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http://www.jimmunol.org/content/179/3/1996
CCR4-Expressing T Cell Tumors Can Be Specifically Controlled via Delivery of Toxins to Chemokine Receptors

Dolgor Baatar,* Purevdorj Olkhanud,* Dianne Newton,† Kenya Sumitomo,* and Arya Biragyn2*2

Expression of chemokine receptors by tumors, specifically CCR4 on cutaneous T cell lymphomas, is often associated with a poor disease outcome. To test the hypothesis that chemokine receptor-expressing tumors can be successfully controlled by delivering toxins through their chemokine receptors, we have generated fusion proteins designated chemotoxins: chemokines fused with toxic moieties that are nontoxic unless delivered into the cell cytosol. We demonstrate that chemokines fused with human RNase Pseudomonas exotoxin 38 are able to specifically kill tumors in vitro upon internalization through their respective chemokine receptors. Moreover, treatment with the thymus and activation-regulated chemokine (CCL17)-expressing chemotoxin efficiently eradicated CCR4-expressing cutaneous T cell lymphoma/leukemia established in NOD-SCID mice. Taken together, this work represents a novel concept that may allow control of growth and dissemination of tumors that use chemokine receptors to metastasize and circumvent immunosurveillance. The Journal of Immunology, 2007, 179: 1996–2004.

Chemokines are a group of small 8- to 15-kDa secreted and structurally related peptides that primarily regulate cell trafficking and diapedesis, although they exhibit a number of additional functions (1, 2, 3). To date, the group consists of ~50 chemokines classified into four superfamilies (two major types CC and CXC and two minor C and CX3C chemokines) on the basis of cysteine residues. They bind and signal through heterotrimeric Gi protein-coupled seven-transmembrane chemokine receptors that are differentially expressed on various subsets of immune cells (4–10). Chemokines and chemokine receptors appear to play a significant role in regulation of growth and metastatic spread of tumors, and their expression is often associated with a poor disease outcome. Since a first report that correlated breast cancer metastasis with expression of CXCR4, CCR7, and CCR10 (11), a number of other chemokine receptors were shown to be differentially expressed on tumors. For example, CXC3 is found expressed on primary melanoma (12), breast cancer (13), and various lymphomas, such as T cell and NK cell lymphomas, chronic lymphocytic leukemia/small lymphocytic lymphoma, and splenic marginal zone B cell lymphoma (14–16). The metastatic migration of tumors to lymphoid organs was associated with their overexpression of CCR7 and CXCR4, whereas tumors homing into the skin were associated with expression of CXCR4 (11, 17–19). Besides migration, chemokine/chemokine receptors also affect the viability and survival of tumor cells through the activation of their prosurvival and proliferation signaling cascades (20, 21). As a result, a prosurvival signal transmitted by the activated CCR1, CCR4, and CXCR4 leads to a greater risk of metastasis and poorer survival in patients with primary melanoma (11, 12) and colorectal cancer (22). Similarly, unfavorable outcome of the disease in patients with adult T cell leukemia/lymphoma (ATLL), mucositis, and Sézary syndrome was also associated with overexpression of CCR4 by malignant CD4+ T cells (1, 2, 19). In contrast, chemokines not only recruit tumors, but they can also induce infiltration of various immunosuppressive cells, such as T regulatory cells (Tregs), immunosuppressive inhibitory macrophages, and NK T cells, leading to escape from immunosurveillance and an unfavorable disease outcome. For example, CCR4+ Tregs were shown to be recruited to cutaneous lymphoma and ovarian cancer sites that expressed high levels of thymus and activation-regulated chemokine (TARC)/CCL17 or macrophage-derived chemokine/CCL22 (19, 23).

Immunotherapeutic interventions that block chemokine receptor signaling expressed by tumors remain an attractive, but insufficiently explored strategy. The CCR4-expressing tumors were successfully controlled in mice treated with Abs to CCR4, inducing NK-mediated Ab-dependent cellular cytotoxicity (1, 24). However, the efficacy of the approach can be affected by the host FcR genetic polymorphism, and its clinical potency in humans remains to be determined. It is tempting to speculate that the strategies that directly kill chemokine receptor-expressing tumors might elicit a higher degree of the disease control. To test this, we generated a formulation designated chemotoxin: chemokines fused with toxic moieties, such as RNases or toxins that are nontoxic unless delivered into the cell cytosol. The work has been inspired by our recent findings that chemokines can deliver exogenous Ags into cytosol to be processed and cross-presented to the MHC class I molecules (25). The process is very efficient because only nM

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1 Laboratory of Immunology, Gerontology Research Center, National Institute on Aging, Baltimore, MD 21224; and 2 Department of Microbiology, SAIC-Frederick, National Cancer Institute, Frederick, MD 21702

Received for publication March 18, 2007. Accepted for publication May 28, 2007.

The costs of publication of this article were defrayed in part by the payment of page charges. This article must therefore be hereby marked advertisement in accordance with 18 U.S.C. Section 1734 solely to indicate this fact.

1 This research was supported by the Intramural Research Program of the National Institute on Aging, National Institutes of Health.

2 Address correspondence and reprint requests to Dr. Arya Biragyn, Laboratory of Immunology, Gerontology Research Center, National Institute on Aging, National Institutes of Health, 5600 Nathan Shock Drive, Box 21, Baltimore, MD 21224. E-mail address: biragyna@mail.nih.gov

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amounts of chemokine-fused Ags are sufficient to induce expression capability of stimulating Ag-specific CTL. In concordance, as shown in this work, chemokotaxis are able to specifically deliver toxins into cytosol of the targeted cells using their chemokine receptors. Our experiments using a mouse model for human cutaneous T cell leukemia/lymphoma demonstrate that the strategy is simple and potent, because treatment with TARC chemotxin almost completely eradicated CCR4-expressing established T cell tumors, while leaving receptor-negative cells untouched. We think that the strategy may have significant clinical value for control of tumors that use chemokine receptors for metastatic spread or to circumvent immunosurveillance.

Materials and Methods

Cell lines and mice

Human acute T-lymphoblastic leukemia cell lines CCRF-CEM (CEM, CCL-119) and MOLT-4 (CRL-1582), and human embryonic kidney (HEK)-293 cells were purchased from American Type Culture Collection. HEK-293 cells stably transfected with human CCR8 (HEK/CCR8) was a gift from Z. Howard (Science Applications International, Frederick, MD). HEK-293 cells were cultured in DMEM (Invitrogen Life Technologies) containing 10% FBS. The same medium, but with 400 µg/ml G418 (Sigma-Aldrich), was used to maintain HEK/CCR8 cells. CEM and MOLT-4 cells were cultured in a standard RPMI 1640 medium (Invitrogen Life Technologies) supplemented with 10% FBS. Female 6- to 8-wk-old NOD/ LtSz-scid/scid (IACR) mice were purchased from The Jackson Laboratory and kept under pathogen-free conditions at the National Institute on Aging animal facility. Animal care was provided in accordance with the procedures outlined in the “Guide for the Care and Use of Laboratory Animals” (National Institutes of Health Publication No. 86-23, 1985).

Plasmid constructs

Schema of constructs is shown in Fig. 1. Mature sequence for TARC/CCL17 was cloned using RT-PCR from human thymus RNA using the following primers: PRhCCL27M-1 (TAT CTC GAG GGA GGT CCC GAG GGC GGC AGC CTG GCC GCG CTGA) and PRhCCL27M-2 (CGT GGT GGT GGT GGT TCT AAA CCG CCG CAG TTC ACT TGG GCT). Eosinophil-derived neurotoxin (EDN) was cloned using the following PCR primers: PREDN-1 (ATA CTC GAG GGA GGT CCC GAG GGC GGC AGC CTG GCC GCG CTGA) and PREDN-2 (CGG GCC TAC GAT GAT ACG GTC CAG ATG AAC CCG AAC) from the EDN-expressing plasmid described elsewhere (26). Bacterial expression vectors with CCR8 antagonist-encoding mature sequence for TARC/CTACK/CCL27 was cloned using primers PRhCCL27M-2 (TAT CTC GAG GGA GGT CCC GAG GGC GGC AGC CTG GCC GCG CTGA) and PRhCCL27-2 (CGG AAC) from the EDN-expressing plasmid described elsewhere (26). A truncated C-terminal of human TARC (TARC-EDN) and inserted into the bacterial I and NcoI fragments encoding for mature sequence of human TARC were cut with GCC GGG CTG GCT). To construct a chemotoxin-expressing vector, PCR fragments encoding for mature sequence for TARC were cut with PstI HaeIII (Osmonics), as described (31, 32). Briefly, the lower chambers were filled with 25 µl of complete RPMI 1640 medium containing titrated amounts of chemotaxtratants, or human TARC/CCL17 (R&D Systems), or binding medium alone (negative control). The top chambers were filled with 50 µl of RPMI 1640 medium (for 1 h at 37°C or at 4°C, and, after extensive washings with PBS, incubated with either anti-c-myc Ab (1/100 dilution, 9E10) or anti-EDN polyclonal rabbit Ab (1/100 dilution). Then cells were incubated with respective secondary Abs conjugated with FITC, such as anti-mouse IgG FITC (Jackson ImmunoResearch Laboratories) and anti-rabbit IgG FITC (Sigma-Aldrich) Abs. The binding/internalization was assessed by flow cytometry on a FACSscan (BD Biosciences) using CellQuest software. Alternatively, chemotoxin binding and internalization were assessed by fluorescent microscopy. Briefly, cells (1 × 10⁶) were treated for 1 h at 37°C or at 4°C with 25 µg/ml fusion proteins in RPMI 1640 containing 10% FBS. Then cells were fixed with 3.7% formaldehyde, permeabilized with 0.2% Triton X-100 (5 min), and incubated with mouse anti-c-myc Ab for 1 h at 37°C, and for 30 min at 37°C with goat anti-mouse IgG Ab conjugated to Alexa Fluor 488 (Molecular Probes). Images were acquired with a ×40 objective on an Axiovert 200 microscope (Carl Zeiss Vision) and using Axiovision software (Carl Zeiss Vision).

Chemokine receptor-binding internalization assays

Expression of CCR4 and CCR8 was tested using FITC-conjugated anti-human CCR4 and CCR8 Abs (R&D Systems), respectively, according to manufacturer instructions. The ligand-binding internalization assays were performed with 1 × 10⁵ cells blocked with 10% mouse serum in PBS containing 2% BSA (PBS-B). Proteins (50 µg/ml) were incubated in cell culture medium for 1 h at 37°C or at 4°C, and, after extensive washings with PBS-B, incubated with either anti-c-myc Ab (1/100 dilution, 9E10) or anti-EDN polyclonal rabbit Ab (1/100 dilution). Then cells were incubated with respective secondary Abs conjugated with FITC, such as anti-mouse IgG FITC (Jackson ImmunoResearch Laboratories) and anti-rabbit IgG FITC (Sigma-Aldrich) Abs. The binding/internalization was assessed by flow cytometry on a FACSscan (BD Biosciences) using CellQuest software. Alternatively, chemotaxtrat secreted from each cell line was detected in the supernatant by ELISA (26). The supernatants were incubated with anti-human CCR8 Ab, 0.05% BSA/PBS, and incubated with either anti-c-myc Ab (1/500 dilution, 9E10) or anti-EDN polyclonal rabbit Ab (1/100 dilution). Then, plates were washed with 0.05% BSA/PBS, incubated with rabbit polyclonal anti-mouse IgG Horseradish Peroxidase (Amersham) for 1 h at 37°C, washed again, and incubated with 1 mg/ml TMB (Sigma-Aldrich) for 10 min at room temperature, and finally, 1 mol/l H₂SO₄ was added and absorbance was read at 450 nm.

Cell viability assays

Cell viability was assessed using cell proliferation reagent WST-1 (Roche Applied Science). A minimum of three independent assays was performed in triplicate for each cell type and protein. Cells (5 × 10⁴ per well), plated in 96-well flat-bottom plates 1 day before the assay, were treated with titrated amounts of proteins in complete RPMI 1640 medium supplemented with 10% FBS (for HEK-293 cells or HEK/CCR8 cells) or 10% FBS (for CEM, CRL-1582, and MOLT-4 cells) and cultured for up to 7 days. Then, cell viability was replaced with 10% WST reagent, and cell viability was assessed after 2–4 h of incubation at 37°C. Results are expressed in percentage of D₅₀ values of PBS-treated cells. For in vitro cell growth assay, 1 × 10⁴ CEM cells were plated in 96-well plate and treated with either PBS or 10 µg/ml proteins, and number of viable cells was evaluated daily for 3 days with WST-1 reagent. Results are expressed as percentage of OD₅₀ values of PBS-treated cells. Cell apoptosis was tested by staining them with Annexin-V-Fluor Staining kit (Roche Applied Science), according to manufacturer instructions, and analyzed by flow cytometry.
CEM tumor growth in vivo and in vitro manipulations

Six- to 8-wk-old female NOD/SCID mice were challenged s.c. with 2 × 10^7 CEM cells. Twelve days later, the mice were checked for the presence of palpable tumors, and mice that did not develop tumors were excluded from the study. The remaining mice were randomized and intratumorally injected with either PBS, TARC-PE38, or TARC-OFA (25 μg/mice, each) daily for 5 days (n = 5 or 6 per treatment group). Tumor size was measured in perpendicular dimensions every other day, and tumor surface area was calculated. Mice were euthanized at day 27 or when tumor area reached 400 mm^2. For histological analysis, tumor was fixed in 10% formalin and embedded in paraffin. Paraffin slides were stained with H&E and analyzed with light microscopy.

To study the sensitivity to TARC-PE38-induced cell death, tumor cells from PBS- and TARC-PE38-treated mice were isolated and cultured ex vivo for 7 days. Then, these tumor-derived cells were seeded in 96-well plates (5 × 10^3 cells/well) and treated for 2 days with 10 μg/ml TARC-PE38. Parental CEM cells (purchased from American Type Culture Collection) were used as control. Cell viability was assessed with WST reagent, as described. Data were expressed as percentage of OD_max values of corresponding control untreated cells. To study the effect of repeated treatments with TARC-PE38 on CCR4 expression and resistance to killing, CEM cells were treated once or five times (with 2-day intervals) with TARC-PE38 (10 μg/ml) and then were cultured for 4 wk to expand remaining cells. Control cells were treated with PBS or TARC-OFA in the same way. Cells were stained with anti-CCR4 Ab and analyzed by FACS. Some cells were plated in 96-well plates and were treated with TARC-PE38 (10 μg/ml), and the cell viability was assessed 2 days thereafter.

To study the effect of chemotoxins after systemic injections, NOD/SCID mice (n = 6) were injected with one million CEM cells via the tail vein. To determine the presence of CEM cells, mice were bled after 16 days of challenge with tumor cells, and the proportion of cells expressing human CD45-FITC Ab (R&D Systems) in peripheral blood was determined by FACS. At day 20 posttumor challenge, mice were randomized and i.v. injected with 10 μg of TARC-PE38 or CTACK-PE38 once daily for 4 consecutive days. At day 25, mice were culled and spleens were removed to assess presence of CEM cells. Spleen-derived mononuclear cells (5 × 10^6) were cultured ex vivo in RPMI 1640 supplemented with 10% PBS for 10 days. Cells were either stained with anti-human CD45 Ab and analyzed by FACS or were stained with trypan blue to evaluate the number of viable cells.

Statistical analysis

All data are expressed as means ± SD. Data were analyzed using a computer-based software system (StatView 5.0.1.; SAS Institute). Differences were tested by Student’s t test or ANOVA, followed by post hoc Scheffe’s F test.

Results

Proof of concept: Chemokines fused with toxic moieties bind their respective chemokine receptors and induce cell killing

We have recently reported that exogenous Ags, if linked (fused) with chemokines, can be efficiently delivered to the cell cytosol and presented to MHC class I molecules using the classical proteasome-mediated pathway (25). We hypothesized that chemokines might also deliver cytotoxic moieties to enable a preferential killing of the chemokine receptor-expressing cells. To test this, we have used toxins that do not bind and kill cells by themselves unless delivered to the cell cytosol. Chemokines were genetically fused with human pancreatic eosinophilic RNase, EDN (MC148-EDN and TARC-EDN), or a truncated form of PE38 (TARC-PE38, MC148-PE38, and CTACK-PE38; Fig. 1). The idea was that MC148 and TARC chemokines would preferentially kill CCR8- and CCR4-expressing cells, respectively. Control constructs expressed MC148 and TARC fusions with irrelevant non-toxic Ags such as mouse plasmacytoma MOPC315 (MC148-VL315) or with tumor Ag OFA (TARC-OFA), or mutant MC148 fused with EDN (MC148-EDN). All constructs contained c-myc and His peptide tags to enable purification and detection (Tag), and a spacer peptide (SP) separating chemokines from Ags to enable proper protein folding.

Several fold less than free EDN. Moreover, it appears that the toxin moiety does not affect the ability of the chemokines to chemotax, because both TARC-EDN and TARC-PE38 specifically attracted CCR4-expressing CEM cells with comparable potency as TARC/CCL17 alone at equal molar ratios (Fig. 2B). Of note: the fusions have ~6 times larger size than free chemokine, and a lesser chemotaxis may be due to the fact that our samples may also contain some amounts of inactive and improperly folded fusions (our unpublished observation). Furthermore, MC148 and TARC chemotoxins could specifically bind CCR8- or CCR4-expressing HEK/CCR8 and CEM cells, respectively (Fig. 2, C–F). Control cells, HEK-293 (Fig. 2C) and MOLT-4 (data not shown), that did not express CCR8 and CCR4 failed to bind chemotoxins. Importantly, chemotoxins were efficiently internalized upon binding with their receptors. Although chemotoxins were mostly detected on the cell surface when incubated at the conditions that sequester receptor internalization (4°C; Fig. 2, D and F), they were internalized within a few minutes of incubation at the permissive temperature (37°C; TARC-EDN; Fig. 2D) and found in the cell cytosol (37°C, TARC-PE38; Fig. 2F). The internalization was through the respective chemokine receptor, because CCR4 was also down-regulated upon incubation with TARC-EDN at 37°C, but not at 4°C (Fig. 2E). Controls, unlinked free toxins such as EDN (Fig. 2E) or toxins fused with mutant chemokine (data not shown), did not bind and affect expression of CCR4.

Taken together, these data indicate that chemotoxins were efficiently delivered into the cytosol using their respective chemokine receptors. Next, we tested whether this would render them cytotoxic. HEK/CCR8 cells were incubated with titrated amounts of MC148-EDN, or control EDN alone and MC148-VL315. As shown in Fig. 3A, HEK/CCR8 cells were only killed when incubated with MC148-EDN, but not with EDN alone or MC148-VL315. The cytotoxicity was specific to the CCR8-expressing cells, because MC148-EDN failed to kill the CCR8-negative parental HEK-293 cells (data not shown). However, despite the fact...
that significant cell death was induced by the treatment with a relatively low amount of EDN chemotoxin (starting from 3 µg/ml; Fig. 3A), it was only detectable after a prolonged incubation up to 7 days.

**Bacterial toxin-containing chemotoxins specifically and efficiently induce cell death in vitro**

To search for chemotoxins with a faster rate of activity, we tested chemokines fused with the exotoxin fragment PE38 from *Pseudomonas aeruginosa* that was reported to kill mammalian cells within a short period of time by inhibiting protein synthesis via the ADP ribosylation of elongation initiation factor 2 (34, 35). Indeed, significant apoptosis was elicited in cells treated overnight with TARC-PE38 (Fig. 3, B and C), leading to mostly dead cells (80%) within 2 days of incubation (Fig. 3, B and D). Similar to EDN chemotoxins, TARC-PE38 also acted through CCR4, because it only killed the CCR4-expressing CEM, but not CCR4-negative MOLT-4 cells (Fig. 4A). Controls, such as TARC-OFA and TGFrα-PE38 (Fig. 4B), or CTACK-PE38 that acts through CCR10 (data not shown), failed to induce cell death, further supporting that the process was specific and required CCR4. TARC-PE38 was a very potent chemotoxin, because a single treatment with as low as 2–8 nM TARC-PE38 (0.1–0.4 µg/ml; Fig. 4A) was sufficient to kill CEM cells. In fact, a single treatment with TARC-PE38 significantly suppressed expansion of fast-growing CEM tumor cells in vitro, leaving only a few viable cells (Fig. 4C). Of note: the remaining cells are presumably derived from a small proportion (3%) of CCR4-negative CEM cells that was present before the treatment. The CCR4-negative cells are no longer susceptible to TARC-PE38 and eventually grow back, as reflected by a slight, but significant increase in OD450 values by day 3 treatment (47 ± 4) when compared with day 2 (20 ± 4, p < 0.01; Fig. 4C).

**TARC-PE38 effectively eradicates established CCR4+ tumors in mice**

CCR4 expression by cutaneous T cell leukemia/lymphomas is an indicator for the patients’ poor clinical outcome (1, 19). Due to the lack of murine models for cutaneous leukemia, we have experimented with s.c. established human T cell lymphoblastoid CEM
tumor cells in NOD-SCID mice. Mice with 12-day growing CEM tumors were injected intratumorally with 25 μg of TARC-PE38 or TARC-OFA, or mock treated with saline once daily for 5 consecutive days. Injections of TARC-OFA or PBS did not affect tumor growth and mice had to be sacrificed by day 27 when tumor size reached ∼400 mm² (Fig. 5, A and E). In contrast, almost complete tumor suppression was observed in mice treated with TARC-PE38 (Fig. 5, B and E), which was also associated with appearance of substantial necrotic lesions at the tumor challenge site (Fig. 5B). The data were supported by histological analysis taken at 27 days after tumor challenge; the TARC-PE38-treated tumors were mostly necrotized (data not shown) and tumor cells were detected only at the margin of necrotic areas (Fig. 5D). In contrast, control-treated samples primarily consisted of tumor cells (cells with large and dense nuclei; Fig. 5C) infiltrated in the dermis and spread into s.c. tissue.

Tumor relapse is due to survival of receptor-negative cells in NOD-SCID mice

Despite the potency of the treatment and the fact that the TARC-PE38 treatment eliminated any palpable signs of tumor in mice, the tumor eventually relapsed and a new tumor growth could be detected at the margins of the necrotic area and surrounding skin. In addition, the tumor no longer responded to the TARC-PE38 treatment, even if treated multiple times by higher doses (100 μg/ml; data not shown). Therefore, we hypothesized that these tumors
might represent an outgrowth of resistant cells that had lost expression of CCR4. In support, the relapsed tumor cells from TARC-PE38-treated mice expressed significantly reduced surface CCR4 compared with tumors grown in mock-treated mice (CCR4 mean fluorescence intensity, 92.5 ± 1.3% vs 147.2 ± 16.4%, respectively, p < 0.05; Fig. 6A). Thus, these data indicate that the TARC-PE38 resistant and relapsed tumors are mostly represented by the cells that either do not signal via CCR4 or lost its surface expression. For example, cells isolated from TARC-PE38-treated and relapsed tumors were significantly less susceptible to additional ex vivo treatments with TARC-PE38 (p < 0.001, Fig. 6B). In contrast, TARC-PE38 killed both control ex vivo cultured cells from PBS-treated mice as efficiently as parental in vitro cultured CEM cells (○ and △, respectively; Fig. 6B). To further assess this, the parental in vitro cultured CEM cells were pretreated either once or five times (with 2-day intervals) with 10 μg/ml TARC-PE38 or TARC-OFA, or PBS. Then, the cells were cultured for 4 wk without any treatments. The majority of control CEM cells pretreated with TARC-OFA or PBS were viable (see also Fig. 4C) and expressed CCR4 (97%; Fig. 6C). In contrast, a single pretreatment with TARC-PE38 yielded dramatic cell death, leaving only a small number of viable cells (<5%, data not shown; also see Fig. 4C) that contained significantly reduced proportion of CCR4-expressing cells (48%, p < 0.001; Fig. 6C). The proportion of CCR4-negative cells was further increased (78%; Fig. 6C) when cells were pretreated with TARC-PE38 for five times. Importantly, these cells not only became CCR4 negative, but also acquired resistance to subsequent treatments with TARC-PE38 (Fig. 6D). Control pretreatments with TARC-OFA or PBS did not affect either the proportion of CCR4-expressing cells (Fig. 6C), or sensitivity to TARC-PE38 (Fig. 6D).

Systemic treatments with TARC-PE38 are safe and effective in mice

Next, we tested whether tumor dissemination can be controlled by the systemic administrations of TARC-PE38. Twenty days after i.v. injection of CEM cells, NOD-SCID mice were treated with 10

**FIGURE 5.** Eradication of CEM tumors established in NOD-SCID mice. A and B, Macroscopic appearance of tumors treated with TARC-OFA (A) or TARC-PE38 (B) at day 27 posttumor challenge. C and D, Microscopic appearance (H&E-stained slides, original magnification, ×200) of tumors treated with TARC-OFA (C) and TARC-PE38 (D, margin of necrotic area) at day 27 posttumor challenge. E, Tumor growth plot of mice treated with TARC-PE38, TARC-OFA, or PBS. *, p < 0.05 is for comparisons between the TARC-PE38 and TARC-OFA groups at the days indicated. Representative data from two independent experiments with comparable data with eight mice per group.

**FIGURE 6.** Mechanism of tumor escape in NOD-SCID mice: TARC-PE38 kills selectively CCR4-positive tumors, but does not prevent growth of escapees that do not express CCR4. A, CCR4 expression in tumor cells isolated from PBS (thin line)- or TARC-PE38 (thick line)-treated mice. Cells were stained with anti-human CD45-PE and anti-human CCR4-FITC Abs and were analyzed by FACS. Histogram shows CCR4 expression in CD45-gated cells. Filled line: isotype-matched FITC-conjugated IgG. B, Effect of TARC-PE38 on viability of tumor cells derived from PBS- or TARC-PE38-treated mice. Parental in vitro cultured CEM cells from American Type Culture Collection were used as control. *, p < 0.01 is for comparisons with the parental CEM cells treated with TARC-PE38 at the indicated doses. C, CCR4 expression on CEM cells treated with 10 μg/ml TARC-PE38 once (TARC-PE38 1×) or five times (TARC-PE38 5×) and cultured thereafter for 4 wk. Controls were treated with TARC-OFA or PBS. D, Effect of TARC-PE38 on viability of CEM cells treated and cultured, as described for C. Cells were overnight treated again with 10 μg/ml TARC-PE38 to test their resistance (viability). *, p < 0.01 is for comparisons with the TARC-OFA treatment. Representative data from at least two independent experiments with comparable data.
μg of TARC-PE38 (i.v., once per day) for 4 consecutive days. Control mice were treated with CTACK-PE38, a PE38 fusion with chemokine that specifically binds CCR10. As shown in Fig. 7A, only <1% of mononuclear cells isolated from spleens of the TARC-PE38-treated mice can be associated with CEM tumors, because they expressed human CD45. In contrast, significantly higher amounts of tumors were found in control CTACK-PE38-treated mouse spleens (9.5%; Fig. 7A). Cells isolated from both TARC-PE38- and CTACK-PE38-treated mice were able to grow ex vivo and became almost 100% positive for human CD45 (data not shown) after 10 days of culture. However, the number of cells expanded from spleens of the TARC-PE38-treated mice was 20-fold less than that of the CTACK-PE38-treated mice (Fig. 7B).

Taken together, our data indicate that chemotoxins can efficiently regress the CCR4-positive T cell tumors established in mice, suggesting their potential usefulness in the control of cutaneous leukemias that express CCR4. Although the CCR4-negative escapees had significantly slower growth rate (data not shown), their impact remains unknown due to the limits of the model used.

Discussion

Chemokines and chemokine receptors expressed by tumors or tumor-containing matrix regulate a number of important functions that eventually affect disease outcome. They can directly induce cell proliferation (MIP1α/CCL3, I-309/CCL11), stromal cell-derived factor 1/CXCL12, TARC/CCL17) (20, 21) and regulate angiogenesis and infiltration of immune cells, etc. (see reviews (3, 36, 37)). Tumor-expressed chemokines attract inhibitory macrophages, NK T cells, and CD25+ CD4+ FoxP3+ Tregs, leading to severe immunosuppression and poor disease outcome in a number of cancers (23, 38). For example, infiltration of suppressive CCR4-expressing CD4+ CD25high Tregs is thought to be caused by CCL17 or CCL22 produced at the tumor site (1, 38). In contrast, tumors by themselves produce various immunosuppressive factors, or even, in the case of ATLL, some of them might originate from transformed immunosuppressive cells (1). These features are thought to negatively affect vaccine-induced antitumor cellular responses, leading to tumor escape despite the presence of tumor-specific cytolytic CD8+ T cells (39, 40). Therefore, alternative strategies that would directly kill malignant cells had been extensively explored. A surge of Ab-mediated therapeutics was initiated by the successful clinical use of rituximab, CD20-targeting Ab (for review, see Ref. 41). Tumor cells from ATLL patients were shown to be killed by incubation with mAb to CCR4, and mice injected with anti-CCR4 Ab were protected from CCR4-expressing tumors (1). This protection was based on induction of the Ab-dependent cellular cytotoxicity-mediated cell killing, and the CCR4-expressing human tumors were not uniformly killed in vitro (1), suggesting that clinical efficiency of anti-CCR4 Abs may be affected by the host FcR polymorphism (42, 43).

In this study, we report a different strategy that uses formulations designated chemotoxins, to preferentially and specifically eliminate chemokine receptor-expressing cells. As we have shown for chemokines fused with tumor Ags, the strategy was based on the ability of chemokines to deliver toxins into the cell cytosol. We recently reported that nonimmunogenic tumor Ags were rendered immunogenic and elicited therapeutic antitumor immunity, if they were targeted to chemokine receptors to use their internalization machinery (25, 33). For example, chemokine fusions that were targeted to CCR6 and CCR8 were internalized into endo/lysoosomal compartments, whereas some of them were degraded and presented to MHC class II molecules, whereas the remaining ligands were able to efficiently escape into cytosol to be degraded by proteasomes for MHC class I cross-presentation (25, 33). Our present data indicate that chemokines could also deliver RNases (EDN) to the cell cytosol and induce the death of cells that expressed the respective chemokine receptor. However, EDN chemotoxin in the present form may not be useful for the treatment of fast-growing malignancies, as its killing dynamics were significantly slow, presumably reflecting the nature of the EDN-induced apoptosis. It remains open to question whether activity of RNase-expressing chemotoxins would be augmented by increasing their RNase activity, protecting them from the intracellular proteases, or improving their delivery into the cytosol. It is tempting to speculate that the reduced RNase activity of chemokine EDN may be improved by generation of EDN-chemokine fusions, so as to free the N terminus of EDN to preserve its activity (our unpublished observation). In contrast, we hypothesized that the efficacy might be improved by use of faster acting moieties such as the exotoxin fragment PE38 of P. aeruginosa. The exotoxin was reported to kill mammalian cells within a short period of time by inhibiting protein synthesis via the ADP ribosylation of elongation initiation factor 2 (34). Importantly, PE38 appears to be nontoxic unless delivered into the cell cytosol (28, 35), and PE38 fusions with Abs to various cell surface Ags, such as CD25, CD22, and IL-4R, were successfully tested for treatment of hematological malignancies (44, 45) (for review, see Ref. 46). Interestingly, although CD22 is expressed on normal B cells and the majority of B cell malignancies, the anti-CD22 immunotoxin treatments only benefited patients with hairy cell leukemia, but not patients with non-Hodgkin’s lymphoma (NHL) and chronic lymphocytic leukemia (45). Similarly, immunotoxins that target CD25 were shown to be moderately effective in patients with NHL and cutaneous T cell lymphoma (46). It is noteworthy that CD25, IL-2Rα, is a necessary and important
survival receptor for cells that depend on IL-2, including immune T cells. Thus, these cells would be also depleted by the anti-CD25 treatment, and beneficial effects of CD25-targeting immunotoxins might be contradicted by the loss of tumor-specific T cells needed for eradication of residual or escaped tumors.

We have wanted to develop a less harmful and specific strategy to eradicate hematological tumors, such as NHL, chronic lymphocytic leukemia, cutaneous T cell lymphoma, and ATLL, delivering toxins via chemokine receptors. Although CCR4 is also expressed by ~20% of human peripheral blood CD4+ T cells (D. Baatar, unpublished data), depletion of CCR4-expressing cells by the systemic injections of TARC-PE38 did not cause apparent and adverse effects in C57BL/6 and BALB/c mice, because they were able to mount Ag-specific humoral and cellular immune responses (our unpublished data). In concordance, others recently reported that the Ab-mediated depletion of CCR4-expressing tumors (and normal cells) was well tolerated in mice (1). Recently, we have shown that the majority of human peripheral blood Tregs expressed CCR4, and their depletion using TARC-PE38 was sufficient to revert the suppressive state of CD8+ T cells (47). Therefore, the TARC-PE38 treatment would not only directly kill the CCR4-expressing leukemia cells, but also would eliminate Tregs to build beneficial conditions for activation of antitumor T cell responses, a necessary step for the elimination of the remaining escapes and residual disease. Of note: the efficacy to bind and kill murine CCR4-expressing cells was comparable regardless of the origin of TARC; both mouse and human TARC fusions with PE38 worked well and human TARC-PE38 effectively depleted murine CCR4+ CD4 T cells in immune-deficient BALB/c or C57BL/6 mice (data not shown and also see Ref. 47). Experimenting with human T cell tumors established in NOD-SCID mice, we demonstrated that TARC-PE38 can efficiently regress the CCR4-expressing cutaneous tumors in mice. We have also tested whether chemotoxins injected systemically can affect dissemination of tumors, although their therapeutic efficacy may be reduced by a relative short plasma t1/2 time. Intravenous injections of TARC-PE38, but not control CTACK-PE38, dramatically reduced the number of spleen-infiltrating CEM tumor cells, indicating that chemotoxins may be also useful for systemic treatment of nonsolid tumors. Because expression of CCR4 is often associated with a poor disease outcome, this may be an attractive strategy to control and combat the disease.

However, despite the fact that the TARC-PE38 treatment eliminated any signs of detectable/palpable tumor, the malignancy eventually relapsed, but with cells that no longer responded to additional and repeated treatments of TARC chemotoxin. Our data indicate that the resistance to TARC-PE38 correlates with the cells that do not express CCR4, although we cannot rule out the possibility that the treatment also selects variants that aberrantly express CCR4 that cannot internalize. Moreover, TARC-PE38 does not induce loss of CCR4 per se, but rather it is unable to affect the growth of the receptor-negative tumor variants, which represented a minor (~3–5%) population that existed before the treatment. Therefore, if a malignancy is heterogeneous, the treatment may lead to outgrowth of receptor-negative tumors, particularly in hosts with aberrant T cell immune responses. It remains to be tested whether escapes can survive and progress in hosts with an adequate immune system that would be activated during chemotoxin-induced massive cell death. Unfortunately, limitations of the model system used in this study do not allow us to address this question. Taken together, we present a novel concept that tumors can specifically be eradicated by delivering toxins through their chemokine receptors. This very simple and potent strategy can presumably be used for treatment of a wide variety of tumors, particularly hematological malignancies, which use chemokine receptors to metastasize or circumvent immunosurveillance. However, a complete eradication of tumors would probably be achieved by use of the strategy in combination with other treatment modalities such as chemotherapy. Finally, chemotoxins may also serve as a useful tool for understanding of biology and functions of various immune cells through their preferential and transient depletion, as it was recently used to elucidate biology of the skin-homing T regulatory cells (47).

Acknowledgments

We are grateful to Drs. Dan Longo and Ashani Weeraratna for helpful comments and suggestions; Ana Lustig (National Institute on Aging/National Institutes of Health) for critical reading of the manuscript; Dr. Ira Pastan (National Cancer Institute/National Institutes of Health) for the gift of PE38 cDNA and anti-TGFα-PE38; and Dot Bertak (National Institute on Aging/National Institutes of Health) for help with immunohistochemistry.

Disclosures

The authors have no financial conflict of interest.

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