CD28 costimulation independence of target organ versus circulating memory antigen-specific CD4+ T cells

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T cell receptor engagement with CD28 costimulation is generally required for naive T cell activation, whereas reactivation of memory cells is less dependent on CD28 costimulation. We studied this process in chronic beryllium disease, in which the frequency of antigen-specific CD4+ T cells in the lung is large and circulating antigen-specific cells are also detectable. In the lung, a large fraction of CD4+ T cells stopped expressing CD28 mRNA and protein, and this change in phenotype correlated with lung inflammation. In the presence of concentrations of CTLA-4 Ig that inhibited the CD28-B7 interaction, beryllium-specific CD4+ T cells in lung were still able to proliferate and secrete IFN-γ in response to beryllium in culture. This functional independence of CD28 costimulation included lung CD28+ effector cells. Although lung CD4+CD28+ cells retained the ability to secrete Th1-type cytokines in response to beryllium, they showed less proliferative capacity and were more susceptible to cell death compared with CD28+ T cells. In contrast to lung cells, inhibition of the CD28-B7 interaction markedly reduced responses of beryllium-specific T cells in blood. Taken together, these findings suggest transition within memory CD4+ T cells from CD28 dependence in central memory cells to functional independence and then loss of CD28 expression in effector cells.


Introduction
The differentiation of naive T cells into memory T cells is a crucial step in the evolution of an immune response. Optimal activation of naive T cells requires two signaling events (1, 2). Signal one is mediated by T cell receptor (TCR) engagement with antigen and MHC molecule on APCs. Signal two, termed costimulation, is most frequently mediated by the engagement of the CD28 molecule on the T cell with its ligands, B7-1 or B7-2, on the APC. In the absence of costimulation, engagement of TCR on naive T cells can result in an unresponsive state of clonal anergy (3, 4), and this clonal inactivation is believed to play an important role in the maintenance of peripheral T cell tolerance (5). In contrast to naive T cells, memory T cells can proliferate and secrete cytokines after TCR engagement without CD28-mediated costimulation (6–8).

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Nonstandard abbreviations used: T cell receptor (TCR); chronic beryllium disease (CBD); bronchoalveolar lavage (BAL); staphylococcal enterotoxin B (SEB); white blood cell (WBC); myelin basic protein (MBP).
Here, we examined the role of CD28-mediated costimulation in antigen-specific T cell activation and survival. The results demonstrate an apparent evolution of independence from CD28-mediated costimulation that correlates with memory cell differentiation. Memory CD4+ T cells in blood continued to require CD28 costimulation for proliferative and cytokine responses to beryllium. In the lung, proliferation and secretion of Th1-type cytokines by effector memory cells were functionally independent of CD28 costimulation, and a proportion of these cells stopped expressing CD28. These CD4+CD28− T cells showed decreased proliferative capacity and an increased rate of apoptosis after stimulation with antigen, suggesting transition to a presenescent state.

Methods

Study population. Twelve patients with a diagnosis of CBD were enrolled in this study. The diagnosis of CBD was established using the following criteria: a history of beryllium exposure, the presence of granulomatous inflammation found as a result of a lung biopsy, and a positive proliferative response of blood and BAL T cells to BeSO4 in vitro (10, 11). A group of four sarcoidosis patients and nine healthy subjects served as negative control subjects. Informed consent was obtained from each CBD patient and control subject, and the protocol was approved by the Human Subject Institutional Review Boards at the University of Colorado Health Sciences Center and National Jewish Medical and Research Center.

Immunofluorescence staining and analysis of CD28 gene expression. PBMCs were isolated from heparinized blood by Ficoll-Hypaque (Amersham Pharmacia Biotech, Piscataway, New Jersey, USA) density-gradient separation, and BAL was performed as described previously (12, 24–26). PBMCs and BAL cells were stained with mAb's to CD4 and CD28 (both from Becton Dickinson Immunocytometry Systems, San Jose, California, USA). A subset of the BAL cells was labeled with CFSE as described (27). The lymphocyte population was identified using forward and 90-degree light-scatter patterns, and fluorescence intensity was analyzed using a FACScalibur cytometer (Becton Dickinson Immunocytometry Systems) as described previously (9).

In some experiments, CD28 gene expression was also analyzed by RT-PCR. BAL cells were stained for CD4 and CD28, and CD4+CD28+ and CD4+CD28− cells were sorted using a MoFlo cell sorter (Cytomation Inc., Fort Collins, Colorado, USA). Equal numbers of cells of each subset were isolated and immediately frozen prior to preparation of total cellular RNA and cDNA. Sterile water and cDNA from a normal subject served as negative and positive controls for the RT-PCR, respectively. PCR amplification of a CD28 fragment was accomplished with the forward primer, 5′-AAACAAGATTGAGAAAGCTGAGG-3′, and the reverse primer, 5′-GTTCTAGAAGGCTTCGAGGAGC-3′, as described (28). As a control for an equal number of cells, a TCR BV351 gene segment was also amplified as described (28). PCR products were visualized electrophoretically on agarose gels.

Cell culture and immunofluorescence analysis for intracellular cytokine expression. Proliferation assays were performed as described previously (9, 12). In brief, BAL cells (10^5 cells/well) and PBMCs (2.5 × 10^5 cells/well) were cultured in 96-well flat-bottom microtiter plates in the presence of 5 μg/ml PHA for 48 hours or 10−5 M BeSO4 with or without the addition of various concentrations of CTLA-4Ig (R&D Systems Inc., Minneapolis, Minnesota, USA) for 4 days. During the last 18 hours, the wells were pulsed with 1 μCi of [3H] thymidine, and incorporation of radioactivity was determined by β-emission spectroscopy. Proliferation assays were performed in triplicate.

For stimulation of cytokine secretion, BAL cells (2.5 × 10^5 to 5 × 10^5 cells) or PBMCs (10^6 cells) were placed in 12 × 75-mm polystyrene tubes (Fisher Scientific Co., Pittsburgh, Pennsylvania, USA) containing 1 ml of RPMI-1640 supplemented with 10% heat-inactivated human serum (Gemini Biological Products, Woodland, California, USA) and one of the following experimental conditions: medium alone, 10 ng/ml staphylococcal enterotoxin B (SEB), or 10−5 M BeSO4 with or without CTLA-4Ig. Cells were incubated for a total of 6 hours with 10 μg/ml brefeldin A added after the first hour of stimulation (9). After stimulation, cells were washed and stained with mAb’s directed against CD4, CD8, and CD28 (all from Becton Dickinson Immunocytometry Systems). Cells were then washed with PBS containing 1% BSA and placed in fixation medium (Caltag Laboratories Inc., Burlingame, California, USA) for 15 minutes at room temperature. Following washing with PBS containing 1% BSA, cells were added to permeabilization medium (Caltag Laboratories Inc.) and stained with a mAb directed against IFN-γ or IL-2 (both from Caltag Laboratories Inc.) for 30 minutes at 4°C (9). Flow-cytometric analysis was performed as described above.

The generation of the BAL T cell clones was performed as described previously (25). In brief, variable numbers of sorted CD4+ T cells and 10^6 irradiated autologous EBV-transformed lymphoblastoid cells were cultured in 96-well microtiter plates (Falcon; Becton Dickinson Labware, Franklin Lakes, New Jersey, USA) in RPMI-1640 media (BioWhitaker Inc., Walkersville, Maryland, USA) supplemented with 10% heat-inactivated human serum (Gemini Biological Products), 20 mM HEPES, 100 U/ml penicillin, 100 μg/ml streptomycin, 2 mM L-glutamine (all from Life Technologies Inc., Gaithersburg, Maryland, USA), and 10−5 M BeSO4. After 12–14 days, T cell colonies were transferred to 1-ml cultures supplemented with 20 U/ml recombinant human IL-2 (R&D Systems Inc.). The T cell clones were maintained in culture by cycles of restimulation every 2–4 weeks.

Apoptosis assay. BAL cells and BAL T cell clones from CBD patients were stimulated with either 20 U/ml of IL-2, 10 ng/ml SEB, or 10−5 M BeSO4 for 24–48 hours. Rates of apoptosis were determined by staining with annexin V and propidium iodide (both from Molecular Probes Inc., Eugene, Oregon, USA), and the cells were immediately analyzed by flow cytometry as described above.
**Results**

**CD28 expression on blood and BAL CD4+ T cells.** Since a subset of CD4+ T cells has been reported to lose CD28 expression in certain other chronic inflammatory conditions (29–32), we initially analyzed CD28 expression on blood and BAL CD4+ T cells from CBD patients. Freshly isolated PBMCs and BAL cells were obtained from CBD patients and control subjects, and the percentage of CD4+ T cells expressing CD28 was determined by immunofluorescence staining and cytofluorographic analysis. As shown in Figure 1a, nearly all CD4+ T cells from the blood of healthy control subjects express CD28 (mean ± SE, 99.8 ± 0.07%). This frequency was significantly decreased in CBD patients (92% ± 2.2%; P = 0.003). The loss of CD28 expression was even more marked in the BAL, and the percentage of CD4+ T cells expressing CD28 ranged from 21% to 94% (mean ± SE, 59% ± 6.5%; P = 0.0003, compared with PBMCs). This loss was not specific to CBD since decreases were also noted in the BAL of sarcoidosis patients (P = 0.09 compared with CBD). Analysis of the level of CD28 expression on negative cells by fluorescence intensity (Figure 1b) suggested that CD28 expression was absent, as opposed to partially downregulated. The essential complete loss of CD28 expression was confirmed by analysis of CD28 mRNA in sorted CD4+CD28+ and CD4+CD28− populations (Figure 1c).

**Correlation between loss of CD28 and degree of inflammation in the lung.** Based on the observation that the frequency of CD4+CD28− T cells in patients with rheumatoid arthritis was associated with the severity of disease (31, 33), we correlated the percentage of BAL CD4+ T cells lacking CD28 expression with total BAL white blood cell (WBC), lymphocyte and macrophage counts in patients with CBD (Figure 2). Significant positive correlations were seen for both total BAL WBC count (r² = 0.495; P = 0.012) and lymphocyte count (r² = 0.482; P = 0.015). On the other hand, no correlation was observed between the frequency of BAL CD4+CD28− T cells and total BAL macrophage count (r² = 0.031; P = 0.52). Thus, the severity of the CD4+ T cell alveolitis appears to be tied to the loss of CD28 expression on CD4+ T cells in the BAL.

**IFN-γ and IL-2 expression by BAL and peripheral blood CD4+ T cells after stimulation with BeSO₄.** Using intracellular IFN-γ expression as a measure of response (9), BAL cells and PBMCs were cultured with BeSO₄ at the optimal concentration of 10⁻⁵ M for a total of 6 hours. Previous studies have shown that PBMCs from healthy individuals and disease controls, as well as BAL cells from sarcoidosis patients, demonstrate no response to BeSO₄ in these assays, but do have the ability to be stimulated by SEB (9). In contrast, cells from CBD patients can show marked responses after the addition of BeSO₄. A representative example of intracellular cytokine expression of BAL CD4+ T cells from CBD patient 1 is shown in Figure 3a. We observed minimal background IFN-γ staining in the absence of stimulation (medium alone). Following BeSO₄ stimulation of fresh BAL cells from patient 1, a greater frequency of CD4+CD28− cells express IFN-γ after BeSO₄ exposure as compared with their CD28− counterparts (21% versus 12%) (Figure 3a). Among the 12 patients studied, the frequency of IFN-γ–producing cells in the CD4+CD28− and CD4+CD28+ subsets was 11.3% ± 3.3% and 18.9% ± 3.6% (P = 0.08), respectively (Figure 3c). Overall, only CBD patients 3 and 10 had a greater percentage of IFN-γ–producing cells in the CD4+CD28− population. On the other hand, SEB stimulated a larger number of cells to secrete IFN-γ, and the CD4+CD28− and CD4+CD28+ subsets each had about the same proportion (mean ± SEM; 51.3 ± 3.0 versus 49.2 ± 2.8) of responsive cells (Figure 3a).

Previous studies have shown that a subset of the IFN-γ–expressing cells also produce IL-2 when stimulated by BeSO₄ (9). Despite the importance of CD28 signaling...
in IL-2 production (34, 35), a significant fraction of CD4+CD28- T cells expressed IL-2, and levels appeared to be similar compared with CD28+-secreting cells (Figure 3b). Among the 10 BAL samples studied for IL-2 production, however, the percentage of CD4+CD28--BAL cells that expressed IL-2 was significantly lower compared with the CD4+CD28+ cells (4.3% ± 1.1% versus 11.8% ± 2.7%; P = 0.02) (Figure 3c), and the ratio to IFN-γ-secreting cells was also lower in this subset (0.46 ± 0.04 versus 0.68 ± 0.04; P = 0.001).

We also analyzed the CD28 phenotype of BeSO4-responsive CD4+ T cells in the peripheral blood of patients with CBD. Previous studies showed that the frequency of peripheral blood IFN-γ expressing CD4+ T cells after stimulation with BeSO4 ranged from undetectable to approximately 1 in 500 cells in CBD patients (9). For the current studies, we selected PBMC samples that proliferated in the presence of BeSO4 and had detectable cytokine-producing cells. One example is shown in Figure 4a. Following short-term culture with BeSO4, approximately 1 in 2,000 CD4+ T cells from patient 1 expressed IFN-γ. The beryllium-responsive cells were localized to the CD28+ T cell population, although this individual had a sizable percentage (5.3%) of CD4+CD28- T cells in the blood. In the example shown, 61% of the CD4+CD28- cells expressed IFN-γ after SEB stimulation compared with 10% of the CD4+CD28+ T cells, yet essentially none (detection limit less than 1:2,000) of the CD4+CD28- cells expressed IFN-γ following BeSO4 stimulation. Few, if any, IFN-γ-producing cells were detected in the CD4+CD28-population of any of the four blood samples from CBD patients (Figure 4b), and the results indicated that this population was not greatly enriched for beryllium-specific effector cells. In these four CBD patients, the frequency of CD4+CD28- T cells ranged from 1.4% to 10% (Figure 4b). Thus, it is possible that the small size of the CD28 population in blood might have prevented the detection of rare responding cells.

Functional dependence on CD28 costimulation. We used inhibitory concentrations of CTLA-4Ig to test for the
requirement of CD28-mediated costimulation when cells from CBD patients are stimulated with beryllium in culture. PBMCs from all CBD patients analyzed demonstrated a positive proliferative response (defined as a stimulation index of 2.5) in the presence of 10^{-5} M BeSO_4 (Figure 5a). For example, PBMCs from CBD patient 1 proliferated in response to BeSO_4, with peak thymidine incorporation of 65,772 ± 2173 cpm compared with background proliferation of 431 ± 165 cpm (Figure 5a). Following the addition of 30 \mu g/ml CTLA-4Ig, beryllium-induced proliferation decreased 93%. Among the seven patients studied, six demonstrated greater than 90% inhibition with this concentration of CTLA-4Ig. The one exception, patient 7, showed the lowest proliferative response, which was inhibited 35% with CTLA-4Ig.

BAL cells from five CBD patients were selected based on positive proliferative responses to 10^{-5} M BeSO_4 in culture (Figure 5b). In contrast to the effects seen in peripheral blood, CTLA-4Ig had minimal, if any, inhibitory effects on the beryllium-induced proliferative response of BAL T cells. Only cells from patient 3 showed a 37% reduction in the proliferative response following the addition of 30 \mu g/ml CTLA-4Ig (Figure 5b). In the remaining four patients, there was either no effect or enhancement of the proliferative response of BAL T cells when CTLA-4Ig was added to the culture.

Figure 5c shows the effects of varying concentrations of CTLA-4Ig on the proliferative responses of PBMCs and BAL cells. These studies again used optimal stimulatory concentrations of BeSO_4 (10^{-5} M). Proliferation of PBMCs was reproducibly inhibited with concentrations of CTLA-4Ig ranging from 0.3 to 30 \mu g/ml (Figures 5, a and c). As shown in Figure 5c, 0.3 \mu g/ml, 3 \mu g/ml, and 30 \mu g/ml CTLA-4Ig inhibited the beryllium-
induced proliferative response 69% ± 12.9%, 85% ± 4.7%, and 92% ± 5.4%, respectively. On the other hand, CTLA-4Ig had little effect on proliferation of BAL T cells, with 0% inhibition when 0.3 µg/ml of CTLA-4Ig was added to the culture conditions. Studies described below show that the BAL T cell proliferative responses occurred in both the CD4+CD28+ and CD4+CD28– populations; therefore, both populations were functionally independent of CD28 costimulation. These studies indicate that the beryllium-induced proliferative response in PBMCs occurs in a CD28-dependent manner, while in the lung, CD28-mediated costimulation is unnecessary.

We also determined what proportion of induced cytokine secretion in BAL CD4+ cells was independent of CD28 costimulation. The addition of 30 µg/ml CTLA-4Ig had no inhibitory effect on IFN-γ production induced in BAL CD4+ T cells following short-term culture with BeSO4 (Figure 6). The lack of inhibition was equally apparent for both the CD4+CD28+ and CD4+CD28– populations; therefore, both populations were functionally independent of CD28 costimulation. These findings suggest that the CD4+CD28– BAL T cells possess either a proliferative and/or a survival advantage following antigen-specific T cell activation.

Proliferative capacity of BAL CD4+ T cells in relation to CD28 expression. In preliminary studies, we noted an increased percentage of CD4+ T cells that expressed CD28 following 14 days of beryllium stimulation of BAL T cells (data not shown). These findings suggested that the CD28– BAL T cells possess either a proliferative and/or a survival advantage following antigen-specific T cell activation. Using CFSE to track T cell division (27), we compared the proliferative capacity of CD4+CD28+ and CD4+CD28– BAL T cells. After 5 days of beryllium exposure, 41% of the CD4+CD28+ T cells underwent at least one cell division compared with 29% of the CD4+CD28– T cells (Figure 7a). Prior to culture, 30% of the CD4+ T

Inhibition of cytokine secretion in stimulated PBMCs was more difficult to test because of the low frequency of antigen-specific cells in blood (9). In the peripheral blood of patient 6, the frequency of beryllium-responsive CD4+ T cells was approximately 1 in 415 (0.24%), and the addition of 30 µg/ml CTLA-4Ig decreased IFN-γ expression after BeSO4 stimulation to background levels (0.07% in the experiment shown in Figure 6). These studies therefore suggest that activation of beryllium-specific CD4+ T cells in blood, whether quantified by cytokine secretion or proliferation, is dependent on CD28 costimulation.

Figure 6
Intracellular expression of IFN-γ after BeSO4 stimulation of BAL T cells and PBMCs from CBD patients with and without the addition of CTLA-4Ig. Representative experiments are shown for the flow-cytometric analysis of BAL and blood CD4+ T cells from CBD patients 3 and 6 stimulated with 10–5 M BeSO4 and with or without 30 µg/ml CTLA-4Ig. Cells were stained for surface CD4 and CD28 and intracellular IFN-γ and then gated for CD expression. The expression of IFN-γ after BeSO4 and CTLA-4Ig exposure in the blood of patient 6 was no greater than background staining (0.06%). The numbers in the upper quadrants of each density plot are the percentages of CD4+CD28+ and CD4+CD28– cells that express IFN-γ.

Figure 7
Differences in beryllium-specific BAL CD4+ T cell proliferation between cells with and without CD28 expression. (a) A representative example of the proliferation in CFSE-labeled BAL CD4+ T cells after 5 days of culture in the presence of 10–5 M BeSO4 is shown. Proliferation is measured by a 50% decrease in fluorescence intensity of CFSE-labeled cells with each division. BAL cells were gated for CD4 expression, and the number in each quadrant of the density plot represents the percentage of CD4+ T cells. (b) Proliferation of sorted CD4+CD28+ and CD4+CD28– cells from CBD patient 3 in medium alone and after the addition of 10–5 M BeSO4 is shown. The data are expressed as the mean counts per minute ± SEM, and the stimulation index for each proliferative response is shown over the corresponding bar.
cells expressed CD28, compared with 34% 5 days after beryllium stimulation. Based on the similar frequencies of CD28+ and CD28− T cells before and after T cell stimulation, it seemed unlikely that beryllium-specific CD28− T cells lost CD28 expression during culture. In separate experiments, BAL T cells were sorted into two populations, CD4+CD28+ and CD4+CD28−, prior to culture with autologous APCs and BeSO4. A strong proliferative response was induced in both populations of T cells (representative example shown in Figure 7b). The proliferative response (stimulated minus background proliferation), however, of the CD4+CD28− cells was 58% of the level of CD4+CD28+ T cells.

Activation-induced apoptosis of BAL CD4+ T cells in relation to CD28 expression. We also studied whether the beryllium-specific CD4+CD28− T cells in lung may be more susceptible to activation-induced cell death compared with their CD28+ counterparts. Fresh BAL cells and BAL-derived T cell clones were stimulated with BeSO4 for 24 hours, and the CD4+CD28− and CD4+CD28+ subsets were stained with annexin V and propidium iodide to determine the fraction of apoptotic cells (representative experiments are shown in Figure 8). Minimal annexin V staining was seen on fresh BAL cells after culture with 20 U/ml of IL-2 for 24 hours (data not shown). After 24-hour stimulation with beryllium, 5.5% and 14% of the CD4+CD28+ and CD4+CD28− subsets, respectively, expressed annexin V (Figure 8a). Using a beryllium-specific T cell clone derived from CBD patient 12, the fraction of apoptotic cells in the CD4+CD28− and the CD4+CD28+ subsets were 2% and 4.9%, respectively, in cultures with IL-2. We observed a dramatic increase in apoptotic cells to 30% of the CD4+CD28− subset after culture with BeSO4, compared with 9.4% of the CD4+CD28− subset (Figure 8b). The increased frequency of apoptotic cells in the CD4+CD28− subset was observed in all three experiments in which it was studied and indicated the increased susceptibility of this subset to activation-induced cell death.

Discussion

We used patients with CBD to investigate the role of CD28-mediated costimulation in human memory CD4+ T cell subsets. Large numbers of beryllium-specific CD4+ T cells are compartmentalized within the lungs of these subjects (9). These cells proliferate and rapidly secrete Th1-type cytokines following recognition of beryllium bound to HLA-DR molecules on the surface of APCs (9, 12, 23, 36). Previous experiments have also shown that these cells have the phenotype of effector-memory cells, including the lack of surface CCR7 and CD62L expression (9). The current studies indicate that activation of these lung cells is independent of CD28-mediated costimulation. Indeed, a large proportion of BAL antigen-specific cells lost expression of CD28 at both the protein and mRNA levels, but continued to respond in culture. In contrast, the proliferative and cytokine responses of beryllium-specific T cells in blood were markedly inhibited by CTLA4-Ig, strongly suggesting a need for CD28-mediated costimulation. Although some CD28− cells were present in blood, few, if any, beryllium-specific cells could be demonstrated in this subset. Taken together, these studies suggest that a decreased reliance on CD28-mediated costimulation is not a universal phenomenon among all memory CD4+ T cells and that functionally distinct memory subsets may be distinguished by CD28 function and expression.

The loss of CD28 on a subset of activated CD4+ T cells has been described in several chronic inflammatory disorders, such as sarcoidosis, rheumatoid arthritis, multiple sclerosis, type 1 diabetes, and chronic infection (29–32, 37). It has also been observed in elderly individuals (38). Although most of these studies have focused on peripheral blood cells, downregulation of CD28 expression has also been observed on BAL cells from patients with sarcoidosis, a pathologically similar lung disease (37). In studies of CD28− cells in rheumatoid arthritis, the decreased surface CD28 was similarly associated with a loss of gene expression (31, 38, 39). What signals the cell to alter its CD28 gene expression is not known. The lungs in both CBD and sarcoidosis are characterized by an accumulation of activated CD4+ T cells that secrete Th1-type cytokines (9, 23, 37). The positive correlation between the loss of CD28 expression on CD4+ T cells and the magnitude of the T cell alveolitis suggests that prolonged Th1-type cytokine exposure might be the underlying mechanism. Support for this idea includes the observation that exposure of CD4+ T cells to increasing concentrations of TNF-α resulted in decreased CD28 expression and even CD28null cells after 8 weeks of continuous TNF-α exposure in culture (40). Furthermore, in the present studies, the loss of CD28 extended to cells that are not responsive to beryllium in our assay systems. Indeed, beryllium-specific cells were only a relatively small fraction of CD28− cells in the lung.
and were absent or rarely present in the blood CD28-cells. On the other hand, the blood CD28- cells were capable of secreting Th1-type cytokines following super-antigen stimulation, indicating that they are not anergic.

The loss of CD28 expression has been extensively described in CD8+ cells, especially in chronic infection and in elderly individuals that harbor clonal expansions (28, 41–45). In these studies, the loss of CD28 more clearly has been shown to be associated with differentiation to effector memory CD8+ cells. For example, in one study of functional subsets within individually expanded CD8+ clones, loss of CD28 was associated with increased perforin expression and loss of proliferative potential (28). We were unable to detect increased perforin expression in the BAL CD4+CD28- cells of CBD patients (data not shown). Studies of CD8+ clonal expansions have also clearly demonstrated that the CD28– cells derive from CD28+ cells (28), and it seems likely that the lung beryllium-specific CD28- cells in CBD patients also arise from CD28+ precursors, especially based on the mechanistic studies described above (40). Interestingly, loss of CD28 expression is apparently not a feature of CD4+ or CD8+ memory T cell differentiation in mice (P. Marrack, personal communication), thus this phenomenon has not been studied in nonhuman systems.

Similar to studies of CD8+ T cells, we also noted decreased proliferation of CD4+CD28- compared with CD4+CD28+ cells after stimulation with antigen, despite maintained effector function. Consistent with this finding, CD28-mediated costimulation lowers the anti-CD3 concentration necessary to induce T cell proliferation in vitro (34) and promotes cell cycle progression and IL-2 production, both of which are required for a sustained T cell response (2). The current studies are the first to suggest that loss of CD28 on human memory CD4+ cells is associated with susceptibility to activation-induced cell death. Our results are, however, consistent with previous studies that have implicated CD28 costimulation in enhanced T cell survival. This effect may be mediated through the enhanced production of IL-2 and increased expression of the antiapoptotic molecule, Bcl-xL (46). CD28-costimulated cells have also been shown to be less dependent on extrinsic growth factor (e.g., IL-2) for their ability to survive in culture (46). Recent studies also suggest that the CD28 signaling pathway affects the regulation of glucose metabolism (47), which also may be important in the prolonged survival of activated T cells. In a target organ characterized by persistent antigen exposure and continuous T cell activation, it is conceivable that the loss of CD28 on a metabolically active T cell limits access to nutrients and marks the differentiation of these cells into a presenescent state.

We also noted that the addition of CTLA-4Ig actually enhanced proliferation and cytokine production of beryllium-specific cells in lung in a number of different experiments. In studies quantifying intracellular cytokine expression, this enhancement occurred in both the CD28- and CD28+ populations. These results seem most likely related to the ability of CTLA-4Ig to block the interaction of CTLA-4 (induced on the T cell after activation) with B7 on the APC, an interaction that usually leads to a downregulatory signal (2, 48, 49). In preliminary experiments, CTLA-4 is upregulated following beryllium stimulation in both CD4+CD28- and CD4+CD28- BAL T cells (A. Fontenot and B. Kortzin, unpublished data). It is noteworthy that our studies show that CTLA-4 may modulate the immediate cytokine production that occurs in these reactions.

Remarkable in the current study was the continued dependence of peripheral blood CD4+ cells on CD28 costimulation for both proliferative and cytokine responses to beryllium in culture. These findings contrast with studies of one group of investigators that have studied responses to the recall antigen, tetanus toxoid, and to certain autoantigens in multiple sclerosis (29) and type 1 diabetes (50). For example, in multiple sclerosis, myelin basic protein–reactive (MBP-reactive) T cells in the blood were found to be less dependent on CD28 costimulation for T cell proliferation as compared with MBP-specific T cells from healthy control subjects (29). Similar findings with respect to glutamic acid decarboxylase 65-reactive T cells in type 1 diabetes have also been described (50). The explanation for the difference in findings is not known. Although Th2- versus Th1-type cells have been described to be more dependent on CD28 costimulation, our cells are almost all Th1 type, as in the above studies. Furthermore, although different reagents were used to test for CD28 dependence, we believe that it is unlikely that the CTLA-4Ig used in our studies could have inhibited through a mechanism other than blocking the CD28-B7 interaction (49). The different findings may relate to the recirculation of effector cells in blood in the different diseases being studied. Our previous studies have shown that the lung effector cells in CBD patients do not circulate in large numbers (9, 25). The remarkable compartmentalization of beryllium-specific cells in the lung may be related to the fact that antigen stimulation is only occurring in the lung. It is possible that differences in where antigen stimulation takes place compared with CBD, or the timing of systemic antigen stimulation in relation to sampling, is key to whether circulating antigen-specific cells are dependent on CD28 costimulation. In this regard, we have noted that patch testing with beryllium leads to a large influx of antigen-specific cells into the blood within a few weeks, and these cells more closely resemble the lung CD4+ effector cells in terms of TCR expression (26). Whether these cells might also be independent of CD28 costimulation is currently unknown.

The current results extend the paradigm that memory CD4+ T cells are relatively independent of CD28-mediated costimulation and have implications for therapies directed at the CD28-B7 interaction, for example, CTLA4-Ig or anti-B7. The results indicate that there is a transition within memory T cell subsets from CD28 dependence in circulating cells, which are mostly central memory cells, to CD28 independence in fully differentiated effector memory cells. The need to block at the level of naive and early memory populations
emphasizes that the accumulation of effector cells in the target organ may preclude therapeutic efficacy with these agents. Furthermore, the ability to target circulating memory cells may be extremely important for this therapeutic strategy to work. Finally, one may need to be cautious when using this therapy in diseases where large numbers of effector cells have accumulated in the target organ, since enhancement of the pathologic response is possible before benefit may take place.

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