Transcription factors RUNX1 and RUNX3 in the induction and suppressive function of Foxp3+ inducible regulatory T cells

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Forkhead box P3 (FOXP3)+CD4+CD25+ inducible regulatory T (iT reg) cells play an important role in immune tolerance and homeostasis. In this study, we show that the transforming growth factor-β (TGF-β) induces the expression of the Runx-related transcription factors RUNX1 and RUNX3 in CD4+ T cells. This induction seems to be a prerequisite for the binding of RUNX1 and RUNX3 to three putative RUNX binding sites in the FOXP3 promoter. Inactivation of the gene encoding RUNX cofactor core-binding factor-β (CBFβ) in mice and small interfering RNA (siRNA)-mediated suppression of RUNX1 and RUNX3 in human T cells resulted in reduced expression of Foxp3. The in vivo conversion of naive CD4+ T cells into Foxp3+ iT reg cells was significantly decreased in adoptively transferred Cbfb−/− CD4−cre naive T cells into Rag2−/− mice. Both RUNX1 and RUNX3 siRNA silenced human T reg cells and Cbfb−/− CD4−cre mouse T reg cells showed diminished suppressive function in vitro. Circulating human CD4+ CD25high CD127− T reg cells significantly expressed higher levels of RUNX3, FOXP3, and TGF-β mRNA compared with CD4+CD25− cells. Furthermore, FOXP3 and RUNX3 were colocalized in human tonsil T reg cells. These data demonstrate Runx transcription factors as a molecular link in TGF-β–induced Foxp3 expression in iT reg cell differentiation and function.

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and can repress the FOXP3 trans-activation process directly in Th2 cells (Mantel et al., 2007). It was further demonstrated that both Th1 and Th2 transcription factors T-bet and GATA3 oppose peripheral induction of Foxp3+ T reg cells in mice through STAT1-, STAT4-, and STAT6-dependent pathways (Wei et al., 2007). Although natural T reg (nT reg) cells that differentiate in the thymus are characterized by their stable Foxp3 expression, the generation of iT reg cells specific for allergens, alloantigens, and autoantigens in the periphery has been associated with a transient Foxp3+ phenotype (Fontenot et al., 2003; Hori et al., 2003). The crucial role of TGF-β in their generation has been demonstrated.

The RUNX gene family (Runt-related transcription factor, acute myeloid leukemia [AML], core-binding factor-α [CBFα], and polypoma enhancer–binding protein–2α [PEBP2α]) contains three members, RUNX1 (AML1/CBFA2/PEBP2αA), RUNX2 (AML3/CBFA1/PEBP2αA), and RUNX3 (AML2/CBFA3/PEBP2αC). They are essential transcriptional regulators of different developmental pathways. RUNX2 is mostly important for bone development and osteoblast differentiation (Komori et al., 1997), RUNX1 plays an important role in hematopoiesis during development, and RUNX3 has important functions in thymogenesis and neurogenesis (Wang et al., 1996; Inoue et al., 2002; Levanon et al., 2002). RUNX1 and RUNX3 also work together in the establishment of lineage specification of T lymphocytes (Taniuchi et al., 2002; Egawa et al., 2007). RUNX1 is a frequent target for chromosomal translocations associated with leukemias (Look, 1997), and RUNX3 methylation and silencing is observed in various human epithelial cancers (Blyth et al., 2005).

RUNX family members share the Runt domain, which is responsible for DNA binding (Ito, 1999). The Runt domain–containing protein constitutes the α-chain partner of the heterodimeric CBF complex. RUNX proteins heterodimerize with the non–DNA-binding partner, CBFβ, which increases the affinity for DNA binding and stabilizes the complex by preventing ubiquitin–dependent degradation (Wang et al., 1993). The CBF complexes regulate the expression of cellular genes through binding to promoters or enhancer elements. The effects of the RUNX–CBFβ complex regulation are clearly cell lineage and stage specific. They include the crucial choices between cell-cycle exit and continued proliferation, as well as between cell differentiation and self-renewal (Blyth et al., 2005).

Because of the involvement of RUNX mutations in different autoimmune diseases and the known interaction with TGF-β, we investigated the impact of RUNX1 and RUNX3 on the expression of FOXP3 and subsequently on the development and function of iT reg cells. This study demonstrates that RUNX1 and RUNX3 induced by TGF-β are involved in the development and suppressive function of Foxp3+ iT reg cells.

**RESULTS**

**The role of RUNX1 and RUNX3 transcription factors in TGF-β–mediated iT reg cell generation**

To investigate the role of RUNX transcription factors in the development of iT reg cells, we cultured naive CD4+ T cells, isolated from human PBMCs, in conditions that enable the development of iT reg cells. Stimulation with anti-CD2/3/28 mAbs or TGF-β alone resulted in a minimal up-regulation of RUNX1 and RUNX3 mRNA (Fig. 1 A). In contrast, the combination of both TGF-β and anti-CD2/3/28 mAbs induced RUNX1 and RUNX3 mRNAs, as well as FOXP3 mRNA, within 48 h in naive CD4+ T cells. This result suggested further experiments to investigate whether the up-regulation of RUNX1 and RUNX3 might be a feature of iT reg cells during their development or even a prerequisite for their induction.

To test this hypothesis, RUNX1 and RUNX3 expression was knocked down in human naive CD4+ T cells by transfection of small interfering RNAs (siRNAs; Fig. 1 B). Deficiency of RUNX1 or RUNX3 resulted in markedly reduced TGF-β–mediated induction of FOXP3 mRNA in naive CD4+ T cells compared with control cells transfected with scrambled siRNA. The level of FOXP3 mRNA was further reduced when both RUNX1 and RUNX3 were knocked down in naive CD4+ T cells during their differentiation to iT reg cells (Fig. 1 B).

The influence of RUNX1 and RUNX3 on the development of other T cell subsets and their specific transcription factor expression was further investigated. Naive CD4+ T cells were cultured under Th1, Th2, T reg cell, and Th17 differentiation conditions and the mRNA expression of the predominant transcription factor for each cell type was subsequently analyzed. We observed no change in GATA3 expression in Th2 cells, T-bet expression in Th1 cells, or RORC2 mRNA expression in Th17 cells in which RUNX1 and RUNX3 were knocked down compared with control cells. On the contrary, FOXP3 mRNA was significantly decreased in RUNX1- and RUNX3-deficient T reg cells compared with control cells (Fig. 1 C).

The effect of RUNX silencing on the expression level of intracellular FOXP3 during naive CD4+ T cell differentiation to iT reg was evaluated by flow cytometry. FOXP3 was only slightly reduced after RUNX1 silencing. Transfection of siRNA for RUNX1 or RUNX3 had a stronger effect. The most striking FOXP3 reduction was observed when both RUNX1 and RUNX3 were knocked down compared with control cells. On the contrary, FOXP3 mRNA was significantly decreased in RUNX1- and RUNX3-deficient T reg cells compared with control cells (Fig. 1 D).

Similar results were obtained in total CD4+ T cells (Fig. S1 and Fig. S2). The increased impact of combined RUNX1 and RUNX3 knockdown implies that RUNX1 and RUNX3 might have redundant functions in the induction of FOXP3. In addition, the levels of IL-4, IL-5, IL-10, IL-13, and IFN-γ in control siRNA-transfected or RUNX1 and RUNX3 siRNA-transfected CD4+ T cells that were cultured with or without anti-CD2/3/28 mAb and TGF-β did not show any significant difference (Fig. S3).

To determine whether RUNX1 and RUNX3 are also expressed in human T reg cells in vivo, we isolated peripheral blood CD4+ CD127CD25+ high T reg cells and compared them with CD4+ CD127CD25+ T cells. Circulating T reg cells expressed significantly higher levels of RUNX3 mRNA compared with CD4+CD25+ cells. As expected, IL-10, TGF-β,
and FOXP3 mRNAs are also expressed in circulating T reg cells (Fig. 2A). There was no difference in RUNX1 mRNA expression between these two cell subsets. We also performed an analysis of human tonsils, which contain high numbers of FOXP3+ T reg cells (Verhagen et al., 2006). Staining of tonsil sections for FOXP3 and RUNX3 demonstrated in vivo transcription of FOXP3 and RUNX3, and the expression of FOXP3 is increased in T reg cells (Verhagen et al., 2006). We also performed a series of experiments to investigate the role of RUNX1 and RUNX3 in the induction of FOXP3 expression in T reg cells. We found that RUNX1 and RUNX3 are involved in the induction of Foxp3 in iT reg cells. (A) RUNX1, RUNX3, and FOXP3 mRNA induction in human naive CD4+ T cells after alone or combined anti-CD2/3/28 mAb and TGF-β stimulation in the presence of IL-2. Real-time PCR of human naive CD4+ T cells after 48 h of culture. Bars show the mean ± SE of three independent experiments. (B) FOXP3 mRNA induction by anti-CD2/3/28 mAb and TGF-β in human naive CD4+ T cells is reduced after siRNA-mediated RUNX1/3 knockdown. Real-time PCR of RNA from human naive CD4+ T cells transfected with RUNX1 and/or RUNX3 siRNA or with a control siRNA and cultured with anti-CD2/3/28, TGF-β and IL-2. Bars show the mean ± SD of three independent experiments. (C) FOXP3 mRNA is down-regulated in iT reg cells after siRNA-mediated knockdown of RUNX1 and RUNX3. Real-time PCR for FOXP3, T-bet, GATA3, and RORC2 from human naive CD4+ T cells, transfected with RUNX1 and RUNX3 siRNA or with scrambled siRNA (control) and cultured under iT reg, Th1, Th2, or Th17-driving conditions for 12 d. Bars show the mean ± SD of three independent experiments. (D) FOXP3 protein induction in iT reg cells is reduced after siRNA-mediated RUNX1/3 knockdown. Human naive CD4+ T cells were transfected with RUNX1 and/or RUNX3 siRNA or with a control siRNA and stimulated with anti-CD2/3/28 and TGF-β in the presence of IL-2. CD4 and intracellular FOXP3 analysis by flow cytometry after 72 h. One of three independent experiments is shown. Statistical differences were verified by the paired Student’s t test. *, P < 0.05; **, P < 0.01.
Figure 2. RUNX1 and RUNX3 expression in Foxp3+ T reg cells and in human CD4+, CD127−, CD25high cells. (A) Real-time PCR analysis of CD4+, CD127−, and CD25high T reg cells and CD4+, CD127−, and CD25− T cells isolated from human peripheral blood showed an increased expression of IL-10, TGF-β, FOXP3, and RUNX3 mRNA in CD25+ compared with CD25− cells. Bars show the mean ± SE of three independent experiments. (B) Human tonsil sections were analyzed by confocal microscopy. Tissue sections were stained for FOXP3, RUNX1, RUNX3, and DAPI or isotype controls. HEK cells RUNX1-transfected or not transfected served as additional control for RUNX1 staining. Data shown are representative from one of the three tissue samples with similar results. Bars, 5 μm. Statistical differences were verified by the paired Student’s t test. *, P < 0.05; **, P < 0.01.
coexpression of these two molecules in a subset of T reg cells, whereas there was low RUNX1 expression in all tonsil cells (Fig. 2 B).

**RUNX1 and RUNX3 bind to the FOXP3 promoter**

Transcription element search system analysis of the human FOXP3 promoter predicted 3 putative RUNX binding sites at 333, 287, and 53 bp upstream of the transcription start site (TSS). All three binding sites are conserved between human, mouse, and rat (Fig. S4). To verify the putative binding sites in the FOXP3 promoter, we transiently transfected HEK293T cells with RUNX1 and RUNX3. After the pull-down with oligonucleotides containing the wild-type binding sequences, but not mutant sequences, RUNX binding to the FOXP3 promoter oligonucleotides was detected by Western blot (Fig. 3 A). To confirm these results and test the ability of single binding site sequences to bind either RUNX1 or RUNX3, we used the promoter enzyme immun assay. Cell lysates were obtained from HEK293T cells that had been transiently transfected with RUNX1 or RUNX3 expression vectors. The biotinylated FOXP3 promoter oligonucleotides were linked to a streptavidin-coated microtiter plate, and bound RUNX1 or RUNX3 was detected by using anti-RUNX antibodies and a peroxidase-labeled secondary antibody. We showed binding of RUNX1 and RUNX3 to the mixture of all three oligonucleotides containing the binding sites, whereas there was no binding detectable when a combination of the mutated oligonucleotides was used in the assay (Fig. 3 B). Although there was a similar and high degree of binding to the −333 and −287 sites, a lower degree of binding was detected when the oligonucleotide containing the −53 site in the Foxp3 promoter was used. This effect was observed both for binding to RUNX1 and RUNX3 (Fig. 3 B). The binding to the two single binding sites at −333 and −287 was comparable to the mixture of all three oligonucleotides. Chromatin immunoprecipitation (ChIP) assay results confirmed the binding of RUNX1 and RUNX3 complexes containing CBFB to FOXP3 promoter during the differentiation of naive T cells toward T reg cells. Here, naive CD4+ T cells were cultured with IL-2, anti-CD2/3/28 mAb, and TGF-β as a Foxp3-inducing stimulation. Amplification of PCR products from the FOXP3 promoter region with the predicted RUNX binding sites showed that RUNX1, RUNX3, and CBFB were immunoprecipitated together with the FOXP3 promoter (Fig. 3 C). Negative control primer targeting open reading frame-free intergenic DNA, IGX1A did not show any significant change in site occupancy.

**Role of CBFB in the induction of Foxp3**

CBFβ, a common cofactor of all RUNX proteins, stabilizes and increases the binding of the runt domain to target DNA sequences. To target all Runx proteins that might be involved in the induction of Foxp3, we used mice in which loxP-fl anked Cbfβ alleles were inactivated in T cells through expression of a CD4-cre transgene. Retinoic acid and TGF-β synergize in the induction of Foxp3 in naive T cells (Kang et al., 2007). To investigate whether Runx-mediated induction of Foxp3 is dependent on the expression of CBFβ, naive CD4+ CD8− T cells from Cbfβfl/fl CD4-cre and control Cbfβfl/fl CD4-cre mice were simultaneously cultured with anti-CD3/28 mAbs, retinoic acid, and increasing concentrations of TGF-β. After 3 d in culture, the cells were restimulated with PMA and ionomycin and analyzed for intracellular Foxp3 and IFN-γ expression. TGF-β induced Foxp3 in Cbfβfl/fl CD4-cre cells in a dose dependent manner, and this was significantly reduced in Cbfβfl/− CD4-cre cells. Retinoic acid enhanced Foxp3 expression even in 20 pg/ml of TGF-β and more than 95% of the CD4+ T cells from Cbfβfl/− CD4-cre mice became Foxp3+ in 100 and 500 pg/ml TGF-β doses. The induction of Foxp3 was again significantly lower in Cbfβfl/− CD4-cre CD4+ T cells even in the presence of retinoic acid, demonstrating that deficiency in Runx binding to DNA affects the TGF-β induction of Foxp3 in T reg cells (Fig. 5 A). There was no difference in the induction of Foxp3 when endogenous IL-4 and IFN-γ were neutralized (Fig. S6).

**Regulation of FOXP3 promoter activity and FOXP3 protein expression by RUNX1 and RUNX3**

To investigate the effect of RUNX1 and RUNX3 binding to the RUNX binding sites in the FOXP3 promoter, we transfected human peripheral blood CD4+ T cells with a FOXP3 promoter luciferase reporter vector and RUNX1 or RUNX3 expression vectors. An increase in luciferase activity was observed only when the FOXP3 promoter (−511 to +176) luciferase construct was cotransfected with RUNX1 or RUNX3 expression vectors (Fig. 4 A). The increase in promoter activity was greater upon cotransfection of RUNX3 compared with RUNX1. Luciferase expression was abrogated when the Runx binding sites in the FOXP3 promoter (−511 to +176) luciferase construct were mutated (Fig. 4 A). In these experiments, the overexpression of RUNX1 and RUNX3 eliminated the need of TGF-β for FOXP3 promoter activation and PMA/ionomycin stimulation was sufficient.

To examine the role of each of the three RUNX binding sites for the FOXP3 promoter activity, we mutated each individually or in combination. No reduction in luciferase activity was observed when the −53 site was mutated and only a slight reduction when either the −287 or −333 site was mutated (Fig. 4 B). However, mutating the −53 site in combination with one of the other two sites led to a significant decrease in luciferase activity, with the greatest reduction observed when all three binding sites were mutated (Fig. 4 B), suggesting that the identified binding sites have redundant functions and RUNX binding to more than one site is necessary for the full activation of the FOXP3 promoter. Supporting these findings, the overexpression of RUNX1 in human primary CD4+ T cells resulted in significantly elevated levels of FOXP3 protein measured by flow cytometry after 48 h. This was achieved without any requirement for anti-CD3, anti-CD28 stimulation, or the presence of TGF-β. Although there was a trend, the transfection of CD4+ T cells with RUNX3 did not lead to statistically significant increase in FOXP3 (Fig. S5).

**Role of CBFβ in the induction of Foxp3**

CBFβ, a common cofactor of all RUNX proteins, stabilizes and increases the binding of the runt domain to target DNA sequences. To target all Runx proteins that might be involved in the induction of Foxp3, we used mice in which loxP-fl anked Cbfβ alleles were inactivated in T cells through expression of a CD4-cre transgene. Retinoic acid and TGF-β synergize in the induction of Foxp3 in naive T cells (Kang et al., 2007). To investigate whether Runx-mediated induction of Foxp3 is dependent on the expression of CBFβ, naive CD4+ CD8− T cells from Cbfβfl/fl CD4-cre and control Cbfβfl/fl CD4-cre mice were simultaneously cultured with anti-CD3/28 mAbs, retinoic acid, and increasing concentrations of TGF-β. After 3 d in culture, the cells were restimulated with PMA and ionomycin and analyzed for intracellular Foxp3 and IFN-γ expression. TGF-β induced Foxp3 in Cbfβfl/fl CD4-cre cells in a dose dependent manner, and this was significantly reduced in Cbfβfl/− CD4-cre cells. Retinoic acid enhanced Foxp3 expression even in 20 pg/ml of TGF-β and more than 95% of the CD4+ T cells from Cbfβfl/− CD4-cre mice became Foxp3+ in 100 and 500 pg/ml TGF-β doses. The induction of Foxp3 was again significantly lower in Cbfβfl/− CD4-cre CD4+ T cells even in the presence of retinoic acid, demonstrating that deficiency in Runx binding to DNA affects the TGF-β induction of Foxp3 in T reg cells (Fig. 5 A). There was no difference in the induction of Foxp3 when endogenous IL-4 and IFN-γ were neutralized (Fig. S6).
Figure 3. Binding of RUNX1 and RUNX3 proteins to the predicted binding sites in the FOXP3 promoter. (A) Mutated and wild-type oligonucleotides are shown. The predicted RUNX binding sites are accentuated (boxed and in green letters) and stars mark mutations introduced into the binding site of the control oligonucleotides. Nuclear extracts from HEK293T cells were incubated with biotinylated oligonucleotides. The precipitated oligonucleotide–transcription factor complexes were separated by SDS-PAGE and identified by Western blotting with anti-RUNX1 and anti-RUNX3 antibodies. A mixture of all three oligonucleotides with the predicted binding sites or with the inserted mutation into the predicted sites was used. Data shown are one representative of three independent experiments with similar results. (B) Promoter enzyme immunoassay using wild-type and mutated oligonucleotides within the FOXP3 promoter. Bars show mean ± SE of three independent experiments. (C) Chromatin immunoprecipitation assay results show binding of RUNX1 and RUNX3 complexes containing CBFβ to the human FOXP3 promoter in naive CD4+ T cells that were cultured with IL-2 together with anti-CD2/3/28 and TGFβ. There was no change in site occupancy in all immunoprecipitations when IGX1A negative control primers were used. The results are normalized to input and isotype control antibody. Bars show mean ± SE of three independent experiments. Statistical differences were verified by the paired Student's t test. *, P < 0.05
Figure 4. Regulation of FOXP3 promoter activity by RUNX1 and RUNX3. (A) Human primary CD4+ cells were transfected with an empty vector (pGL3 Basic), a vector containing the wild-type or mutated FOXP3 promoter region (FOXP3 -511/+176) fused to the luciferase reporter gene together with a GFP, RUNX1, or RUNX3 expression vector. Bars show the mean luciferase activity ± SE measured as arbitrary light units of three independent experiments. (B) Human primary CD4+ cells were transfected with an empty vector (pGL3 Basic), a vector containing the putative FOXP3 promoter region (FOXP3 -511/+176) fused to the luciferase reporter gene, or with a vector containing the putative FOXP3 promoter region (FOXP3 -511/+176) with single RUNX binding sites mutated (53, 287, or 333) or with the combination of two or three RUNX binding sites mutated (53, 287, or 333) fused to the luciferase reporter gene. Bars show the mean luciferase activity ± SD measured as arbitrary light units of three independent experiments.

DISCUSSION

This study demonstrates that RUNX transcription factors 1 and 3 play an important role in the generation of FOXP3+ iT reg cells by TGF-β. TGF-β mediates RUNX induction and FOXP3 is efficiently up-regulated by RUNX1 and RUNX3.
Figure 5. Diminished capacity of Cbfb-deficient CD4-cre mice T cells in the generation of Foxp3+ CD4+ T cells. (A) FACS-purified naive CD4+ CD8− T cells from CbfbF/F CD4-cre and control CbfbF/+ CD4-cre mice were activated in vitro with anti-CD3/28 mAb, 50 U/ml IL-2, ± 10 nM retinoic acid (RA), and increasing concentrations of TGF-β. After 3 d in culture, the cells were restimulated with PMA + ionomycin, and then analyzed for intracellular Foxp3 and IFN-γ expression. One of five experiments is shown. (B) Naive CD4+ T cells from Cbfb CD4-cre or control mice (harboring a Foxp3-IRES-GFP allele) were adoptively transferred into Rag-deficient mice (5 × 10⁶ cells per transfer). 6 wk later, TCRβ+ CD4+ cells from the spleen, mesenteric lymph node (MLN), and lamina propria of the small intestine (LP) were analyzed for Foxp3-GFP expression. Results from one of four CbfbF/F CD4-cre and control CbfbF/+ CD4-cre mice with same findings are shown. The data from four sets of mice is shown in C. Statistical analysis was performed with Mann-Whitney U test. *, P < 0.05 between groups.
in human CD4+ T cells. There are three putative RUNX binding sites in the proximal FOXP3 promoter. One binding site was predicted as a binding site for RUNX2. Promoter enzyme immunoassay results showed that binding of RUNX1 and RUNX3 also occurred at this site (as well as at the other two), which were initially identified as RUNX1 binding sites. This finding is not surprising because RUNX proteins bind to promoter or enhancer elements of their target genes via the runt domain, which is conserved between members of the RUNX family. The RUNX protein that actually induces the expression of FOXP3 might therefore be dependent on the availability of the specific RUNX family member at certain stages of T cell development. RUNX proteins are able to increase or inhibit transcriptional activity of their target genes depending on the cell type and the target gene (Ott et al., 2003). Mutation of only one of the three binding sites had only a little effect on the promoter activity; however, when two binding sites were mutated, the FOXP3 promoter activity dropped to a greater extent. The most striking effect was observed when all three binding sites were mutated. We therefore assume that these binding sites have partially redundant functions, but binding to at least two sites seems to be necessary for full promoter activation.

TGF-β promotes or inhibits the proliferation, differentiation, and survival of a wide array of different cells. It is also produced in activated T cells and it inhibits T cell proliferation (Kehr et al., 1986; Siegel and Massagué, 2003). It was shown that TGF-β is mandatory for the maintenance of peripheral T reg cells and their expression of Foxp3 (Marie et al., 2005; Rubtsov and Rudensky, 2007). RUNX transcription factors are targets of the TGF-β superfamily and they are involved in the TGF-β pathway. They interact directly with regulatory SMADs (Miyazawa et al., 2002; Ito and Miyazono, 2003). TGF-β can activate RUNX genes at the transcriptional level, and at the posttranscriptional level through activation or stabilization of RUNX proteins (Jin et al., 2004). It was shown that RUNX2 regulates the expression of TGF-β type I receptor (Ji et al., 2001), suggesting that other mechanisms for their function could be involved. The fusion proteins RUNX1-EVII and RUNX1-ETO block TGF-β inhibition of leukemic cell growth. RUNX3 plays an important role in TGF-β–mediated growth control in epithelial cells, as loss of RUNX3 leads to decreased sensitivity to TGF-β and hyperproliferation of the gastric mucosa (Blyth et al., 2005). The present study demonstrates that RUNX3 expression is more dominant in circulating human T reg cells and tonsl T reg cells compared with RUNX1. This could be dependent on the stage of the cells and organ from which they were isolated.

We observed that single siRNA interference of either RUNX1 or RUNX3 alone shows a slight decrease in Foxp3+ T reg cell induction, which could be caused by redundancy of these proteins. For this reason, we decided to use Cbfb+/− CD4+ T reg mice. Foxp3 induction by TGF-β is reduced in CD4+ T cells of Cbfb+/−/− CD4+ T reg mice compared with Cbfb+/+ CD4+ T reg mice. Retinoic acid is secreted by a subset of dendritic cells in the gut-associated lymphoid tissue. It inhibits the IL-6–driven induction of Th17 cells and facilitates the differentiation of naive T cells to Foxp3+ T reg cells (Mucida et al., 2007). We observed an increased number of Foxp3+ cells by retinoic acid and TGF-β compared with TGF-β treatment alone in Cbfb+/− CD4+ T reg cells. In addition, we showed a defective in vivo generation of T reg cells from Cbfb-deficient CD4+ T reg cells in Rag2−/− mice. These data in mice confirm the human data that RUNX proteins play an important role for TGF-β–dependent FOXP3 induction, as well as in the suppressive capacity of T reg cells. As an additional support for this concept, the overexpression of RUNX1 induced increased FOXP3 protein expression without any requirement of TGF-β and anti-CD3 and anti-CD28 stimulation in human primary CD4+ cells. In both human and mouse systems, reduced Foxp3 expression was associated with reduced T reg cell suppressive activity.

In a recent study, the role of Runx–Cbfβ was investigated in nT reg cell development in the thymus (Rudra et al., 2009). It was reported that Foxp3 expression in nT reg cells is unstable in the absence of Runx–Cbfβ complexes. Cbfb-deficient nT reg cells progressively lose Foxp3 upon division, and there is no evidence of increased death of Cbfb-deficient nT reg cells in that study. The experiments in Cbfb-deficient CD4+ T reg cells in mice and the knockdown experiments in humans in this study suggest that the induction of Foxp3 expression is a major contributing factor in the in vivo conversion experiment. Here, we observed that there is a twofold increased Foxp3+ iT reg cell generation in vivo. This is in the same range with previously published studies targeting different mechanisms in Foxp3 induction (Maynard et al., 2007; Sun et al., 2007). Whether the diminished capacity of Cbfb-deficient CD4+ T cells in the generation of Foxp3 may be caused by peripheral expansion of Cbfb-deficient non–T reg cells or survival problems faced by Cbfb-deficient iT reg cells after Foxp3 induction remains to be elucidated.

The involvement of RUNX proteins in autoimmune diseases has been previously suggested (Alarcón-Riquelme, 2003). A mutation in the RUNX1 binding site in the promoter of programmed cell death 1 gene (PDCD-1) has been implicated in systemic lupus erythematosus pathogenesis (Prokunina et al., 2002). Polymorphisms that alter RUNX1 binding to other genes have also been described in rheumatoid arthritis linkage at 5q31 in Japanese patients (Tokuhiro et al., 2003) and in a psoriasis linkage at 17q25 (Prokunina et al., 2002; Helms et al., 2003). RUNX3-deficient mice spontaneously develop inflammatory bowel disease and hyperplastic gastritis–like lesions (Brenner et al., 2004). These disease symptoms resemble those occurring after depletion of Foxp3-expressing T reg cells (Sakaguchi, 2004). Derepression of Th2 cytokines might also account for some of the observed disease symptoms, as it was shown that T-bet first induces Runx3 in Th1 cells and then partners with Runx3 to direct lineage-specific gene activation. Runx3/Cbfb are both required for the activation of the Ifng gene and silencing of the Il4 gene in Th1 cells (Djuric et al., 2007; Naoe et al., 2007).
Figure 6. *CbfβF/F CD4-cre* mouse cells and iT reg cells generated from human naive CD4+ T cells undergoing siRNA-mediated RUNX1 and RUNX3 knock down show a diminished suppressive activity. Experimental setup (A) and results of the mouse suppression assay (B), FACS-purified
Runx proteins also play an essential role during T lymphocyte differentiation in the thymus (Taniuchi et al., 2002). Runx1 regulates the transitions of developing thymocytes from the CD4+ CD8- double-negative stage to the CD4+ CD8+ double-positive stage and from the DP stage to the mature single-positive stage (Egawa et al., 2007). Runx1 and Runx3 deficiencies caused marked reductions in mature thymocytes and T cells of the CD4+ helper and CD8+ cytotoxic T cell lineages. In addition, inactivation of both Runx1 and Runx3 at the double-positive stages resulted in a severe blockage in the development of CD8+ mature thymocytes. These results indicate that Runx proteins have important roles at multiple stages of T cell development and in the homeostasis of mature T cells, and suggest that they may play a role in T reg cell development, which remains to be elucidated. Furthermore, it was shown that Runx1 activates IL-2 and IFN-γ gene expression in conventional CD4+ T cells by binding to their respective promoter. RUNX1 interacts physically with Foxp3 protein, and it was demonstrated that this interaction might be responsible for the suppression of IL-2 and IFN-γ production and up-regulation of T reg cell–associated molecules (Ono et al., 2007).

It has been shown that Foxp3 also influences Th17 differentiation. Specifically, Foxp3 physically interacts with RORγt, and this interaction inhibits RORγt function (Zhou et al., 2008). This relationship of RORγt and Foxp3 and probably yet unknown mechanisms might be the basis of the observation that the differentiation of Th17 cells and T reg cells is often reciprocal (Bettelli et al., 2006). Recently, data suggests that Runx1 may also be involved in regulating IL17 transcription, functioning in complex with RORγt to activate transcription (Zhang et al., 2008).

The Runx3-deficient mice develop spontaneous Th2-dominated autoimmune colitis and asthma (Brenner et al., 2004; Fainaru et al., 2005). Cbfb+/- Cd4- mice also show a spontaneous Th2 dominated disease, with increased serum IgA, IgG1, and IgE titers and lymphocyte and eosinophil infiltration of the lung (Naoe et al., 2007). All these phenotypes were previously attributed to a loss of Th2 silencing whereas our findings additionally suggest that loss of T reg function plays a role. We have shown a link between Foxp3 induction in iT reg cells and RUNX1 and RUNX3. RUNX proteins play a central role in pathways regulating cell growth and differentiation, and their interaction with the TGF-β pathway is of particular interest.

Foxp3 protein interacts not only with RUNX proteins but also with several other transcriptional partners, such as NFAT and possibly NF-κB; with histone acetyl transferases, such as TIP60; and histone deacetyl transferase (HDAC) complexes, such as HDAC7 and HDAC9 (Wu et al., 2006; Sakaguchi et al., 2008). NFAT forms a complex with AP-1 and NF-κB and regulates the expression of IL-2, IL-4, IFN-γ, and CTLA4 in conventional T cells, which leads to the activation and differentiation to effector T cells (Dolganov et al., 1996; Hu et al., 2007). The NFAT–AP-1 complex also binds to the Foxp3 promoter after TCR triggering and regulates its gene expression positively (Mantel et al., 2006). It was shown that NFAT and Smad3 cooperate to induce Foxp3 expression through its enhancer (Tone et al., 2008), but no TGF-β response element was identified in the Foxp3 gene or in the surrounding regions. The initial induction of RUNX1 and RUNX3 and the subsequent binding of these transcription factors to the Foxp3 promoter that we showed here might explain the relatively late induction of Foxp3 mRNA that peaks 24–48 h after stimulation. The interaction of Foxp3 and NFAT is dependent on their cooperative binding to DNA (Wu et al., 2006). RUNX1 alone, or together with its interacting partners p300 and CREB-binding protein, may cooperate with the NFAT transcription complex to activate the IL-2 promoter (Sakaguchi et al., 2008). Similar to this interaction, NFAT may also cooperate with RUNX1 or RUNX3 to activate Foxp3, but further studies are necessary to elaborate on this concept.

In conclusion, our findings elucidate the role of RUNX proteins in iT reg cell development and function. The induction of the transcription factors RUNX1 and RUNX3 by TGF-β and the subsequent up-regulation of Foxp3 play a role in iT reg cell generation and its suppressive capacity.

**MATERIALS AND METHODS**

**Mice.** Cbfb+/+ CD4+ and Foxp3+/- mice have previously been described (Bettelli et al., 2006; Naoe et al., 2007). Cd44.1 and Rag2−/− mice were purchased from Jackson ImmunoResearch Laboratories and Taconic, respectively. For the in vivo Foxp3 conversion assay, naive CD4+ T cells from Cbfb Cd44- or control mice (harboring a Foxp3-IRES-GFP allele) were adaptively transferred into Rag-deficient mice. 5 × 10^6 cells were used per transfer. 6 wk later, TCRβ+ CD4+ gated cells from the spleen, mesenteric lymph node (MLN), and lamina propria of the small intestine were analyzed for Foxp3-GFP expression. All analyses and experiments were performed on animals at 6–8 wk of age. Animals were housed under specific pathogen–free conditions at the animal facility of the Skirball Institute, and experiments were performed in accordance with approved protocols for the New York University Institutional Animal Care and Usage Committee.
Isolation of PBMCs, CD4+ T cells, and culture conditions. Human PBMCs were isolated by Ficoll (Bioschir) density gradient centrifugation and CD4+ T cells were then isolated using the Dynal CD4+ Isolation kit (Invitrogen) according to the manufacturer’s instructions. The purity of CD4+ T cells was initially tested by flow cytometry and was ≥95%. Cells were stimulated with the following combination of mAbs to T cell surface molecules (Meiler et al., 2008): anti-CD2 (clone 4B2 and 6G4; 0.5 μg/ml), anti-CD3 (clone OKT3; 0.5 μg/ml), and anti-CD28 mAb (clone B7G5; 0.5 μg/ml; all from Sanquin) and cultured in serum-free AIM-V medium (Life Technologies) with the addition of 1 nmol/liter IL-2 (Roche). TGF-β (R&D Systems) was used at 5 ng/ml if not stated otherwise. A combination of PMA (25 ng/ml) and ionomycin (1 μg/ml; Sigma-Aldrich) was used.

Mouse CD4+ CD127+ CD45RO− and CD4+ CD127+ CD25+ cells were purified by flow cytometry using anti-CD127, anti-CD25-PC5 (Beckman Coulter), and anti-CD4-FITC antibodies (Dako).

Mouse naive (CD62L+/44+25-) CD4+ T cells were purified by flow cytometry and activated in vitro with 5 μg/ml plate-bound anti-CD3 and 1 μg/ml soluble anti-CD28 antibodies (dBiotech) in RPMI supplemented with 10% FCS, 5 mM β-mercaptoethanol, and antibiotics. Neutralizing anti–IFN-γ and anti–IL-4 mAbs (BD) were used at 1 μg/ml concentrations when indicated.

In vitro T cell differentiation. CD4+ CD45RA+ magnetically sorted (CD45RO depletion with AutoMACS; Miltenyi Biotec) cells were stimulated with immobilized plate-bound anti-CD3 (1 μg/ml; OKT3; IgG1) and anti-CD28 (2 μg/ml). For Th1 differentiation conditions, cells were stimulated with the following: 40 ng/ml IL-2, 5 μg/ml anti–IL-4, and 25 ng/ml IL-12 (R&D Systems). For Th2 conditions, cells were stimulated with the following: 40 ng/ml IL-2, 25 ng/ml IL-4, and 5 μg/ml anti–IL-12 (R&D Systems). For Th reg cell conditions, cells were stimulated with the following: 40 ng/ml IL-2, 5 μg/ml TGF-β, 5 μg/ml anti–IL-12, 5 μg/ml anti–IL-4. For Th17 conditions, cells were stimulated with the following: 40 ng/ml IL-2, 20 ng/ml IL-6, 5 μg/ml TGF-β, 20 ng/ml IL-23 (Alexis Biochemicals Corp.), 10 ng/ml IL-1β, 5 μg/ml anti–IL-4, and 5 μg/ml anti–IL-12 were used. Proliferating cells were expanded in medium containing IL-2. The cytokine profile of these cells demonstrated that IFN-γ is the predominant cytokine in Th1 cells, IL-4 and IL-13 in Th2 cells, and IL-17 in Th17 cells (Akdí et al., 2000; Burgler et al., 2009).

Immunohistochemistry. Human tonsils were obtained from tonsillec- tomy samples of hypertrophic and obstructive tonsils without a current infection. Ethical permission was obtained from the Cantonal Ethics Commission, and informed consent was obtained from patients. Paraformaldehyde-fixed tonsil cryosections were stained with unconjugated rabbit IgG polyclonal antibody to human RUNX1 (Santa Cruz Biotechnology, Inc.) or unconjugated mouse IgG1 mAb to human RUNX3 (Abcam). After a washing step, the sections were stained with the corresponding secondary antibodies. RUNX1–binding antibodies were detected by using Alexa Fluor 633–conjugated goat anti–rabbit IgG and RUNX3–binding antibodies were detected by using Alexa Fluor 633–conjugated mouse anti–human RUNX3 (dBiotech) or the corresponding isotype control. Tissue sections were stained with DAPI for the demonstration of nuclei and mounted with Prolong antifade (Life Technologies) and three Silencer Pre-designed siRNAs for RUNX3 (Applied Biosystems) and three putative RUNX binding sites in the FOXP3 promoter region was introduced using the QuickChange kit (Stratagene), according to the manufacturer’s instructions and confirmed by sequencing the DNA. The following primers and their complementary strands were used: foxp runx-333, forward 5′-CACCCTTTCCTTTAAAACCTGTCCTTTCTCTAGCCCGTATTATCTC-3′; foxp runx-333 reverse 5′-GATAATAGGGCTCTAGGAAAGAGCAGTCTTTTAAAAACAGGTG-3′; foxp runx-287 forward 5′-CCTCTCCACCTGCTTGGAGGGAAGAAATC-3′; foxp runx-287 reverse 5′-GATTTCCTCCTCCCTCAGGACAGGTGAGAGG-3′; foxp runx-33 forward 5′-GCTTCCACACCGGTACACGGTTCTTCTCTTCTCCTGATATAAGAG-3′; foxp runx-53 reverse 5′-CTTTTATACGCAGAAGAAAAAGGACGCTGTACGGTGTGGAAGC-3′.

The human RUNX1 fragment from the Addgene plasmid 12504 (Bigs et al., 2006) pFlagCMV2-AML1B was sub-cloned in the pEGFPN1 vector (Clontech Laboratories). The RUNX1 vector pCMV human RUNX3, which was a gift from K. Ito (Institute of Molecular and Cell Biology, Proteins, Singapore) was subcloned into pEGFPN1 vector (Clontech Laboratories; Yamamura, et al., 2006).

Transfections and reporter gene assays. T cells were rested in serum-free AIM-V medium overnight. 3.5 μg of the FOXP3 promoter luciferase reporter vector or a combination together with the RUNX1, RUNX3 pEGFPN1 vector, and 0.5 μg phRL-TK were added to 3×10^6 CD4+ T cells resuspended in 100 μl of Nucleofector solution (Lonza) and electroporated using the program U-15. After a 24-h culture in serum-free conditions and stimuli as indicated in the figures, luciferase activity was measured by the dual luciferase assay system (Promega) according to the manufacturer’s instructions. PMA/ionomycin was used to stimulate the cells, because the transfection was only transient and the luciferase assay required a strong and fast stimulation of the cells. To evaluate the effect of overexpression of RUNX1 or RUNX3 on FOXP3 protein levels, CD4+ T cells were pretreated with 2 μg/ml phytohemagglutinin (Sigma-Aldrich) in serum-free AIM-V medium in the presence of 1 nmol/liter IL-2 (Roche) for 12 h, and then transfected with the vector pEGFPN1 containing the RUNX1 or RUNX3 fragment using the Nucleofector system (Amusa Biosys- tems) and the program T-23. FOXP3 expression was evaluated by flow cytometry after 48 h of culture in AIM-V medium containing 1 nmol/liter IL-2.

RNA interference. CD4+ or naive CD4+ T cells were resuspended in 100 μl of Nucleofector solution (Lonza) and electroporated with 2 μM siRNA using the Nucleofector technology program U-14 (Lonza). Five different Silencer or Silencer Select Pre-designed siRNAs for RUNX1 (Applied Bio- systems) and three Silencer Pre–designed siRNAs for RUNX3 (Applied Biosystems) were tested, and the best was selected for all further experiments.
The Silencer Negative Control #1 siRNA (Applied Biosystems) was used for normalization. Cells were then left unstimulated or were stimulated after 12 h with anti-CD2, anti-CD3, and anti-CD28. Cells were cultured in serum-free AIM-V medium with the addition of 1 nmol/L-2 (Roche). Cells were harvested for mRNA detection of the target genes after 24 h and for protein detection after 48 h.

**RNA isolation and cDNA synthesis.** RNA was isolated using the RNeasy Mini kit (Qiagen) according to the manufacturer’s protocol. Reverse transcription of human samples was performed with reverse-transcription reagents (Fermentas) with random hexamers according to the manufacturer’s protocol.

**Real-time PCR.** PCR primers and probes were designed based on the sequences reported in GenBank with the Primer Express software version 1.2 (Applied Biosystems) as follows: FOXP3 forward primer, 5’-GAAACAGC-ACATTCGCAAGTC-3’; FOXP3 reverse primer, 5’-ATGGCCCGAGGATGG-3’; EF-1a forward primer, 5’-CTGAACCATCCAGGCCAAAT-3’; and EF-1a reverse primer, 5’-GCCGTGTTGCAATCCT-3’, as previously described (Amato et al., 2007). GATA3 forward primer, 5’-CCGGGCCTCATCACAATAGGTA-3’; and GATA3 reverse primer 5’-GCTCTCTGTGGCTGACGAGC-3’ (Amato et al., 2007). T-bet forward primer, 5’-GATTGGCCAGAAGTTGTC-3’; T-bet reverse primer, 5’-GACAATTCTCATCGTGCTCAT-3’; RUNX2 forward primer, 5’-CAGTCATGAGAACACAAATTGAAGTG-3’; and RUNX2 reverse primer 5’-CAGGTGATAACCCGTAGTTGAT-3’. The prepared cDNAs were amplified using SYBR green PCR master mix (Fermentas) according to the recommendations of the manufacturer in an ABI PRISM 7000 Sequence Detection System (Applied Biosystems).

RUNX1 and RUNX3 mRNA was detected by using TaqMan Gene Expression Assays from Applied Biosystems and used according to the manufacturer’s instruction using TaqMan master mix using a 7000 real-time PCR system (Applied Biosystems). PCR amplification of the housekeeping gene encoding elongation factor (EF)-1α or by using the 18S rRNA Gene Expression Assay (Applied Biosystems) was performed to allow normalization between samples. Relative quantification and calculation of the range of confidence was performed using the comparative ∆∆CT method (Applied Biosystems). The percentage of FOXP3 mRNA in siRNA-mediated RUNX knockdown cells was calculated in relation to cells, which were transfected with scrambled control siRNA. Arbitrary units show the 2^−ΔΔCT values multiplied by 10,000 incorporating the ct values of the gene of interest and the housekeeping gene.

**Flow cytometry.** For analysis of human FOXP3 expression on the single-cell level, cells were first stained with the monoclonal CD4 mAb (Beckman Coulter), and after fixation and permeabilization, they were incubated with biotinylated double-stranded oligonucleotides containing the wild-type or mutated RUNX binding sites, and polydeoxyinosinic-deoxycytidylic acid (Sigma-Aldrich). A combination of all three oligonucleotides containing the mutated or the wild-type binding sites was used in the assay. DNA-bound proteins were collected with streptavidin-agarose beads (GE Healthcare), incubated with biotinylated double-stranded oligonucleotides containing the wild-type or mutated RUNX binding sites, and polydeoxyinosinic-deoxyoxyctydlylic acid (Sigma-Aldrich). A combination of three oligonucleotides containing the mutated or the wild-type binding sites was used in the assay. DNA-bound proteins were collected with streptavidin-agarose beads, washed with HKMG buffer, and finally resuspended in NuPAGE loading buffer (Invitrogen Life Technologies), heated to 70°C for 10 min, and separated on a NuPage 4-12% Bis-Tris gel (Invitrogen Life Technologies). The proteins were electroblotted onto a PVDF membrane (GE Healthcare) and after fixation and permeabilization, they were incubated with horseradish peroxidase-labeled mAb (Cell Signaling Technology). For analysis of human FOXP3 expression on the single-cell level, cells were cultured either with IL-2 only or with IL-2, anti-CD2/3, and anti-CD28. Cells were cultured in serum-free AIM-V medium with the addition of 1 nmol/L-2 (Roche). Cells were harvested for mRNA detection of the target genes after 24 h and for protein detection after 48 h.

**Pull-down assay.** HEK293T cells were transfected with RUNX1 or RUNX3 using the Lipofectamine 2000 reagent (Invitrogen) according to the manufacturer’s instruction. Cells were lysed by sonication in HKMG buffer (10 mM Hepes, pH 7.9, 100 mM KCl, 5 mM MgCl2, 10% glycerol, 1 mM DTT, 0.5% Nonidet P-40) containing a protease inhibitor cocktail (Roche Diagnostics). The cell lysate was preclared using streptavidin-agarose beads (GE Healthcare), incubated with biotinylated double-stranded oligonucleotides containing the wild-type or mutated RUNX binding sites, and polydeoxyinosinic-deoxyoxyctydlylic acid (Sigma-Aldrich). A combination of three oligonucleotides containing the mutated or the wild-type binding sites was used in the assay. DNA-bound proteins were collected with streptavidin-agarose beads, washed with HKMG buffer, and finally resuspended in NuPAGE loading buffer (Invitrogen Life Technologies), heated to 70°C for 10 min, and separated on a NuPage 4-12% Bis-Tris gel (Invitrogen Life Technologies). The proteins were electroblotted onto a PVDF membrane (GE Healthcare) and detected using RUNX1 or RUNX3 antibodies described in the previous section.

**Promoter enzyme immuno assay.** As performed in the pull-down assay, HEK293T cells were transfected with RUNX1 or RUNX3 and subsequently lysed. Insoluble material was removed by centrifugation. 384-well plates, precoated with streptavidin (Thermo Fisher Scientific) were washed 3 times with washing buffer (PBS and 0.05% Tween 20). Biotinylated FOXP3 promoter/oligonucleotides probes containing the RUNX binding sites were added (1 pmol per well; 50 fmol/μl) and incubated for 1 h at room temperature. Either a combination of all three oligonucleotides containing the mutated or the wild-type binding sites was used or single oligonucleotides were used in the assay. After 3 washing steps with washing buffer, the nuclear extract was added (concentration > 0.2 μg/μl) and incubated overnight at 4°C. The lysates were incubated with 10 μg of poly-deoxyinosinic-deoxyoxyctydlylic acid (Sigma-Aldrich). The plate was washed with HKMG buffer, secondary antibody (anti-rabbit IgG-HRP, 1:3,000 in HKMG buffer), Cell Signaling Technology) was added, and the plate was incubated for 1 h at 4°C. The wells were washed 4 times with HKMG buffer before adding the substrate reagent (RoD Systems). The colorimetric reaction was stopped by adding 2 M H2SO4. Absorbance at 450 nm was measured using a microplate reader (Berthold Technologies).

**ChIP.** Human naive CD4+ T cells were cultured either with IL-2 only or with IL-2, anti-CD2/3/28, and TGFB-β for 72 h, and protein–DNA complexes were fixed by cross-linking with formaldehyde in a final concentration of 1.42% for 15 min. Formaldehyde was quenched with 125 mM glycine for 5 min, and cells were subsequently harvested. The ChIP assay was performed as described in the fast chromatin immunoprecipitation method (Nelson et al., 2006). Cells were lysed with immunoprecipitation buffer (150 mM NaCl, 50 mM Tris-HCl, pH 7.5, 5 mM EDTA, NP-40 [0.5% vol/vol]) containing phosphatase (Roche) and protease inhibitors cocktails (Roche), the nuclear pellet was washed, the chromatin was sheared by sonication and incubated with antibodies for RUNX1 (H-65 X; Santa Cruz Biotechnology, Inc.), RUNX3 (H-50 X; Santa Cruz Biotechnology, Inc.), CBFBP (PEBP2β; FL-182 X; Santa Cruz Biotechnology, Inc.), and as controls normal rabbit IgG (Santa Cruz Biotechnology, Inc.), anti-human RNA polymerase II antibody, and mouse control IgG (both from SA Biosciences). The cleared chromatin was incubated with protein A agarose beads and, after several washing steps, DNA was isolated with 10% (wt/vol) Chelex 100 resin. Samples were treated with protease K at 55°C for

**HRP-labeled mAb (Cell Signaling Technology) and visualized with a LAS-1000 gel documentation system (Fujiﬁ lm). To conﬁ rm sample loading and transfer membranes were incubated in stripping buffer and reblotted for 1 h and reprobed using anti-GAPDH (6CS; Abcam) and developed using an anti–mouse IgG HRP-labeled mAb (Cell Signaling Technology).**

**Western blotting.** For human RUNX1 and RUNX3 analysis on the protein level, 106 cells were lysed and loaded next to a protein-mass ladder (Invitrogen) on a NuPAGE 4–12% Bis-Tris gel (Invitrogen). The proteins were electroblotted onto a PVDF membrane (GE Healthcare). Unspecific binding was blocked with 3% milk in TBS Tween, and the membranes were subsequently incubated with a 1:1,000 dilution of rabbit anti-RUNX1 (ab11903; Abcam) or 1:200 dilution of rabbit anti-RUNX3 (H-50; Santa Cruz Biotechnology, Inc.) in blocking buffer containing 3% milk in TBS Tween overnight at 4°C. The blots were developed using an anti-rabbit IgG HRP-labeled mAb (Cell Signaling Technology) and visualized with a LAS-1000 gel documentation system (Fujiﬁ lm). To conﬁ rm sample loading and transfer membranes were incubated in stripping buffer and reblotted for 1 h and reprobed using anti-GAPDH (6CS; Abcam) and developed using an anti–mouse IgG HRP-labeled mAb (Cell Signaling Technology).
30 min. The proteinase K was then inactivated by boiling the samples for 10 min. The purified DNA was used in a real-time PCR reaction. Specific primers for the FOXP3 promoter, spanning the region from −87 to −3, FOXP3 promoter forward primer 5′-AGAGGTCTGCGGCTTCCAC-3′, FOXP3 promoter reverse primer 5′-GGAACAGTCTGACATCAAAAA-CAA-3′, or control GAPDH primer (SA Biosciences) for the RNA poly- nase II were used. A negative control PCR for each immunoprecipitation using IGX1A negative control primer targets ORF-free intergenic DNA (SA Biosciences) was used. The fold enrichment in site occupancy was calculated incorporating IgG control values and input DNA values using the ChomptonChIP qPCR data analysis file (SA Biosciences).

Quantification of cytokine levels. IL-4, IL-5, IL-6, IL-10, IL-13, IL-17, and IFN-γ secretion was assessed using fluorescent bead-based technology. The Bio-Plex-hu Cytokine Panel, 17-Plex Group 1 was used according to the manufacturer’s instructions (Bio-Rad Laboratories). Fluorescent signals were read and analyzed using the Bio-Plex 200 System (Bio-Rad Laboratories).

Online supplemental material. Fig. S1 shows the induction of RUNX1, RUNX3, and FOXP3 mRNA in human CD4+ T cells after anti-CD2/3/28 mAb and TGF-β stimulation. Fig. S2 shows decreased RUNX1 and RUNX3 mRNA and protein expression after siRNA-mediated knockdown. Fig. S3 shows the quantification of IL-4, IL-5, IL-10, IL-13, and IFN-γ levels in control siRNA transfected or RUNX1 and RUNX3 siRNA transfected human CD4+ T cells. Fig. S4 shows the putative RUNX binding sites in the FOXP3 core promoter sequence of human, mouse, and rat. Fig. S5 shows the induction of FOXP3 protein after overexpression of RUNX1 and RUNX3 in human CD4+ T cells. Fig. S6 shows that endogenous IL-4 and IFN-γ do not effect Foxp3 expression in naive CD4+ T cells of Cby+/ Cby+ Cby+Cby control mice, which were stimulated with anti-CD3 and anti-CD28 mAbs, IL-2 and TGF-β in the absence or presence of anti-IL-4 and anti-IFN-γ neutralizing mAbs. Fig. S7 shows similar cell death (A) and proliferation (B) of Foxp3− cells and Foxp3+ cells in Cby+/ Cby+ Cby+Cby control mice cultures. Online supplemental material is available at http://www.jem.org/cgi/content/full/jem.20090596/DC1.

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