particularly striking in three of the four mice, each with health scores of 3, that received 200,000 Treg cells and were evaluated 13–16 wk posttransfer (Fig. 7C). Collectively, these results indicate that there is substantial TCR repertoire reshaping after adoptive transfer of Treg cells into IL-2Rβ−/− mice and at sufficient input cell numbers (>100,000 Treg cells), control of autoimmunity is often still achieved.

**Discussion**

Although the Treg cell repertoire is highly diverse (7–11), the relevance of this high TCR diversity in suppressing potentially autoreactive T cells that escape thymic negative selection is poorly understood. One major finding in this report is that control of autoreactive T cells is readily achieved by only a fraction of the total TCR diversity expressed by Treg cells. We show that a fixed population of input donor Treg cells is selected after adoptive transfer into IL-2Rβ−/− mice to express a restricted portion of the total Treg cell TCR repertoire and such Treg cells effectively suppress a continual source of autoreactive polyclonal T cells that escape thymic negative selection. This was strikingly shown when WT Treg cells were adoptively transferred into IL-2Rβ−/− mice. All such recipients were judged to be autoimmune-free, but the donor Treg cells exhibited readily measurable limitations on their TCR diversity when compared with the input donor pretransferred Treg cells. These limitations were broadly seen over multiple Vo and Vβ CDR3s and were particularly striking for allogeneic donor Treg cells. Moreover, control of autoimmunity also often occurred after transfer of an optimal number of 2 × 10^5 Treg cells with a defined limitation on their TCR diversity through expression of a single TCR β-chain, although the durability of suppression appears lower than WT Treg cells.

Posttransferred WT Treg cells showed limitations of their TCR diversity that was more striking when compared with CD4+ T cells from autoimmune untreated IL-2Rβ−/− mice, suggesting that Treg cells may require less TCR diversity than the target autoreactive cells that they suppress. In support of this idea, donor Treg cells with a single TCRβ also readily suppressed autoimmunity in IL-2Rβ recipients. We assume that the large majority of the Treg cells that persist in these recipients are dedicated to suppress autoreactive T cells. However, it is possible that the therapeutic Treg cells are a subset of the persistent engrafting cells. If this proves to be true, even more limited TCR diversity than we have measured in this study is sufficient for suppression of peripheral polyclonal autoreactive T cells. Of note, the calculation for diversity for all Vo2 CDR3 sequences from posttransferred TCRβ Tg+ Treg cells derived from the entire group of recipients that received a common pool of Treg cells was greater than measured for Treg cells from a single recipient. Thus, the sum of Treg cell specificities in the donor population used to suppress autoreactive T cells is greater than found in an individual recipient.

Why is the Treg cell repertoire highly diverse when only a fraction of this diversity is required to effectively suppress self-reactive T cells? One obvious answer to this question and supported by our results is that high Treg cell TCR diversity guarantees that self-tolerance is readily established toward a random conventional TCR repertoire with unpredictable and distinctive self-specificities that are not deleted as a course of central tolerance. Such a mechanism likely reflects an essential step during the evolution of the adaptive immune system to ensure self-tolerance at a very high frequency for individuals of a species. High TCR diversity by Treg cells also likely ensures their participation in regulating immune responses to various nonself-Ags. Indeed, there is considerable data that Treg cell suppressive activity is not limited to autoaggressive T cells but also downregulates many conventional immune responses to foreign and tumor Ags (27–29).

A second major finding is that there is substantial peripheral reshaping of the Treg TCR repertoire in IL-2Rβ−/− recipients and such reshaping varies for each recipient. This notion was suggested by spectratyping of posttransferred WT Treg cells as the resulting profiles varied for TCR Vβ subgroups when isolated from distinct recipients. Vo2 sequences of pre- and posttransferred TCRβ Tg+ Treg cells indicate that this reshaping often involves selection of rare and distinct specificities within the input Treg population by each recipient, often at the expense of the dominant specificities found on the pretransferred Treg cells. Thus, distinct pools of input Treg cells showed greater similarity to each other than when compared with virtually all posttransferred donor cells from individual recipients.

Peripheral reshaping of Treg TCR repertoire has been previously noted (8–10) and represented by a lowering in the periphery of those specificities that were dominant within the thymus. Although we often noted a decrease in the dominant specificity of the pretransferred Treg cells, the peripheral reshaping within IL-2Rβ recipients was much more extensive and characterized by selection and expansion of minor specificities. It is intriguing to speculate that this latter process may exaggerate peripheral reshaping that normally occurs where flattening of the dominant Treg TCR specificity must lead to some favoring of other less represented specificities. An important distinction of IL-2Rβ−/− mice is that they lack mature Treg cells (30). After transfer of CFSE-labeled Treg cells into IL-2Rβ−/− mice, a minor fraction of the initial donor inoculum of 2 × 10^5 Treg cells successfully engrafted, rapidly expand to normal levels, where the CFSE-label was nearly fully diluted 7 d posttransfer (17). These donor cells provide a lifelong pool of Treg cells through extensive homeostatic proliferation that function to prevent autoimmunity (17, 18, 30). Therefore, this intense homeostatic pressure likely influences the selection of a persistent pool of Treg cells. Our data rule out that the selection and reshaping of the Treg TCR repertoire is strictly a random process of homeostatic proliferation by the donor Treg cells. In this case, we expected that the TCR repertoire of posttransferred Treg cells to closely mirror the input cells. Furthermore, simply the presence of a normal proportion of Treg cells within IL-2Rβ recipients that received TCRβ Tg+ Treg cells did not predict control of autoimmunity. Thus, specificities associated with the Treg cells clearly matter to prevent autoimmunity. Consistent with this notion, the more limited TCR diversity of engrafted allogeneic Treg cells when compared with syngeneic Treg cells suggests that the former cells were selected on allo-Ag. Considering these points, repertoire reshaping, although influenced by homeostasis, likely also includes indexing in some manner to self-Ag(s) and/or the autoreactive T cells, whose specificities appear to vary between individual IL-2, IL-2Rα (24), and IL-2Rβ−/− mice (Fig. 1).

In conclusion, our data favor a model where Treg TCR specificities are actively selected and reshaped in the periphery to favor...
specifications to optimally suppress autoreactive T cells. This mechanism does not normally represent a risk, but rather a benefit to maintain self-tolerance, because of the high diversity of the Treg cell TCR repertoire. Thus, peripheral Treg cell TCR repertoire reshaping represents a feature of adaptive immunity to maintain tolerance as thymic output wanes or during insults to the immune system. In the latter situations, the immune system must be continuously rebalanced after infections, bone marrow transplantation, or the use of drugs for immunosuppression and tumor chemotherapy. In settings where the diversity of Treg TCRs is limited, such as Treg cell transfers into IL-2Rβ<sup>−/−</sup> mice, these selective pressures may result in a key specificity to become underrepresented or absent, leading to autoimmune attack. In an analogous fashion, a similar risk for autoimmune disease may result from pressures may result in a key specificity to become underrepresented or absent, leading to autoimmune attack. In an analogous fashion, a similar risk for autoimmune disease may result in a high reliance on Treg cell specificities of the existing repertoire and from nonautoreactive T cells by endogenous TCR expression. T cells from autoreactive T cells simultaneously with their negative selection in the thymus and from nonautoreactive T cells by endogenous TCR expression.

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Disclosures

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