Characterization of Protective Human CD4+CD25+ FOXP3+ Regulatory T Cells Generated with IL-2, TGF-β and Retinoic Acid

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Abstract

Background: Protective CD4+CD25+ regulatory T cells bearing the Forkhead Foxp3 transcription factor can now be divided into three subsets: Endogenous thymus-derived cells, those induced in the periphery, and another subset induced ex-vivo with pharmacological amounts of IL-2 and TGF-β. Unfortunately, endogenous CD4+CD25+ regulatory T cells are unstable and can be converted to effector cells by pro-inflammatory cytokines. Although protective Foxp3+CD4+CD25+ cells resistant to proinflammatory cytokines have been generated in mice, in humans this result has been elusive. Our objective, therefore, was to induce human naïve CD4+ cells to become stable, functional CD25+ Foxp3+ regulatory cells that were also resistant to the inhibitory effects of proinflammatory cytokines.

Methodology/Principal Findings: The addition of the vitamin A metabolite, all-trans retinoic acid (atRA) to human naïve CD4+ cells suboptimally activated with IL-2 and TGF-β enhanced and stabilized FOXP3 expression, and accelerated their maturation to protective regulatory T cells. AtRA, by itself, accelerated conversion of naïve to mature cells but did not induce FOXP3 or suppressive activity. The combination of atRA and TGF-β enabled CD4+CD45RA− cells to express a phenotype and trafficking receptors similar to natural Tregs. AtRA/TGF-β-induced CD4+ regs were anergic and low producers of IL-2. They had potent in vitro suppressive activity and protected immunodeficient mice from a human-anti-mouse GVHD as well as expanded endogenous Tregs. However, treatment of endogenous Tregs with IL-1β and IL-6 decreased FOXP3 expression and diminished their protective effects in vivo while atRA-induced iTregs were resistant to these inhibitory effects.

Conclusions/Significance: We have developed a methodology that induces human CD4+ cells to rapidly become stable, fully functional suppressor cells that are also resistant to proinflammatory cytokines. This methodology offers a practical novel strategy to treat human autoimmune diseases and prevent allograft rejection without the use of agents that kill cells or interfere with signaling pathways.

Introduction

CD4+ regulatory T cells (Tregs) bearing the Forkhead Box P3 (Foxp3) transcription factor are required to maintain immunologic homeostasis and prevent autoimmunity [1,2]. Mutations of the Foxp3 gene result in immune dysregulation and multiorgan autoimmunity [3]. Both CD4+ cells and CD8+ cells can express Foxp3 [4,3], but the former have received the most attention. Because abnormalities in the numbers and function of Tregs can lead to autoimmunity, allergy and graft rejection, manipulation of these cells to correct these defects offers a novel treatment strategy [6]. Endogenous CD4+Foxp3+ cells can be divided into thymus-derived, natural regulatory T cells (nTregs) which constitutively express high levels of CD25, the IL-2 receptor alpha chain and those induced in the periphery from CD4+CD25+Foxp3− precursors by a TGF-β dependent mechanism (iTregs). In mice and humans these two subsets have been indistinguishable phenotypically until recently [7], and may have separate or synergistic roles in vivo [8,9]. In humans CD4+FOXP3+ Tregs express high levels of CD25 and low levels of CD127, the IL-7 receptor alpha chain [10].

In addition to endogenous Foxp3+ Tregs, substantial evidence exists that the combination of IL-2 and TGF-β can induce naïve CD4+CD25− cells to become FOXP3+ iTregs in both mice and humans. In mice, suboptimal polyclonal TCR stimulation of naïve CD4+ cells with IL-2 and TGF-β can induce iTregs that have protective effects in autoimmune diabetes [11], experimental autoimmune encephalitis[12] and myasthenia gravis [13]. Because
of decreased numbers and/or function of FOXP3+ Tregs in human autoimmune diseases [14], the transfer of iTregs generated ex-vivo could be therapeutic to subjects with these diseases.

In humans CD4+CD25− cells activated by either superantigens or alloantigens with IL-2 and TGF-β developed potent in vivo suppressive activity [15,16], and these alloantigen-induced FOXP3+ iTregs could also induce other CD4+CD25− cells to become TGF-β-dependent suppressor cells [17]. One group recently also reported that polyclonal TCR stimulation of naïve CD4+ cells with TGF-β could result in FOXP3+ suppressor cells [18]. However, the generation of fully functional polyclonal human FOXP3+ iTregs ex vivo is controversial. First, TCR activation without TGF-β can induce naïve CD4+ T cells to transiently express FOXP3 [19]. Secondly, although we and others have observed that TGF-β can greatly increase FOXP3 expression and stability, after one week in vitro suppressive activity of these human CD4+ cells was not greater than control cells [20,21]. Moreover, unlike nTregs which are anergic in response to TCR stimulation, these human CD4+ cells primed with TGF-β produced IL-2 and proliferated robustly following re-stimulation. Interestingly, however, repeated stimulation of TGF-β primed CD4+ cells did result in anergy, membrane-expression of TGF-β, and in vitro suppressive activity similar to that described with nTregs [20,22]. We concluded that human TGF-β primed CD4+ cells one week after culture were partially differentiated cells and required a much longer time to mature than similar mouse FOXP3+ iTregs [21]. Thus, agents that accelerate cell differentiation might be useful for a more rapid generation of human iTregs ex-vivo.

Retinoic acid (RA), a vitamin A derivative, has an important role in the development of various organs including the immune system. RA metabolites strongly contribute to the maintenance of immunologic tolerance. All-trans retinoic acid (atRA), an active metabolite of retinoic acid, markedly enhances TGF-β-induced Foxp3 expression and stability in mice [23], and the expansion of these iTregs by either direct cytokine-dependent [24] or cytokine independent mechanisms [25]. In human CD4+ cells, atRA has been shown to induce histone acetylation at the FOXP3 gene promoter and expression of the FOXP3 protein [26]. Recently, atRA has been shown to enhance the stability and expansion of TGF-β induced iTreg and endogenous nTreg cells [27]. Here we have extensively characterized the phenotype and functional properties of iTregs induced by TGF-β and atRA. We report that atRA markedly accelerates the differentiation of naïve cells to fully functional suppressor cells. Unlike CD4+ cells generated in one week with IL-2 and TGF-β the presence of atRA during this time enabled naïve CD4+ cells to demonstrate strong protective suppressive effects not only in vivo, but also in vitro when transferred to immunodeficient mice. Moreover, unlike endogenous nTregs, atRA-induced iTregs were resistant to the inhibitory effects of IL-1β and IL-6. Thus, we have demonstrated that it is possible to induce naïve human CD4+ cells to rapidly become iTregs that have protective effects in vivo and that are resistant to the inhibitory effects of proinflammatory cytokines.

Results

The addition of atRA to TGF-β enhanced FOXP3 stable expression and accelerated the maturation of naïve CD4+ cells to memory/effector regulatory cells. Naïve CD4+ cells were activated with suboptimal anti-CD3/28 beads titrated to numbers needed for the cells to express CD25. While IL-2 and TGF-β increased the percentage of CD4+CD25+ cells that expressed FOXP3 after 5 days of culture, the addition of atRA to TGF-β markedly enhanced this effect (Figure 1A and B). Time course studies revealed that adequate levels of IL-2 can sustain TGF-β induced FOXP3 [28] (result not shown). However, with less IL-2, TGF-β induced Foxp3 also decreased after culture for 7 to 9 days, while FOXP3+ cells induced by atRA and TGF-β remained stable (Figure 1B). This finding is in agreement with Wang et al [27].

Thus, the combination of atRA and TGF-β rapidly induced naïve CD4+ cells to express FOXP3, and the stability of this transcription factor is less IL-2 dependent than FOXP3 induced by TGF-β alone.

Although IL-7 is an important growth and survival factor for certain T cell subsets, CD127, the α chain of the IL-7 receptor, is down-regulated on nTreg cells. These cells are CD25+CD127dim [10]. Figure 2A shows that following activation for 6 days, CD127 displayed by naïve CD4+ cells was moderately down-regulated. While TGF-β enhanced this down-regulation of CD127, atRA resulted in more than 90% of the CD4+ cells becoming CD127dim. However, only a small fraction expressed FOXP3. Because >80% of CD4+ cells activated with atRA and TGF-β expressed FOXP3, presumably most were also CD127dim. Similarly, although atRA alone also accelerated transition from CD45RA+ to CD45RO+ cells, most of these cells were FOXP3+. Only with the combination of atRA and TGF-β do most CD45RO+ cells also express FOXP3 (Figure 2B).

Figure 3A shows ten markers characteristically expressed by nTreg cells [22,29]. Of these, naïve CD4+ cells express only L-selectin (CD62L) and CCR7 (see below). The figure shows that following TCR activation with atRA and TGF-β added separately or together, naïve CD4+ cells acquire other Treg-related receptors and retain CD62L and CCR7. These cells expressed GITR, and CTLA-4, although these markers are also expressed by control activated T cells. Besides FOXP3 and CD127 shown above, the combination of atRA and TGF-β increased the intensity of CD122, PD-1, and TNFRII (Tumor necrosis factor receptor II) staining. Although some activated T-Mac expressed TNFRII, atRA markedly enhanced this effect. These cells, however, were FOXP3+ . Only with both atRA and TGF-β did most of these cells display FOXP3. TNFRII expression has been described on mouse nTregs, but to date not on human Tregs [29].

Finally, activated nTreg cells express membrane-bound TGF-β [30]. We observed that adding atRA to TGF-β greatly increased the number of iTreg cells expressing membrane TGF-β (Figure 3B). Unlike most markers that are stained at 4°C, membrane-bound TGF-β is maximal at 37°C. Thus, the addition of atRA to TGF-β increases the conversion of activated CD4+ cells to the effector/memory cells and accelerates their phenotypic differentiation to FOXP3+ Treg cells.

Activation of naïve CD4+ cells with atRA added to TGF-β enables them to retain or acquire receptors needed to recirculate from blood to lymphoid organs. Naïve and central memory CD4+ cells constitutively express the lymphoid homing receptors CD62L and CCR7 that enable them to circulate from blood to secondary lymphoid tissues [31]. CD4+CD45RO+ effector cells generally lack these receptors and express others that enable them to migrate to extravascular sites. Even though most nTregs are CD45RO+, they continue to express CD62L and CCR-7 which facilitate their trafficking to lymphoid organs. CCR7 has been reported to be needed for nTreg homing and function in lymphoid tissues [32].

Figure 3A shows that following activation of naïve CD4+ CD45RA+ cells, down-regulation of CD62L is decreased by TGF-β. Similarly, activation with atRA decreases CCR7 down-regulation. Accordingly, when naïve CD4+ cells were activated with both TGF-β and atRA, expression of both CD62L and CCR7 was retained, even though the cells they underwent transition from CD45RA to CD45RO (Figure 2B). Thus, similar
to nTregs, most atRA/TGF-β-induced iTregs are CD45RO+ effector/memory cells that continue to express CCR7 and CD62L. Human nTregs also express CCR4, another lymphoid homing receptor [33]. Interestingly, TGF-β also induced naïve CD4+ cells to express CCR4 (Figure 3A). CD103 (αE integrin) is another
homing receptor especially important in the mucosal immune system [34]. Human nTregs and iTregs express CD103 [18]. Here we observed that the combination of TGF-β and atRA induced higher levels of CD103 than either agent used alone (Figure 3A).

In other experiments, atRA/TGF-β-induced iTregs and expanded nTregs were restimulated with anti-CD3/28 beads and low dose IL-2 for 3 days. We observed that CCR4 and CCR7 expression remained high on iTregs, but began to decrease on nTregs (Figure S1).

Tregs generated with atRA and TGF-β produce low levels of proinflammatory cytokines, are hypoproliferative in vitro, and develop potent suppressive activity in vitro and in vivo. Unlike naïve CD4⁺ cells primed with TGF-β, the addition of atRA to TGF-β resulted in functional activities similar to those of nTregs within 5 to 7 days following activation. As shown in Figure 4, atRA/TGF-β-induced Tregs (iTregs) produced only small amounts of intracellular IL-2 and IFN-γ. They were non-responsive following restimulation with anti-CD3/28 coated beads and remained hyporesponsive (Figure 5A). Interestingly, the addition of anti-TGF-β to the cultures restored their proliferative response, a finding suggesting that membrane-bound TGF-β may contribute to anergy in vitro.

As has been reported previously, human polyclonally-stimulated CD4⁺ cells primed with IL-2 and TGF-β did not acquire marked

Figure 3. Phenotypic characterization of human atRA/TGF-β induced CD4⁺ iTregs at day 5. Naïve CD25 depleted CD4⁺ cells were stimulated with CD3/CD28 beads and the additives indicated above for 5 days. (A) The staining intensity of each marker and the isotype control is shown in comparison with the staining of unstimulated CD4 naïve cells. (B) The cells were rested in fresh medium containing 10% FCS for 24 hours and restimulated with anti CD3/CD28 beads (1:1) and IL-2 (20 U/ml) for 48 hours. FACS analysis of CD25 and mTGFβ1 expression by the various CD4⁺ conditioned subsets is shown. The results are representative of three separate experiments.

doi:10.1371/journal.pone.0015150.g003
Figure 4. AtRA/TGF-β-primed CD4+ cells produce less IL-2 and IFN-γ. Naïve CD4+ cells were stimulated with the additives described above for 5 days, washed and the beads were removed. The cells were rested for 24 hours and then restimulated with anti CD3/CD28 beads for 48 hours. PMA and ionomycin was added for the last 5 hours, and brefeldin A for 4 hours, and intracellular IL-2 and IFN-γ cytokine expression was then assessed by flow cytometry. The results shown are representative of three separate experiments. doi:10.1371/journal.pone.0015150.g004

Figure 6A. Examination of the blood and spleen of mice at 14 days revealed activated with IL-2 (Tcon) to PBMC resulted in similar demise. Nonetheless, they had developed the human anti-mouse graft-versus-host disease (GVHD) described by others [35,36,37]. The percentage of human CD4+ cells present in the blood was small, but comprised >80% of the mononuclear cells isolated from the spleen, liver and lung. In all animals studied engraftment of CD4+ cells was greater than CD8+ cells. An even greater predominance of CD4+ cells was observed in the blood and liver of animals that received iTreg cells. Relatively more NK cells found in spleen, liver, lung and peritoneum compared to blood and bone marrow. The kidney was the only organ with >10% B cells. Thus, to increase the survival of these immunodeficient mice, it is likely that iTreg cells inhibit the magnitude of engraftment, the percentage of human cells found, and the proportion of the mononuclear cells trafficking to various organs, but not the pattern of human engraftment. In addition to using suppression of T cell-dependent IgG production. By two weeks after transfer of human PBMC >2 mg/ml of human IgG was detected in mouse serum (Figure 7C). The addition of iTregs or nTregs to PBMC markedly suppressed human IgG production.
Figure 5. CD4+CD25+Foxp3+ cells induced by atRA and TGF-β are anergic and have potent suppressive effects in vitro. (A) The various primed CD4+ cell subsets indicated were prepared as described above. After culture for 5 days, the cells were washed, rested for 48 hours, and...
Because of the possible therapeutic potential of Treg cells generated and expanded ex-vivo, they should be not only functional, but also resistant to T effector cell conversion in vivo. The plasticity of Foxp3+ nTregs has become evident with proinflammatory cytokines such as IL-1β and IL-6 converting human nTreg cells to Th17 cells [38,39]. These cytokines also inhibit nTreg suppressor cell activity [40]. Accordingly, both atRA-induced iTregs and expanded nTregs were treated similarly with these cytokines. They were restimulated with anti-CD3/28 beads, low dose IL-2, IL-1β and IL-6 for three days (see methods). These treated Treg cells were mixed with autologous PBMC and injected into NOG mice. The xeno-GVHD observed in two experiments was even more aggressive than the previous study (maximum survival 14 vs. 18 days). Nonetheless, Figure 7D shows that both iTregs and nTregs had significant protective effects (p = 0.004). However, after treatment with IL-1β and IL-6, the protective activity of the atRA-induced iTregs was significantly greater than nTregs (Figure 7E). While the protective activity of cytokine-treated nTregs decreased in comparison with control nTregs, the protective activity of cytokine-treated iTregs was modestly greater than control cells (Figures 7F, G).

Phenotypic analysis of these Treg subsets after restimulation using low dose IL-2 revealed decreased expression of Foxp3 by nTregs compared with iTregs. PD-1, GITR, CD103, CTLA-4, and CD62L expression by nTregs also decreased and was weaker than iTregs (Figure S4). This was probably a consequence of the strong TCR stimulation and high IL-2 dose used for Treg expansion. After iTregs and nTregs were restimulated with IL-1β and IL-6, Foxp3 expression by the nTregs decreased even further. Surprisingly, expression of PD-1, GITR, was greater on iTregs restimulated with IL-1β and IL-6 than iTregs restimulated without these cytokines. Thus, atRA-induced iTregs were not only resistant to the inhibitory effects of IL-1β and IL-6, but these proinflammatory cytokines appeared to have a mild positive effect in stabilizing their phenotype and functional activity.

Discussion

We have shown that within one week the addition of atRA to IL-2 and TGF-β can induce polyclonally activated human naive CD4+ cells to become CD25+FOXP3+ Tregs that resemble nTregs phenotypically and functionally. These atRA/TGF-β iTregs have strong suppressive activity both in vitro and in vivo. While IL-2 and
TGF-β can induce polyclonally activated human CD4+ cells to express FOXP3, with one exception [18], most workers have found that these cells lack the functional profile of iTreg [20,21]. However, the addition of atRA to IL-2 and TGF-β completes the maturation of these partially differentiated cells and enables them to protect immunodeficient mice from a xeno-GVHD at least as well as expanded human nTreg cells. Two experiments were combined so that each group contained six to eight mice. A) Survival: TatRA/TGF-β cells significantly enhanced the survival of NOG mice (P<0.01, Log Rank test), B) prevented weight loss (P<0.01); C) Suppressed human IgG production (P<0.01). Panels D–G, Effects of IL-1β and IL-6 on Treg protective effects. Induced Tregs and expanded nTregs were restimulated with IL-1β and IL-6 (See Methods) and 4 million cells were mixed with 20 million PBMC and injected IV into NOG mice. D) Equivalent effects of restimulated iTregs and nTregs; E) Significantly decreased protective effects of nTregs after treatment with cytokines; F) Comparison of nTregs restimulated ± IL-1β and IL-6. G) Comparative effects of iTregs restimulated ± IL-1β and IL-6. Results are a combination of two experiments with 6 mice per group. P values shown were calculated using the Log Rank test.

Figure 7. AtRA/TGF-β iTregs have equivalent protective effects in vivo as expanded nTregs and are also resistant to the inhibitory effects of IL-1β and IL-6. A rapidly fatal xenogenic GVHD was induced by the transfer of human PBMC to NOG mice (See materials and methods). Various conditioned CD4+ cells cultured for 5 to 6 days were rested for 24 h. Then 5 million were added to 20 million human PBMC and transferred IV to sublethally irradiated NOG mice. Two experiments were combined so that each group contained six to eight mice. A) Survival: TatRA/TGF-β cells significantly enhanced the survival of NOG mice (P<0.01, Log Rank test), B) prevented weight loss (P<0.01); C) Suppressed human IgG production (P<0.01). Panels D–G, Effects of IL-1β and IL-6 on Treg protective effects. Induced Tregs and expanded nTregs were restimulated with IL-1β and IL-6 (See Methods) and 4 million cells were mixed with 20 million PBMC and injected IV into NOG mice. D) Equivalent effects of restimulated iTregs and nTregs; E) Significantly decreased protective effects of nTregs after treatment with cytokines; F) Comparison of nTregs restimulated ± IL-1β and IL-6. G) Comparative effects of iTregs restimulated ± IL-1β and IL-6. Results are a combination of two experiments with 6 mice per group. P values shown were calculated using the Log Rank test.

doi:10.1371/journal.pone.0015150.g007
The principal effect of atRA, however, was to accelerate the maturation of naïve CD4+ cells to become effector/memory cells. Uprégulation of CD45RO and down-regulation of CD127 was markedly accelerated. Although these CD127dim cells were FOXP3−, in combination with TGF-β, most now co-expressed FOXP3. Moreover, expression of membrane-bound TGF-β was maximal in the presence of atRA and TGF-β. Most suppressive FOXP3+ cells are CD127dim [10]. When added together, atRA and TGF-β also enhanced expression of PD-1 and CD103. Thus, atRA and TGF-β have induced human naïve CD4+ cells to become phenotypically fully mature Tregs in one week.

Both TGF-β and atRA have well established effects on CD4+ cell trafficking. TGF-β induces CD103, αE integrin [41,42], and atRA induces CCR9 and integrin β7 [26]. We found the combination of atRA and TGF-β enhanced CD103 expression by human CD4+ cells, as previously observed in mice [42]. We also found that TGF-β induced CCR4 and that atRA enhanced this effect. While human CD4+CD25+FOXP3+ cells expressing CCR4 are found in certain tumors and in rheumatoid synovial tissue [43], to our knowledge this is the first description of CCR4 expressed by iTregs induced ex vivo.

Naïve CD4+ cells constantly recirculate from the blood to lymphoid organs and these cells express CD62L (L-selectin) and CCR7 for this function. CCR7 enables CD4+ cells to enter lymph nodes through high endothelial venules [32]. Following strong TCR stimulation, these receptors are down-regulated as CD4+ cells become CD45RO+ effector/memory cells. However, although almost all nTregs have become CD45RO+, they retain both CD62L and CCR7. TGF-β inhibited the loss of CD62L and atRA blocked the downregulation of CCR7 so that the combination of both agents resulted in iTregs similar to nTregs that were effector/memory cells that had not downregulated expression of both of these homing receptors.

In addition to phenotypic similarities, there were several other similarities between iTregs and nTreg cells: 1) As has been described with activated mouse nTregs [44], activated iTregs also express membrane-bound TGF-β. Recently, others reported that human macrophage-induced FOXP3+ iTregs express membrane-bound TGF-β [45]. Previously, we had observed that CD4+ cells primed with IL-2 and TGF-β had to be repeatedly stimulated before they expressed membrane-bound TGF-β [21]; 2) Both nTregs and iTregs produce much less IL-2 and IFN-γ than conventional CD4+ cells. 3) Both Treg subsets proliferate poorly in response to TCR stimulation in vivo. However, this property of iTregs was abolished by antagonizing TGF-β, a result suggesting that the membrane-bound TGF-β contributed to this effect. Reversal of nTreg anergy by neutralizing TGF-β is unusual. The anergy experiments also revealed that atRA and TGF-β do not have to be added together for the cells to become hyporesponsive. Although CD4+ primed with TGF-β respond robustly to re-stimulation, adding atRA to these cells resulted in anergy.

This is the first report showing that the addition of atRA to IL-2 and TGF-β enabled polyclonally TCR-stimulated naïve CD4+ cells to acquire protective suppressive activity in vivo within one week. Whether human TGF-β induced CD4+CD25+FOXP3+ cells develop suppressive activity has been controversial [18,20]. As stated above, we and others could not induce TGF-β primed human naïve CD4+ cells to resemble nTregs in one week. However, since we observed that they did acquire these properties following repeated stimulation, we suspected that at one week they were only partially differentiated cells. With the addition of atRA to accelerate maturation, here we document almost complete suppression of T cell proliferation with only 1 iTreg added to 32 T responder cells, and in vivo suppressive activity at least as strong as nTreg cells. We also assessed the ability of iTreg cells to block a xeno-GVHD and prevent human T cell-dependent IgG production in NOD SCID IL-2R common γ chain−/− immunodeficient deficient (NOD) mice. Others have used RAG−/− SCID cγ chain−/− mice to induce a human xeno-GVHD, and demonstrated protective effects of expanded endogenous nTregs [36,37]. To induce GVHD these

**Table 1.** Human naïve CD4+ cells polyclonally activated with TGF-β and retinoic acid rapidly become CD25+ cells phenotypically similar to nTreg cells.

| Relative expression by naïve CD4+ cells following TCR activation with: | nTreg markers | TGF-β | atRA | Both |
|---|---|---|---|---|
| nTreg markers | medium only | TGF-β | atRA | Both |
| Foxp3 | +/- | | +/- | ++ |
| CD127dim | +/- | + | + | ++ |
| CD45R0 | + | +/- | + | ++ |
| CD122 | +/- | + | +/- | ++ |
| CTLA-4 | + | ++ | + | ++ |
| GITR | +/- | + | + | ++ |
| Membrane-bound | - | + | - | ++ |
| TGF-β | | | | |
| CD6L | Decreased | Sustained | Decreased | Sustained |
| CD103 | +/- | + | +/- | ++ |
| CCR4 | + | ++ | + | ++ |
| CCR7 | Decreased | Decreased | Sustained | Sustained |
| Naïve CD4+ cells were stimulated with suboptimal numbers of anti-CD3/28 beads with IL-2 (50 U/ml) in serum-free medium without APCs for 5 days with the indicated additives. The markers were assessed by flow cytometry.
| - no significant change.
+ minimal to modest enhancement.
++ moderate enhancement.
+++ marked enhancement.

doi:10.1371/journal.pone.0015150.t001
mice must be irradiated and given toxic chlorodronate liposomes to deplete macrophages. As documented by others, to develop GVHD in NOG mice, the use of toxic liposomes is not necessary [35]. Thus, possibly confounding toxic effects contributing to the early death of these mice is avoided. In anticipation of further studies with other epigenetic agents to induce iTregs, we desired to have a rapid readout for our suppressor cell assay. Therefore, we transfected a large dose of human CD25− depleted human PBMC to have a rapid demise of the mice studied.

The addition of CD4+ cells conditioned with atRA and TGF-β, but not TGF-β by itself, delayed the onset of weight loss and extended their survival for two additional months. Other groups had reported that expanded human nTregs cells had protective effects in this mouse [35,36,37]. We learned that the protective effects of iTregs were at least as strong as nTregs. It is not surprising that the protective effect of these Treg cell subsets in NOG mice was not permanent. These Treg cells require human IL-2 to maintain Foxp3 expression [46], and production of this cytokine by the human cells engrafted in these mice will decrease with time. The result will be a corresponding decrease in the suppressive effects of the Treg cells.

Some workers have suggested that FOXP3 expression by TGF-β induced iTregs is only transient [47]. Others have reported that iTregs cannot suppress acute GVHD [48]. It has become evident that the methodology used to prepare mouse iTregs affects both the stability of Foxp3 expression and the protective effects of these cells in vivo [12,49]. The methodology used to prepare human iTregs in this study resulted in suppressive activity that was equivalent to that of nTregs in protecting immunodeficient mice from a rapidly fatal xeno-GVHD. To address the mechanism of the protective effect of iTregs, we provide evidence that they may have suppressed the numbers of human mononuclear cells trafficking to various organs, as others have reported for expanded nTregs [36,37].

The last and most important new finding of this study is that, in contrast to expanded nTregs, atRA-induced iTregs were resistant to the inhibitory effects of the pro-inflammatory cytokines IL-1β and IL-6 on an in vivo protective activity. It has become evident that Foxp3+ Tregs are not stable. These cytokines can down-regulate Foxp3 expression and convert these Tregs to Th17 effector cells [38,39]. This effect can have adverse consequences in established chronic immune-mediated diseases where these cytokines are very abundant. Since retinoic acid in mice can stabilize Foxp3 [23,24,25] and confer T cells resistance to Th17 conversion [25,26,50,51], we expected that human atRA-induced iTregs would be resistant to the inhibitory effects of IL-1β and IL-6. In mice where IL-2 and TGF-β are sufficient to induce iTregs, we found that these cytokines enabled these iTregs to be resistant to Th17 conversion by IL-6 [52].

Because of the well described protective effects of Tregs in immunologic diseases and allograft rejection, it is possible that these cells can be exploited as a therapeutic modality. Efforts are currently underway to use expanded endogenous CD4 Tregs for this purpose. However, because of the small numbers of these cells in the blood, the technical difficulties to expand them and their instability after extended expansion [33], this procedure may be impractical for commercial development. Alternatively, personalized iTreg therapy may be more practical because: 1) large numbers of CD4+CD45RA+ cells can be obtained following pheresis; 2) the procedures to obtain these cells would utilize present methodologies used to isolate stem cells; 3) it is likely that these cells will have proliferative potential in vivo following transfer, and 4) atRA induced iTregs are resistant to Th17 conversion. Thus, the generation of Tregs ex vivo is a promising therapeutic strategy to treat autoimmune diseases and prevent allograft rejection.

Materials and Methods

Mice

NOD/scid/IL2r common γ chain−/− (NOG) mice were obtained from Jackson Laboratory (Bar Harbor, ME). The mice were bred and housed under specific pathogen-free conditions in microisolate cages and given unrestricted access to autoclaved food and sterile water. Animals of both sexes were used for experiments at 8–12 weeks of age. The mice received a single dose of 200 cGy gamma irradiation from a linear accelerator before injection of human PBMC on the same day. Some mice were irradiated but did not receive human PBMC. All experiments were performed according to the guidelines of the Institutional Animal Committee of the University of Southern California.

Monoclonal antibodies and cytokines used

The following FITC, PE, or Cy5 conjugated human antibodies were used for flow cytometric analysis: CD4 (RM4-5), CD25 (PC61), CD45RA (L48), CD45RO (UCHL1), CD122 (Mik-β3), CD127 (hIL-7R-M21), CD103 (Ber-Act3), CD28 (CD28.2), CC4 (1G1), CCR7 (3D12), CTLA-4 (BNI3), PD-1 (MH4), Foxp3 (FJK-16), TNF-αR1 (2B7/97) and TGF-β (4E3). All reagents were purchased from BD Pharmingen (San Diego, CA) and eBioscences (San Diego, CA). Other agents purchased included: OKT3 from Ortho Biotech Products (Bridgewater, NJ), all-trans retinoic acid (atRA) and RPMI medium from Sigma-Aldrich (St. Louis, MO), Biotin-conjugated anti-GITR, recombinant human TGF-β1 and IL-2 from R&D Systems Inc. (Minneapolis, MN), IL-1β and IL-6 from HumanZyme (Chicago, IL), anti-human CD3/CD28-conjugated Dynabeads and carboxyfluorescin succinimidyl ester (CFSE), and AIM-V serum-free medium from Invitrogen (Carlsbad, CA), rapamycin from Calbiochem® EMD Chemicals (Gibbstown, NJ).

Isolation of human nTregs and generation of human iTregs ex vivo

PBMC were prepared from heparinized venous blood of healthy adult volunteers by Ficoll-Hypaque density gradient centrifugation. All protocols that involved human blood donors were approved by the IRB at the University of Southern California. T cells were prepared by negative selection as described previously to a purity of >95% [8]. The CD4+CD25high cells were obtained by fluorescence-activated cell sorting and FOXP3 expressed by these cells was >90%. These endogenous Treg cells are a mixture of nTregs and iTregs induced in vivo, but for simplicity they will be called nTregs. They were expanded for two weeks by activation with anti-CD3/CD28 beads in the presence of IL-2 (500 U/ml) and rapamycin (100 nM); CD4+CD45RA cells were isolated from the CD4+CD25− cells by negative selection and activated with anti-human CD3/CD28 beads 1:10 (one bead to 10 cells) in AIM-V serum-free medium containing Hepes buffer (10 mM), sodium pyruvate (1 mM), glutamine, non-essential amino acids and penicillin and streptomycin. This complete medium was supplemented with IL-2 (50–100 U/ml) ± TGF-β1 (5 ng/ml) ± atRA (100 nM). The dose of TGF-β1 was determined from testing concentrations from 1–20 ng/ml, and atRA from testing 0.01–1000 nM. The populations studied were: naïve CD4+CD45RA+ cells activated with IL-2 (Tno AL): 2) CD4+ cells activated with IL-2 and TGF-β1 (T-rgf-b); 3) CD4+ cells activated with IL-2 and atRA (T-rgf-b); and 4) CD4+ cells activated with IL-2, TGF-β and atRA (T-rgf-b or iTreg cells). The cells were stimulated for 5 days in 24 or 48 well plates, washed, and transferred to new wells with fresh culture medium containing IL-2 (50–100 U/ml) unless stated otherwise. Depending upon cell density, they were split and fresh culture medium with the corresponding additives replaced every 3 days.
Treatment of iTregs and nTregs with IL-1β and IL-6

Nave CD4+ cells that had been stimulated with anti-CD3/28 beads with IL-2 (50 U/ml), TGF-β and atRA for 7 days and nTregs that had been expanded with anti-CD3/28 beads and IL-2 (300 U/ml) for 1 to 2 weeks were prepared. Foxp3 expression by iTregs was between 70 to 75% and expanded nTregs was between 75 to 80%. After the cells were harvested, the beads were removed and each preparation restimulated with anti-CD3/28 beads (1:10), IL-2 (12.5 U/ml), IL-6 (20 ng/ml) for 5 days. This low dose of IL-2 was chosen since others have reported this amount is required for IL-1β to convert nTregs to Th17 cells [39]. High dose IL-2 was avoided because this dose would stabilize Foxp3 expression [34], and thus mask the inhibitory effects of IL-1β and IL-6 on Foxp3+ Treg cells. Neither additional atRA nor TGF-β was added to the iTregs.

Suppressive assays of CD4+ Treg cells in vitro and in vivo

The T cells were labeled with CFSE as previously described[9]. Various ratios of CD4+ conditioned T cells were added to CD25 depleted T cells (T responder cells) and stimulated with soluble OKT3 (20 ng/ml) for 96 hours in the presence of irradiated (30 Gy) non-T cells (1:1 ratio). Cell division was monitored by levels of CFSE dilution. The model to assess suppressor activity in vivo was to protect mice from a rapidly fatal GVHD as described previously[36]. Twenty million/0.2 ml CD25 depleted human PBMC were injected IV into NOG mice sublethally irradiated with 200cGy. Five×10^6 conditioned Treg or T control CD4+ cell subsets stimulated with or without IL-1β and IL-6 were mixed with 20×10^6 PBMC and transferred to the mice. Other mice received 5×10^6 naive CD4+CD45RA+ cells + PBMC. The animals were examined and weighed every two days for evidence of GVHD. The mice were bled 2 weeks after cell injection and human IgG in recipient sera was measured by an ELISA using a human immunoglobulin assay kit (Bethyl, Montgomery, IL).

Histological examination of human mononuclear cell engraftment in NOG mice

Since animals that received PBMC ± non-Treg CD4+ cells died between 14 and 16 days, another series of mice given similar cells were all sacrificed at 15 days for a comparative histologic evaluation of mice that received Treg or non-Treg cells. Peripheral blood obtained by cardiac puncture, spleen, liver kidney, lung, intestine and skin were harvested from recipient mice. Samples were either fixed in formalin for histologic analysis, or collagenase digested and subjected to Ficol/Hypaque centrifugation to study engrafted human mononuclear cells. After formaldehyde fixation, paraffin sections were stained with hematoxylin and eosin. The sections were scanned with a Duoscan T2000XL microscope, and photos were taken with a Nikon 80i digital Camera.

Flow cytometric analysis

Single cells suspensions were stained with conjugated anti-human lymphocyte antibodies indicated above. Percentages of human CD4, CD8, NK and B cells in mouse tissues were determined by gating on human anti-CD45r cells. All analytic flow cytometry was done on a modified dual laser LSRSscan [BD Immunocytometry Systems, San Diego, CA]. For the membrane bound TGF-β staining, each previously primed CD4+ cell subset was restimulated with anti-CD3/CD28 beads (1:1) for 72 h, and stained with anti-TGF-β or isotype control at 37°C for 4 h.

Statistical analysis

Differences in animal Kaplan-Meier survival curves were analyzed by the log-rank test. Differences in proliferation and phenotypes of T cells, FOXP3 expression, and serum IgG levels were analyzed using the 2-tailed Student t test using Prism 4 software (San Diego, CA).

Supporting Information

Figure S1 Stability of homing receptors on Tregs induced with atRA and TGF-β. AtRA/TGF-β-iTregs and expanded nTregs were rested for 2 days and restimulated with anti-CD3/28 beads for 3 days. The cells were then stained for CCR4 and CCR7 and examined by flow cytometry for expression of these chemokine receptors. This result was observed in three separate experiments.

Figure S2 Suppressive activity by Tregs induced with atRA and TGF-β can be abolished by anti-TGF-β antibody. The various primed T cell subsets shown were tested in an in vitro suppressive assay as described in Figure 5. In this experiment the suppressive activity was abolished by anti-TGF-β.

Figure S3 Engraftment human cells in NOG mice 15 days after transfer CD4+ cells activated with TGF-β and expanded nTregs. Hematoxylin and cosin sections of organs from the mice indicated were prepared as described above and compared with sections from control mice injected with PBS. The result shown is representative of studies in three mice.

Figure S4 Effect of IL-1β and IL-6 on the phenotype of iTregs and expanded nTregs. A) Histograms of Foxp3 expression by iTregs and nTregs at the conclusion of the primary cultures and other markers after the Tregs were re-stimulated for 3 days ± IL-1β and IL-6. This experiment was repeated twice with similar results.

Author Contributions

Conceived and designed the experiments: DAH SGZ. Performed the experiments: LL NZ JW. Analyzed the data: LL NZ JW SGZ DAH. Wrote the paper: DAH.

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