Targeted Elimination of Prostate Cancer by Genetically Directed Human T Lymphocytes

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Abstract

The genetic transfer of antigen receptors is a powerful approach to rapidly generate tumor-specific T lymphocytes. Unlike the physiologic T-cell receptor, chimeric antigen receptors (CARs) encompass immunoglobulin variable regions or receptor ligands as their antigen recognition moiety, thus permitting T cells to recognize tumor antigens in the absence of human leukocyte antigen expression. CARs encompassing the CD3ζ chain as their activating domain induce T-cell proliferation in vitro, but limited survival. The requirements for genetically targeted T cells to function in vivo are less well understood. We have, therefore, established animal models to assess the therapeutic efficacy of human peripheral blood T lymphocytes targeted to prostate-specific membrane antigen (PSMA), an antigen expressed in prostate cancer cells and the neovasculature of various solid tumors. In vivo specificity and antitumor activity were assessed in mice bearing established prostate adenocarcinomas, using serum prostate-secreted antigen, magnetic resonance, computed tomography, and bioluminescence imaging to investigate the response to therapy. In three tumor models, orthotopic, s.c., and pulmonary, we show that PSMA-targeted T cells effectively eliminate prostate cancer. Tumor eradication was directly proportional to the in vivo effector-to-tumor cell ratio. Serial imaging further reveals that the T cells must survive for at least 1 week to induce durable remissions. The eradication of xenogeneic tumors in a murine environment shows that the adoptively transferred T cells do not absolutely require in vivo costimulation to function. These results thus provide a strong rationale for undertaking phase I clinical studies to assess PSMA-targeted T cells in patients with metastatic prostate cancer. (Cancer Res 2005; 65(19): 9080-8)

Introduction

Prostate cancer is the most frequent cancer in males in the United States and the cause of nearly 31,000 deaths per year (1). When diagnosed early, local adenocarcinomas can be effectively treated by radical extirpation or radiation (2, 3). Postsurgical residual disease requires radiation and/or hormonal therapy, which may prevent tumor progression and metastasis, but there is at present no curative treatment for hormone refractory, metastatic prostate cancer (4). Immunotherapy is a targeted therapy that is in principle an ideal approach for the treatment of minimal residual disease and micrometastases. The induction of cellular immunity requires that tumor antigens be presented in an immunogenic context, a far greater challenge for tumor antigens than for foreign molecules, such as viral antigens (5, 6). Successful immunization using dendritic cells or plasmid DNA vaccines (7–9) requires the presentation of human leukocyte antigen (HLA) class I–restricted peptides by both the antigen-presenting cell and the tumor itself. Immunogenic epitopes have been identified, at least for some HLA alleles, in antigens found in normal and transformed prostate cells, which include acid phosphatases, prostate-secreted antigen (PSA), prostate-specific membrane antigen (PSMA), and prostate stem cell antigen (10–13). Nonetheless, several obstacles remain to induce tumor immunity, which requires the expansion of cytotoxic T lymphocytes to numbers sufficient to mediate tumor rejection. Furthermore, the potency of the T-cell response may be undermined by several tumor escape mechanisms, which include antigen loss and HLA down-regulation, both of which deprive the T-cell receptor of its specific ligand on the surface of tumor cells (14).

The genetic engineering of T cells is a novel strategy designed to accelerate the generation of tumor-specific T cells and remedy the biological limitations that constrain the antitumoral functions of normal T cells. Chimeric antigen receptors (CARs) are essential constituents of this new armamentarium (15–18). Unlike the physiologic T-cell receptor, CARs encompass immunoglobulin variable regions or receptor ligands as antigen-recognition elements, thus permitting T cells to recognize cell surface tumor antigens in the absence of HLA expression (15–18). The signaling function of CARs has been questioned based on findings in transgenic mice and transduced primary T cells, indicating that antigen ligation was not always sufficient to activate resting naïve mouse T cells (19) or elicit interleukin 2 (IL-2) secretion in retrovirally transduced human peripheral blood T cells (20). We and others subsequently showed that ζ chain–based CARs could induce strong activation capable of sustaining T-cell proliferation and permitting secondary antigenic restimulation in vitro provided that antigen was presented in the context of CD82–mediated costimulation (21–23). It is not known, however, whether T cells expanded in this manner, particularly human T cells, could mediate tumor eradication in vivo, and whether further in vivo costimulation would be needed to sustain their function.
Here, we investigate the in vivo function of Pz1, a CAR-targeting human PSA. PSA is a highly attractive target antigen that is found in most prostate cancer cells, including hormone refractory metastatic disease (24, 25). The Pz1 receptor encompasses the \( \xi \) chain of the CD3 complex as its activation domain and specificallyredirects in vitro cytolysis against PSA-positive tumor cell lines (20). To investigate whether expanded Pz1+ T cells are active in vivo and require costimulation after adoptive transfer, we established three tumor models in severe combined immunodeficient (SCID)-bg/bg mice—orthotopic, s.c., and pulmonary. Direct administration of Pz1+ T cells in orthotopic and s.c. human prostate tumors eliminated a majority of the tumors. In a systemic tumor model in which mice rapidly succumbed to pulmonary disease, i.e., administration of Pz1-transduced T cells induced objective responses in all mice and cured a substantial fraction of them. Altogether, our results strongly support the feasibility of targeting prostate cancer with autologous T lymphocytes directed against PSA by a transduced \( \xi \) chain–based receptor.

**Materials and Methods**

**Cell Lines**

Retroviral producer cells were maintained in DMEM (Invitrogen, Carlsbad, CA) supplemented with 10% FCS. LNCaP and LNCaP/B7 cells were maintained in RPMI (Invitrogen) with 10% fetal bovine serum (FBS; Hyclone, Logan, UT). NIH3T3 fibroblasts (American Type Culture Collection, Rockville, MD) were maintained in DMEM with 10% donor calf serum (Hyclone). RM1, RML.PGLS, and EL4 were grown in high-glucose DMEM supplemented with sodium pyruvate, 10% FBS, and 2 mmol/L glutamine (Invitrogen). All media contained penicillin (100 U/mL) and streptomycin (100 \( \mu \)g/mL).

**Vectors**

The oncoretroviral vectors encoding Pz1, human PSA, 19z1, and CD80/1640 and seeded at a density of \( 1 \times 10^6 \)/mL with 1 \( \mu \)g/mL phytohemagglutinin (PHA). Isolated cells were analyzed in an independent observer.

**Peripherial Blood Lymphocyte Collection and Retroviral Transduction**

An average volume of 100 mL peripheral blood was drawn from healthy donors with informed consent. Peripheral blood mononuclear cells were isolated with Accu-Prep Lymphocytes (Accurate Chemical and Scientific Company, Westbury, NY) by low-density centrifugation and cultured in RPMI 1640 and seeded at a density of \( 2 \times 10^6 \)/mL with 1 \( \mu \)g/mL phytohemagglutinin. After 48 hours, T cells were exposed to gibbon ape leukemia virus envelope-pseudotyped Pz1 or 19z1 vector as previously described (20, 23, 26).

**Pz1+ T-cell Expansion**

LNCaP cells were transduced with the SFG-CD80 vector (20) and sortedbased on CD80 expression. The sorted cells, termed LNCaP/B7, were found to stably express CD80, obviating the need to periodically repurify CD80-expressing cells. Three days after peripheral blood lymphocyte (PBL) transduction, Pz1+ T cells were expanded on LNCaP/B7 cells in six-well tissue culture plates in the presence of 50 units/mL IL-2 (supplemented every 2-3 days). This cycle was done once before adoptive transfer or repeated on day 7 for in vitro restimulation studies. 19z1+ T cells were expanded on fibroblast-derived artificial antigen-presenting cells expressing CD19 and CD80 as described (26).

**Cytotoxicity Assays**

Transduced T-cell effector function was measured in standard \( ^{51} \)Cr release assays using EL4, EL4PSMA (23), RM1, RML.PGLS, and LNCaP as targets. The T cells were incubated with \( ^{51} \)Cr-labeled target cells for either 4 or 12 hours. Specific \( ^{51} \)Cr release was calculated using the following formula: percentage of specific lysis = \( (cpm \text{ experimental release} - cpm \text{ spontaneous release}) / (cpm \text{ maximum release} - cpm \text{ spontaneous release}) \) \times 100. The SD of triplicates was in all cases <5%. The effector-to-target cell ratio was based on the determination of CD8+ Pz1+ or CD8+ 19z1+ T cells by fluorescence-activated cell sorting (FACS) analysis. For the antibody blocking experiment, targets cells were preincubated for 30 minutes (24-26) with 10 \( \mu \)g/10^6 cells of anti-MHC class I (clone W6/32; Axxora, San Diego, CA) or purified mouse IgG1 as control.

**Flow Cytometry**

Expanded PBLs were stained for the expression of differentiation markers immediately before injection into SCID beige mice. Pz1 and 19z1 cell surface expression was measured using a cross-reactive goat anti-mouse IgG phycoerythrin conjugate (CN Biosciences, La Jolla, CA). The following antibodies were used for subset identification: anti-CD8-APC, anti-CD27-phycoerythrin (PE), anti-CD4-APC, anti-CD4-PE-Cy5, anti-CD28-PE (Caltag, La Jolla, CA), anti-CCR7-PE (R&D Systems, Minneapolis, MN), and anti CD45RA-PE-Cy5 (eBioscience, San Diego, CA), and, as isotype controls, PE-conjugated mouse IgG2b (BD Biosciences) or mouse IgG1-PE. Intracellular perforin expression was assessed after fixation with CYTOFIX buffer and permeabilized with PERM/WASH buffer (both from BD Biosciences) using a PE-conjugated mouse anti-human perforin monoclonal antibody (mAb; BD Biosciences).

**Mouse Models**

**Orthotopic surgery.** Male SCID beige mice between 5 and 8 weeks old (Taconic, Haverford, NY) were anesthetized with i.p. injection of ketamine and xylazine solution. The dorsolateral aspect of the prostate was injected with 1 \( \times 10^6 \) LNCaP cells in 25 mL RPMI + 10% FCS buffer. Tumors were allowed to grow for 10 days to 3 weeks before injection with T cells. Serum PSA-positive mice were randomized to receive either Pz1 or 19z1 cells. Pz1 and 19z1-transduced T cells were injected directly into the prostate tumors at equal doses of 20 \( 10^6 \) total T cells (6-8 million CD8+ Pz1+ T cells) given in two injections over 48 hours. Mice were then followed until death or sacrificed if they showed any signs of distress, as determined by an independent observer.

**Subcutaneous model.** One million LNCaP/C4-2 cells were injected s.c. Mice were sacrificed when the tumors reached a volume of 1 cm^3.

**Lung model.** Five thousand RM1.PGLS cells were administered i.v. on day 0. The RM1 cells were retrovirally transduced to express human PSA and Egfp/Luc and sorted to high purity (99% PSA*, Egfp/Luc*). Pz1+ CD8+ PBLS (5 \( \times 10^5 \)-\( 8 \times 10^6 \)) were administered i.v. at 24, 48, and 72 hours after tumor injection. For the 1:10 ratio (Pz1:19z1) experiment (Fig. 5C), we injected i.v. 20 \( 10^6 \) 19z1 T cells on days 2, 4, and 5, and 10 \( 10^6 \) 19z1 and 5 \( 10^6 \) CD8+ Pz1+ T cells on days 1 and 2; and 5,000 RML.PGLS cells were administered i.v. on day 0.

**Lung T-cell retrieval.** Single cell suspensions of tissues from sacrificed animals were made by macerating tissues through a 40 μm nylon cell strainer (Becton Dickinson, Franklin Lakes, NJ). Isolated cells were analyzed by FACS for expression of human CD3 and CD45 using PE-labeled anti-human CD3 and FITC-labeled anti-human CD45 mAbs (Caltag).

**Magnetic Resonance Imaging**

**Orthotopic model.** Mice selected at random from the two treatment groups were anesthetized with a single i.p. injection of ketamine (90 mg/kg) and xylazine (9 mg/kg). Thirteen to 15 mice were then positioned together within a custom-built three-turn parallel wound foil solenoid. The studies were done on a GE Signa 1.5 T clinical scanner (GE Medical Systems, Milwaukee, WI) operating at 63.9 MHz. Axial prostate images were acquired within a custom-built three-turn parallel wound foil solenoid. The studies were also done on a GE Advantage Workstation 4.0_03 software.

**Lung model.** Two mice from each group were selected at random for magnetic resonance imaging (MRI) on days 7 and 13 posttumor inoculation. Each mouse was anesthetized using 2% isoflurane (Forane, Baxter Healthcare Corp, Deerfield, IL) and positioned within a four-turn parallel wound foil solenoid. These studies were also done on a GE Signa 1.5 T
clinical scanner. Axial images were acquired using a spoiled gradient echo sequence with TR = 100 milliseconds, TE = 6.3 milliseconds, flip angle = 20°, NEX = 14, FOV = 4 × 4 cm, slice thickness = 1.0 mm, slice gap = 1.5 mm, matrix size = 256 × 256. To better quantify tumor burden and distribution at day 7, a mouse was positioned within a Bruker Biospec spectrometer (Bruker, Billerica, MA) operating at 7.0T. Contiguous axial images were acquired using a spoiled gradient echo sequence with TR = 179 milliseconds, TE = 5.2 milliseconds, flip angle = 30°, NEX = 48, FOV = 3 × 3 cm, slice thickness = 0.8 mm, slice gap = 1 mm, matrix size = 256 × 256. The tumor volumes were quantified using Bruker Paravision 3.0.1 software.

Microcomputed Tomography Imaging
Noncontrast small-animal X-ray computed tomography (CT) imaging was done using the MicroCAT II scanner (ImTek, Inc., Knoxville, TN), with an X-ray source having a 50 μm tungsten anode and operated at 55 kVp and 800 μA and a phosphor detector optically coupled to a 1,828 × 3,488 charge-coupled device. A total of 360 conebeam projections were acquired with an exposure time of 750 milliseconds each. Images were reconstructed three-dimensionally in a 512 × 512 × 1,024 matrix using the Feldkamp conebeam filtered back-projection algorithm. Total CT acquisition time was ~10 minutes and the spatial resolution (full-width half-maximum) was 100 μm. Visualization and analyses of CT images were done using the Amira (Konrad-Zuse-Zentrum für Informationstechnik Berlin and Indeed Visual Concepts GmbH) software.

Immunohistochemistry
Fresh tissue samples were embedded in optimal cutting temperature compound and frozen in isopentane cooled in liquid nitrogen before storing at −70°C. Sections of 8 to 10 μm were cut on a cryostat at −20°C. The sections were laid onto Vectobond slides (Sigma, St. Louis, MO) and dried in air for 15 minutes. The sections were fixed in cold acetone for 2 minutes and then immersed in 0.3% hydrogen peroxide in 0.1% (w/v) sodium azide for 10 minutes. Slides were washed with PBS and incubated with 5% bovine serum albumin (BSA) in PBS for 60 minutes at room temperature. Following the removal of BSA, 5 μg/ml of anti-PSMA J591 mAb* (generously provided by Dr. Neil Bander, Weill Medical College of Cornell University) was applied followed by thorough washing with PBS before application of horseradish peroxidase-conjugated goat anti-human IgG mAb (Jackson Immunoresearch, West Grove, PA) at a 1:1,000 dilution for 60 minutes. After washing with PBS, 0.06% diamobenzidin tetrachloride and 0.01% hydrogen peroxide in PBS were applied onto the specimens for 3 minutes. The sections were then washed, lightly stained with Carazzi’s hematoxylin, dehydrated, and mounted.

Quantitative Bioluminescence
Bioluminescence images were collected on a Xenogen IVIS Imaging System (Xenogen, Alameda, CA) using a cryogenically cooled charge-coupled device camera. Living Image software (Xenogen) was used to acquire and quantitate the bioluminescence imaging data sets. Ten to 15 minutes before the time of imaging, a single i.p. injection of 150 mg/kg D-luciferin (Xenogen) in PBS was administered to each mouse. Subsequently, the animals were induced using 2% isoflurane (Forane, Baxter Healthcare) and positioned within the imaging chamber under anesthesia. Two to five mice were imaged simultaneously with exposure times ranging from 0.5 to 3 minutes. Ten, 15, or 25 cm field of view and low, medium, or high binning levels were applied to maximize sensitivity and spatial resolution. Both dorsal and ventral images were obtained for each animal. The dorsal and ventral signals were separately quantified through region of interest (ROI) analysis over the thoracic region. The resulting signal summations (in units of photons/s) were normalized to the ROI area so that all measurements are given in photons/second/cm². For each image, normalized background signal from similarly sized ROIs was subtracted.

Statistics
Statistical analyses and curve fittings were done using GraphPad Prism version 4.00 for Windows (GraphPad Software, San Diego, CA). Survival curves were created using the product limited method of Kaplan and Meier and compared using the log-rank test. Linear regression analysis was used to fit the quantitated bioluminescence data to the FACS-determined tumor cell counts, whereas nonlinear regression analysis was used to fit the quantitated bioluminescence data to the FACS-determined T-cell counts.

Results
Specific expansion of prostate-specific membrane antigen–specific primary T lymphocytes. We previously showed that Pz1-transduced human primary T cells efficiently lyse PSMA-positive target cells in vitro, including LNCaP cells and PSMA-transduced PC3 cells, EL4 cells, and NIH 3T3 fibroblasts (20, 23). However, these Pz1+ PBLs failed to expand in response to repeated antigenic stimulation in the absence of CD28-mediated costimulation (23). As shown in Supplementary Fig. S1A, Pz1-transduced PBLs likewise failed to sustain proliferation in the presence of LNCaP cells, but expanded 8 to 10-fold in a single round of in vitro stimulation with CD80/B7.1-expressing LNCaP cells (LNCaP/B7). Both CD4+ and CD8+ Pz1 T cells present in these PBL cultures expanded, starting from a 2:1 ratio and typically reaching a 1:2 ratio by day 14. When cultured separately, the CD4+ Pz1 T cells expanded in response to PSMA + CD80 in two consecutive stimulations, but not the CD8+ Pz1 T cells, which showed diminished expansion upon restimulation despite the addition of exogenous IL-2 (Supplementary Fig. S1B). Thus, absolute expansion was consistently the highest when CD4 and CD8 T cells were cocultured in the presence of PSMA and CD80. For the in vivo studies shown below, the transduced PBLs were exposed once to LNCaP/B7 cells before adoptive transfer. T cells expanded in this manner were highly specific for xenogeneic PSMA+ tumor cells and efficiently lysed hematopoietic and epithelial mouse tumor cells expressing human PSMA (Fig. 1A-C). Pz1+ T cells also lysed LNCaP cells (Fig. 1D). Lysis of LNCaP cells was not reduced in the presence of blocking antibody against HLA class I (Supplementary Fig. 1D), indicating that alloreactivity did not significantly contribute to target cell recognition and further demonstrating that tumor cell lysis occurred in an HLA-independent manner. T cells transduced with a control CAR specific for CD19 showed little or no cytotoxic activity against the same target cells (Fig. 1).

Expanded Pz1+ T cells comprise activated and effector memory T cells. Based on their cell surface phenotype, CD8+ T cells can be assigned to five distinct differentiation stages (28): naïve (CCR7+; CD45RA+), central memory (T CM; CCR7+; CD45RA+), early effector T cells (CCR7+; CD45RA+; CD28+); effector memory T cells (T EM; CCR7–; CD45RA+; CD28+); and CD45RA+ effector memory cells (T EMRA; CCR7–; CD45RA+; CD28–) (28). The expanded Pz1+ CD8+ T cells were characterized on the day of injection, typically 5 to 7 days after initiating cocultivation with LNCaP/B7 cells. The vast majority of the Pz1+ CD8+ T-cell population was CCR7+ CD45RA–, about half of which were positive for CD28, thus corresponding to early effector and effector memory phenotypes, respectively (Supplementary Fig. S2A-C). The expanded Pz1+ T cells markedly up-regulated perforin expression compared with untransduced T cells (Supplementary Fig. S2D), consistent with their high lytic potential (Fig. 1).

Pz1+ peripheral blood lymphocytes eradicate established orthotopic LNCaP tumors. To assess the in vivo function of Pz1-transduced human PBLs, we first established an orthotopic model using the PSA-secreting prostate adenocarcinoma cell line LNCaP.C4-2 (29). Following surgical implantation of 1 million cells, PSA serum levels and tumor size were monitored by RIA and MRI, respectively. The tumors were ~0.02 to 0.03 cm³ when serum PSA became unequivocally positive (>0.1 ng/ml) and increased to a median value of 0.75 cm³ within 5 weeks, by which time tumor-bearing animals had to be sacrificed. The correlation between tumor weight estimated by MRI and tumor weight measured after necropsy was linear ($r^2 = 0.96$, $P = 0.002$). Ten days after tumor...
inoculation, serum PSA-positive mice were randomized to two treatment groups, receiving either Pz1 or control 19z1-transduced PBLs. Ten million transduced PBLs were administered locally on days 11 and 12 after tumor implantation. Fifty-four mice were treated (n = 26 for Pz1 and n = 28 for 19z1). In half of the mice, tumor progression was followed weekly by MRI (Fig. 2A and B).

The median survival in the Pz1 group was 211 days (range 28-272 days), in contrast to 53 days in the 19z1 group (range 23-147 days; P < 0.0001). Fifteen mice treated with Pz1-transduced PBLs (57%) remained without evidence of disease by the end of the study on day 272, which was corroborated by absence of detectable serum PSA (data not shown) and by MRI. No mouse survived in the 19z1 control group (Fig. 2C). The 19z1-treated animals showed the same growth rate as untreated mice (data not shown). Interestingly, the tumors in Pz1-treated mice initially showed modest growth, reaching 0.08 cm³ on average (range 0.01-0.6 by day 18) followed by shrinkage and tumor elimination in the long-term survivors.

**Pz1-transduced peripheral blood lymphocytes eradicate established s.c. LNCaP tumors.** To test T-cell efficacy at other tumor sites and assess the tumor response in relation to tumor size, we established a s.c. tumor model. Serum PSA was detectable 7 to 14 days after implantation of 5 × 10⁶ LNCaP/C4-2 cells and tumors were palpable by days 10 to 14. As in the orthotopic model, the relationship between serum PSA and tumor volume was linear 15 and 30 days after tumor implantation (data not shown).

A first set of 22 mice with an average tumor size of 0.07 cm³ (mean PSA, 1.7 ng/mL) was treated with 10 million human PBLs (either Pz1 transduced or untransduced) by direct intratumoral injection on 2 consecutive days. The tumors progressed to >1 cm³ in 11 of 11 mice treated with control PBLs, correlating with a rapid rise in serum PSA. In contrast, the tumors were successfully eradicated in 6 of 11 (55%) of the mice receiving Pz1-transduced lymphocytes as evidenced by impalpable tumor and undetectable serum PSA for at least 180 days (P < 0.0001; Fig. 3A). Moreover, the tumors remaining at day 60 in the Pz1 + T cell–treated mice were smaller than the tumors in the control cohort (mean 0.23 versus 1.61 cm³, P = 0.02).

![Figure 1. In vitro cytolytic activity of Pz1-transduced PBLs. Target cell lysis by Pz1-redirected PBLs.](http://www.aacrjournals.org)
A second set of animals with a larger tumor burden (mean size, 0.28 cm³; mean PSA, 3.6 ng/mL) was treated with the same T-cell dose ($n = 8$ with Pz1 PBLs and $n = 8$ with mock transduced PBLs). Half (four of eight) of the established tumors were eradicated by the Pz1-transduced T cells, remaining free from both local and systemic recurrence for at least 180 days, whereas none of the eight tumors were eradicated by the control T cells ($P = 0.0043$; Fig. 3B).

**Systemically administered Pz1 peripheral blood lymphocytes can eradicate pulmonary tumors.** To investigate the bioactivity of systemically delivered T cells, we established a lung metastasis model. The murine prostate cell line RM1 (30) was retrovirally transduced to express human PSMA as well as the GFP/Luciferase fusion protein (RM1.PGLS) to facilitate tumor tracking and the quantification of tumor burden. RM1.PGLS cells were detectable in the lung within 24 hours of i.v. administration and remained localized in the lung parenchyma, developing as nodules detectable by MRI and CT scan within 7 days (Fig. 4A). The tumors gradually invaded the lung parenchyma, causing major airway obstruction by day 13, at which time the animals were sacrificed (Fig. 4B). T cells transduced with either Pz1 or 19z1 were infused i.v. on days 1, 2, and 3 at a dose of $5 \times 10^6$ to $8 \times 10^6$ transduced PBLs per day. All mice treated with Pz1 ($n = 16$) responded to treatment (Fig. 5A). One third remained tumor-free (survival >90 days without evidence of tumor), whereas the others showed significantly delayed tumor progression (mean survival time doubling from 12 to 23 days; Fig. 5A). Control 19z1 + T cells did not delay tumor progression relative to untreated mice (Fig. 5A), underscoring the specificity imparted by the CAR. Conversely, Pz1 + T cells were without effect on PSMA/C0, CD19 + Raji cells, which in turn were eliminated by the 19z1-transduced T cells (Fig. 5B). To assess whether the lymphodepleted milieu of the recipient SCID mice facilitated tumor eradication by Pz1 + T cells, we did an additional experiment in which mice received either Pz1 + T cells or Pz1 + T cells plus 19z1 + T cells at a 1:10 ratio. As shown in Fig. 5C, there was no significant survival difference between mice treated with Pz1 T cells ($n = 8$) and mice treated with Pz1 T cells and 19z1 T cells ($n = 12$; Fig. 5C). No tumor was detectable by quantitative bioluminescence in long-term survivors with Pz1 T cells or Pz1’19z1 T cells (data not shown).

**Prostate-specific membrane antigen-specific T cells take over a week to eradicate established tumors.** To analyze the kinetics of tumor elimination, we measured tumor progression by quantitative bioluminescence in all Pz1-treated mice ($n = 16$).
as well as 19z1-treated mice \((n = 9)\) and untreated mice \((n = 8)\). Tumor progression was indistinguishable between the two latter groups (Fig. 6), indicating that non-PSMA targeted T cells had no effect on the tumor. Eleven of the Pz1-treated mice showed partial responses or complete remission followed by relapse (Fig. 6A). Of these, two were negative in two consecutive imaging sessions (days 10 and 13), followed by delayed, fatal tumor progression. Examination of lung tissue by immunohistochemistry showed that the progressing tumors remained by and large positive for PSMA, thus excluding antigen down-regulation as the mechanism of tumor escape. Importantly, we found that mice brought into transient complete remission \((n = 2; \text{Fig. 6A})\) as well as mice eventually cured by Pz1 treatment \((n = 5; \text{Fig. 6B})\) showed continued tumor progression in the 3 to 10 days following T-cell administration before tumors started regressing. These data established that the T cells were not immediately effective and implied that therapeutic T cells must persist for at least 1 week to eradicate the tumor. This conclusion was further supported by measuring residual T-cell numbers in mice responding to therapy. We enumerated lung T cells and quantified tumor mass in seven residual T-cell numbers in mice responding to therapy. We enumerated lung T cells and quantified tumor mass in seven mouse experiments, mice with orthotopic \((n = 19)\), s.c. \((n = 26)\), and lung metastases—we show that Pz1 but not 19z1-transduced T cells induce durable remissions and cures in a substantial fraction of the treated animals. In all of our experiments, mice with orthotopic \((n = 26)\), s.c. \((n = 19)\), or pulmonary tumors \((n = 16)\) received a total of 18 to 20 million Pz1−T cells administered in either two (Figs. 2 and 3) or three (Figs. 4 and 5) consecutive injections. Complete responses and long-term survival were achieved in 55% of the direct infusions in prostate or skin \((n = 45)\). In the partial responders, survival was extended up to 200 days in mice with orthotopic tumors (average tumor size of 0.02 cm$^3$; Fig. 2), to 110 days in mice with 0.04 cm$^3$ s.c. tumors (Fig. 3A), and to 65 days in mice with 0.28 cm$^3$ s.c. tumors (Fig. 3B). These results indicate that the therapeutic response rate is proportional to the intratumoral effector-to-target cell ratio.

Significantly, Pz1-transduced human T cells could not engage their endogenous T-cell receptor when recognizing PSMA on RM1.PGLS cells, thus excluding any role for the endogenous T-cell receptor and establishing that tumor elimination was strictly mediated by the Pz1 receptor. The 19z1+ T cells had no effect on tumor progression in prostate or in lung, which underscores the targeting specificity afforded by Pz1 and excludes nonspecific inflammation as a mechanism sufficient to account for tumor regression, thus further underscoring the targeting specificity afforded by Pz1.

Antigen-induced \textit{ex vivo} expansion of the transduced cells required costimulation, which was provided here by expressing CD80 in LNCaP tumor cells (Fig. 1). A critical question for immunotherapy is whether adoptively transferred T cells require \textit{in vivo} costimulation to achieve tumor protection. This important question is difficult to assess because of the difficulty of pinpointing whether the infused T cells interact with antigen cross-presented by dendritic cells or other antigen-presenting cells at any time before their contact with the tumor, irrespective of the costimulatory ligands presented by the latter (31–34). Here, we took advantage of non–MHC-restricted antigen recognition and a xenogeneic setting to address this question, establishing \textit{in vivo} models in which cross-presented LNCaP antigens could not stimulate the Pz1 antigen receptor. To
address whether the “immunologic space” available in recipient SCID mice contributed to this therapeutic response, we did an experiment in which recipient mice were “filled” with a 10:1 excess of irrelevant 19z1+ T cells (100 million T cells per recipient, which alone had no effect on tumor progression). As shown in Fig. 5C, this massive infusion of nontargeted T cells did not reduce tumor eradication, establishing that Pz1+ T cells were still fully effective under such competitive conditions.

Our results also show that the adoptively transferred T cells eliminated established tumor cells in the absence of in vivo costimulation, as shown by the eradication of parental (CD80-negative) LNCaP cells as well as nonhuman, PSMA-positive RM1.PGLS tumors. This finding thus establishes that human T cells redirected through a ζ chain–based CAR are able to mediate tumor rejection without further in vivo costimulation. This observation does not rule out a role for in vivo costimulation of adoptively transferred T cells in tumor eradication, but establishes that, if T cells are provided in sufficient numbers and if they are adequately activated ex vivo before infusion, further in vivo costimulation is not necessary. Costimulation may nonetheless be important to promote T-cell proliferation, survival, and/or memory formation (15), thereby increasing or sustaining the effector-to-tumor cell ratio.

We found that tumor elimination was indeed closely correlated with short-term T-cell persistence after T-cell transfer. Quantitative bioluminescence imaging was used to monitor tumor burden in the RM1.PGLS lung model. Serial imaging studies showed that the tumors decreased to undetectable levels within 7 days in 7 of 16 Pz1-treated mice (Fig. 6). Among these seven complete responders, five mice showed continued tumor progression over the course of 1 week following the last T-cell infusion. This finding establishes that the therapeutic impact of T cells is neither immediate nor even very rapid, and further implies that, in this model, the T cells have to survive and remain functional for at least 7 days to exert their therapeutic effect. This time lag may represent a period during which tumor-specific T cells expand until a sufficient effector-to-target cell ratio is reached, or, alternatively, the period needed for T cells to recirculate and gradually reach each and every tumor site, which is required to achieve a cure. Two of seven mice in complete remission relapsed and showed progressive disease by day 18 after having scored negative on at least two consecutive imaging sessions on days 10 to 14 (Fig. 6A). Collectively, our measurements of tumor progression (Fig. 6) and correlation of T-cell count and tumor burden suggest that the T cells eliminated most, but not all, of the tumor cells by day 10 in these two animals and that the number of remaining T-cells did not impede subsequent tumor relapse from minimal disease because of an insufficient T cell-to-tumor ratio.

There is presently no effective treatment for disseminated prostate cancer. Approximately 25% of newly diagnosed prostate cancer cases present with metastatic disease (35) and 30% to 50% of patients who undergo definitive therapy for prostate cancer will show evidence of metastatic disease (36). Metastatic prostate cancer involves the bone and lung in over 90% and 46% of cases, respectively. Other frequent sites of metastases include the liver, pleura, and adrenal glands (37). Whereas metastatic disease is initially hormone sensitive, the natural history is progression to a hormone refractory state where the median survival is 12 to 18 months (38). Current systemic therapies have limited impact on survival (37).

Figure 5. RM1.PGLS lung model. Survival after i.v. administration of Pz1- versus 19z1-transduced PBLs. A, RM1.PGLS lung model: Pz1+ PBLs (n = 16, ■), 19z1+ PBLs (n = 9, ●), or no treatment (n = 8, ○). Pz1-transduced T cells eliminate established RM1.PGLS, but not 19z1+ PBLs (P < 0.0001, Pz1 versus 19z1 or untransduced T cells). B, Raji medullary model (25): Pz1- PBLs (n = 10, □) and 19z1+ PBLs (n = 20, ■). 19z1-transduced T cells eliminate established Raji, but not Pz1- PBLs (P < 0.0001). C, RM1.PGLS lung model: Pz1+ PBLs (n = 8, ◊), Pz1+ PBLs in combination with a 1:10 ratio of 19z1+ PBLs (n = 12, ▲) or 19z1+ PBLs (n = 4, ■). Survival is not statistically different between the two former groups.
Our studies show that Pz1 T cells are highly specific for PSMA and capable of mediating tumor elimination in vivo. Having established an efficacious method for expanding functional, PSMA-specific human T cells, we show antitumor function in three tumor models, including a systemic disease model. The data presented here establish that genetically modified human T cells targeted against PSMA not only suppress tumor growth, but markedly increase long-term survival. The genetic targeting of T cells is a strategy designed to rapidly generate populations of tumor-specific T lymphocytes (15–18). T cells may be engineered with physiologic T-cell receptors that are HLA-restricted or CARs that recognize cell surface antigens independent of HLA, as we do here. This latter approach is attractive for its potential applicability to any patient, irrespective of HLA haplotype, but is restricted to cell surface antigens. These include carcinoembryonic antigen (39), folate-binding protein (40, 41), erbB2-3-4 (42, 43), and Tag-72 (44), which have been targeted in vivo by genetically modified murine T lymphocytes (reviewed in ref. 45).

Human T cells were shown to mediate in vivo tumor elimination in two recent studies targeting s.c. (42, 46) or medullary prostate tumors (47). In another tumor model, in which tumors localize in the medullary cavity (26), we showed that CAR-modified T cells could effectively eliminate bone marrow tumors. In aggregate, our results show that CAR-modified T cells are active in prostate, skin, lung, and bone marrow. PSMA is an attractive target for novel therapeutic applications because it is expressed in all prostate cancers and its expression levels increase in more poorly differentiated prostate cancers, including hormone refractory disease. Bander et al. (48) recently reported on the targeting of metastatic prostate cancer with a radiolabeled PSMA-specific mAb. The J591 mAb was found to accurately target bone or soft tissue lesions in at least 94% of evaluable tumor sites. Toxicity was limited to dose-related, reversible myelosuppression thought to be secondary to the antibody radiolabel. No patients developed anti-idiotypic antibodies (49). Thus, whereas PSMA is also expressed in normal prostate, the kidney proximal tubule, and

Figure 6. Quantitative bioluminescence analysis of tumor response to PSMA-targeted adoptive T-cell therapy. In (A and B), the two thick lines represent the mean thoracic optical signal measured in mice receiving either no T cells or 19z1-transduced T cells (average of eight and nine mice, respectively; the SDs are very small and not apparent). All other lines represent serial imaging in single mice treated with Pz1 T cells. In (A), partial or complete responder mice that rapidly progressed or relapsed after complete remission \((n = 11)\). In (B), quantitative bioluminescence imaging in five mice that were ultimately cured by Pz1+ PBLs (see Fig. 5). In several mice, continued tumor progression was noted in the imaging sessions following the T-cell infusions (vertical arrows), before the tumors eventually began regressing. In (C) and (D), T-cell and tumor cell counts were obtained by FACS assays done on whole lungs removed from another set of seven animals immediately following bioluminescence imaging on day 17. Regression analyses of bioluminescence tumor signal intensity versus lung RM1.PGLS tumor counts or lung Pz1+ T-cell counts are shown in (C) and (D), respectively. C, linear relationship between bioluminescence imaging signal intensity and the number of tumor cells found in the lung 16 to 17 days after the injection of RM1.PGLS cells \((y = 4.2e4 + 1.25e6x, r^2 = 0.92)\). D, decreasing monoexponential relationship between bioluminescence imaging signal intensity and the relative number of Pz1 receptor–positive T cells found in the lung at the same time points \((y = 1e7exp(-0.077x)), r^2 = 0.93\).
the jejunal brush border (25), and putatively in the central nervous system (50), no related toxicities have been observed in man.

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