ABSTRACT

Background: CD8+ T suppressor cells may play a role in immunoregulation. Recent studies have characterized this population by the lack of the CD28 molecule. These CD8+CD28– T cells differ phenotypically and functionally from CD8+CD28+ T cells. Little is known about CD8+CD28– cells in atopy. Our aim was to analyze the phenotype and functional properties of CD8+CD28– T cells in atopic and non-atopic individuals.

Methods: Peripheral blood mononuclear cells (PBMC) were obtained after density gradient centrifugation. CD8+CD28– and CD8+CD28+ T cells were isolated using immunomagnetic beads. Relative percentages of these cells and expression of several phenotypic markers were analyzed by flow cytometry. Proliferation was assessed by thymidine incorporation in isolated populations and in co-cultures with PBMC using Dermatophagoides pteronyssinus as stimulus. Cytokine synthesis was evaluated in culture supernatants by cytometric bead array.

Results: The relative percentages of CD8+CD28– T cells and their phenotypic expression in atopic and non-atopic volunteers were not significantly different. However, CD8+CD28– T cells showed greater proliferation than did CD8+CD28+ T cells when stimulated with D. pteronyssinus, although cytokine synthesis patterns were similar. CD8+CD28+ co-cultures with PBMC showed greater proliferation than CD8+CD28– T cell co-cultures, but cytokine synthesis patterns were not different.

Conclusions: Our data confirm phenotypic and functional differences between CD28+ and CD28– T cells, irrespective of atopic status. Purified human CD8+CD28– T cells, freshly isolated from peripheral blood, do not have suppressor properties on allergen-specific proliferation or on cytokine synthesis in PBMC.

Key words: Allergy; Atopy; CD8+CD28– T cells; Human.

INTRODUCTION

T cells that use non-cytolytic mechanisms to downregulate the immune response (suppressor T cells or Ts) are thought to play an essential role in controlling reactivity to foreign antigens and inducing tolerance to self antigens1. Nevertheless, more than three decades since their discovery, an understanding of the mechanisms whereby suppressor cells exert their activity is still incomplete.
Phenotypically, CD8+ T cells are characterized by the lack of the CD28 co-stimulatory molecule. CD8+CD28− T cells are thought to arise from CD8+CD28+ T cells that have proliferated several times since they have shorter telomeres and share oligoclonal expansions. In vitro, CD8+CD28− T cells arise from CD8+CD28+ T cells repeatedly stimulated in the presence of interleukin (IL)-2. In contrast, IL-4 can block this differentiation.

Recent evidence has been accumulating to show that CD8+CD28− T cells can inhibit T helper cell activation and proliferation in mitogen and antigen-driven responses. In vitro, antigen-specific CD8+CD28− T suppressor cells have been generated by multiple rounds of stimulation of human CD8+CD28− T cells with APC either from an allogeneic or a xenogeneic donor. The generated cells expressed Foxp3, a gene related with regulatory function, and could inhibit proliferation of CD4+ T cells interacting directly with the APC used for priming.

Non-antigen specific CD8+CD28− T cells with suppressor activity have also been generated from CD8+CD28− T cells in the presence of IL-2 and IL-10. These suppressor cells inhibited both antigen-specific CD4+ T cell proliferation and cellular cytotoxicity by secreting cytokines such as IFN-γ, IL-6 and IL-10. Defects in this antigen-nonspecific suppression have been described in multiple sclerosis and in systemic lupus erythematosus patients, and are primarily seen in chronic progressive situations.

Atopic allergic diseases are immune disorders caused by abnormal, Th2 cell-dominated responses to otherwise innocuous substances, such as proteins from house dust mite or grass or tree pollen. Many authors believe that atopy and allergic diseases are the result of inadequate or impaired inhibition of allergen-specific T helper-type responses by the regulatory T cells, or even a consequence of decreased frequencies of those cells in atopic individuals. There is a growing body of evidence which suggests that CD8+ T cells play an important part in regulating the IgE response to non-replicating antigens. Nonetheless, the presence of CD8+ suppressor T cells in human atopic patients is not yet confirmed.

**MATERIALS AND METHODS**

**Subjects**

Peripheral blood was obtained from 30 non atopic and 32 atopic (with allergic rhinitis) adult volunteers (Caucasian, non-smokers), matched for age and gender. Atopic volunteers were recruited from the allergy clinic of the Cova da Beira Hospital and non atopic volunteers were recruited among the staff of the Hospital and the University.

Atopy was assessed by positive skin prick tests and specific IgE levels to Dermatophagoides pteronyssinus (Der p). Volunteers who received immunotherapy or were on systemic medication were excluded. Pregnant or breastfeeding women and all volunteers with disease affecting the immune system were also excluded. Informed written consent was signed by all the volunteers. The study was approved by the Ethics Committee of Centro Hospitalar Cova da Beira.

**Monoclonal antibodies**

The following monoclonal antibodies (mAbs) were used: anti-CD3 mAb conjugated with APC was purchased from Pharmingen (San Diego, CA, USA), anti-CD8 mAb conjugated with PerCP was purchased from Becton Dickinson (San José, CA, USA), anti-CD28 mAb conjugated with FITC was purchased from Pharmingen (San Diego, CA, USA), anti-CD25, -CTLA-4, -HLA-DR, -CD56, -CD94, -CD158a, -CD161, -NK1B1 mAb conjugated with PE were all purchased from Pharmingen, anti-VA24 mAb conjugated with PE was purchased from Serotec (Oxford, UK). Isotype-matched, PE-conjugated control mAbs IgG1, IgG2a, and IgM were purchased from Pharmingen.

**Blood preparation and antibody staining**

Freshly collected peripheral blood mononuclear cells (PBMC) were stained either from whole blood after lysis of the erythrocytes (10 mM Tris, 0.15 M NaCl, pH = 7.4). Staining was performed at 4 °C for 30 min in staining solution (PBS, 0.2 % bovine serum albumin (BSA), 0.1 % NaN3) in round-bottomed microtiter plates (Greiner, Nürtingen, Germany) with ≈ 0.5 x 106 cells/well. After staining, the cells were washed, resuspended in 500 μl PBS and acquired in a FACSCalibur Flow cytometer (Becton Dickinson).

**Flow cytometry analysis**

Data were collected on 20,000 cells/sample using FACSCalibur flow cytometer equipped with an argon ion laser and a red diode laser, for quantification of CD8+ and CD28+ T cells. Anti-CD8, -CD3, and -CD28 monoclonal mAb were used to define the CD3+CD8+CD28+ and CD8+ T cell populations. T lymphocytes were gated for analysis based on light scatter.
tering properties and on CD3 staining. Positively and negatively stained populations were calculated by quadrant dot plot analysis determined by isotype controls.

For phenotypic analysis of the defined subpopulations, 30,000 events were collected per sample. Fluorescence dot plots and histograms were analysed using cytological software (Cell Quest Pro, Becton Dickinson). Phenotypic analysis of the TCRαβ and TCRγδ was also performed on the isolated fractions of CD3+CD8+ and CD28+ cells.

Cell isolation

Peripheral blood mononuclear cells (PBMC) from the volunteers were separated from peripheral blood (120 ml) by centrifugation over Lymphoprep. Lymphocytes were further purified by allowing the cells to adhere to plastic for 1 h at 37 °C, 5 % CO₂, CD8⁺ T cells were isolated from the supernatant by negative selection using magnetic beads (Miltenyi Biotec, Bergisch Gladbach, Germany). To separate CD28⁺ and CD28⁻ T cells from CD8⁺ cell suspensions, cells were incubated with CD28-FITC mAb (Pharmingen) and then with anti-FITC microbeads (Miltenyi Biotec). Purity of the fractions was greater than 85 % as evaluated by flow cytometry analysis. The adherent fraction was collected as well and used as APC in cell cultures after mitomycin treatment.

Cell Cultures

Responder cells were cultured at a concentration of 0.5 × 10⁶ cells/ml for 6 days in RPMI 1640 complete medium (RPMI 1640, 2 mM L-glutamine, 1 % antibiotic/antimicotic (Sigma-Aldrich, St Louis, MO, USA), and 10 % fetal calf serum (Biochrom, Berlin, Germany)) in 96-well flat-bottomed plates (Nunc, Roskilde, Denmark) with or without APC and 10 μg/ml purified Der p extract (a kind gift from Dr. Joost Van Neerven, The Netherlands). For positive control, cells were incubated with 0.5 μg/ml OKT3 (eBiosciences, San Diego USA) or with 5 μg PHA (Sigma-Aldrich) for 3 days in the same conditions as for Der p. Fourteen hours before the end of the culture [³H] thymidine was added to the wells (1 μCi/well). [³H] thymidine (Amersham Biosciences, Uppsala, Sweden) incorporation was determined by scintillation counting in a TopCount (Perkin Elmer, MA, USA) and results were expressed as counts per minute (cpm). Mean cpm of the triplicate cultures and standard error of the mean were calculated.

Culture conditions were as follows: CD8⁺CD28⁺ cells alone, CD8⁺CD28⁻ cells alone, CD8⁺CD28⁺ cells + APC + Der p, and CD8⁺CD28⁻ cells + APC + Der p.

Cell co-cultures

PBMC (0.5 × 10⁶ cells/ml) were incubated in different proportions (1:1, 2:1, 4:1) with freshly isolated CD8⁺CD28⁻ and CD8⁺CD28⁺ T cells for 6 days in 96-well flat-bottomed plates with and without 10 μg/ml Der p extract. Tritiated thymidine (1 μCi/well) was added 14 hours prior the end of the culture. Cells were harvested on fiber filters, and incorporated thymidine was determined by scintillation counting.

Cytokine production

For soluble cytokine detection in the culture supernatants, the human Th1/Th2 cytokine bead array kit was used, according to the manufacturer’s instructions (Becton Dickinson).

Statistical analysis

Statistical analysis of the results was performed using Minitab 14 statistical Software. Mann Whitney U test was used to assess differences between atopic and non atopic volunteers. Wilcoxon signed rank test was used to assess differences within the same population. A p value of less than 0.05 was considered significant.

RESULTS

First, we evaluated the frequency of CD8⁺CD28⁻ T cells in whole blood from 22 atopic and 18 non atopic volunteers (age 22-42) by flow cytometry. We observed that the relative percentages were not significantly different (p = 0.80), with 41 ± 14.3 and 40 ± 13.4 % of CD8⁺CD28⁻ T cells in atopic and non atopic volunteers respectively. CD8⁺ T cells were larger, morphologically more complex and expressed CD3 with higher mean fluorescence intensity (MFI) than CD8⁻ T cells (1048 vs 978).

We then evaluated the expression of several phenotypic markers on CD3⁺CD8⁺ and CD28⁻ PBMC subpopulations in whole blood from ten atopic and ten non atopic volunteers (fig. 1). Phenotypic expression was similar in both of these groups. However, taking all the data together, more CD8⁺CD28⁻ T
cells than CD8+CD28+ T cells expressed CD25 (5% vs. 1.7%) and CTLA-4 (5.5% vs. 4%). The reverse was true for HLA-DR expression (7.7% vs. 29.7%).

Next, we examined the expression of TCR\^\text/H9251/H9252, TCR\^\text/H9253/H9254 and TCRV\^24 in CD8+CD28+ and CD28− T cells. As expected, the majority of CD8+ T cells expressed a TCR\^\text/H9251/H9252 (87% of CD8+CD28+ and 83% of CD8+CD28− T cells). Interestingly, when the expression of TCR\^\text/H9253/H9254 was examined, we found a higher percentage of CD28− T cells than CD28+ T cells expressing that receptor (11% vs. 4%).

Expression of chemokine receptors was also different, with the CXCR3 relative percentage being higher on CD28+ than on CD28− cells (69.6% vs. 49.1%). However, no difference was observed in terms of CCR5 expression between these two cell subsets. On the other hand, a higher percentage of CD28− than CD28+ T cells expressed the CCR4 chemokine receptor (21.3% vs. 18.4%).

In order to assess the proliferative response to allergen of CD8+CD28+ and CD28− T cells, we set up cultures of isolated fractions with and without APC and Der p. As shown in figure 2, contrary to what occurred with unstimulated cells, CD8+CD28− T cells proliferated when stimulated with Der p, displaying a significantly higher level (p = 0.001) of proliferation as compared with CD8+CD28+ T cells in atopic and non atopic volunteers. However, the proliferation level was not significantly different between atopic and non atopic volunteers. We also stimulated some fractions with OKT3 and, in this case, CD8+CD28− T cells proliferated less significantly (p = 0.003) than CD8+CD28+ T cells (data not shown).
T cells synthesise IL-10, thus mediating T cells in allergy. Previous studies on the functional properties of the CD8+CD28+ T cells and their implication in the development of atopy.

We observed that CD8+CD28+ T cells are not phenotypically different between atopic and non atopic individuals. This may imply that atopy is not associated with a specific CD8+ T cell phenotype. However, it is important to mention that our data on phenotypic differences between CD28+ and CD28– subpopulations is in line with other reports in health and disease25,26. Moreover, the increased expression of the NK cell related receptors in CD8+CD28+ T cells, in a context where co-stimulation is not present, may be important towards limiting T cell cytolytic responses, and act as a form of “regulation”.

Classical antigen-presentation studies showed that MHC class I molecules present peptides derived from proteins synthesized within the cell, whereas MHC class II molecules present exogenous proteins captured from the environment. Emerging evidence indicates, however, that dendritic cells have a specialized capacity to process exogenous antigens into the MHC class I pathway25. According to our expectations, unstimulated cells did not proliferate. Interestingly enough, CD8+CD28– but not CD8+CD28+ T cells proliferated in response to Der p, in the presence of APCs. This fact suggests that CD28+ T cell proliferation is not impaired in spite of the absence of CD28 co-stimulation26 and that CD8+CD28+ T cells respond to common aeroallergens. This results should be considered with caution as the two subsets of CD8+ T cells studied here are oligoclonal with re- spective TCR, and the use of this allergen might stimulate an insignificant minority of clones and lead to inconclusive results. For this reason, we performed cell cultures for 3 days with OKT3 to confirm whether multi-clonal cellular responses were different in both subpopulations. Results show a proliferation impairment in the CD8+CD28+ T cells which corroborates previous results26. This implies that our results must be confirmed by performing the same studies with another allergen.

When cytokine production was analysed, no detectable IL-5, IL-4 or IL-2 production was induced by the allergen in CD28+ or CD28– T cells. On the other hand, IFN-γ, TNFα and IL-10 were synthesized at similar levels by both isolated populations. Other investigators mention that a subpopulation of suppressor CD8+CD28+ T cells synthesises IL-10, thus mediating the suppressive effect14. Moreover, Seneviratne et al. linked low levels of IL-10 with severe atopic disease15. Since both our subsets produced IL-10, we performed the co-culture studies in order to evalu-
ate the “regulatory” capacity of these cells. The pro-
duction of cytokines varied between donors, but fol-
lowed similar patterns. The failure to detect the oth-
er cytokines (IL-5, IL-4, IL-2) in the supernatants does
not necessarily imply that they are not synthesized
but may also suggest that they are immediately used
as autocrine factors.

Finally, we performed co-culture studies and obser-
vanced that freshly isolated CD8\(^{+}\)CD28\(^{-}\) or CD8\(^{+}\)CD28\(^{-}\)
T cells did not show suppressive properties, and were
not able to inhibit allergen-driven PBMC proliferation or
cytokine synthesis. These results are in line with previ-
ous reports namely by Suciu-Foca and Filaci groups\(^3\)
where suppressor CD8\(^{+}\)CD28\(^{-}\)T cells were only gener-

Figure 3.—Cytokine synthesis is similar for both isolated populations. Isolated CD8\(^{+}\)CD28\(^{-}\) and CD8\(^{+}\)CD28\(^{+}\) T cells from 4 atopic and 3 non
atopic volunteers were incubated in 96-well flat-bottomed plates with and without 10 \(\mu\)g/ml Der p extract. On day 3, culture supernatants
were collected and cytokine synthesis (IFN-
\(\gamma\), TNF-
\(\alpha\), IL-10, IL-5, IL-4, IL-2) was evaluated by cytometric bead array, following the manu-
facturer’s protocol. Values shown are mean ± sem. Mann-Whiney U test was used for comparison between atopic and non atopic and
Wilcoxon signed rank test was used for comparison between conditions.

Figure 4.—CD8\(^{+}\)CD28\(^{-}\) co-cultures with PBMC proliferate more than CD8\(^{+}\)CD28\(^{+}\) co-cultures. Isolated CD8\(^{+}\)CD28\(^{-}\) and CD8\(^{+}\)CD28\(^{+}\) T cells
from 8 atopic and 6 non atopic volunteers were incubated in different proportions with PBMC for 6 days in 96-well flat-bottomed plates with
and without 10 \(\mu\)g/ml Der p extract. Tritiated thymidine (\(\mu\)Ci/well) was added 14 h prior to the end of the culture. Cells were harvested
on fiber filters, and incorporated thymidine was determined by scintillation counting. Results show thymidine incorporation (cpm,
mean ± s.e.m.). Wilcoxon signed rank test was used for comparison between conditions.
ated after multiple rounds of stimulation of PBMCs with allogeneic 13, xenogeneic 8 or antigen-pulsed 32 autologous APC.

However the presence of CD8⁺CD28⁻ suppressor T cells in vivo was observed in transplanted patients without rejection 33. These facts may imply a need for an elevated and sustained contact with the antigen (hence the multiple rounds of stimulation) in order to generate antigen-specific suppressor cells. It would be interesting to further study suppressive properties of CD8⁺CD28⁻ T cells, by using allergen-specific T cell lines developed from atopic and non atopic volunteers.

In summary, in the present study we have shown that atopy is not associated with altered relative percentages or specific phenotypes in CD8⁺CD28⁺ or CD28⁻human T cells. Freshly immunomagnetically isolated CD8⁺CD28⁺ or CD28⁻human T cells have distinct phenotypes and, although sharing similar cytokine production patterns, they proliferate at different levels to common stimuli. Both subpopulations show similar proliferation capacity in atopic and non atopic individuals and do not have any suppressor capacity.

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Figure 5.—Cytokine synthesis is similar in all the co-cultures stimulated with Der p. Isolated CD8⁺CD28⁺ and CD8⁺CD28⁻ T cells from 4 atopic and 3 non atopic volunteers were incubated in a proportion of 1:1 with PBMC in 96-well flat-bottomed plates with and without 10 μg/ml Der p extract. On day 3 and day 5, culture supernatants were collected and cytokine synthesis (IFN-γ, TNF-α, IL-10, IL-5, IL-4, IL-2) was evaluated by cytometric bead array, following the manufacturer’s protocol. Values shown are mean ± s.e.m. Mann-Whiney U test was used for comparison between atopic and non atopic volunteers and Wilcoxon signed rank test was used for comparison between conditions.


