Effector CD8\textsuperscript{+} T-cell Engraftment and Antitumor Immunity in Lymphodepleted Hosts Is IL7R\textalpha Dependent

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Abstract

Adoptive cellular therapy, in which activated tumor-reactive T cells are transferred into lymphodepleted recipients, is a promising cancer treatment option. Activation of T cells decreases IL7 responsiveness; therefore, IL15 is generally considered the main driver of effector T-cell responses in this setting. However, we found in lymphodepleted mice that CD8\textsuperscript{+} T cells activated with IL12 showed enhanced engraftment that was initially dependent on host IL7, but not IL15. Mechanistically, enhanced IL7 responsiveness was conferred by elevated IL7R\textbeta expression, which was critical for antitumor immunity. Elevated IL7R\textalpha expression was achievable without IL12, as polyclonal CD8\textsuperscript{+} T cells activated with high T-cell receptor (TCR) stimulation depended on T-cell IL7R\textalpha expression and host IL7 for maximal engraftment. Finally, IL12 conditioning during the activation of human CD8\textsuperscript{+} T cells, including TCR-modified T cells generated using a clinically relevant protocol, led to enhanced IL7R\textalpha expression. Our results demonstrate the importance of the donor IL7R\textalpha/host IL7 axis for effector CD8\textsuperscript{+} T-cell engraftment and suggest novel strategies to improve adoptive cellular therapy as a cancer treatment.


Introduction

The cytokines IL7 and IL15 are both critical for T-cell homeostasis (1–5). In the context of adoptive T-cell therapy (ACT), involving transfer of effector T cells into lymphodepleted hosts, the relative importance of each cytokine for T-cell support has not been fully elucidated; however, several lines of evidence suggest IL15 is more critical. First, activated T cells downregulate IL7R\textalpha (CD127) and upregulate IL2/15R\textbeta (CD122), leading to a gain in IL15 responsiveness but concomitant loss in IL7 responsiveness (6–8). Second, IL15 has been shown to be more important for antitumor efficacy than IL7 in a preclinical ACT model (6, 9). Third, memory CD8\textsuperscript{+} T cells predominantly require IL15 for proliferation in lymphodepleted hosts (10, 11). Next, multiple studies have demonstrated that IL7 and/or IL7R\textalpha are not critical for the accumulation of effector CD8\textsuperscript{+} T cells at the peak of an antiviral immune response (12–14). Finally, IL15 more potently and specifically maintains effector CD8\textsuperscript{+} T-cell numbers at the culmination of infection compared with IL7 (15). On the basis of these studies, IL15 would be predicted to be more relevant than IL7.

Priming activated T cells with the Th1/Tc1 polarizing cytokine IL12 (16, 17) dramatically improves the persistence and antitumor efficacy of CD8\textsuperscript{+} T cells after adoptive transfer (18–20). As IL7 and IL15 are elevated after lymphodepletion (21–23), this enhanced persistence may be due to an increase in the expression of IL2R\textalpha and/or IL7R\textalpha induced by IL12 (8, 24). Although IL2R\textbeta has consistently been shown to be increased by IL12 (25, 26), data concerning IL7R\textbeta are conflicting. Several studies have found that IL12 exposure decreased IL7R\textbeta levels (27–30), although in other settings IL12 increased IL7R\textbeta on activated CD8\textsuperscript{+} T cells (25, 26, 31). Thus, the impact of IL12 on the ability of CD8\textsuperscript{+} T cells to respond to the homeostatic cytokines IL7 and IL15 warrants further consideration.

In this study, we investigated the cytokine requirements of effector CD8\textsuperscript{+} T cells in murine lymphodepleted hosts. We initially focused on CD8\textsuperscript{+} T cells conditioned with IL12 because these cells expand robustly in a lymphodepleted host without a requirement for exogenous cytokines or vaccination (18). This strategy revealed that activated CD8\textsuperscript{+} T cells require host IL7, but not IL15, for maximal initial expansion in a lymphodepleted host. Accordingly, the persistence and antitumor activity of these cells was dependent on IL7R\textbeta. These findings are generalizable and translatable, as polyclonal CD8\textsuperscript{+} T cells activated in the absence of IL12 were also dependent on IL7/IL7R\textalpha for initial engraftment, and human T cells cultured with IL12 acquired superior IL7 responsiveness. These findings have direct implications for the design of future adoptive cellular therapy trials for cancer therapy.
**Materials and Methods**

**Mice**

C57BL/6 (B6), B6.PL (Thy1.1), pmel-1 T-cell receptor (TCR) transgenic (32), β2-microglobulin−/− (B2m−/−), and IL7Rα−/− mice were obtained from The Jackson Laboratory. IL15−/− mice were purchased from Taconic. H3T TCR transgenic mice were generated as previously described (33). Pmel-1 mice were maintained by crossing a pmel-1 (male) to a Thy1.1 (female) generating hemizygous offspring. IL7Rα−/− heterozygous mice were generated by crossing an IL7Rα−/− male to either a Thy1.1/1.1 homozygous B6 female (generating B6 IL7Rα−/− Thy1.1/1.2 mice) or a pmel−/− Thy1.1/1.1 homozygous female (generating pmel−/− IL7Rα−/− Thy1.1/Thy1.2 mice). All mice were used between 6 and 16 weeks of age. Mice were housed under specific pathogen-free conditions in accordance with institutional and federal guidelines at the Medical University of South Carolina (MLISC, Charleston, SC).

**Cell cultures**

B16-F1 tumor cells were obtained from the ATCC and immediately expanded and frozen down into a large number of aliquots. Cells were verified to be *Mycoplasma* free and one aliquot was briefly expanded for each experiment using culture conditions as previously described (18). All T cells were grown in RPMI-1640 complete media as described previously (18). For generation of mouse gp100-reactive T cells, pmel-1 TCR transgenic splenocytes (1 × 10^6 cells/mL) were stimulated with 1 μg/mL H-2D^d-restricted human gp100_25-32 peptide (KFPKNQDWL; American Peptide Company) for 3 days with or without IL12 (10 ng/mL; Shenoadoah Biotechnology) to generate Tc1 or Tc0 T cells, respectively. For generation of mouse tyrosinase-reactive T cells, h3T TCR transgenic splenocytes (1 × 10^6 cells/mL) were stimulated with 1 μg/mL anti-CD3 mAb (37.51) directly or by transduction of a previously described protocol (REP) by incubation in a cytokine-containing media to maintain a concentration of 1 × 10^5 cells/mL. After 3 days of activation, cytokine responsiveness and phenotype were assessed. In some experiments, activated cells were maintained in cytokines as indicated for 2 weeks. Every 2 to 3 days cells were counted and given fresh cytokine-containing media to maintain a concentration of 0.8 × 10^6 cells/mL. For generation of TCR-modified human T cells, we used a modification of a previously described protocol (34). On day 1, human PBMCs were stimulated with soluble anti-CD3 mAb (OKT3, NCI preclinical repository) for 48 hours. Beginning on day 3, cells were cultured with IL2 (300 IU/mL) and IL15 (100 ng/mL), and maintained between 1 and 2 × 10^6 cells/mL. Also on day 3, activated T cells were transduced by coculture with 50% retroviral supernatant from PG13 packaging cells transfected with the TIL1383ITCR/CD34t construct (35). Transduction was done with retroinfectin-coated plates and spinoctination (2,000 × g for 2 hours at 32°C). On day 8, cells underwent a rapid expansion protocol (REP) by incubation in a G-Rex 100 flask (Wilson Wolf Manufacturing) of 1 × 10^6 transduced T cells with 2 × 10^6 irradiated (50 Gy) allogeneic feeder cells from human donors. Soluble anti-CD3 mAb (OKT3, 30 ng/mL) and hIL15 (100 ng/mL) were added to culture every 2 to 3 days and tumor surface area (mm^2) was calculated as length × width.

**Cytokine responsiveness**

Cytokine responsiveness was assessed by washing cells three times in PBS, then repatinating cells at 0.8–1 × 10^6/mL with the indicated cytokine (mouse cytokines from Shenoadoah Biotechnology). After overnight incubation, cells were either fixed/permeabized for phosflow analysis per the manufacturer’s instructions (Phosflow; BD Bioscience) or 10 μmol/L bromodeoxyuridine (BrdU) was added for 1 hour at 37°C and cells were processed according to the manufacturer’s protocol (BrdU Flow Kit; BD Bioscience). Note that the percentage of cells that were pSTAT5−15 minutes after restimulation was not significantly different from values obtained after overnight incubation (data not shown).

**Flow cytometry**

For flow-cytometric analysis, cells were processed as previously described (18) and analyzed on either an LSRII or Accuri C6 flow cytometer (BD Bioscience). Data were processed using FlowJo (TreeStar) or C6 software (BD Bioscience). Mouse antibody clones used in this study include: CD4 (GK1.5), CD8 (53-6.7), CD25 (PC6.15), CD62L (Mel-14), CD122 (TM-β1), IL7Rα (SB/199 or A7R34), Eomes (Dan1imag), granzyme B (GB12), IFNγ (XMG1.2), pAKT S473 (D9E), pSTAT5 (47/Stat5 pY694), p86 (D57.2.2E), Thy1.1 (OX-7 or HB51), TNF-α (TN3-19.12), and Tbet (4810). Human antibody clones used are CD8 (OKT8 or SK1) and IL7Rα (eBioBDR5 or A019DS). These were purchased from BD Bioscience, BioLegend, Invitrogen, eBioscience, and/or Cell Signaling Technology.

**Tumor challenge, lymphodepletion, and adoptive T-cell transfer**

For tumor experiments, B6 mice were injected subcutaneously (s.c.) with 2.5 × 10^5 B16-F1 tumor. Tumor growth was measured by an observer blinded to treatment groups with calipers two to three times per week and tumor surface area (mm^2) was calculated as length × width. Mice were sacrificed when tumors reached >400 mm^2. Total body irradiation (TBI) was administered at 6 Gy the day before adoptive transfer. Mice were excluded from analysis if they developed i.p. tumor spread within the first 4 weeks after injection.

**In vivo cytokine neutralization**

All neutralizing antibodies were purchased from BioXCell except for JES6-1A12 (UCSF monoclonal antibody core). Unless otherwise indicated, the following amounts of mAb were injected i.p. on days 0, 2, 5, 8, 12, and 17 following adoptive transfer: 32G7 (25 μg), IL7Rα (A7R34, 500 μg), IL7Rα (250 μg each of S4B6 and JES6-1A12 injected together), and mgL2b isotype control (MPC-11, 200 μg).

**Measurement of IFNγ**

Day 3 culture supernatants were analyzed for mIFNγ via ELISA per the manufacturer’s instructions (BioLegend).

**Experiments involving human PBMCs**

Deidentified human PBMCs were isolated from a leukapheresis pack obtained from Research Blood Components and experiments were performed in accordance with MUSC Institutional Review Board (IRB) guidelines. For *in vitro* stimulation, cells were thawed and rested in 100 IU/mL hIL2 overnight. The next day, 0.5 μg/mL soluble αCD3 (OKt3, NCI repository) was added to culture plus 10 ng/mL hIL12. After 3 days of activation, cytokine responsiveness and phenotype were assessed. In some experiments, activated cells were maintained in cytokines as indicated for 2 weeks. Every 2 to 3 days cells were counted and given fresh cytokine-containing media to maintain a concentration of 0.8 × 10^6 cells/mL. For generation of TCR-modified human T cells, we used a modification of a previously described protocol (34). On day 1, human PBMCs were stimulated with soluble αCD3 mAb (OKT3, NCI preclinical repository) for 48 hours. Beginning on day 3, cells were cultured with hIL2 (300 IU/mL) and hIL15 (100 ng/mL), and maintained between 1 and 2 × 10^6 cells/mL. Also on day 3, activated T cells were transduced by coculture with 50% retroviral supernatant from PG13 packaging cells transfected with the TIL1383ITCR/CD34t construct (35). Transduction was done with retroinfectin-coated plates and spinoctination (2,000 × g for 2 hours at 32°C). On day 8, cells underwent a rapid expansion protocol (REP) by incubation in a G-Rex 100 flask (Wilson Wolf Manufacturing) of 1 × 10^6 transduced T cells with 2 × 10^6 irradiated (50 Gy) allogeneic feeder cells from human donors. Soluble anti-CD3 mAb (OKT3, 30 ng/mL)
was also added to the cultures. On REP day 14, cultures were harvested, washed, and replated for IL7R analysis 3 days later.

**Statistical analysis**
Statistical analysis was done with GraphPad Prism 6 software. One-way ANOVA with a Tukey multiple comparisons correction or a two-sided two-sample t test was used to evaluate statistical significance of means between groups. When variances were unequal, the Welch t test was used. Data expressed on a ratio scale (e.g., fold change) were first log-transformed to normalize the distribution, then analyzed by the t test or one-way ANOVA, as appropriate. For survival data, the log-rank test was used. Unless otherwise indicated, summary statistics in figures are presented as mean ± SEM.

**Results**
The enhanced initial engraftment of IL12-conditioned effector CD8+ T cells (Tc1) transferred into lymphodepleted hosts is dependent on IL7 but not IL15
We previously demonstrated that the persistence and antitumor abilities of IL12-conditioned pmel-1 CD8+ T (Tc1) cells were enhanced by cyclophosphamide, a lymphodepleting agent (18). Similarly, lymphodepletion with 6-Gy TBI before adoptive transfer of Tc1 significantly delayed the growth of established B16 tumors, while transfer of Tc1 alone or transfer of cells activated without IL12 (Tc0) into irradiated hosts did not (Fig. 1A and B). The persistence of Tc1 cells was also strikingly enhanced relative to Tc0 cells, with the peak of expansion seen about 1 week after transfer (Fig. 1C and D). This enhanced persistence with multiple

![Figure 1](https://example.com/figure1.png)

The enhanced persistence of IL12-conditioned CD8+ T cells (Tc1) in lymphodepleted hosts is dependent on IL7. A and B, B6 mice were injected with B6 melanoma tumor s.c. on day 0 and then irradiated on day -1. On day 0, mice were adoptively transferred with 2 x 10^6 3-day activated pmel-1 CD8+ T cells with IL12 conditioning (Tc1) or without (Tc0). A, survival curves (n = 8; ****, P = 0.001 for Tc1 vs. control, P < 0.0001 for Tc1 vs. Tc1 + TBI), and B, individual tumor growth curves. C and D, 5 x 10^6 Tc1 or Tc0 cells were transferred into mice with or without 6 Gy TBI and Thy1.1+ donors were tracked in the (C) peripheral blood over time (n = 5) or D, in the spleens 7 days after transfer (n = 5; ****, P < 0.0001). E, as in D, except cells were transferred into WT B6 or IL15-/ mice with or without all7-neutralizing mAb (clone M25; n = 5; ****, P < 0.0001). All results are representative of at least two independent experiments. n.s., not statistically significant; SA, surface area.
forms of lymphodepletion but without the need for IL2 or vaccination establishes the feasibility of using our Tc1 model to investigate the host cytokine requirements of effector CD8+ T cells.

Because IL7 and IL15 are thought to be the dominant cytokines for T-cell homeostatic expansion (1–3), and they are elevated post-lymphodepletion (21–23), we assessed their importance for the expansion of Tc1 cells. We transferred Tc1 cells into irradiated wild-type (WT) or IL15−/− mice with or without an IL7-neutralizing mAb (clone M25). We then harvested spleens at day 7 after transfer, as this correlated with the peak of their expansion (Fig. 1C). Surprisingly, Tc1 cells exhibited a significant expansion defect at day 7 in WT mice treated with IL7-neutralizing antibody against IL7Rα (A7R34; Supplementary Fig. S1A). Like IL15, IL2 was not critical, as a combination of neutralizing IL2 antibodies (JES6-1A12 and S4B6; ref. 36) did not significantly affect Tc1 cell expansion (Supplementary Fig. S1B). In addition, the absence of host IL2, IL7, and/or IL15 did not significantly impair the ability of Tc1 cells to secrete IFNγ and TNFα after ex vivo restimulation (Supplementary Fig. S2). In summary, Tc1 cells are dependent on host IL7 alone for their initial expansion.

Certain T-cell subsets require TCR engagement for homeostatic maintenance (3, 4). Because pmel-1 T cells have engineered specificity against gp100, a self-antigen, we transferred Tc1 cells into β2m−/− mice, which are devoid of HLA-C presentation. Tc1 cells persisted equally well in WT B6 and β2m−/− B6 mice, indicating that Tc1 did not require TCR engagement for effector expansion (Supplementary Fig. S3A). To confirm our results in a second model, we used the h3T TCR transgenic mouse, whose T cells recognize tyrosinase in an HLA-A2-restricted manner (33). h3T T cells activated in the presence or absence of IL12 showed...
similar persistence when transferred into irradiated WT B6 or HLA-A2 transgenic mice (Supplementary Fig. S3B). Thus, activated Tc1 cells do not require contact with cognate MHC-I for maximal effector expansion in irradiated hosts.

IL7 and IL15 are required for maximal antitumor efficacy of Tc1 cells

The results above were obtained in tumor-free animals. Therefore, we assessed the cytokine requirements for optimal expansion of effector CD8^+ T cells adoptively transferred into B6 mice bearing 12-day established B16 tumors. In a manner similar to tumor-free mice, the initial engraftment of Tc1 cells was dependent on IL7 but not IL15 (Fig. 2A). Consistent with our early expansion data (Fig. 2A), Tc1 cells required IL7 for maximum antitumor efficacy (Fig. 2B and C). In contrast with these data, Tc1 cells also needed IL15 for maximal antitumor efficacy (Fig. 2B and C). This result is likely because IL15 is required for the long-term persistence and memory formation of Tc1 cells (Fig. 2D), although IL15-dependent host cells may be relevant. Thus, Tc1 cells require IL7 for initial expansion but both IL7 and IL15 for maximal antitumor efficacy.

Figure 3.

IL12 conditioning during CD8^+ T-cell activation leads to elevated IL7 responsiveness and IL7Rα expression in vitro. A–C, Pmel-1 T cells were activated for 3 days with (Tc1) or without (Tc0) IL12, washed and replated in the indicated cytokines (A, top). Representative histograms depicting pSTAT5 and pS6 levels after reculture without cytokine or with IL7 (A, bottom). Mean pSTAT5 and pS6 levels after reculture in 100 ng/mL of the indicated cytokine (n = 4; *, P < 0.05; ***, P < 0.001; ****, P < 0.0001). B, BrdU was added for the final hour after overnight culture in the indicated cytokine (n = 10; ****, P < 0.0001). C, cells were counted on days 0, 1, 2, and 3 after reculture in the indicated concentration of IL7 in ng/mL (results are from one experiment with two replicates and are representative of at least three independent experiments). D–F, Tc0 and Tc1 cells were analyzed for the indicated cytokine receptors via flow cytometry. D, representative histograms and E, MFI ratios (****, P < 0.0001; ***, P < 0.001; ***, P < 0.01; *P < 0.05; P values represent statistically significant difference from Tc0, which is indicated by the dashed line). F, the percentages of cells expressing each cytokine receptor are shown (n = 11 independent experiments; ***, P < 0.001 via the Welch t-test). G, the percentage of cells expressing IL7Rx on days 2 and 3 after stimulation (n = 7; ****, P < 0.0001 for all comparisons with Tc1 day 3; not statistically for others).
Tc1 cells show superior IL7 responsiveness and elevated IL7Rα levels in vitro

Because Tc1 cells exhibited IL7-dependent expansion in irradiated hosts, we assessed the in vitro IL7 responsiveness of Tc1 cells compared with Tc0 cells. We also assessed IL2 and IL15 signaling as controls. We first cultured Tc0 cells and Tc1 cells in high doses (100 ng/mL) of IL2, IL15, or IL7 overnight and then assessed phosphorylation of STAT5 and ribosomal S6 (Fig. 3A), both of which are downstream of IL2/7/15 cytokine signaling (4, 36). As expected, IL2 and IL15 led to high levels of phosphorylation in both Tc0 and Tc1 cells. However, when cultured with IL7, only Tc1 cells robustly phosphorylated STAT5 and S6 (Fig. 3A). These enhanced signaling events translated into increased proliferation of Tc1 cells after reculture in IL7 as determined by BrdU incorporation (Fig. 3B). In contrast, Tc0 and Tc1 cells proliferated extensively in IL2 or IL15, as over half of the cells had incorporated BrdU in 1 hour (Fig. 3B). The enhanced proliferation rate after overnight culture led to about a 5-fold expansion of Tc1 over Tc0 cells after 3 days of culture in IL7 (Fig. 3C). Remarkably, even 100-fold lower levels of IL7 (1 ng/mL) led to an increased concentration of Tc1 cells after 3 days, while Tc0 cells at the highest dose barely maintained their numbers (Fig. 3C). These signaling and proliferation events were inhibited by JAK-STAT and PI3K inhibitors, but not mTOR inhibitors (Supplementary Fig. S4), indicating that IL7 was engaging established pathways for cytokine-mediated T-cell proliferation (38–40). In summary, these findings demonstrate the ability of IL12 conditioning to induce IL7 responsiveness in effector CD8+ T cells.

We next sought to delineate the mechanism(s) responsible for the enhanced IL7 responsiveness of Tc1 cells by evaluating IL7Rα as well as IL2Rα and IL7Rα expression on Tc0 and Tc1 cells. The expression of all three receptors was increased by the addition of IL12 (Fig. 3D and E), although the magnitude of these increases varied (Fig. 3E). When expressed as a proportion of cells staining positive for the receptor rather than the magnitude of expression, a striking difference was seen with IL7Rα and IL2Rα expression on Tc1 cells. A large proportion of Tc1 cells expressed IL7Rα while Tc0 cells had almost none, in contrast with high levels seen with IL2Rα and IL7Rα on Tc0 and Tc1 cells. Because Tc1 cells exhibited IL7-dependent expansion in irradiated hosts (6 Gy), and the absolute number of donor cells in host spleens 7 days later is displayed (data are combined from three independent experiments; ****, P < 0.0001), E-G, on day 32 B6 tumor-bearing mice were injected with 2 x 10^6 T cells the day after irradiation. E, the percentage of donor cells in the peripheral blood on day 8 after transfer (*, P < 0.05). F, survival curves (****, P < 0.0001) for TBI only vs. IL7Rα+/+ Tc1; **, P < 0.01 for IL7Rα+/+ Tc1 vs. Tc1; combined from two independent experiments for total n = 14–17). G, growth curves. SA, surface area.

Figure 4.

IL7Rα expression is required for maximal expansion and antitumor efficacy of Tc1 cells. A, representative histogram of IL7Rα levels in Tc0, Tc1, and IL7Rα+/+ Tc1 cells. B, pSTAT5 and C, pS6 levels of Tc1 and IL7Rα+/+ Tc1 cells after reculture in 100 ng/mL of the indicated cytokine (n = 4–6; **, P < 0.01). D, 3 to 5 x 10^5 pmel Tc1 or IL7Rα+/+ Tc1 cells were transferred into irradiated hosts (6 Gy); and the absolute number of donor cells in host spleens 7 days later is displayed (data are combined from three independent experiments; ****, P < 0.0001). E-G, on day 32 B6 tumor-bearing mice were injected with 2 x 10^6 T cells the day after irradiation. E, the percentage of donor cells in the peripheral blood on day 8 after transfer (*, P < 0.05). F, survival curves (****, P < 0.0001) for TBI only vs. IL7Rα+/+ Tc1; **, P < 0.01 for IL7Rα+/+ Tc1 vs. Tc1; combined from two independent experiments for total n = 14–17). G, growth curves. SA, surface area.
(Fig. 3F). We next investigated the kinetics of IL7Rα expression. As expected, IL7Rα was initially decreased on both cell types after T-cell activation, but Tc1 cells showed increased expression by 72 hours after stimulation (Fig. 3G). Thus, IL2 promotes IL7Rα reexpression in Tc1 cells, a finding that may explain the enhanced IL7-mediated persistence of effector CD8+ T cells (Tc1) cells after transfer into lymphodepleted hosts.

IL7Rα upregulation is responsible for the enhanced IL7 responsiveness and subsequent in vivo persistence of Tc1 cells

To directly test whether IL7Rα was critical for the enhanced IL7 responsiveness of Tc1 cells, we generated pmel-1 IL7Rα−/− mice. As expected, Tc1 cells generated from IL7Rα−/− and IL7Rα+/− pmel-1 mice expressed similar levels of IL2Rα, IL2Rα, granzyme B (GzB), Tbet, Eomes, and CD62L (Supplementary Fig. S5A), and produced equivalent levels of IFNγ by 3-day culture (Supplementary Fig. S5B). In contrast, IL7Rα levels in the IL7Rα−/− Tc1 cells were about half that of Tc1 cells (Fig. 4A and B). This decreased IL7Rα expression translated to reduced IL7-induced STAT5 and S6 phosphorylation for IL7Rα−/− Tc1 compared with WT Tc1, despite having similar levels when maintained in IL2 or IL15 (Fig. 4B and C). BrdU incorporation also trended lower with IL7 cultures of IL7Rα−/− Tc1 relative to Tc1 (Supplementary Fig. S5C).

These in vitro results indicate that IL7Rα−/− Tc1 cells can be used to evaluate the functional importance of IL7Rα, given that they appeared identical to WT Tc1 in all aspects tested except for IL7Rα expression and IL7 responsiveness. Therefore, we transferred WT and IL7Rα+/− Tc1 cells into irradiated hosts. On day 7 after transfer into irradiated hosts, there were about half as many IL7Rα+/− Tc1 cells as WT Tc1 cells in the spleens of recipient mice (Fig. 4D). Similar results were observed in the peripheral blood of tumor-bearing mice 7 days after transfer (Fig. 4E).

Importantly, this decreased initial expansion of Tc1 cells also led to significantly reduced antitumor activity in IL7Rα+/− Tc1 cells relative to WT pmel-1 Tc1 cells (Fig. 4F and G). Together, these results indicate that elevated IL7Rα expression is critical for driving the initial engraftment and subsequent antitumor activity of Tc1 cells.

Host IL7 and donor IL7Rα are required for maximal persistence of polyclonal CD8+ T cells in lymphodepleted hosts

Next, we investigated the importance of IL7Rα for the initial engraftment of effector CD8+ T cells activated without IL12. As shown in Fig. 1C and D, pmel-1 T cells stimulated with hgp100 alone (Tc0) persisted poorly, presumably due to low IL7Rα expression (Fig. 3F). Therefore, we sought IL12-independent activation conditions that would elevate IL7Rα appreciably and thereby generate effector cells capable of persisting in lymphodepleted hosts. Because TCR strength has been shown to modulate IL7Rα levels in human CD4+ T cells (41), we activated pmel-1 T cells over a broad range of hgp100 concentrations. Although
higher peptide concentrations increased IL7Rα expression, the receptor levels did not reach those achieved with IL12 (Fig. 5A). To further increase the strength of TCR stimulation, we next activated T cells nonspecifically with soluble or plate-bound anti-CD3 mAb with or without anti-CD28 mAb. Consistent with reports demonstrating elevated TCR signaling with immobilized anti-CD3 mAb (42) and costimulation with anti-CD28 mAb (43), IL7Rα levels were increased in plate-bound conditions and even higher when anti-CD28 mAb was added (Fig. 5B). In fact, plate-bound anti-CD3 mAb and anti-CD28 mAb (PB CD3/CD28) were statistically indistinguishable from Tc1 cells (hgp100 + IL12; Fig. 5B).

Having established that higher TCR signals increase IL7Rα expression in the pmel-1 model, we evaluated this relationship in Fig. 5B). Higher peptide concentrations increased IL7Rα expression, the receptor levels did not reach those achieved with IL12 (Fig. 5A). To further increase the strength of TCR stimulation, we next activated T cells nonspecifically with soluble or plate-bound anti-CD3 mAb with or without anti-CD28 mAb. Consistent with reports demonstrating elevated TCR signaling with immobilized anti-CD3 mAb (42) and costimulation with anti-CD28 mAb (43), IL7Rα levels were increased in plate-bound conditions and even higher when anti-CD28 mAb was added (Fig. 5B). In fact, plate-bound anti-CD3 mAb and anti-CD28 mAb (PB CD3/CD28) were statistically indistinguishable from Tc1 cells (hgp100 + IL12; Fig. 5B).

Human T cells conditioned with IL12 display enhanced IL7Rα expression and IL7 responsiveness. A–D, human PBMCs were activated with soluble anti-CD3 mAb (0.5 µg/mL, Okt3 clone) with or without hIL12 (10 ng/mL) for 3 days. A, IL7Rα expression after 3-day activation (***, P < 0.001; "ND" is normal donor). B and C, 3-day activated T cells were washed and then replated in the indicated cytokines (300 IU/mL IL2 + 100 ng/mL IL15; IL7, 100 ng/mL). B, pSTAT5 staining via flow cytometry after overnight culture (n = 8 from two independent experiments with four normal donors; ***, P < 0.0001). C, cells were counted and given fresh media every 2 to 3 days (n = 6 from two independent experiments with three normal donors). D, as in C except activated cells were recultured in IL2 + IL15 on day 3 and then assessed for IL7Rα expression at the indicated time points (n = 6–9 from two independent experiments with four normal donors; ***, P < 0.001; ****, P < 0.0001 via the Welch t test). E, overview of the clinical transduction protocol to generate TCR-transduced melanoma-reactive human T cells. Shown is the timing of IL12 addition and 3-day reculture in IL2 (300 IU/mL) + IL15 (100 ng/mL). F, IL7Rα expression at day 26 of above timeline of human T cells initially grown with or without hIL12. This result is representative of two independent experiments.
Human T cells conditioned with IL12 display enhanced IL7Rα expression and IL7 responsiveness

Given the importance of donor IL7Rα and host IL7 for the persistence of effector CD8+ T cells in mice, we next tested the ability of IL12 to enhance IL7Rα expression in activated human CD8+ T cells. CD8+ T cells from day 3 activated human peripheral blood mononuclear cells (PBMC) exhibited higher IL7Rα expression with IL12, although the magnitude of this effect was not as large as our murine data (Fig. 6A compared with mouse data in Fig. 3D). In contrast with this small change in IL7Rα expression, human T cells were only able to phosphorylate STAT5 robustly in response to IL7 if they were activated with IL12 (Fig. 6B). When these activated T cells were washed and recultured in vitro, only those activated with IL12 expanded in the presence of IL7 (Fig. 6C). Given the discordance between initial IL7Rα levels (Fig. 6A) and IL7 responsiveness (Fig. 6B and C), we assessed IL7Rα levels after reculture of cells. We speculated that the ability to reexpress IL7Rα after withdrawal of TCR stimulation might explain the observed differences in IL7 responsiveness. Consistent with this hypothesis, the presence of IL12 during the first 3 days of activation led to a striking enhancement in IL7Rα expression that lasted for at least 1 week after reculture (Fig. 6D). Finally, we sought to evaluate the translatability of our findings from 3-day cultures in a clinically relevant scenario by using the retroviral transduction protocol depicted in Fig. 6E, in which IL12 was added or withheld during the REP. We found that the inclusion of IL12 did not significantly increase IL7Rα levels at the end of the REP. As was the case in our 3-day cultures, however, the transduced T cells that underwent the REP in the presence of IL12 possessed higher IL7Rα expression 3 days after reculture (Fig. 6F). These results suggest that the addition of IL12 to human T-cell cultures during the REP is a feasible strategy to augment IL7Rα levels, and this may be applicable in a number of clinically used protocols (44–46).

Discussion

In this study, we evaluated the host cytokines required for the initial engraftment of effector CD8+ T cells transferred into lymphodepleted hosts. Contrary to our expectations, IL7 was initially required, whereas IL15 was not. Because multiple methodologies for the activation of CD8+ T cells, including IL12 conditioning or strong TCR stimulation, demonstrated IL7 and IL7Rα dependence, our results are likely generalizable to a variety of T-cell activation methodologies.

Our results indicate that transferred effector T cells should be IL7 responsive for maximal engraftment in a lymphodepleted host without exogenously provided cytokine. In our murine models, CD8+ T cells required IL7Rα for maximal engraftment after adoptive transfer; however, in a clinical setting, expression of IL7Rα on donor T cells was one of 45 markers that failed to differentiate persisting T-cell clones from those that failed to engraft (47). In this prior study, T cells were not conditioned with IL12. Our results with human T cells suggest that reexpression of IL7Rα after cessation of TCR stimulation and extended culture corresponds most directly with IL7 responsiveness (Fig. 6). We therefore predict that assessing IL7Rα levels after extended reculture may have more clinical utility than determining IL7Rα levels at the predetermined point of infusion.

An intriguing result from this work is that IL15 does not initially play a role in the support of effector Tc1 cells. These data are in contrast with results from prior studies with memory phenotype

CD8+ T cells transferred into lymphopenic hosts (10–12). Because IL15 is known to be elevated in the lymphodepleted host (21), these differences are potentially explained by distinct trafficking of activated versus resting T cells.

That in vitro IL12 priming increases IL7Rα expression appears to be discordant with the well-described phenomenon that enhanced IL7/IL12/IFN-γ/IL-12 expression during effector responses in vivo leads to more terminally differentiated CD8+ T cells with decreased IL7Rα expression (28, 30, 48). A potential explanation is that the programming for terminal differentiation has not yet occurred after 3 days of activation in the presence of IL12, a theory supported by the increased IL7Rα and CD62L expression observed with IL12 priming on day 3 (25). The kinetics of IL7Rα reexpression we observed further support this idea, as IL7Rα transcription appears to be initiated on day 2 of culture. Given that the expression of IL7Rα is modulated by the transcription factors Gfi-1 and GABPRα, the relationship between IL12 and these transcription factors warrants further investigation (49).

In summary, our results suggest a model in which effector CD8+ T cells are dependent on host IL7 for maximal persistence and antitumor efficacy in a lymphodepleted host. This represents a shift in the current paradigm that considers IL15 as the critical cytokine capable of modulating effector CD8+ T-cell durability and efficacy in this increasingly relevant clinical setting. In practical terms, our results demonstrate that a direct and feasible way to produce IL7Rα-expressing, IL7-responsive effector T cells is ex vivo IL12 conditioning.

Disclosure of Potential Conflicts of Interest

No potential conflicts of interest were disclosed.

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Conception and design: C.B. Johnson, D.J. Cole, M.P. Rubinstein
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Analysis and interpretation of data (e.g., statistical analysis, biostatistics, computational analysis): C.B. Johnson, B.P. Riesenberg, K.F. Staveley-O’Carroll, E. Garrett-Mayer, D.J. Cole, M.P. Rubinstein
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