Induction of Cytotoxic T Lymphocyte Antigen 4 (CTLA-4) Restricts Clonal Expansion of Helper T Cells

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Abstract

Cytotoxic T lymphocyte antigen (CTLA)-4 plays an essential role in immunologic homeostasis. How this negative regulator of T cell activation executes its functions has remained controversial. We now provide evidence that CTLA-4 mediates a cell-intrinsic counterbalance to restrict the clonal expansion of proliferating CD4+ T cells. The regulation of CTLA-4 expression and function ensures that, after ~3 cell divisions of expansion, most progeny will succumb to either proliferative arrest or death over the ensuing three cell divisions. The quantitative precision of the counterbalance hinges on the graded, time-independent induction of CTLA-4 expression during the first three cell divisions. In contrast to the limits imposed on unpolarized cells, T helper type 1 (Th1) and Th2 effector progeny may be rescued from proliferative arrest by interleukin (IL)-12 and IL-4 signaling, respectively, allowing appropriately stimulated progeny to proceed to the stage of tissue homing. These results suggest that the cell-autonomous regulation of CTLA-4 induction may be a central checkpoint of clonal expansion of CD4+ T cells, allowing temporarily and spatially restricted growth of progeny to be dictated by the nature of the threat posed to the host.

Key words: lymphocyte • CTLA-4 • cell cycle • CD4+ • CCR7

Introduction

Proliferation and differentiation of CD4+ T cells are regulated by numerous extrinsic signaling pathways. Variables such as the amount of antigen, inflammation, costimulatory ligands, and cytokine milieu all serve to multiply the potential outcomes of an immune response (1). Despite the seemingly limitless possibilities, there are certain behaviors of activated CD4+ T cells that suggest a level of invariant, autonomous control (2, 3). One strikingly uniform feature in the activation of helper T cells is the finding that clonal expansion seems invariably accompanied by clonal contraction (4). Activation-induced cell death is generally regarded as the major mechanism for lymphocyte homeostasis during an immune response (5–7). CD4+ T cells can be induced to undergo apoptosis through a mechanism that is linked to their activation, but how the processes are coupled is not well understood. Whether nonapoptotic mechanisms exist to limit lymphocyte expansion is also uncertain (8).

The phenotype of cytotoxic T lymphocyte antigen 4 (CTLA-4) deficiency in mice (9–11) suggests that this receptor plays a central role in homeostasis. Together with other gain- and loss-of-function experiments, CTLA-4 has been implicated as a regulator of cell cycle (12–14), energy, tolerance (15–17), autoimmunity (18), transplantation (18), effector choice (19–21), and immunity to tumors (22) and foreign pathogens (23–25). One explanation for how negative regulation by CTLA-4 might be involved in many disparate immunologic reactions is that its regulation and function are coupled to the proliferative program. It has become evident that the cell cycle organizes some changes in gene expression in CD4+ T cells that are critical for effector subset choice (26–28) and effector-memory homing (29, 30). This suggests that many immunologic reactions, despite being cued by extrinsic variables, have levels of cell-autonomous control that are determined by the lineage relationship of proliferating cells (2–4).

*Abbreviations used in this paper: CCR, CC chemokine receptor; CFSE, carboxy-fluorescein diacetate succinimidyl ester; CTLA, cytotoxic T lymphocyte antigen; HPRT, hypoxanthine guanine phosphoribosyl transferase; RT, reverse transcription.
We wished to determine the precise lineage relationship between cells that were engaged in a proliferative response when well-characterized variables of activation were experimentally manipulated. Using the dilution of carboxyfluorescein diacetate succinimidyl ester (CFSE) to assess cell division (31), in conjunction with quantitation of cellularity and apoptosis, CD4+ T cells were found to have a cell-intrinsic mechanism to control their proliferation. The control mechanism can be ascribed to the graded, cell cycle–coupled regulation of the expression and function of the CTLA-4 receptor. These findings suggest that vastly different immunologic outcomes mediated by CD4+ T cells result from the integration of extrinsic stimuli with a cell-autonomous program of gene regulation that is linked to cell division.

Materials and Methods

Mice. Wild-type C57BL/6 and BALB/c mice were obtained from The Jackson Laboratory. CTLA-4−− (9), CD28−− (32), Bcl-xL (under the control of the CMV promoter and E. coli enhancer) transgenic (33), and DO11.10 TCR transgenic (34) mice were generated as described. CTLA-4−− DO11.10 TCR transgenic mice were used to delay onset of disease in CTLA-4−− animals (17, 35), and wild-type DO11.10 TCR transgenic, CD28−− DO11.10 TCR transgenic, and Bcl-xL transgenic DO11.10 TCR transgenic mice were used as their controls. All mutant mice were backcrossed to the BALB/c or B10.D2 background six times before intercross with mice from the same background. Animals were genotyped using PCR and flow cytometry. All animal work was done in accordance with guidelines of the University of Pennsylvania.

Cell Culture. In experiments with nontransgenic mice, splenocytes were depleted of CD8+ cells using magnetic beads (PerSeptive Biosystems), and stimulated (2 × 10^6 cells/ml) using soluble anti-CD3 mAb (2.0 μg/ml; BD Pharmingen) as described (26), unless indicated. Anti-CD28 mAb (BD Pharmingen), human rIL-2 (Life Sciences), murine rIL-4 (Roche), rIL-12 (BD Pharmingen), anti-IL-4 mAb (BD Pharmingen), and anti–IL-12 mAb (BD Pharmingen) were used at concentrations indicated in the figure legends. Stimulations of DO11.10 transgenic splenocytes were performed as described for wild-type cells, except without prior CD8 depletion. All cells were labeled with CFSE (Molecular Probes) as described previously (26). Briefly, cells (2 × 10^7 cells/ml) were incubated in PBS with CFSE (5 μM) for 9 min at room temperature. Labeling was quenched with an equal volume of FCS, and then cells were washed twice with HBSS supplemented with 10% FCS.

Flow Cytometry. Flow cytometric analysis was performed on fixed cells as described previously (26). Briefly, cells were washed with PBS and fixed in 4% paraformaldehyde for 11 min at room temperature. Fixed cells were stained in permeabilization buffer (PBS with 0.2% saponin, 1% FCS, and 0.1% sodium azide). Phycoerythrin-conjugated mAbs against CTLA-4, CD25, CD44, and allopurinol-conjugated mAbs against IFN-γ and IL-4 were used where indicated. For intracellular cytokine staining only, cells were restimulated with PMA (50 ng/ml; Sigma-Aldrich) and ionomycin (500 ng/ml; Sigma-Aldrich) for 4 h, with brefeldin A (2.0 μg/ml; Sigma-Aldrich) added for the final 2 h, before fixation.

Live, unfixed cells were washed in HBSS 1% FCS before staining. All antibodies were obtained from BD Pharmingen, unless indicated. Flow cytometry was performed using a Becton Dickinson FACSCalibur™ instrument and CELLQuest™ software. Throughout the article, data represent only CD4+ events, based on specific staining with fluorochrome-conjugated anti-CD4 mAbs (Caltag). Unless specified, analysis includes only live (as determined by forward and side light scatter), CD4+ events. Gates for specific staining of surface or intracellular proteins were determined using fluorochrome-conjugated species-matched and, where possible, isotype-matched, control mAbs.

Cell Death and DNA Content. Binding of PE-conjugated Annexin V (BD Pharmingen) was performed on unfixed cells according to the manufacturer's instructions. Propidium iodide (P.I.) exclusion was performed by resuspending unfixed cells in PBS with P.I. (1.0 μg/ml; Sigma-Aldrich) 5 min before flow cytometry. ToPro-3 (Molecular Probes) was used to assess DNA content in fixed, permeabilized cells. Cells were first stained with mAbs, and then washed, fixed, and incubated in PBS with saponin (0.3%), RNase A (50 μg/ml), EDTA (5 mM), and ToPro-3 (1.0 μM) for 30 min at 4°C before analysis by flow cytometry.

Reverse Transcription PCR of CC Chemokine Receptor 7 Expression. CD8-depleted, CFSE-labeled cells were stimulated for 4 d in the cytokine conditions indicated in the figure legend, and then stained with fluorochrome-conjugated anti–CD4 mAb, before sorting of live-gated, CD4+ events into separate generations by flow cytometry, using a MoFlo instrument (Cytomation). Cells were washed and total RNA was extracted using Trizol Reagent (Life Technologies) and reverse transcribed using random hexamer primers (Amersham Pharmacia Biotech), as described (36). cDNA levels were equalized by reverse transcription (RT)-PCR, using a competitive template of hypoxanthine guanine phosphoribosyl transferase (HPRT) cDNA as an internal standard, as described (36). The competitive template, because of its higher molecular weight, migrates slower than authentic HPRT cDNA during electrophoresis. PCR for CC chemokine receptor (CCR)7 was performed on equalized cDNA samples. Each reaction was performed at least three times. The following primer sets were used: CCR7 sense CTACACGCCCCCAGGACCAT, CCR7 antisense GAAGGGGAAATGAGGAAAAAG, HPRT sense GTTGGAGACGCCAGACTTTGTTG, and HPRT antisense GAGGTTAGGCTGCGGCTATAGGGCT.

Results

Limits on Clonal Expansion Are Regulated by the Cell Division Cycle. CFSE labeling was used to resolve individual cell divisions by the quantitative dilution of green fluorescence that accompanies cytokinesis (31). CD8-depleted splenocytes were stimulated with soluble anti-CD3 mAb, using endogenous accessory cells to cross-link the anti-CD3 mAb and provide a natural source of ligands for CD28 and CTLA-4. We first performed a quantitative analysis of relative cellularity as a function of cell division in an asynchronously dividing population that encompassed at least five cell divisions. By comparing the relative number of live CD4+ T cells in each successive division, we found that the dynamics of clonal expansion follow a highly reproducible pattern over a range of conditions (Fig. 1). The greatest number of cells was typically found in the popula-
tion having completed the third or fourth cell division. Although many cells continued through 1–3 more cell divisions, cellular expansion was limited not only by proliferative arrest, but apoptosis (see below). As a result, fewer than 5% of the viable cells in culture completed the sixth cell division.

Changes in mitogenic dose of anti-CD3 mAb (0.05–5.0 μg/ml), duration of culture (3–6 d), and genetic background of T cells did not alter the pattern of clonal expansion (Fig. 1 A), suggesting that this regulatory mechanism is independent of both responder frequency and time. Addition of varying concentrations of anti-CD28 mAb (0.3–10.0 μg/ml) under the same mitogenic conditions shifted the point of maximal cellularity to a later cell division (Fig. 1 A). The increase in cell expansion that resulted from costimulation, however, did not result from expansion of cells past the fifth division, and, again, fewer than 5% of cells successfully completed the sixth division.

The distribution of cell death within discrete cell divisions was examined by analyzing the binding of Annexin V to apoptotic membranes (Fig. 1 B). As expected, there was significant cell death among activated, proliferating cells. We consistently observed, however, that the majority of apoptosis occurred after the third cell division, and increased substantially in later cell divisions (Fig. 1 B). Identical results were obtained using exclusion of propidium iodide as a parameter of survival (see below). Thus, over a wide range of conditions, clonal expansion of CD4⁺ T cells is limited by a combination of cell cycle arrest and apoptosis.

*CTLA-4 Limits Clonal Expansion.* The fact that each experiment followed the same fundamental pattern was quite unexpected since the incorporation of radioactive thymidine yielded orders-of-magnitude differences between the experimental variables we employed (data not shown). Although CD28 costimulation could increase the percentage of cells that complete the fifth division before arresting and dying, there appeared to be an absolute barrier past which cells did not proceed, or if they did, underwent apoptosis. This reproducible pattern under different stimulatory conditions and at different time points suggests the possibility that clonal expansion is limited by a division-dependent counterbalance that is metered in a cell-autonomous manner.

We observed both a failure to proceed past five divisions and increasing cell death in later divisions even in cells that were deficient in FAS (data not shown), suggesting that FAS, although potentially important to subsequent cell elimination, was not required for the termination of proliferative expansion. CTLA-4⁻/⁻ mice were bred to DO11.10 TCR transgenic mice, to delay the onset of disease seen in CTLA-4-deficient animals (17, 35). Although CTLA-4⁻/⁻ cells from mice of this age may contain up to fivefold more activated cells than wild-type cells (9), we used only younger CTLA-4⁻/⁻ DO11.10 animals and excluded from analysis any mice with grossly enlarged spleens or lymph nodes. CTLA-4⁻/⁻ DO11.10 TCR transgenic cells were compared with those from DO11.10 TCR transgenic mice, and CD28⁻/⁻ DO11.10 TCR transgenic mice. CD4⁺ T cells from CD28⁻/⁻ mice had a profound proliferative defect compared with normal cells (Fig. 2 A). CD28⁻/⁻ cells did not achieve significant cellular expansion, despite many precursors undergoing at least one division, because many of the cells in culture underwent apoptosis (Fig. 2 A).

In contrast, T cells from CTLA-4⁻/⁻ mice divided at least 2–3 divisions more than wild-type cells, and did not show any evidence of reaching a clear limit of proliferative cell division (Fig. 2 A). The proliferative arrest of wild-type cells was accompanied by substantial apoptosis of cells that completed multiple rounds of division, manifest as an increase in propidium iodide uptake in the later cell divisions (Fig. 2 A). In contrast, the unchecked clonal expansion of CTLA-4⁻/⁻ cells correlated with decreased cell death in the later divisions of proliferating cells (Fig. 2 A). The apex of cellularity in CTLA-4⁻/⁻ cells occurred at even later divisions of cells in the cell division having the largest number of cells (y-axis) within each experimental condition. Top panel includes 10 experiments (with each line representing one experimental condition and time point) using varying concentrations of both anti-CD3 (0.01–2.0 μg/ml) and anti-CD28 (0.3–10.0 μg/ml) mAbs, all analyzed on day 4. Cells from both C57BL/6 and BALB/c mice were tested. Dose responses of anti-CD3 and anti-CD28 were tested at least twice. (B) CD8-depleted, CFSE-labeled splenocytes were stimulated with anti-CD3 mAb (1.0 μg/ml), anti-CD28 mAb (2.0 μg/ml), and rIL-2 (2 U/ml) for 3 d, and stained with fluorochrome-conjugated Annexin V and anti-CD4 mAb. Binding of Annexin V (% Apoptotic, y-axis) among CD4⁺ cells of indicated division number (x-axis) is depicted in bar graph (left panel), and flow cytometric data of CD4⁺ events is displayed (right panel) as cell division (x-axis) versus Annexin V binding (y-axis). Polygons gate indicates upper limit of background fluorescence. Results are representative of four separate experiments.
CD4⁺ T cells expressing a Bcl-\(x_I\) transgene had substantially reduced apoptosis compared with normal cells, but no significant enhancement of proliferation (Fig. 3 A). In contrast, we could correct the proliferative defect in CD28⁻/⁻ cells by adding rIL-2 (Fig. 3 A), however, few viable cells remained that could complete the sixth cell division. Thus, CTLA-4 antagonizes both proliferative and survival signals mediated by costimulation from CD28, which can be separately attributed to the actions of IL-2 and Bcl-\(x_I\), respectively (37). In addition, we tested the effect of a relative decrease in B7 ligands by stimulating highly purified CD4⁺ cells with immobilized anti-CD3 plus anti-CD28 and rIL-2. In the absence of antigen-presenting cells, we found that division-dependent changes in apoptosis were substantially attenuated (Fig. 3 B).

**Induction of CTLA-4 Is Coupled to the Cell Cycle.** The precise, invariant regulation of the limits on clonal cell division (Figs. 1 and 2) suggested the existence of a cell-autonomous mechanism controlling the action of CTLA-4. Kinetics of CTLA-4 induction (38, 39), together with the phenotype of CTLA-4-haplo-insufficiency (Fig. 2 B), further suggested that this counterbalance might be regulated at the level of receptor dosage. We, therefore, examined the total cellular levels of CTLA-4 in stimulated, proliferating CD4⁺ T cells. We found that the levels of CTLA-4 were low in undivided cells, and that levels increased progressively within the initial cell divisions, achieving maximal expression after ~3 cell divisions (Fig. 4 A). Addition
of anti-CD28 and rIL-2, positive regulators of CTLA-4 expression (39), did not influence the pattern of induction per cell division but did accelerate its induction in relation to time, by causing more cells to be represented in later divisions over a fixed time (data not shown).

To further discriminate between time- and division-dependent controls of expression, we used inhibitors of cell cycle and monitored CTLA-4 expression in proliferating or arrested cells cultured for the same length of time. Cells arrested in G1 using mimosine (40) had defective induction of anti-CD3 mAb and anti-CD28 mAb (2.0 μg/ml) plus human rIL-2 (5 U/ml) (right, “no APCs”). (B) In a separate experiment, splenocytes from 6-wk-old C57BL/6 mice were either CD8-depleted (left) or subjected to positive selection (right panel), human rIL-2 (50 U/ml) was added. (B) In a separate experiment, splenocytes from 6-wk-old C57BL/6 mice were either CD8-depleted (left) or subjected to positive selection (>98% purity) of CD4+ cells (right) before labeling with CFSE and stimulation with either soluble anti-CD3 mAb and anti-CD28 mAb (2.0 μg/ml) plus human rIL-2 (50 U/ml) was added. In a separate experiment, splenocytes from 6-wk-old BALB/c (from left to right) wild-type, Bcl-xL-transgenic, and CD28−/− DO11.10 TCR transgenic mice were stimulated with anti-CD3 mAb (2.0 μg/ml) plus human rIL-2 (5 U/ml) (right, “no APCs”). All results are representative of at least two separate experiments.

The division-dependent induction of CTLA-4 in naive cells, and its more rapid reiteration in immunologically experienced cells (42), suggested that gene expression might be regulated by epigenetic mechanisms of repression. To test this, we cultured cells in the presence of sodium butyrate, an inhibitor of histone deacetylation (43) that is capable of derepressing some silent genes. The graded pattern of total cellular CTLA-4 expression in the initial cell divisions was substantially increased (Fig. 4, C and D) by the addition of butyrate. Trichostatin A, a more specific inhibitor of histone deacetylation, and 5-aza-2-deoxycytidine, an inhibitor of cytosine methylation, also caused increases in the level of CTLA-4 induction within the first three cell divisions (data not shown). Thus, chromatin structure might be a rate-limiting factor in the induction of CTLA-4.

The augmentation of CTLA-4 expression with agents that modify chromatin structure and the requirement of cell cycling to induce CTLA-4 expression suggested that activation of this gene might be coupled to DNA replication. To dissect the contributions of cytoplasmic from nuclear division, cells were cultured with cytochalasin B, an inhibitor of actin polymerization (44). Cytochalasin B–treated cells were unable to undergo cytokinesis (Fig. 4 E), but, in stark contrast to cells arrested in G1 (Fig. 4 A), could be to express near-maximal levels of CTLA-4 (Fig. 4 E). When we analyzed which cells had undergone gene induction, we found that the level of CTLA-4 expression was directly linked to DNA content. Lowest levels of CTLA-4 expression (4 geometric mean-fluorescence-intensity units above background) were found in cells with 2N DNA content, and highest levels (61 geometric mean-fluorescence-intensity units above background) were found in cells with 8N DNA content (Fig. 4 E). It is unlikely this 15-fold increase is due simply to the quadrupling of alleles as, in a separate analysis, we found that levels of CD25 and CD44 increased only 4-fold while levels of CTLA-4 increased 13-fold. Thus, CTLA-4 gene expression may be progressively induced in the first three cell divisions because of a mechanism coupled to DNA replication. That such small changes in protein expression between the initial cell divisions might indeed mediate significant biological effects is further suggested by the observation that the differences in levels of CTLA-4 expression in CTLA-4+/+ and CTLA-4−/− cells (Fig. 4 F) were found to closely parallel their differences in survival and cell division (Fig. 2 B).

Growth and Maturation Signals Can Counteract the Limits on Clonal Expansion. During analysis of proliferating CTLA-4+/+ and CTLA-4−/− cells, we noted that CTLA-4 functions in activated cells to limit their size as well as their proliferation (data not shown). Comparison of cells at day 4 demonstrated that the over-representation of
Figure 4. Cell cycle–coupled induction of CTLA-4. (A–E) C57BL/6, CD8-depleted, CFSE-labeled splenocytes were stimulated with anti-CD3 (2.0 μg/ml) for 3 d. All flow cytometric plots depict only live-gated, CD4+ events. Polygonal gates are drawn around the upper limit of background staining, as illustrated in part A. (A) After culture in the absence (left) or presence (right) of mimosine (300 μM), cells were fixed, permeabilized, and stained (see Materials and Methods) with anti-CD4 mAb and either fluorochrome-conjugated hamster control mAb or hamster anti-mouse CTLA-4 mAb (lower row), before analysis of cell division (x-axis) versus control staining (top) or total cellular CTLA-4 expression (bottom) (y-axis). (B) Cells were washed and stained (without prior fixation or permeabilization) with either anti-CD25, anti-CD44, or control mAb, before analysis of cell division (x-axis) versus surface expression (y-axis) of CD25 (left panels) and CD44 (right panels). (C) Cells were washed, fixed, and either stained directly (“surface,” left panel) or permeabilized before staining (“total,” right panel) as above. (D) In the same experiment as C, a group of cells was stimulated in the presence of sodium butyrate (600 μM), to inhibit histone deacetylases, and analyzed for total cellular CTLA-4 expression (y-axis). (E) Cytochalasin B (3.5 μg/ml) was used to prevent cytokinesis in stimulated cells (left panel) before analysis of both DNA content (x-axis, right panel) and CTLA-4 expression (y-axis). (F) Splenocytes from 4 week-old BALB/c, CTLA-4−/− (filled symbols) and CTLA-4+/− (open symbols) littermate mice were stimulated and analyzed as in part A. Geometric mean fluorescence intensity of CTLA-4 expression (y-axis) for each cell generation (x-axis) is displayed. All experiments were performed at least twice.

CTLA-4−/− cells in later cell divisions was accompanied by increased cell size, as measured by forward light scatter, in comparison to control cells (Fig. 5 A). This suggested that CTLA-4 function was restricting not only proliferation but also cell growth.

Extrinsic growth factor signaling can permit nutrient utilization in T lymphocytes (45). We, therefore, used the parameters of cell division and cell size to determine whether the clonal growth arrest mediated by CTLA-4 was a general checkpoint in the maturation of CD4+ T cell immunity. Because IL-2 expression begins to be downregulated after approximately three cell divisions (26), we tested whether saturating levels of rIL-2 could alter the pattern of growth arrest. Indeed, cells cultured in 200 U/ml of rIL-2 could remain larger and divide more than those cultured in 2 U/ml of rIL-2 (Fig. 5 B). Thus, in the course of an immune response, the presence of survival signals might regulate the outcome of clonal expansion.

IL-12 (46, 47) and IL-4 (48, 49) promote growth and maturation of committed Th1 and Th2 subsets, respectively. We, therefore, examined the effects of IL-12 and IL-4 on cell division and cell size during polarized immune responses. In the presence of rIL-12, newly differentiating, IFN-γ–positive, Th1 cells were able to undergo more division and grow to substantially larger size than undifferentiated, IFN-γ–negative cells within the same culture (Fig. 5 C). In the presence of rIL-4, newly differentiating, IL-4–positive, Th2 cells were able to undergo more division and grow to substantially larger size than undifferentiated, IL-4–negative cells in the same culture. A large proportion of differentiated cells progressed past the fifth cell division, while only a small fraction of undifferentiated cells could do so (Fig. 5 C). Thus, in subsets of cells that are specifically responsive, growth signals can successfully counteract CTLA-4–mediated proliferative and growth-arrest.

In human CD4+ T cells, expression of CCR7 protein begins to be downregulated after five cell divisions (30), giving progeny that achieved greater cell division the potential to migrate from lymphoid organs to peripheral tissues (29). To determine whether the pattern of expression in relation to cell division was similar in the mouse and, further, whether this was the result of transcriptional repression, we analyzed mRNA levels of CCR7 using RT-PCR among individual sorted cell generations of CD4+ T cells that had been stimulated in Th1- and Th2-polarizing conditions (Fig. 5 D). In either cytokine milieu, the expression of CCR7 was highest in cells that had divided less than five times. After ~5 cell divisions levels of CCR7 mRNA became low-to-undetectable in both cytokine conditions (Fig. 5 D). Therefore, as is the case in human cells (30), expression of murine CCR7 in Th cells is regulated by cell division. Thus, cytokines that mediate polarization of immunity might function to ensure that subsets of effector T cells can overcome clonal arrest and selectively
achieve division numbers (Fig. 5 C) that allow their emigration from lymph nodes (2, 29).

Discussion

The present results suggest that the pattern of clonal expansion of CD4+ T cells has a remarkable, and previously uncharacterized, uniformity in response to a broad range of stimuli (Fig. 6). After nonpolarizing stimulation, CD4+ T cells clonally expand, reaching an apex of cellularity at approximately the third cell division. Less than 5% of viable cells are likely to complete the sixth cell division. This control over proliferation appears dependent on CTLA-4, and in its absence, CD4+ T cells display unbridled expansion, due to both excessive cell division and enhanced survival. Although death receptors may be the final executioner of those cells undergoing attrition (5–7), our genetic experiments indicate that CTLA-4 alone is sufficient to discriminate the divisional history of a cell and mark generations for death or arrest.

Signal transduction through CD28 is likely to be an important element in this control mechanism for at least two reasons. First, CD28-deficient T cells, despite a limited ability to divide, do not undergo clonal expansion in the presence of an intact CTLA-4 system (Fig. 2 and 3). Second, genetic experiments reveal that a deficiency of B7 ligands, which bind both receptors, can correct the autoimmune phenotype of CTLA-4 deficiency (50). Thus, the gene duplication that gave rise to this pair of receptors represents an evolutionary “yin-yang” (51), in which the locus encoding the receptor pair is wholly epistatic.

The unique regulation of CTLA-4 gene expression allows the CD28 receptor, the nondominant member of the
pair for binding B7 (52), to become engaged initially without complete interference from CTLA-4. Counterbalance from CTLA-4 is then allotted in a precisely controlled manner by the cell cycle and its ability to permit gene induction in proportion to proliferation. We are still determining whether there is actual repression in cis at the CTLA-4 locus, in trans at the locus of an activator of the CTLA-4 promoter, or some other explanation for the graded, division-dependent gene induction. Nonetheless, cell cycle–coupled regulatory and epigenetic effects might be the mechanistic explanation for what others have recently recognized as a level of cell-autonomous control in the way lymphocytes respond to extrinsic signals (2, 4).

There are surely some redundant or cooperative negative controls for CD4+ T cells since CTLA-4−/− cells do not behave as though immortalized. These are likely to be the death receptors and the in vivo limitations or niches that permit survival or growth, such as nonpolarizing and polarizing cytokines. The phenotype of cells with CTLA-4 deficiency that have been stimulated in vitro is strikingly similar to the behavior of wild-type Th1 cells cultured in IL-12 or wild-type Th2 cells cultured in IL-4. These results support a model in which programming of selective growth factor reception (46–49) is a proximate event in effector lineage commitment (53). Early in differentiation, Th1 cells become uniquely programmed for IL-12Rβ2 expression (53–55), and Th2 cells become uniquely competent in IL-4R signal transduction (56, 57). Subsets of differentiated cells might, therefore, receive selective signaling from the cytokine milieu to progress to a CCR7-negative stage and thereby mediate immunity in tissues (Fig. 6). This checkpoint, thus, sharpens polarity from cytokines, and also restricts important fate transitions in the effector and memory response (2, 29, 30).

Our results may provide insight into the elusive basis of long-lived immunologic memory, as well as the unusual auto-aggressive phenotype of CTLA-4−/− mice, wherein peripheral tissues are the primary targets of inflammation. The limitation on the number of divisions a cell undergoes, provided by CTLA-4−/− (Fig. 6), may ensure a long-lived remnant of CCR7-positive clonal progeny in central lymphoid compartments, which can be rapidly mobilized during rechallenge (58, 59). If T cells are fated to linear, terminal differentiation as they divide, a proliferative arrest of some progeny may be the only way to maintain self-renewal of clonally selected cells. The substantial increase in division number that is readily achieved by CTLA-4−/− deficient cells would, therefore, be compatible with predominantly peripheral tissue inflammation (Fig. 6).

In double-positive thymocytes, variations in signal transduction through the antigen receptor are integrated into vastly different outcomes by intrinsic rheostats, such as Grb2 (60). In the periphery, where TCR signaling is coupled to proliferation, and perhaps obligatory for survival (61–63), CTLA-4 gene induction and function appears to be an intrinsic rheostat that may be a central checkpoint of antigen-driven clonal expansion (Fig. 6). Further maturation and expansion might then become dependent on nonantigenic extrinsic signals, such as those provided by inflammatory cytokines, which can mediate avoidance of clonal arrest.

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