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Th17 cells are refractory to senescence and retain robust antitumor activity after long-term ex vivo expansion

Jacob S. Bowers,1,2,3 Michelle H. Nelson,1,2,3 Kinga Majchrzak,1,2,3,4 Stefanie R. Bailey,1,2,3 Baerbel Rohrer,5,6 Andrew D.M. Kaiser,7 Carl Atkinson,1 Luca Gattinoni,8 and Chrystal M. Paulos1,2,3

1Department of Microbiology and Immunology, 2Department of Dermatology, 3Department of Surgery, Medical University of South Carolina, Charleston, South Carolina, USA. 4Department of Physiological Sciences, Faculty of Veterinary Medicine, Warsaw University of Life Sciences, Warsaw, Poland. 5Department of Ophthalmology, Medical University of South Carolina, Charleston, South Carolina, USA. 6Ralph H. Johnson VA Medical Center, Charleston, South Carolina, USA. 7Center for Cancer Research, 8Experimental Transplantation and Immunology Branch, National Cancer Institute (NCI), NIH, Bethesda, Maryland, USA.

Adoptive immunotherapy for solid tumors relies on infusing large numbers of T cells to mediate successful antitumor responses in patients. While long-term rapid-expansion protocols (REPs) produce sufficient numbers of CD8+ T cells for treatment, they also cause decline in the cell’s therapeutic fitness. In contrast, we discovered that IL-17–producing CD4+ T cells (Th17 cells) do not require REPs to expand 5,000-fold over 3 weeks. Also, unlike Th1 cells, Th17 cells do not exhibit hallmarks of senescence or apoptosis, retaining robust antitumor efficacy in vivo. Three-week-expanded Th17 cells eliminated melanoma as effectively as Th17 cells expanded for 1 week when infused in equal numbers into mice. However, treating mice with large recalcitrant tumors required the infusion of all cells generated after 2 or 3 weeks of expansion, while the cell yield obtained after 1-week expansion was insufficient. Long-term-expanded Th17 cells also protected mice from tumor rechallenge including lung metastasis. Importantly, 2-week-expanded human chimeric antigen receptor–positive (CAR+) Th17 cells also retained their ability to regress human mesothelioma, while CAR+ Th1 cells did not. Our results indicate that tumor-reactive Th17 cells are an effective cell therapy for cancer, remaining uncompromised when expanded for a long duration owing to their resistance to senescence.

Introduction

Adoptive cell therapy (ACT) is a powerful means for augmenting a patient’s immunity to cancer via the ex vivo selection, expansion, and enhancement of autologous tumor-reactive T cells. An advantage of ACT is the potential for complete and durable responses executed by memory T cell populations (1). In patients with metastatic melanoma, this therapy achieves remarkable outcomes for patients, with objective responses of 54% and complete remissions of 24% (2, 3).

To accomplish durable results in patients, ACT requires the infusion of large numbers of tumor-reactive T cells (~10^10 cells) (4–10). Most protocols for generating vast T cell numbers require their expansion for up to 3 months with at least 1 CD3-dependent reactivation (8–10). However, ex vivo expansion works in direct opposition to maintaining antitumor efficacy because the cytotoxic CD8+ T cells used in clinical trials lose their potency when extensively expanded ex vivo. This loss of antitumor efficacy is due to a reduced capacity to persist in vivo once CD8+ T cells reach terminal differentiation (11). Additionally, this effect occurs rapidly, as even highly potent IL-12–primed CD8+ T cells lose antitumor efficacy in as little as 1 week of expansion (12). Rapid-expansion protocols (REPs) also compromise the integrity of human tumor-infiltrating lymphocytes (TILs), while TILs that undergo shorter expansion ex vivo persist longer (1, 13–15).

Although investigators have developed strategies to prevent T cell differentiation ex vivo (16, 17), another potential avenue is to circumvent this dilemma using a T cell subset relatively refractory to senescence. We used CD4+ T cells with a TCR specific for tyrosinase-related protein-1 (TRP-1, a shared melanocyte/melanoma antigen) that were polarized to a Th17 phenotype (CD4+ T cells that express the transcription
factor RORγt and secrete IL-17). In murine models of melanoma, TRP-1 Th17 cells offer an alternative to melanoma-reactive pmel-1 CD8+ T cells, as they mediate a potent response against aggressive melanoma in mice even without antigen vaccination and in vivo IL-2 support required in pmel-1 CD8+ ACT models (18–20). This remarkable Th17 antitumor response occurs in part by enhancing CD8+ T cell responses against transformed cells (21–24) and via their ability to directly lyse tumors (25, 26). In both cases, Th17 cells exert superior antitumor immunity compared with Th1 and other CD4+ T cell subsets (18), as well as superior enhancement of CD8+ T cells (23). Also, since Th17 cells show durable efficacy in vivo and have stem memory properties (27, 28), we posited that unlike CD8+ T cells, Th17 cells would retain their antitumor capacity even after long-term ex vivo expansion to clinically relevant doses.

We found that, even without a REP, Th17 cells were capable of robust growth for 21 days ex vivo, yielding ~5,000 times the initial CD4+ T cell numbers. Additionally, while exhibiting a CD44hiCD62Llo effector memory phenotype, Th17 cells gained nuclear Tcf7 throughout in vitro culture, indicating active and preserved stem memory signaling. Unlike Th1 or CD8+ T cells, Th17 cells retained their ability to eliminate melanoma tumors over 2 weeks of expansion. Th17 cells were resistant to senescence indicated by retention or upregulation of CD127, CD28, and CD27 along with nominal expression of KLRG1 and TIM3. In contrast, Th1 cells expressed exhaustion markers and higher levels of checkpoint receptors and were far more apoptotic than Th17 cells at any time point of expansion. Importantly, Th17 cells expanded for either a 7- or 21-day duration ex vivo were equally able to persist in the tumor-bearing host after transfer. Despite similar antitumor potency between 7- and 21-day Th17 cells on a per-cell basis, we hypothesized that mediating curative responses in mice with large, nearly terminal tumors would require the full yield of Th17 cells generated from 3 weeks of culture. We found that treatment with 2- or 3-week-expanded Th17 cells rapidly and completely eradicated aggressively growing tumors, while the full yield of 1-week-expanded Th17 cells was insufficient to control these tumors. Of clinical significance, human Th17-polarized CD4+ cells also exhibit marked durability in antitumor potency compared with Th1-polarized cells when expanded for either 7 or 12 days. Our analysis of all parameters of Th17 cell ex vivo preparation for ACT (in vitro time of expansion, cell yield, T cell subset, and size of tumor) indicates that tumor-reactive Th17 cells represent a cell product that can be reliably applied to treat highly aggressive solid tumors.

**Results**

*ACT with Th17 cells still requires transferring large numbers of T cells to clear tumors.* The effectiveness of ACT is thought to depend on the transfer of large numbers of tumor-specific T cells (11). However, because Th17 cells mediate potent antitumor responses in mice and have durable stem memory properties, we hypothesized that we could titrate down the number of Th17 cells infused into mice without impairing tumor clearance. To test this idea, TRP-1 CD4+ T cells were polarized to a Th17 phenotype and expanded ex vivo for 1 week. This protocol generated cells with expression of the Th17 transcription factor RORγt, high IL-17 production, and nominal Tbet (the Th1 transcription factor) and IFN-γ (Figure 1, A and B). We infused decreasing doses of Th17 cells (4.5, 1.5, or 0.5 × 10^6 cells/mouse) into C57BL/6J mice with established B16F10 melanoma preconditioned with nonmyeloablative (5 Gy) total body irradiation (Figure 1C). While the highest number of Th17 cells (4.5 × 10^6) eliminated tumors in most mice, tumor responses and overall survival declined in mice treated with fewer Th17 cells (Figure 1, C and D). Thus, despite their enhanced antitumor efficacy and a stem memory profile, Th17 cells require a sufficient number of transferred cells for successful ACT.

*REP decreases the antitumor capacity and recall responses of Th17 cells.* Since large numbers of Th17 cells are required to eradicate tumors, we tested whether extending the in vitro culture time for another week using a REP with TCR reactivation would improve their yield. To test this, as presented in Figure 2A, Th17 cells were expanded for 2 weeks, with the REP culture undergoing TCR reactivation using TRP-peptide-pulsed feeder cells at 1 week. Interestingly, Th17 cells only activated once (primary expansion) could effectively expand for 2 weeks to similar yields as REP Th17 cells (Figure 2B). By day 14, both Th17 and REP Th17 cells were mainly CD44hiCD62Llo effectors (Figure 2C). However, REP Th17 cells were slightly compromised in their ability to regress tumors and promote survival in mice (Figure 2, D and E). This was not due to increased levels of apoptosis, KLRG1, PD1, or TIM3 on the REP Th17 cells compared with Th17 cells (Supplemental Figure 1, A–C; supplemental material available online with this article; https://doi.org/10.1172/jci.insight.90772DS1). However, while the frequency of donor Th17 cells in the spleen and tumors of surviving mice was similar to the frequency of REP Th17 cells 40 days after treatment (Figure 2, F...
and G), memory recall against TRP-1 antigen by mice treated with REP donor Th17 cells was significantly impaired compared with mice treated with Th17 cells in an in vivo cytotoxicity assay (Figure 2H). Thus, REPs appear to compromise the antitumor efficacy and antigen-recall responses of donor Th17 cells.

Th17 cells become less activated but not terminally differentiated during ex vivo expansion. We next sought to determine how long Th17 cells could be expanded in vitro without losing antitumor efficacy. We found that Th17 cells could expand for 3 weeks without TCR reactivation (Figure 3A). While tyrosinase-specific (Vβ14+) CD4+ T cells have a naive phenotype (CD44loCD62Lhi) when isolated from the spleens of TRP-1 transgenic mice, Th17 cells exhibited an effector memory phenotype with high CD44 and low CD62L expression as early as 1 week after activation, which prevailed throughout culture (Figure 3B). This phenotype corroborates previous studies with human and murine Th17 cells (18, 27). However, Th17 cells expressed Tcf7 (a downstream product of active β-catenin signaling), throughout culture (Figure 3, C and D; see complete unedited blots in the supplemental material), indicating sustained capacity for self-renewal and persistence (29).

Because we were not using a REP to expand the Th17 cells, we expected that the cells would gradually become quiescent and downregulate activation markers as they were expanded for 21 days in vitro. As expected, activation markers CD69 and CD25 (IL-2Rα) were initially high 1 week after stimulation but progressively declined (Figure 3E). Similarly, Th17 cells initially expressed high levels of ICOS, which is an important costimulatory molecule that supports their function and maintenance (19, 30), while the costimulatory receptor 4-1BB was only increased moderately above preactivation levels at the early Th17 time point. In contrast, CD127 (IL-7Rα) was maintained on Th17 cells expanded for 3 weeks, suggesting a persistent responsiveness to IL-7 (31). The costimulatory ligand CD28 was similarly retained. CD27, which is associated with adaptive memory responses (32), increased between week 1 and week 3 (Figure 3E). Importantly, the retention of CD127 and CD28, and the increased expression of CD27 during expansion, may indicate that Th17 cells are resistant to the senescence that naturally occurs in other T cell subsets (33).

Long-term-expanded Th17 cells retain antitumor potency and persist after transfer. Based on evidence that Th17 cells maintained active Wnt/β-catenin signaling (Figure 3, C and D), and the pattern of chemokine and costimulatory receptor expression we observed throughout culture, we posited that Th17 cells (in contrast to CD8+ T cells; see ref. 11) would retain antitumor activity as they expanded long term in vitro. To compare Th17 cell antitumor efficacy over ex vivo culture time on a per-cell basis, Th17 cells were expanded for 3 weeks, with a portion of cells cryopreserved at either (a) early (day 6 or 7), (b) intermediate (days 12–14), or (c) late (days 18–21) time points in culture. Th17 cells were then transferred at equal numbers (2 × 106 Th17 cells/mouse) from each time point into lymphodepleted mice bearing established melanoma. Note that for late Th17 culture, the infusion product represented a mere 1/2,500 of their total cell count. However, regardless of the duration of in vitro expansion, Th17 cells killed tumors (Figure 3F) and supported long-term survival (Figure 3G) in the majority of mice.

Figure 1. High numbers of transferred Th17 cells are required for successful antitumor response. (A) RORγt and Tbet expression by Th17 cells after 7 days of culture; representative of 3 independent experiments. (B) IL-17A and IFN-γ expression by Th17 cells after 7 days of culture; representative of 5 independent experiments. (C) Th17 cells cultured ex vivo for 9 days were administered to mice with B16F10 melanoma after nonmyeloablative (5 Gy) total body irradiation 1 day prior to treatment with the indicated Th17 cell doses; n = 6–7 mice/group, representative of 3 independent experiments. (D) Percentage survival of mice treated with 4.5, 1.5, or 0.5 × 106 Th17 cells/mouse. Kaplan-Meier curves compared by log-rank test. *P < 0.05.
T cell persistence correlates with enhanced treatment outcome in melanoma patients given ACT (1). Thus, the similar antitumor efficacy mediated by early- or late-culture Th17 cells in mice suggested that these cells — regardless of time in vitro — would engraft and persist in mice at similar levels. To test this idea, we followed donor TRP-1 CD45.2+ Th17 cells expanded for 7 (early) or 21 days (late) in CD45.1+ recipient mice over 28 days. Early and late Th17 cells were present at equivalent numbers after treatment at 7, 14, and 28 days in the spleen and after 7 days in the tumor (Figure 3H). The eye is a common off-site target for T cell models specific for melanocyte-associated antigens (i.e., tyrosinase) because of melanocytes in the pigment epithelium (34). Here, too, we detected early and late Th17 cells at comparable amounts in melanoma-bearing mice (Figure 3H).

In ACT clinical trials, a donor T cell with longer telomeres persists better in melanoma patients and is associated with improved survival (13, 14). We hypothesized that preserved telomere length during expansion both in vitro and in vivo might be a mechanism by which Th17 cells, at any stage of expansion, preserved their antitumor efficacy after in vitro expansion. Telomere length only slightly decreased in Th17 cells as they expanded in vitro (Figure 3I). However, Th17 cells from both early and late cultures maintained comparable telomere length 4 weeks after ACT (Figure 3J). Thus, along with Tcf7, another partial explanation for the durable antitumor response of Th17 cells may be their maintenance of telomere length in vitro and in vivo. Collectively, our data reveal that Th17 cells expanded for 3 weeks retain antitumor efficacy compared with 1-week-old counterparts, and that this may be due to persistent stem signaling and telomere length which sustain the in vivo persistence of these cells.

Th17 cells undergo polyfunctional conversion during expansion but do not change their interaction with host immune cells. We noticed that even without inflammatory signaling such as IL-12, which drives Th17 cell conversion to a Th1-like phenotype (termed Th17 plasticity; see refs. 35–37), Th17 cells became progressively polyfunctional. As seen by ELISA performed 18 hours after peptide-dependent reactivation, IL-17A and IL-17F decreased in concert throughout culture, IL-22 was lost rapidly after 1 week, and IL-2 and IFN-γ production increased during ex vivo expansion (Supplemental Figure 2A). This correlated with reduced levels of nuclear RORγt and STAT3, although both molecules persisted throughout culture (Supplemental Figure 2, B and C). Additionally, Th17 cells maintained nuclear STAT1, while Tbet was not detected (Supplemental Figure 2, B and C). We suspected that the increased polyfunctionality of
3-week-expanded Th17 cells compared with 1-week-old counterparts would increase the recruitment of host immune cells in the tumor-bearing mouse to inflammatory sites. In contrast, we found that mice treated with 21-day-expanded Th17 cells actually had slightly fewer (though not statistically significant) host CD4+ and CD8+ cells in the spleens and eyes of recipient CD45.1+ mice, indicated by their higher ratio of donor Th17 to host cells (Supplemental Figure 2, D and E).

These differences between early and late Th17 cells, as well as previous data showing that Th17 cells enhance tumor-specific CD8+ T cell immunity (23), led us to question if early or late Th17 cells utilize endogenous CD8+ T cells to a different degree in their antitumor response. We transferred early or late Th17 cells and antibody-depleted CD8+ T cells to address this question. Interestingly, all groups had similar antitumor responses and survival rates (Supplemental Figure 2, F and G), suggesting that Th17 cells are capable of tumor lysis independent of CD8+ T cells. However, one should be conservative with this conclusion, as other reports suggest depleting host T cells might be beneficial to ACT by removing competition for homeostatic cytokines (38), which may have confounded the results. Thus, while long-term-expanded donor Th17 cells undergo functional conversion, they recruit and interact with host immune cells in tumor-bearing mice comparably to infused Th17 cells expanded for 7 days in vitro.
Th17 cells but not Th1 cells resist senescence and apoptosis in vitro. We next tested whether TRP-1 Th1 cells, like TRP-1 Th17 cells, could retain their antitumor efficacy after long-term ex vivo expansion. While Th17 cells cleared tumors similarly after 1 or 2 weeks of expansion, 2-week-expanded Th1 cells lost their ability to control tumor growth (Figure 4, A and B). This is similar to the loss of antitumor efficacy by CD8+ T cells, which we found occurs in IL-12–primed pmel-1 CD8+ T cells even without REP. Expanding these cells for only 10 days still caused a dramatic reduction in antitumor activity compared with 4-day-expanded CD8+ T cells (Supplemental Figure 3, A and B). Note that the CD8+ T cells lost control of tumors even after in vitro activation (a substitute for antigen vaccination; see ref. 39) and post-transfer injections of high-dose IL-2, which were not used in either our Th17 or Th1 therapies.

To understand why Th17 and not Th1 cells had durable antitumor efficacy during expansion, we surveyed expression of memory and other surface markers as well as levels of apoptosis in culture over time. Th1 cells had a significant cohort of central memory cells at 1 week of culture compared with Th17 cells, as indicated by their CD62L expression (32% on Th1 vs. 4% on Th17). However, by the second week, CD62L decreased on Th1 cells by half (Figure 4, C and D). Thus, CD62L expression on Th1 cells appears to correlate with their antitumor potential, similar to CD8+ T cells.

Unlike Th17 cells, Th1 cells maintained CD69 and CD25 expression between day 7 and day 14. Importantly, however, levels of CD25 on early Th1 cells were lower than on early Th17 cells, indicating a reduced capacity to respond to the homeostatic cytokine IL-2. Similarly, Th1 cells also had less CD127 than Th17 cells (Figure 4D). We assayed the levels of other markers that, along with homeostatic cytokine receptor levels,
can indicate T cell fitness (33, 40). We observed that levels of CD28 were lower on early Th1 cells compared with early Th17 cells, and CD27 expression decreased on Th1 cells between day 7 and day 14, in contrast to Th17 cells (Figure 4E). Additionally, KLRG1 and the checkpoint inhibitor TIM3 were higher at each time point on Th1 cells versus Th17 cells. Interestingly, PD1 expression on early Th17 and Th1 cells was similar, but expression of this inhibitory molecule declined by day 14 on Th17 cells, while it increased on Th1 cells (Figure 4E).

To determine if expression of exhaustion/checkpoint molecules was merely an artifact of activation, we activated day-14 Th17 and Th1 cells using irradiated splenocytes bearing TRP-1 peptide. While CD69 expression increased substantially on Th17 cells, KLRG1 did not, and TIM3 and PD1 only increased marginally. However, expression of all these markers was higher on Th1 cells compared with Th17 cells with or without activation and was not affected by activation status (Supplemental Figure 4A).

An important difference between Th1 and Th17 cells is their tendency to undergo apoptosis and necrosis in culture. At day 7 ex vivo, 18% of Th1 cells were single positive for annexin V (indicating early entry into apoptosis), while 42% stained positive for propidium iodide (PI) (indicating either the end stages of apoptosis or necrosis). On the other hand, only 2% of Th17 cells were annexin V single positive or PI positive (Figure 4F). By day 14, 18% of Th1 cells were annexin V single positive and 3% were PI positive, while Th17 cells had 4% annexin V–single-positive and no PI-positive cells (Figure 4F). However, Th1 cells demonstrated much more sensitivity to activation-induced cell death than Th17 cells. Twelve hours after TCR engagement with an anti-CD3 antibody we observed that up to 55% of Th1 cells were annexin V single positive and 8% PI positive. Yet, Th17 cells remained resistant to apoptosis upon reactivation, demonstrated by only a slight increase in annexin V staining from 4% to 8% on these cells (Figure 4F). We observed a similar pattern of increased annexin V+ cells in Th1 cells but not Th17 cells after TRP-1 and irradiated-splenocyte activation (Supplemental Figure 4B). These data suggest that Th17 cells maintain durable memory responses after long-term expansion due in part to their resistance to senescence and apoptosis, which may explain their improved ability to regress tumors compared with Th1 or CD8+ T cells.

### Large numbers of Th17 cells cure mice with aggressive tumors

Up to this point, we tested the antitumor capacity of Th17 cells expanded for 1 to 3 weeks on a per-cell basis. This ignored the vast expansion of these cells by the end of 3 weeks of ex vivo expansion. We hypothesized that more Th17 cells (>1 \times 10^9) would be required to ablate large tumors and therefore would necessitate long-term expansion.

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**Figure 5. Successful Th17 cell adoptive cell therapy against large tumors requires the cell numbers achieved by long-term ex vivo expansion.**

(A) Mean (± SEM) number of Th17 cells administered to mice with large (median 136 mm^2) B16F10 tumors from parallel cultures expanded to either early, intermediate, or late culture time points and divided among 12 mice/culture. Boxes represent the 25th to 75th quartiles and bars represent the minimum and maximum values; n = 2 cultures. (B) Tumor control in mice treated with early-, intermediate-, or late-culture Th17 cells; n = 12 mice/group, representative of 3 independent experiments. (C) Mean time (± SEM) in days after treatment until tumors reached peak size (mm^2); n = 12 mice/group. Mean time to peak tumor area compared by 1-way ANOVA with multiple comparisons; ***P < 0.001. (D) Mean (± SEM) peak tumor area (mm^2) in each mouse before onset of tumor control; n = 12 mice/group. Mean tumor areas compared by 1-way ANOVA with multiple comparisons; *P < 0.5. (E) Percentage survival of mice treated with early-, intermediate-, or late-culture Th17 cells; n = 12 mice/group, representative of 3 independent experiments. Kaplan-Meier curve compared by log-rank test; ***P < 0.001. (F) Onset of vitiligo in treated mice for each group; n = 12 mice/group. Kaplan-Meier curve compared by log-rank test; **P < 0.01, ns, not significant.
We therefore capitalized on the high yield of Th17 cells, which still had remarkably preserved effector potency, to regress large, recalcitrant melanoma. To obtain differentially aged Th17 cell products and test their potential against large tumors, parallel cultures were grown from one Th17 polarization to early (day 7), intermediate (day 14), or late (day 18) time points, cryopreserved, and then split evenly among 12 mice/group. This protocol yielded mean treatment doses of $1.3 \times 10^6$, $9.9 \times 10^6$, or $1.3 \times 10^7$ cells/mouse, respectively, to treat mice with large melanomas (~136 mm$^2$) (Figure 5A). As anticipated, we found that tumors regressed rapidly in mice treated with the high numbers of Th17 cells from either the intermediate or late cultures (Figure 5B). This was evidenced by the tumors in these mice reaching peak size earlier and having a smaller peak tumor size than mice treated with early Th17 cells (Figure 5, C and D). Conversely, mice treated with $1.3 \times 10^6$ early Th17 cells were unable to control tumor growth, with most mice succumbing to disease (Figure 5, B and E). The reduced antitumor efficacy of early Th17 cells was paralleled by a delay in onset of autoimmune vitiligo (Figure 5F). However, all mice given Th17 cells (at any amount or at any age) eventually developed a similar degree of vitiligo (Supplemental Figure 5A).

Infusing more Th17 cells from long-term expansion sets up long-term donor residence with enhanced memory response against antigen and tumor rechallenge. As expected, higher frequencies of donor Th17 cells were detected in the spleen and tumor draining lymph nodes (dLN) of surviving mice 40 days after treatment; $n = 5$ mice/group. Mean frequencies compared by 1-way ANOVA with multiple comparisons; *$P < 0.05$, **$P < 0.01$. TRP, CD4+ T cells with a TCR specific for tyrosinase-related protein-1. (C) Memory recall assessed by in vivo lysis of peptide-pulsed splenocytes 40 days after treatment compared with control mouse (mean percentage specific lysis ± SEM); $n = 5$ mice/group. Mean percentage specific lysis compared by 1-way ANOVA with multiple comparisons; **$P < 0.01$. (D) Macropscopic images of lungs and microscopic images of en-bloc sections from intermediate or late Th17-treated mice compared with control mice 21 days after rechallenge with $0.2 \times 10^6$ B16F10 cells injected i.v.; $n = 2$-6 mice/group. (E) Quantification of percentage of lung taken up by tumor as determined by microscopic examination (mean ± SEM); $n = 2$-6 mice/group. Mean percentage area compared by 1-way ANOVA with multiple comparisons; ****$P < 0.0001$. 

Infusing large numbers of Th17 cells after 2 or 3 weeks of expansion sets up long-term donor residence with enhanced memory response against antigen and tumor rechallenge. (A and B) Mean frequency (± SEM) of donor Th17 cells in (A) spleens and (B) draining lymph nodes (dLNs) of surviving mice 40 days after treatment; $n = 5$ mice/group. Mean frequencies compared by 1-way ANOVA with multiple comparisons; *$P < 0.05$, **$P < 0.01$. TRP, CD4+ T cells with a TCR specific for tyrosinase-related protein-1. (C) Memory recall assessed by in vivo lysis of peptide-pulsed splenocytes 40 days after treatment compared with control mouse (mean percentage specific lysis ± SEM); $n = 5$ mice/group. Mean percentage specific lysis compared by 1-way ANOVA with multiple comparisons; **$P < 0.01$. (D) Macroposcopic images of lungs and microscopic images of en-bloc sections from intermediate or late Th17-treated mice compared with control mice 21 days after rechallenge with $0.2 \times 10^6$ B16F10 cells injected i.v.; $n = 2$-6 mice/group. (E) Quantification of percentage of lung taken up by tumor as determined by microscopic examination (mean ± SEM); $n = 2$-6 mice/group. Mean percentage area compared by 1-way ANOVA with multiple comparisons; ****$P < 0.0001$. 

To determine if there was a difference between treating mice with intermediate or late Th17 cells, we administered a second dose of subcutaneous B16F10 to the cured animals. All animals from both treatment groups were protected from rechallenge (Supplemental Figure 5B), underscoring the therapeutic ability of Th17 cells to mediate durable tumor immunity. To further test the potency of this therapy, we rechallenged the mice again with a more aggressive metastatic model. B16F10 tumor cells were administered intravenously to mice to induce lung pseudometastases, which are far more challenging to treat than the subcutaneous tumors. Despite causing aggressive tumor outgrowth in the lungs of control mice, the lungs of all mice previously...
treated with either intermediate or late Th17 cells were protected from metastasis (Figure 6D and Supplemental Figure 5C). We performed histological analysis of the lungs to confirm our macroscopic observations of no metastatic tumor growth. Stepped sections were cut through whole-lung blocks (which included the heart and surrounding structures) from each group to determine the presence of any macroscopic or microscopic metastatic tumor deposits. In concordance with macroscopic findings, clear tumor nodules were noted in control lungs, while no melanoma deposits were seen in any of the other treatment groups (Figure 6D and Supplemental Figure 5C). In control mice, tumors occupied ~61% ± 4.5% of the total lung volume (Figure 6E). These results highlight the capacity of long-term-expanded Th17 cells to eradicate tumors and provide durable protection from tumor metastasis.

Human Th17-polarized, but not Th1-polarized cells retain antitumor activity after expansion. As an important step toward translation of this work to clinical trials, we sought to determine if human Th17 cells possess the same undiminished capacity to clear tumors after long-term expansion as murine Th17 cells. We expected that human Th17 cells would remain highly therapeutic, while Th1 cells would decline in efficacy after expansion, similar to our observation with murine Th17 versus Th1 cells. To examine function, phenotype, and antitumor efficacy of human Th17 and Th1 cells, human CD4+ T cells from the peripheral blood of normal donors were polarized to either a Th17 or Th1 phenotype (see Methods). These 2 subsets were then expanded for 2 or more weeks after activation with anti-CD3 beads coated with ICOS and CD28, and redirected to express a chimeric antigen receptor (CAR) that recognizes mesothelin and signals through CD3ζ and 4-1BB (MesoCAR). We also expanded bulk CD8+ T cells with CD3/CD28/ICOS beads and IL-2, and engineered them with the same MesoCAR construct. During expansion, Th17 cells became more polyfunctional with increased capacity to secrete IFN-γ and IL-17A (Figure 7A). Interestingly, at all time points human Th17-polarized cells secreted more IFN-γ than Th1-polarized cells. This may be because among the cells derived from the peripheral blood were mature T cells that may have had prior functional profiles before polarization compared to the mainly naive T cells from TRP-1 transgenic mice.

However, Th17 polarization did induce high expression of the Th17-associated CCR6 chemokine, which was largely absent on Th1-polarized cells (Figure 7B). However, Th1 and Th17 cells both expressed similar levels of CXCR3, a chemokine receptor associated with Th1 cells. Thus, a large portion of Th17 cells became double positive for CCR6 and CXCR3 by the end of culture, a phenotypic marker of a highly functional nonclassical Th17/Th1 hybrid cell (Figure 7B) (41). Additionally, Th17 cells with the highest CCR6 expression also expressed CCR5 and CCR10, 2 chemokine receptors important for trafficking to inflamed tissues (42, 43) (Supplemental Figure 6, A and B). While present, these chemokine receptors were at lower concentrations on Th1 cells. Similar to murine CD4+ T cells, human Th1 cells expressed more CCR6L2 at the beginning of culture than human Th17 cells, but by day 12, both subsets had less (though still ample) CD62L expression, indicating a progressive shift to an effector memory phenotype (Supplemental Figure 6C) — though not to the extent seen in murine Th17 cells (Figure 4A).

To test whether human Th17-polarized cells preserved antitumor activity during ex vivo expansion, we infused 7- or 12-day-expanded MesoCAR+Th17 cells (2 × 10⁷/mouse) with 12-day-expanded MesoCAR+CD8+ T cells (2 × 10⁶/mouse) into NSG mice with established M108 mesothelioma tumors. As a control, we also treated mice with 7- or 12-day-expanded MesoCAR+ Th1 cells (2 × 10⁷/mouse) with 12-day-expanded MesoCAR+CD8+ T cells (2 × 10⁶/mouse). We found that MesoCAR+ Th17 cells eradicated human mesothelioma regardless of expansion time (Figure 7C). Conversely, treatment with 12-day-expanded MesoCAR+ Th1 mediated poor antitumor activity compared with treatment with 7-day-expanded MesoCAR+ Th1 cells (Figure 7C). Importantly, Th17 cells could not mediate tumor immunity if engineered with a CAR against CD19 (Supplemental Figure 6D). This is not surprising, as mesothelioma tumors do not express CD19 antigen. Also, we tested the benefit of antigen-specific MesoCAR+CD8+ T cells in our human model by cotransferring MesoCAR+ Th17 cells with CD19CAR+CD8+ T cells. While antigen-specific Th17 cells with CD19CAR+CD8+ T cells controlled tumor growth well (and better than MesoCAR+CD8+ T cells transferred with CD19CAR+ Th17 cells), we only observed curative responses when both the Th17 cells and CD8+ T cells were antigen specific against the correct target, mesothelin (Supplemental Figure 6D). Collectively, our data reveal that human Th17 cells retain their antitumor capacity when expanded long term ex vivo. Our human data confirm our findings that durable Th17 cells generated from ex vivo expansion (Figure 3), which dramatically eliminated large and metastatic melanoma tumors in mice (Figures 5 and 6), translates to human T cells with major implications for improving ACT clinical trials (Figure 7).
Discussion

ACT mediates objective response rates in ~50% of cancer patients. This therapy relies on the infusion of large numbers of T cells ($10^{11}$) to achieve this goal. Although CD8$^+$ T cells are often used for ACT, they lose efficacy as they expand ex vivo. Herein, we discovered that Th17 cells have an undiminished capacity to regress tumors as they expand. Transferring $2 \times 10^6$ Th17 cells expanded for 3 weeks ex vivo mediated antitumor immunity in mice similar to that mediated by infusing $2 \times 10^6$ younger Th17 cells expanded for 1 week. In contrast, neither Th1 nor CD8$^+$ T cells retained antitumor efficacy after only 2 weeks of expansion. Thus, on a per-cell basis, the antitumor capacity of Th17 cells is more effective than Th1 or CD8$^+$ T cells after long-term ex vivo expansion.

We hypothesized that the retention of antitumor efficacy by Th17 cells, coupled with their expansion to large numbers over 3 weeks, would allow us to treat large recalcitrant melanoma tumors (~175 mm$^2$) by treating mice using the entire cell product. Indeed, this high-dose Th17 therapy eradicated tumors in mice and mediated curative responses for 6 or more months. Also, our cured animals were protected from a subsequent tumor rechallenge both subcutaneously and intravenously to model aggressive lung metastases. In contrast, the few Th17 cells expanded for only 1 week were insufficient to clear even primary tumors. To our knowledge, this is the first report that concomitantly investigates 4 factors critical for ACT: (a) T cell age, (b) T cell yield, (c) T cell subset, and (d) level of tumor burden. Our findings are relevant, as they provide key insight into the potency of Th17 cells, which expand robustly without diminishing in quality.

We and other investigators have reported beneficial traits of Th17 cells for ACT (18, 27, 28). Unlike Th1 cells, Th17 cells do not express KLRG1 or TIM3 during expansion, while cytokine receptors and costimulatory molecules are more abundant on Th17 cells than Th1 cells. Moreover, although resistance to apoptosis in Th17 cells has been reported (27, 44), we provide evidence that their resistance to apoptosis is continuous throughout in vitro culture. We suspect that these attributes also permit Th17 cells to persist in the harsh tumor microenvironment.
Effector memory CD8+ T cells (CD62L-) are far less effective at regressing tumors when infused into mice or humans compared with their less differentiated counterparts, such as stem and central memory CD8+ T cells (11, 16, 45). This observation begs the question: how can early or late effector memory Th17 cells that express nominal CD62L persist and still regress tumors in vivo? While previous studies indicate that Th17 cells have active β-catenin signaling after short-term in vitro expansion (28), our Western blot analysis of tumor-reactive Th17 cells found the transcription factor Tcf7 present in the nucleus at all time points during expansion. This observation is insightful, as Tcf7 is an essential protein in the Wnt/β-catenin pathway that is critical for stem memory T cell self-renewal and was very recently found to be critical for formation of memory daughter cells through asymmetric T cell division (28, 29). Therefore, we posit that constitutive Tcf7 signaling may partially explain why Th17 cells expand robustly in a Petri dish and persist long term in vivo. This finding might also explain why Th17 cells maintain their telomere length in vitro and in vivo, a property directly associated with T cell persistence in patients with enhanced survival in ACT trials (1, 13, 14).

We found that Th17 cells are capable of directly lysing tumors. Both early- and late-expanded donor Th17 cells regressed melanoma independently of host CD8+ T cells, as antibody depletion of CD8+ T cells did not compromise treatment outcome. This is in alignment with a report by the Antony lab (46) that found tumor-reactive CD4+ T cells clear tumors in a class II–restricted manner in vivo. However, in our human CAR model, we found that infusion of both antigen-specific Th17 cells and CD8+ T cells was needed to mediate optimal antitumor responses, as MesoCAR+ Th17 cells were unable to eradicate the tumors when coinfused with nonspecific CD8+ T cells (i.e., CD19CAR+CD8+ T cells) (Supplemental Figure 6D). This finding agrees with a study by the Dong laboratory (23), which found that depletion of endogenous CD8+ T cells diminished the antitumor response of Th17 cells, though it did not abrogate it completely. It is also important to appreciate that a possible confounding variable of depleting host CD8+ T cells in our TRP-1 ACT melanoma model would be the creation of space for the donor Th17 cells to engraft and better consume homeostatic cytokines (38, 47). Collectively, our data support a multilateral model where donor Th17 cells directly lyse tumors as well as cooperate with other immune cells, such as CD8+ T cells, to protect the host from tumor relapse long term.

Based on recent evidence supporting the benefit of shorter preparation time for ACT cell products (11, 48), cancer immunotherapists are devoting significant effort to reduce T cell expansion time ex vivo before infusion. A clinical trial at the NCI reported that they could generate young TILs that could mediate improved regression of metastatic melanoma. Yet, these TIL products were still expanded for a month or more ex vivo before being returned to the patient (49). Additionally, it is important to note that many parameters besides time contribute to CD8+ T cell exhaustion/loss of efficacy including: the number of times the T cells are restimulated; the growth factor cytokines used (IL-2 vs. IL-15 vs. IL-21); and the type of costimulatory signals (CD28, ICOS, or 4-1BB) delivered to expand the cells, all of which can alter their cellular bioenergetics (19, 50–54). In particular, IL-2 is the staple growth factor used to expand TIL and CAR-T cells for ACT. However, while this cytokine has potent effects on proliferation, IL-2 also differentiates CD8+ T cells to an effector or terminal status (55). These considerations give further credence to the use of T cells with stemness, such as Th17 cells, or pharmacologically altered CD8+ T cells (16, 17, 28, 56), both of which expand long term without reaching terminal differentiation.

Of clinical importance, human CAR+ Th17-polarized cells also retain their antitumor potency after long-term expansion, while human CAR+ Th1-polarized cells do not. The antitumor activity of human CAR+ Th17 cells in our mouse xenograft model of mesothelioma was superior to that of CAR+ Th1 after either 7 or 12 days of ex vivo expansion. These findings recapitulate those in our TRP-1 melanoma model where murine Th17 cells mediate greater tumor immunity than TRP-1 Th1 cells. Of note, while the shortest reported TIL expansion protocols still require 1 month of ex vivo expansion, our data with 12-day-expanded human Th17 CD4+ T cells is particularly relevant in the context of CAR therapies where many T cells can be quickly generated from peripheral blood within 1 or 2 weeks after apheresis.

In closing, Th17 cells offer promise for next-generation ACT trials, as they can: (a) expand rapidly in vitro for 3 weeks without requiring reactivation, (b) repopulate the host after infusion regardless of ex vivo expansion time, and (c) mediate potent destruction of large tumors when the total number of cells generated after 2 or more weeks of ex vivo culture is used. Th17 cell durability may also result in more consistent success in generating a therapeutic cell product, owing to a larger window of opportunity for securing potent T cells after ex vivo expansion. The resilience of Th17 cells in vitro may also simplify clinical preparation, rendering them more feasible to prepare for clinical trials in many cancer centers around the world, thereby extending ACT treatment to a wider patient population.
Methods

Mice and tumor lines
C57BL/6J (B6), CD45.1 congenic B6, TRP-1 TCR transgenic mice, and NOD/scid/γ chain knockout (NSG) mice were purchased from Jackson Laboratories and housed and bred in the Medical University of South Carolina (MUSC) Hollings Cancer Center comparative medicine department. NSG mice were housed under specific pathogen–free conditions in microisolator cages and given autoclaved food and acidified water. Housing and experiments were conducted in accordance with MUSC's IACUC procedures. B16F10 (H-2b) melanoma tumor, a gift of the Nicholas Restifo lab at the NCI surgery branch, was maintained in culture media (RPMI 1640 with L-glutamine; 10% FBS; 1% penicillin/streptomycin, nonessential amino acids, and Na pyruvate; and 0.1% 2-ME and Hepes). M108 xenograft tumors were a gift from the June lab at the University of Pennsylvania Philadelphia, Pennsylvania, USA. M108 were cultured and engrafted as described previously (57).

T cell cultures

**TRP-1 cells.** TRP-1 splenocytes were activated using 10-Gy-irradiated B6 splenocytes (feeder cells) pulsed with 1 μM TRP-1 peptide and polarized to a Th17 phenotype at 3 × 10⁶ cells/2 ml of cell media in 1 well of a 24-well plate with the following cocktail: 100 ng/ml recombinant human (rh) IL-6 (NIH repository), 100 ng/ml rhIL-21 (Shenandoah), 30 ng/ml rhTGF-β1 (Biologend), 10 ng/ml rhIL-1β (NIH Repository), 10 μg/ml each of anti–mIFN-γ (clone XMG1.2), anti–mouse IL-4 (clone 11B11), and anti–mIL-2 (clone JES6-1A12) (all BioXCell). Th1 polarization occurred under peptide activation with irradiated feeder cells with the following cocktail: 3 ng/ml recombinant mouse (rm) IL-12 (Shenandoah), 100 IU/ml rhIL-2 (NIH repository), and 10 μg/ml anti–mIL-4 (clone 11B11). Cultured cells were supplemented with new media containing 100 IU/ml rhIL-2 throughout expansion, then cryopreserved at time points indicated in 10% DMSO in FBS. One day before transfer, cells were thawed and plated in cell media overnight and then transferred into mice.

**Pmel-1 T cells.** Pmel-1 splenocytes were activated using 1 μM human gp100 peptide + 100 IU rhIL-2/ml and primed on day 1 with 40 ng/ml of rmIL-12. Cells were supplemented with culture media containing 100 IU rhIL-2/ml and expanded as indicated. CD8+ T cell cultures were in vitro activated with feeder cells and peptide 12 hours before transfer as described (39).

**Human normal donor peripheral T cells.** To generate mesothelin-specific T cells, human CD4+ T cells were activated with CD3/CD28/ICOS–coated beads and programmed to a Th17 or Th1 phenotype and then transduced with a CAR that contained either an anti-mesothelin or anti-CD19 single-chain variable fragment (scFv) fusion protein containing the TCRζ signaling domain and 4-1BB that was generated as described previously (57) (gifts from the June lab). CD4+ T cells were polarized to the Th17 phenotype as follows: 10 ng/ml rhIL-1β, 10 ng/ml rhIL-6, 20 ng/ml rhIL-23, 10 μg/ml anti–hIL-4 (clone 11B11), and anti–hIFN-γ (clone H22, eBioscience). Experiments were conducted with fetal calf serum containing endogenous sources of TGF-β. Th1 cells were polarized with 10 ng/ml rhIL-12 and 100 IU/ml rhIL-2. Cell cultures were maintained with 100 IU/ml of rhIL-2 and cells were expanded for up to 2 weeks. Likewise, human CD8+ T cells were activated with CD3/CD28/ICOS beads and redirected with either the anti-MesoCAR or anti-CD19CAR constructs.

ACT

**B6 mice.** B6 mice were given 4.5 × 10⁵ B16F10 cells subcutaneously and tumors were allowed to establish between 5 and 12 days before ACT. One day before therapy, mice received nonmyeloablative 5 Gy total body irradiation. T cells were infused via tail vein. In the experiments with IL-12–primed CD8+ T cells, starting on the day of ACT, mice were given intraperitoneal injections of 600,000 IU rhIL-2 every other day for a total of 6 doses.

**NSG mice.** NSG mice were given 5 × 10⁶ M108 suspended in matrigel subcutaneously. Tumors were allowed to establish for 40 days prior to adoptive therapy. In all experiments, mice were randomized to treatment groups and tumor burden was monitored in blinded fashion using perpendicular caliper measurements and reported as tumor area (mm²).

Antibody depletion in vivo

For cell depletion assays, mice were injected intraperitoneally with 100 μg of either anti-CD8 depleting antibody (clone 53-6.72) or isotype control (clone 2A3) (both BioXCell) beginning the day before ACT and continuing weekly throughout the duration of the experiment.
Vitiligo autoimmunity score
Severity of autoimmune vitiligo was scored after the phenotype stabilized. Depigmentation was rated based on a graduated scale from 1 to 5 with 1 being no loss of pigment (all black) and 5 being total loss of pigment (all white). Mice were scored in blinded fashion in triplicate.

Tissue distribution assays
Spleens from treated mice were harvested and mechanically disrupted using the tip of a syringe plunger. Cells were filtered through a wire mesh, red blood cells lysed with RBC lysis buffer (Biolegend), and then resuspended in cell media for analysis. Eyes and tumors were sectioned, and then incubated in 1 mg/ml collagenase type II (Life Technologies) at 37°C for 1 hour. Digested tissue was filtered, resuspended in cell media, and plated for assay. Before probing with antibodies, FC block (Biolegend) was applied to cells at 1 μg/100 μl. Cell suspensions were then analyzed by flow cytometry.

Flow cytometry, ELISA, and in situ flow
Flow cytometry was performed with a BD FACsVerse instrument. Antibodies used were: anti-mCD3e-Fluor450 (clone 17A2), anti-h/mRORγt-PE (clone AFKJS-9) (both eBioscience); anti-m4-1BB-Biotin (clone 17B5), anti-hCCR6-PECy7 (clone G034E3), anti-hCCR10-PE (clone 6588-5), anti-mCD25-FITC (clone 7D4), anti-mCD27-PECy7 (clone LG.3A10), anti-mCD28-PerCP-Cy5.5 (clone 37.51), anti-h/mCD44-PerCP-Cy5.5 (clone IM7), anti-mCD69-PECy7 (clone HI.2F3), anti-mCD127-PE (clone A7R34), anti-hCXCR3-APC-F467 (clone G025H7), anti-mLAG3-PE (clone C9B7W), anti-mPD1-PerCP-Cy5.5 (clone 29F.1A12), Streptavidin-BV421, anti-mTIM3-PE (clone B8.2C12) (all Biolegend); anti-hCCR5-V450 (clone 2D7/CCR5), anti-mCCR7-PE (clone 4B12), anti-hCD4-APC/PE (clone RM4-5), anti-mCD45.1-APC-Cy7 (clone A20), anti-mCD45.2-V500 (clone 104), anti-mCD62L-APC (clone ME1-14), anti-mCD95-PECy7 (clone Jo2), anti-miCOS-Biotin (clone 7E.17G9), anti-mKLRG1-APC (clone 2F1), anti-mNK1.1-PE (clone PK136), anti–h/mTbet-AF647 (clone 4B10), and anti-mVB14-FITC (clone 14-2) (all BD). Apoptosis assays were performed with Biolegend’s FITC Annexin V Apoptosis Detection Kit with PI, per the manufacturer’s instructions. ELISAs for IL-17A, IL-17F, IL-22, IFN-γ, and IL-2 were performed using DuoSet ELISA kits (R&D Systems) per the manufacturer’s instructions. Telomere length was measured by in situ hybridization/flow cytometry using the Telomere PNA Kit/FITC via Flow Cytometry (Dako) per the manufacturer’s instructions.

In vivo cytotoxicity assay
C57BL/6 splenocytes were incubated with either TRP peptide or OVA peptide residues 323–339 (OTII) (Invivogen) and then labeled with a low level (0.5 μM) or high level (5 μM) of cell trace violet (CTV) (Thermo Fisher Scientific), respectively. TRP- and OTII-pulsed splenocytes were mixed at an equal ratio and then injected intravenously into mice previously treated with Th17 cells or untreated control mice. After 12 hours, spleens were harvested from mice and the ability of donor cells to lyse TRP-pulsed splenocytes versus OTII-pulsed splenocytes was assessed via flow cytometry and reported as percentage specific lysis using the following equation: % specific lysis = (1 – [ratio of no-T-cell-control mice] / [ratio of ACT mice]) × 100, where the ratio = % OTII/% TRP.

Western blot
Nuclear protein from Th17 cells cultured for 7, 14, or 21 days was extracted via the NE-PER Nuclear and Cytoplasmic Extraction kit (Thermo Fisher Scientific) and then analyzed by Western blot. The following primary antibodies were used: rabbit anti–h/mTCF7 (Abcam, clone EPR2035); rabbit anti–h/mHistone H3 (clone D1H2), rabbit anti–h/mSTAT1 polyclonal (catalog 9172), rabbit anti-h/mSTAT3 (clone D32Z2G) (all Cell Signaling Technology); rat anti–h/mRORγt (clone AFKJS-9), and mouse anti–h/mTbet (clone 4B10) (both eBioscience).

Histology
For lung histology and morphometry, lungs were inflated en bloc and fixed with 4% buffered paraformaldehyde at a constant hydrostatic pressure of 25 cm and further immersed in fixative for 48 hours. The inflated lungs were embedded in paraffin and 4-μm sagittal stepped sections were stained with H&E for histological evaluation. In total, 10 stepped sections from each lung were analyzed to calculate the volume of metastatic...
tumor in each group using an automated motorized stage Olympus BX61 microscope to generate lung superimages using Visiopharm software. The total area of lung and tumor metastases was calculated using Visiopharm image acquisition software by an observer blinded to the experimental groups.

Statistics
Kaplan-Meier survival curves were assessed for significance using a log-rank test between treatment groups. Comparisons between 2 groups were analyzed using a 2-tailed Student’s t test with Welch’s correction for parametric distribution or Mann-Whitney signed-rank test for nonparametric distribution. For comparisons between multiple groups, a 1-way ANOVA was performed followed by multiple comparisons. P values less than 0.05 were considered significant.

Study approval
Studies were approved by the IACUC of the MUSC Animal Resource Center (number 3039). De-identified human peripheral blood mononuclear cells were collected under approval of the MUSC IRB (pro13570). Human T cells were engineered via approval from the MUSC institutional biosafety committee (number 2335).

Author contributions
JB and CP were responsible for conception and design of this project. Methodology was developed by JB, BR, CA, AK, LG, and CP. Data were acquired by JB, AK, CA, and CP. Interpretation and analysis of data were performed by JB, M. Nelson, KM, SB, CA, and CP. The following participated in writing, review, and revision of the manuscript: JB, MN, KM, SB, BR, LG, CP. CP was responsible for administrative, technical, and material support as well as supervision of this study.

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Address correspondence to: Jacob S. Bowers or Chrystal M. Paulos, Hollings Cancer Center, Medical University of South Carolina, 86 Jonathan Lucas Street, Room 606B MSC509, South Carolina 29425, USA. Phone: 843.792.8926; E-mail: bowersjs@musc.edu (J.S. Bowers). Phone: 843.792.3210; E-mail: paulos@musc.edu (C.M. Paulos).

ADMK’s present address is: Miltenyi Biotec, Bergisch Gladbach, Germany.
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