Differential Activation of CD8+ Tumor-Specific Tc1 and Tc2 Cells by an IL-10-Producing Murine Plasmacytoma

CHRISTOPH SPECHT*, HANS-GERD PAUELS, CHRISTIAN BECKER and ECKEHART KOLSCH

Institute for Immunology, University of Münster, Domagkstr. 3A, D-48129 Münster, Germany

The involvement of counteractive CD8+ T-cell subsets during tumor-specific immune responses was analyzed in a syngeneic murine plasmacytoma model. CD8+ Tc cells against the immunogenic IL-10-producing BALB/c plasmacytoma ADJ-PC-5 can be easily induced by immunization of BALB/c mice with X-irradiated ADJ-PC-5 tumor cells in vivo and in vitro. However, the failure of recipient mice to mount a protective Tc response against the tumor during early stages of a real or simulated tumor growth is not due to immunological ignorance, but depends on the induction of tumor-specific tolerance, involving a population of tumor-induced CD8+ T cells that are able to inhibit the generation of tumor-specific Tc cells in a primary ADJ-PC-5-specific MLTC, using IFN-γ as a suppressive factor. Whereas most long-term cultivated CD8+ ADJ-PC-5-specific Tc lines produce type-1 cytokines on stimulation, at least two of them, which were derived from a primary MLTC, display a type-2 cytokine spectrum. Furthermore, the primary in vivo Tc response against ADJ-PC-5 cells shows characteristics of a Tc2 response. The Tc response is strictly depending on tumor-derived IL-10. CD8+ Tc cells that are induced in a primary MLTC do not produce IFN-γ, and the tumor-specific Tc response is enhanced by IL-4 but suppressed by IFN-γ or IL-12. In contrast, ADJ-PC-5-specific CD8+ Tc cells from immunized mice are IFN-γ-producing Tc1 cells. Since the primary in vitro Tc response against the tumor is suppressed even by the smallest numbers of irradiated ADJ-PC-5-specific Tc1 cells via IFN-γ, these Tc1 cells behave similar to the suppressive CD8+ T cells that are induced during early stages of ADJ-PC-5 tumorigenesis.

Keywords: CD8+ T cells, cytokine profiles, IFN-γ, plasmacytoma, T-cell subsets, tumor immunology

INTRODUCTION

In most cases, tumors grow in their primary hosts or syngeneic recipients without being hampered by the immune system, even if the tumor cells express immunogenic neoantigens. The lack of apparent immunogeneity of tumors in situ might be due to special properties of the tumor cells, for example lack of costimulatory molecules, downregulation of MHC molecules, or production of immunosuppressive
factors (Browning and Bodmer, 1992; Chen et al., 1992; Sulizeanu, 1993), or due to intrinsic tolerance mechanisms of the immune system (Mengersen et al., 1975; Naor, 1979; North, 1985). Under special experimental conditions, however, it is possible to mount protective immune responses against tumor cells, involving specific Tc cells or nonspecific macrophages and NK cells. Activation of Tc cells against tumor-specific transplantation antigens is necessary for any kind of vaccinization strategy or for adoptive immunotherapy against cancer. Thus, in the context of such therapies, it is necessary to analyze phenotypes and activation modalities of tumor-specific Tc cells and their fate in situ, in the presence of a growing tumor. Using the highly malignant BALB/c plasmacytoma ADJ-PC-5 as a model tumor, we studied the regulation of tumor-specific Tc responses, especially during early stages of tumorigenesis. Syngeneic BALB/c mice can be protectively immunized against this tumor by four weekly i.p. injections with $10^7$ X-irradiated ADJ-PC-5 cells (Cihak et al., 1981), which leads to the activation of tumor-specific CD8$^+$ Tc cells. Specific CD8$^+$ Tc cells against the tumor can also be induced in vitro, in a primary syngeneic mixed lymphocyte tumor cell culture (MLTC) (Haubeck and Kölsch, 1982). However, the growth of ADJ-PC-5 tumor cells in syngeneic BALB/c mice is accompanied by the induction of a subset of CD8$^+$ regulatory T cells, which are characterized by their ability to suppress the induction of tumor-specific CD8$^+$ Tc cells in a primary MLTC in vitro (Haubeck and Kölsch, 1982, 1986). These suppressive T cells are induced during early stages of tumorigenesis, when the tumor burden is small, and possibly hold down a protective cytotoxic T-cell (Tc) response against the tumor, even when the tumor mass reaches an immunogenic level. Recent data indicate that the T-cell-mediated suppressive effect in vitro involves IFN-$\gamma$ as a suppressor factor, but does not depend on IL-2 consumption (Pauels et al., 1996). The finding that IFN-$\gamma$ is a suppressive factor for Tc induction in vitro prompted us to study phenotypes and activation modalities of ADJ-PC-5-specific Tc cells in more detail.

RESULTS

Simulation of ADJ-PC-5 Tumor Growth in the Peritoneal Cavity is Accompanied by the Induction of IFN-$\gamma$-Producing Suppressive CD8$^+$ T Cells

In previous studies, an in vitro protocol for the induction of ADJ-PC-5-specific Tc cells was developed, which is based on a primary syngeneic MLTC (Haubeck and Kölsch, 1982). As shown in Figure 1(a), tumor-induced suppressive T cells can be isolated from the peritoneal cavity of syngeneic BALB/c mice that have been subjected to a simulated ADJ-PC-5 tumor growth by daily i.p. injections of exponentially increasing numbers of 60-Gy X-irradiated ADJ-PC-5 cells. These T cells are characterized by their ability to suppress the induction of a tumor-specific Tc response from naive BALB/c spleen cells in a primary syngeneic ADJ-PC-5-specific MLTC in vitro. The respective suppressive T cells have been shown previously to be CD8$^+$ and to display no substantial cytolytic activity against ADJ-PC-5 cells (Haubeck and Kölsch, 1982, 1986). Such suppressive cells are not found in the peritoneal cavity of naive mice. The dominant role played by IFN-$\gamma$ as a suppressive factor in this experimental system is proven by two lines of evidence: (1) Addition of recombinant IFN-$\gamma$ to a primary ADJ-PC-5-specific MLTC substantially reduces Tc induction against the tumor cells; see Figure 1(c). (2) Addition of IFN-$\gamma$-specific monoclonal antibodies to an MLTC containing PEC from tumor-treated mice can almost completely restore Tc induction, whereas addition of control monoclonal antibodies has no such effect; see Figure 1(d).

Characterization of the ADJ-PC-5-Specific Tc Response in a Primary MLTC

Since the primary in vitro Tc response against ADJ-PC-5 plasmacytoma cells is suppressed by tumor-induced CD8$^+$ regulatory T cells via IFN-$\gamma$, the effects of other exogenous and endogenous cytokines on a primary MLTC as well as the phenotype of the resulting Tc cells were analyzed in more detail.
a) Suppression of Tc induction by PEC from tumor-treated mice

b) IFN-γ concentrations in MLTC supernatants

c) Suppression of Tc induction by recombinant IFN-γ

d) Restorage of Tc induction by IFN-γ-specific mAb

FIGURE 1 Suppression by PEC from ADJ-PC-5-treated mice is due to IFN-γ. (a)Suppressive T cells were generated by repeated i.p. injections of exponentially increasing numbers of X-irradiated ADJ-PC-5 cells (according to Materials and Methods) and added to an ADJ-PC-5-specific MLTC on day 0. The resulting tumor-specific cytotoxicity was measured on day 6 by a 31Cr-release assay. The cytotoxicity in the presence of nonadherent PEC from tumor treated mice (▼—▼) was compared with the cytotoxicity in the presence of PEC from naive mice (■—■). Control cultures were set up without PEC (□—□) or without PEC and ADJ-PC-5 cells (〇—〇). (b) Cytokine kinetics in MLTC supernatants corresponding to part (a). Substantial amounts of IFN-γ can only be detected in MLTC supernatants containing suppressive T cells from tumor-treated mice (tum.). Samples of culture supernatants were taken at the indicated time points and assayed for IFN-γ according to Materials and Methods. (c) MLTC cultures were set up with 2 × 10⁷ naive BALB/c spleen cells and 1 × 10⁶ X-irradiated ADJ-PC-5 cells in the presence of different concentrations IFN-γ: □: no IFN-γ added; ■: 50 U/ml; and ▼ 500 U/ml. Specific cytotoxicity was measured on day 6 of the culture period. (d) MLTC cultures were set up with (▼—▼) or without (□—□) nonadherent PEC from tumor-treated mice in the presence of 20 μg/ml of an IFN-γ-specific monoclonal rat IgG antibody (α-IFN-γ) or in the presence of 20 μg/ml of an irrelevant IgE-specific rat IgG monoclonal antibody (α-IgE).

Beside the suppressive effect of IFN-γ on Tc induction in vitro, addition of IL-4 to a primary ADJ-PC-5-specific MLTC has an enhancing effect on Tc induction, whereas addition of IL-12 is slightly suppressive in this context (Becker et al., 1997). Another striking difference between the primary in vitro Tc response against ADJ-PC-5 cells and a typical Tc response, as seen in the case of a primary allogenic MLC, is the absolute dependence of the tumor-specific Tc response on tumor-derived IL-10. As shown in Figure 2, addition of IL-10-specific monoclonal antibodies to a primary MLTC can completely eliminate the ADJ-PC-5-specific Tc response, whereas an allogenic control MLC of CBA/J spleen cells against irradiated BALB/c spleen cells is not affected. In addition, cytokine production of CD8⁺
T cells from a primary ADJ-PC-5-specific MLTC and a primary allogenic MLC was compared on a single-cell level by flow cytometry. As shown in Figure 3, the percentage of CD8+ T cells increases significantly in both types of cultures as compared with unstimulated control cultures, indicating that an antigen-specific stimulation of CD8+ T cells must have taken place in both types of cultures. Despite the development of a cytotoxic phenotype, CD8+ T cells from an ADJ-PC-5-specific MLTC differ from typical alloantigen-induced CD8+ Tc cells by a lack of IFN-γ production. Production of IL-4 could neither be

---

**FIGURE 2** Induction of ADJ-PC-5-specific Tc cells in a primary MLTC depends on tumor-derived IL-10. MLTC cultures (BALB/c α ADJ) were set up with 2 × 10^7 BALB/c spleen cells and 1 × 10^6 X-irradiated ADJ-PC-5 cells in the absence (closed bars) or presence (open bars) of a mix of two IL-10-specific monoclonal antibodies (SXC-1 and SXC-2; 10 μg/ml each). Allogenic MLC cultures (CBA/J α BALB/c) were set up in the presence or absence of IL-10-specific antibodies with 2 × 10^7 CBA/J spleen cells and 1 × 10^6 X-irradiated BALB/c spleen cells. The resulting cytotoxicity was measured on day 6 against ADJ-PC-5 (ADJ) cells and a syngeneic control B-cell tumor (A20).
detected in MLTC cultures nor in MLC cultures. However, it must be mentioned that at least some CD8+ T-cell lines that were derived from a primary MLTC by repeated \textit{in vitro} stimulation with irradiated ADJ-PC-5 cells produce substantial amounts of IL-4 but no IFN-\(\gamma\) and acquire a full Tc2 phenotype (see what follows).

All three features of the primary tumor-specific Tc response, that is, the suppression by IFN-\(\gamma\) and IL-12, the dependence on IL-10, and the lack of IFN-\(\gamma\) production by the resulting tumor-specific Tc cells, point toward a type-2 nature of the \textit{in vitro} Tc response against the tumor.

### Characterization of Long-Term Cultivated ADJ-PC-5-Specific CD8+ Tc Lines

In order to phenotypically characterize ADJ-PC-5-specific Tc cells further, CD8+ Tc lines were generated from spleen cells of ADJ-PC-5-immunized BALB/c mice or from spleen cells of naive BALB/c mice that were stimulated with irradiated ADJ-PC-5 cells in a primary syngeneic MLTC. Nine of initially 24 CD8+ ADJ-PC-5-specific Tc lines could be cultivated over a period of more than 3 months; among them, seven developed a quite homogeneous phenotype. As shown in Table I, all of the nine lines specifically lyse ADJ-PC-5 cells but not cells of

![Flow cytometric detection of intracellular cytokines on a single-cell level.](image)

*FIGURE 3* Flow cytometric detection of intracellular cytokines on a single-cell level. Living cells were isolated from MLTC or allogeneic MLC cultures after day 5 of the culture period and stained for intracellular cytokines according to Materials and Methods. Both cytokine-specific MoAb are of the same isotype and were used at the same concentrations. Thus, data for an isotype control antibody are not shown. Percentages of the respective populations are indicated in the upper right quadrant of each plot.*
| Line | Preimmunization* | Target E:T | % CD8<sup>+</sup> | V<sub>β</sub> usage<sup>a</sup> | Cytokine production<sup>a</sup> | IFN-γ | IL-4 |
|------|------------------|-------------|------------------|-----------------------------|------------------------|-------|------|
| BTc2 | 10<sup>7</sup> (7×) | ADJ 78 | 63 97 6 98 0 | 8,439 0 998 | Meth.A | 13 5 |
| BTc3 | ADJ 56 | 33 88 6 99 0 | 1,717 0 22,579 | Meth.A | 0 0 |
| BTc4 | 5 × 10<sup>5</sup> (1×) | ADJ 42 | 20 90 6 94 0 | 31,129 0 1,030 | Meth.A | 0 0 |
| BTc5 | 5 × 10<sup>5</sup> (1×) | ADJ 58 | 47 68 6 90 0 | 13,311 0 692 | Meth.A | 18 11 |
| BTc6 | 5 × 10<sup>5</sup> (1×) | ADJ 72 | 56 95 6 34 0 | 27,760 0 977 | Meth.A | 13 10 |
| BTc7 | - | ADJ 41 | 30 40 8.1/8.2 63 | 974 0 23,893 | Meth.A | 5 3 |
| BTc8 | - | ADJ 55 | 28 95 8.1/8.2 76 | 3,500 0 0 | Meth.A | 1 0 |
| BTc9 | - | ADJ 57 | 32 94 8.1/8.2 78 | 3,1200 0 3,1200 | Meth.A | 0 0 |
| BTc14 | - | ADJ 60 | 27 95 8.1/8.2 62 | 3,300 0 0 | Meth.A | 0 0 |

*Tc lines were generated from spleen cells of naive or ADJ-PC-5 preimmunized BALB/c mice. Cytotoxicity of Tc lines was measured against 10<sup>4</sup> ADJ-PC-5 or Meth A cells in a 6-hr 5<sup>1</sup>Cr-release assay at the indicated effector:target ratios. Total percentage of CD8<sup>+</sup> within the respective line. Dominant V<sub>β</sub>TcR type and percentage of T cells expressing this V<sub>β</sub> type among CD8<sup>+</sup> T cells. T cells were cultivated at a density of 1 × 10<sup>6</sup>/ml for 2 days and stimulated with either immobilized αCD3 antibodies or left unstimulated.

Table II shows the results from a three-color fluores-
TABLE II  Flow Cytometric Analysis of in vivo Induced ADJ-PC-5-Specific Tc Cells

| Tc induction | % CD8* | % V$\beta$6/CD8* | % IFN-γ/CD8* | % IFN-γ/CD8* V$\beta$6* | % IL-4*/CD8* |
|--------------|--------|-----------------|--------------|-------------------------|-------------|
| In Vivo      |        |                 |              |                         |             |
| Spleen naive | 13.5   | 9.5             | 0.3          | 0.3                     | 0.06        |
| Spleen immune| 14.9   | 9.3             | 3.8          | 3.1                     | 0.08        |
| PEC naive    | 8.0    | 10.1            | 0.8          | 0.9                     | 1.24        |
| PEC immune   | 9.8    | 12.0            | 7.1          | 12.0                    | 1.67        |

Spleen cells or PEC were pooled from three individual ADJ-PC-5 immune or naive BALB/c mice and analyzed by flow cytometry according to Materials and Methods.

ence analysis of spleen cells and PEC from naive and ADJ-PC-5 immune BALB/c mice stained for CD8, V$\beta$6, IFN-γ, and IL-4. Repeated immunization of BALB/c mice with the tumor does not substantially alter the total number of splenic CD8* or CD8*V$\beta$6* T cells, which is a main phenotype of ADJ-PC-5-specific Tc lines, but induces IFN-γ production in a small but significant fraction of CD8* and CD8*V$\beta$6* T cells. The data demonstrate that the actual percentage of ADJ-PC-5-specific Tc cells among CD8* or CD8*V$\beta$6* T cells must be low, even in immune mice. However, production of IFN-γ by the respective cells identifies them as Tc1 cells. Production of IL-4 is only detectable in a small fraction of CD8+ spleen cells from immune mice, but not in CD8+ cells. Peritoneal T cells from immune and naive BALB/c mice display similar properties, as seen for splenic T cells. As for spleen cells, immunization with the tumor increases the percentage of IFN-γ-producing CD8* or CD8*V$\beta$6* T cells by a factor of about 10. A relatively high percentage of peritoneal CD8+ T cells produces IL-4. However, this population of IL-4-producing peritoneal CD8+ T cells does not expand during immunization, indicating that the tumor-specific Tc response in the peritoneal cavity is a Tc1 response, too.

ADJ-PC-5-Specific Tc1 Cells Are Potent Suppressors of Tc Induction in a Primary ADJ-PC-5-Specific MLTC In Vitro

Since IFN-γ is a suppressive factor for Tc induction in a primary ADJ-PC-5-specific MLTC, we analyzed whether this suppressive effect is also exerted by ADJ-PC-5-specific Tc1 cells. As shown in Table III, a substantial amount of Tc suppression in a primary MLTC is seen in the presence of even the lowest numbers of X-irradiated ADJ-PC-5-specific Tc1 cells. The suppressive effect in this case must be due to IFN-γ, since addition of IFN-γ-specific monoclonal antibodies to the cultures can in part avoid suppression (Table III, Exp. III). Irradiation of the Tc1 cells is necessary, because otherwise the cultures would be overgrown by restimulated Tc1 cells within a few days, making an analysis of the primary in vitro Tc response impossible. Suppression of the primary Tc response by a rapid elimination of ADJ-PC-5 stimulator cells can be excluded in the case of low numbers of irradiated Tc1 cells, since at any time point almost equal numbers of ADJ-PC-5 stimulator cells could be visually detected in MLTC cultures, irrespective if irradiated Tc1 cells were added or not. In contrast to low numbers of irradiated Tc1 cells, suppression by high numbers of irradiated Tc1 cells is not influenced by IFN-γ-specific mAb (data not shown). Suppression of the primary in vitro Tc response in this case, or in the presence of high numbers of irradiated Tc2 cells, can be entirely related to a rapid elimination of ADJ-PC-5 stimulator cells within the first hours of the culture period. The data demonstrate that Tc1 cells can interfere with the generation of ADJ-PC-5-specific Tc cells in a primary syngeneic MLTC using IFN-γ as a suppressor factor. Thus, these ADJ-PC-5-specific Tc1 cells behave similarly to tumor-induced suppressive T cells from ADJ-PC-5 tumor-bearing mice (Pauels et al., 1996).

DISCUSSION

Although the BALB/c plasmacytoma ADJ-PC-5 is highly immunogenic, tumor growth in syngeneic
### TABLE III  Suppression of ADJ-PC-5-Specific Tc Induction in a Primary MLTC by ADJ-PC-5-Specific Type-1 Tc Lines

| Stimulator | Anti-IFN-γ | Type-1 Tc | % specific cytotoxicity against ADJ-PC-5 |
|------------|------------|----------|----------------------------------------|
|            |            |          | 100:1 50:1 25:1                        |
| Exp. I     |            |          | 0 0 0                                  |
| ADJ        |            |          | 46 40 15                               |
| ADJ        | BTc 2 (1 × 10⁶) | 0 0 0 |
| ADJ        | BTc 2 (5 × 10⁵) | 0 0 0 |
| Exp. II    |            |          | 0 0 0                                  |
| ADJ        |            |          | 29 19 10                               |
| ADJ        | BTc 4 (1 × 10⁵) | 0 0 0 |
| ADJ        | BTc 4 (5 × 10⁵) | 0 0 0 |
| Exp. III   |            |          | 70 61 54                               |
| ADJ        |            |          | 70 64 40                               |
| ADJ        | BTc 8 (1 × 10⁶) | 8 10 1 |
| ADJ        | BTc 8 (1 × 10⁵) | 36 26 16 |
| ADJ        | BTc 8 (1 × 10⁵) | 4 9 8 |

Thirty-Gy X-irradiated type-1 Tc cells were added on day 0 at the indicated cell numbers to a primary ADJ-PC-5-specific MLTC containing different concentrations of IFN-γ-specific monoclonal antibodies. The resulting ADJ-PC-5-specific cytotoxicity was measured on day 6 in a ^⁵¹Cr-release assay at the indicated effector:target ratios.

BALB/c mice does not elicit a protective tumor-specific Tc response, but induces a state of tumor-specific tolerance. We could previously demonstrate that the tumor induces a population of CD8⁺ T cells during early stages of tumorigenesis, which are able to suppress a primary ADJ-PC-5-specific Tc response in vitro through IFN-γ (Pauel et al., 1996).

Recent data concerning Tc cells imply a functional and phenotypical dichotomy among CD8⁺ T cells (LeGros and Erard, 1994; Sad et al., 1995), similar to the one observed in CD4⁺ Th1 and Th2 cells. Since suppressive T cells from tumor-tolerant mice and Tc cells from tumor-immune mice are both CD8⁺, we asked whether the failure of BALB/c mice to mount a protective Tc response against a growing ADJ-PC-5 tumor might reflect the involvement of counteractive CD8⁺ Tc1 and Tc2 cells. To answer this question, the phenotypes and cytokine profiles of ADJ-PC-5-specific T cells had to be analyzed.

Long-term cultivated ADJ-PC-5-specific Tc cells are CD8⁺ and use TcR of the Vβ6 and Vβ8 types. The exclusive CD8⁺ phenotype of the lines is in accordance with the finding that ADJ-PC-5 cells are MHC class II negative (Becker et al., 1997). The majority of the Tc lines produce high amounts of IFN-γ but no or only marginal amounts of IL-4 on stimulation, and can thus be classified as Tc1 cells (Sad et al., 1995). Nevertheless, at least two of five ADJ-PC-5-specific CD8⁺ T-cell lines, which were derived from a primary MLTC by repeated restimulation with X-irradiated tumor cells, are IL-4-producing Tc2 cells. Whereas in other experimental systems, the addition of high amounts of IL-4 and blocking antibodies against IFN-γ is a prerequisite for the generation of antigen-specific Tc2 lines (Sad et al., 1995), the exogenous addition of immunomodulating cytokines or antibodies against cytokines is not necessarily required for the induction of tumor-specific Tc2 cells against ADJ-PC-5 plasmacytoma cells. The driving force for Tc2 differentiation in this case seems to be endogenous IL-10 produced by the tumor cells themselves, since addition of blocking antibodies against IL-10 to a primary MLTC can completely abrogate the induction of tumor-specific Tc cells (see what follows). Since the cytokine phenotype of long-term cultivated T-cell lines is strongly influenced by the culture conditions, the observed numbers of Tc1 and Tc2 lines must not necessarily reflect the participation of both T-cell subsets during tumor-specific Tc responses in situ. For this reason, we examined the involvement of either of these T-cell subsets among CD8⁺ and CD8⁺ Vβ6⁺ T cells during ADJ-PC-
5-specific Tc responses in vivo and in vitro using intracellular cytokine staining (CD8^+Vβ6^+ is a major phenotype of ADJ-PC-5-specific Tc lines).

ADJ-PC-5-specific Tc cells from immunized mice are Tc1 cells. This is proven by the exclusive generation of ADJ-PC-5-specific Tc lines from preimmunized mice. Furthermore, immunization with the tumor significantly increases the percentages of IFN-γ-producing CD8^+ and CD8^+Vβ6^+ T cells among spleen cells and PEC, as compared with naive mice.

In contrast to in vivo induced ADJ-PC-5-specific Tc cells, the in vitro Tc response against the tumor shows Tc2 characteristics. This is demonstrated by a lack of IFN-γ production of the tumor-reactive Tc cells, which are induced during a primary MLTC. It is also confirmed by means of supplementation experiments with exogenous cytokines, since the primary in vitro Tc response against ADJ-PC-5 cells is enhanced by IL-4, but suppressed by IFN-γ and IL-12. Furthermore, the primary in vitro Tc response against ADJ-PC-5 is suppressed by irradiated Tc1 cells via IFN-γ.

In this context, the Tc1 cells behave similar to the previously described CD8^+ suppressive T cells from tumor-tolerant mice (Pauels et al., 1996). However, these suppressive CD8^+ T cells did never show any significant cytotoxicity against ADJ-PC-5 cells, whereas the Tc1 lines are potent tumor-specific Tc cells. It is presently not clear if the suppressive T cells in tumor-tolerant mice are Tc1 cells that in situ occur at frequencies too low to cause any substantial cytotoxicity or if they are incompletely activated Tc1 cells, which produce suppressive cytokines but are not sufficiently activated for cytotoxicity.

IL-10 has been described as a cytotoxic T-cell differentiation factor (Chang and Zlotnik, 1991), as well as an inhibitor of CD8^+ cytotoxic T-cell responses in primary allogenic MLC cultures (Bejarano et al., 1992). Thus, the effects of IL-10 on CD8^+ T cells seem to depend on the kind of CD8^+ T-cell response studied. In our experimental system, tumor-derived IL-10 is an essential differentiation factor for the induction of CD8^+ ADJ-PC-5-specific Tc cells, since blocking of IL-10 in MLTC cultures by monoclonal antibodies can completely abrogate the tumor-specific Tc response.

Production of IL-10 is a general property of malignancies of the B-lymphocyte lineage (Bost et al., 1995). It has previously been shown for CD4^+ T cells that IL-10, which is produced by Th2 cells, is a potent inhibitor of Th1 functions (Fiorentino et al., 1989), whereas Th1 cells are able to suppress Th2 cells via IFN-γ (Gajewski and Fitch, 1988). It is also known that Th1 functions are drastically decreased in mice bearing IL-10-producing plasma-cell tumors (Ruzek and Mathur, 1995). Therefore, it is likely that production of IL-10 by ADJ-PC-5 cells strongly influences the CD8^+ Tc-cell response against the tumor. In this context, high cell numbers of ADJ-PC-5 cells, which are present in a MLTC produce sufficiently high amounts of IL-10 to dictate a Tc2-dominated response against the tumor in vitro. On the other hand, low numbers of tumor cells, which are present during early stages of tumorigenesis in vivo, could result in a Tc1-dominated response due to a lack of substantial amounts of IL-10. The same seems to be true for mice that were weekly immunized with high numbers or X-irradiated ADJ-PC-5 tumor cells. In this case, the IL-10 might be rapidly cleared from the surrounding of the tumor cells without being further produced by the tumor cells that die from irradiation.

Suppression via IFN-γ of the IL-10-driven Tc2 response against ADJ-PC-5 cells is the main reason for the downregulation of the tumor-specific Tc response in a primary MLTC by CD8^+ T cells from tumor-treated mice. However, the question remains to be answered, how an early response of IFN-γ-producing CD8^+ T cells during early stages of tumorigenesis can omit the onset of a Tc1-dominated response during the course of a subsequent immunization with high numbers of tumor cells.

We are presently favoring the following model: During early stages of tumorigenesis, when no substantial amounts of IL-10 are present, the growing tumor induces a transient tumor-specific Tc1 response. This early Tc1 response, which is not strong enough to cause a regression of the tumor, is subsequently stopped by tumor-derived IL-10, when
the tumor mass increases. These Tc1 cells must be in some state of anergy, because a subsequent immunization with high numbers of X-irradiated ADJ-PC-5 tumor cells can no longer induce a protective tumor-specific Tc1 response, as it would be the case in naive mice. The Tc1 cells are likely to be anergized, but not deleted by the tumor, since CD8+ T cells from tumor-bearing mice, or from mice that have been subjected to a simulated tumor growth, can still suppress a Tc2-dominated primary MLTC via IFN-γ.

MATERIALS AND METHODS

Mice and Tumors
Female BALB/c mice were aged 8 to 12 weeks at the beginning of each experiment. The animals were either purchased from Charles River (Sulzfeld, Germany) or bred at our own animal facility. The nonsecreting variant of the BALB/c plasmacytoma ADJ-PC-5 (ADJ-PC-5-NS) (Blatt and Haimovich, 1977), originally obtained from J. Haimovich (Weizmann Institute, Rehovot, Israel) was used as a model tumor. The BALB/c fibrosarcoma Meth A (Old et al., 1962), the DBA/2 plasmacytoma ULMC (Bosslet et al., 1979), and the BALB/c B-cell lymphoma A20 (Kim et al., 1979) were used as control tumors in some experiments.

Media
Dulbecco’s modified Eagle’s medium, containing penicillin (50 U/ml), streptomycin (50 μg/ml), L-glutamine (2 mM), 2-mercaptoethanol (5 × 10−5 M), nonessential amino acids, 10% FCS, and additionally 3.5 g/l glucose (referred to as MLC medium), was used for most experiments. Hank’s balanced salt solution (BSS) was used as washing medium.

Monoclonal Antibodies
Monoclonal rat antibodies specific for murine IFN-γ (R4-6A2, IgG1; Spitalny and Havell, 1984), Vβ6 (44-22-1, IgG2a; Acha-Orbea et al., 1985), and Vβ8.1 (KJ 16, IgG2a; Haskins et al., 1984) were used as hybridoma supernatant or as affinity-purified material using Protein G sepharose (Pharmacia, Uppsala, Sweden). Rat monoclonal antibodies against IL-10 (SXC-1 and SXC-2, IgM; Mosmann et al., 1990) were purified from hybridoma supernatants by ammonium sulphate precipitation followed by ion-exchange chromatography on hydroxylapatite columns.

Generation of Tumor-Induced Suppressive T Cells by Tumor Growth Simulation
The in vivo induction of suppressive T cells by tumor growth simulation has been previously described (Haubeck and Kölsch, 1982). Briefly, BALB/c mice received daily intraperitoneal injections of exponentially increasing numbers of 60-Gy X-irradiated ADJ-PC-5 cells starting with eight cells on day 0. According to the generation time of the tumor, doses of injected tumor cells were doubled every day, until mice had received 10⁶ cells on day 16. Eight days after the last treatment with ADJ-PC-5 cells, peritoneal exudate cells (PEC) were collected by a peritoneal lavage and fractionated into adherent and nonadherent cells by overnight incubation on plastic dishes at 37°C in MLC medium. Mice in which tumor growth was simulated by the preceding protocol are subsequently denoted “tumor-treated.”

Generation and Culture of ADJ-PC-5-Specific Tc Lines
Tumor-specific Tc lines were generated by weekly in vitro restimulations of 1 × 10⁶ spleen cells or peritoneal exudate cells from naïve or ADJ-PC-5 preimmunized BALB/c mice with 5 × 10⁶ X-irradiated BALB/c spleen cells and 2 × 10⁵ X-irradiated ADJ-PC-5 cells in 10 ml MLC medium. After the second to fifth restimulation, most Tc cultures were supplemented with 5% of an IL-2/IFN-γ containing supernatant from Con-A-stimulated rat spleen cells (CM) or with 5 U/ml recombinant human IL-2.
Mixed Lymphocyte Tumor Cell Culture (MLTC)

ADJ-PC-5-specific Tc cells were generated in a primary syngeneic MLTC from spleen cells of naive mice as previously described (Haubeck and Kölsch, 1982). Briefly, $2 \times 10^7$ BALB/c spleen cells were cultured with $1 \times 10^6$ 60-Gy X-irradiated ADJ-PC-5 cells in 10 ml of MLC medium for 6 days. Thereafter, cells were harvested and tested in a 6-hr $^{51}$Cr-release assay for specific cytotoxicity (Haubeck and Kölsch, 1982). In some experiments, the medium used for Tc induction was supplemented with recombinant IL-4, IL-12, or IFN-γ or monoclonal antibodies against IL-10.

In Vitro Assay for T-Cell-Mediated Suppression

To test their suppressive capacity, nonadherent PEC from naive and tumor-treated mice, or 30-Gy X-irradiated ADJ-PC-5-specific Tc cells were added to an ADJ-PC-5-specific MLTC (as described earlier) at day 0 of culture. After 6 days, specific cytotoxic activity against ADJ-PC-5 cells was measured in a 6-hr $^{51}$Cr-release assay and compared with the cytotoxicity from cultures without added T cells. For some experiments, IFN-γ-specific monoclonal antibodies were added to the cultures at various concentrations.

Flow Cytometric Analysis

For flow cytometric analysis of surface markers on tumor-specific T cells, cells were harvested and washed with BSS. For two-parameter analysis of Vβ expression on CD8+ ADJ-PC-5-specific Tc lines, cells were incubated with FITC-coupled, Vβ-specific, and PE-coupled CD8-specific monoclonal antibodies (Pharmingen, San Diego). All washing and incubation steps were performed with ice-cold MLC medium containing 0.1% NaN₃. Labeled cells were analyzed using a FACSComp flow cytometer (Becton-Dickinson, Heidelberg). Prior to flow cytometric analysis, samples were incubated with 10 μg/ml propidium iodide to discriminate dead cells from living cells.

Sources and Measurement of Cytokines

Recombinant mouse IL-4 and recombinant rat IFN-γ were purchased from Hycult Biotechnology (Uden, The Netherlands), whereas recombinant mouse IL-12 was kindly provided by Frank J. Podlasky (Hoffmann-LaRoche, Nutley, NJ). IL-4, IFN-γ, and IL-10 were determined by means of a sandwich ELISA using 2A5.7 (Sander et al., 1993), and biotinylated XT1 monoclonal anti-bodies for IL-10, R4-6A2 (Sipitalny and Havell, 1984), and biotinylated XMG 1.2 for IFN-γ, 11B11 (Ohara and Paul, 1985), and biotinylated BVD6-24G2 for IL-4. All biotinylated monoclonal antibodies were purchased from Pharmingen (San Diego).

Detection of Intracellular Cytokines

Staining of intracellular cytokines was performed according to the saponin/monensin method, as described elsewhere (Jung et al., 1993). Briefly, spleen cells, peritoneal exudate cells, or cells from primary MLTC or MLC cultures were purified by centrifugation over a discontinuous Ficoll gradient (Pharmacia, Uppsala, Sweden). Purified cells were stimulated with 10 ng/ml PMA (Serva, Heidelberg) and 1 μM ionomycin (Sigma, Deisenhofen, Germany) in the presence of 1 μM monensin (Sigma) for 4 hr. Subsequently, cells were fixed with 4% paraformaldehyde in BSS for 15 min on ice. Intracellular cytokines were stained with biotinylated mAb against murine IFN-γ or murine IL-4 (both from Pharmingen) followed by Streptavidin-PE. All incubation steps were performed for 30 min at 4°C. The medium used for incubation and washing steps was BSS containing 5% FCS and 0.1% saponin (Fluka, Neu-Ulm, Germany) to cause reversible pore formation. Immediately before flow cytometric analysis, cells were extensively washed with BSS without saponin to reverse pore formation and counterstained for 30 min with CyChrome-coupled mAb against CD8 and FITC-coupled mAb against Vβ6 (both from Pharmingen). Cytokine expression was evaluated with a FACSComp flow cytometer (Becton-Dickinson).
Acknowledgments

This work was supported by the Wilhelm Sander-Stiftung through grant 91.024 and the DFG through SFB 310. C.B. was recipient of a fellowship from the Konrad-Adenauer-Stiftung.

References

Acha-Orbea H., Zinkernagel R. and Hengartner H. (1985). Cytotoxic T cell clone-specific monoclonal antibodies used to select clonotypic antigen-specific cytotoxic T cells. Eur. J. Immunol. 15:31-36.

Bejarano M.T., deWaal-Malefyt R., Abrams J.S., Bigler M., Becker C., Kölsch E. and Pauels H.G. (1997). CD8+ tumor-specific Konrad-Adenauer-Stiftung. SFB310. C.B. was recipient of a fellowship from the Stiftung through grant 91.024 and the DFG through Bosslet K., Schirrmacher V. and Shantz G. (1979). Tumor Blatt C. and Hämovich J. (1977). Cellsurface immunoglobulins of Bost K.L., Bieligk S.C. and Jaffe B.M. (1995). Lymphokine mRNA expression by transplanted murine B lymphocytic malignancies. Tumor-derived IL-10 as a possible mechanism for modulating the anti-tumor response. J. Immunol. 154:718-729.

Bejarano M.T., deWaal-Malefyt R., Abrams J.S., Bigler M., Becker C., Kölsch E. and Pauels H.G. (1997). CD8+ tumor-specific Konrad-Adenauer-Stiftung. SFB310. C.B. was recipient of a fellowship from the Stiftung through grant 91.024 and the DFG through Bosslet K., Schirrmacher V. and Shantz G. (1979). Tumor Blatt C. and Hämovich J. (1977). Cellsurface immunoglobulins of Bost K.L., Bieligk S.C. and Jaffe B.M. (1995). Lymphokine mRNA expression by transplanted murine B lymphocytic malignancies. Tumor-derived IL-10 as a possible mechanism for modulating the anti-tumor response. J. Immunol. 154:718-729.

Bejarano M.T., deWaal-Malefyt R., Abrams J.S., Bigler M., Becker C., Kölsch E. and Pauels H.G. (1997). CD8+ tumor-specific Konrad-Adenauer-Stiftung. SFB310. C.B. was recipient of a fellowship from the Stiftung through grant 91.024 and the DFG through Bosslet K., Schirrmacher V. and Shantz G. (1979). Tumor Blatt C. and Hämovich J. (1977). Cellsurface immunoglobulins of Bost K.L., Bieligk S.C. and Jaffe B.M. (1995). Lymphokine mRNA expression by transplanted murine B lymphocytic malignancies. Tumor-derived IL-10 as a possible mechanism for modulating the anti-tumor response. J. Immunol. 154:718-729.