

Development of CAR T Cells Expressing a Suicide Gene Plus a Chimeric Antigen Receptor Targeting Signaling Lymphocytic-Activation Molecule F7

Christina Amatya,¹ Melissa A. Pegues,¹ Norris Lam,¹ Danielle Vanasse,¹ Claudia Geldres,¹ Stephanie Choi,¹ Stephen M. Hewitt,² Steven A. Feldman,³ and James N. Kochenderfer¹

¹National Institutes of Health, National Cancer Institute, Center for Cancer Research, Surgery Branch, Bethesda, MD 20892, USA; ²National Institutes of Health, National Cancer Institute Laboratory of Pathology, Bethesda, MD 20892, USA; ³Stanford University School of Medicine, Stanford, CA, USA

Chimeric antigen receptors (CARs) are fusion proteins that contain antigen-recognition domains and T cell signaling domains. Signaling lymphocytic-activation molecule F7 (SLAMF7) is a promising target for CAR T cell therapies of the plasma cell malignancy multiple myeloma (MM) because SLAMF7 is expressed by MM but not normal nonhematopoietic cells. We designed CARs targeting SLAMF7. We transduced human T cells with anti-SLAMF7 CARs containing either CD28 or 4-1BB costimulatory domains. T cells expressing CD28-containing CARs or 4-1BB-containing CARs recognized SLAMF7 *in vitro*. SLAMF7-specific cytokine release was higher for T cells expressing CARs with CD28 versus 4-1BB domains. In murine solid tumor and disseminated tumor models, anti-tumor activity of T cells was superior with CD28-containing CARs versus 4-1BB-containing CARs. Because of SLAMF7 expression on some normal leukocytes, especially natural killer cells that control certain viral infections, the inclusion of a suicide gene with an anti-SLAMF7 CAR is prudent. We designed a construct with a CD28-containing anti-SLAMF7 CAR and a suicide gene. The suicide gene encoded a dimerization domain fused to a caspase-9 domain. T cells expressing the anti-SLAMF7 CAR plus suicide-gene construct specifically recognized SLAMF7 *in vitro* and eliminated tumors from mice. T cells expressing this construct were eliminated on demand by administering the dimerizing agent AP1903 (rimiducid).

INTRODUCTION

Multiple myeloma (MM) is an almost-always incurable malignancy of plasma cells; new therapies are needed for MM.^{1,2} Chimeric antigen receptors (CARs) are artificial fusion proteins with antigen-recognition domains and T cell signaling domains.^{3,4} T cells expressing CARs targeting the MM-associated antigen B cell maturation antigen (BCMA) have proven clinical efficacy.^{5–10} However, variable BCMA expression and loss of BCMA expression from MM after anti-BCMA CAR T cell treatment have been documented,^{5,8} which makes development of CAR T cell therapies targeting other antigens an important goal.

Signaling lymphocytic-activation molecule F7 (SLAMF7), also known as CD319, CD2 subset-1 (CS1), and CD2-like receptor-activating cytotoxic cells (CRACCs), is a highly glycosylated protein that belongs to the SLAMF family of cell-surface receptor proteins.^{11,12} SLAMF7 is a promising CAR target for MM therapy due to high and uniform expression on MM cells and lack of expression on normal nonhematopoietic cells.^{13–15} Although the role of SLAMF7 in MM is unclear, SLAMF7 expression is speculated to enhance growth, survival, and adhesion of MM cells to bone marrow stromal cells.^{16,17} Additionally, SLAMF7 is also known to be expressed on a variety of other cells.^{15,18,19} A CAR targeting SLAMF7 has potential to eliminate SLAMF7-expressing leukocytes, especially natural killer (NK) cells and CD8⁺ T cells. Thus, despite being a promising therapeutic target for MM in preclinical studies,^{20–23} anti-SLAMF7 CAR T cells might cause cytopenias, especially NK cell deficiency that might lead to a high risk for severe viral infections, such as cytomegalovirus infections.^{24,25} A suicide gene could be a useful strategy to eliminate anti-SLAMF7 CAR T cells in case of severe toxicities.

The inducible caspase-9 (iCasp9 or IC9) suicide gene has been used to eradicate T cells in humans with reversal of clinical toxicity.^{26,27} The IC9 gene contains part of the human caspase-9 protein fused to a dimerization domain from the human FK-506-binding protein-12 (FKBP12).²⁸ Activation of the caspase-9 domain of IC9 is dependent on dimerization of FKBP12 domains; this dimerization occurs only upon administration of the small molecule drug rimiducid (AP1903).^{26,27}

An important consideration in CAR design is the selection of costimulatory domains. Costimulatory domains of most clinically used CARs are from either CD28 or 4-1BB.²⁹ CD28 costimulatory domains have been reported to provide faster and stronger functional activity to CAR T cells, and

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Correspondence: James N. Kochenderfer, National Institutes of Health, National Cancer Institute, Center for Cancer Research, Surgery Branch, 10 Center Drive, CRC Room 3-3330, Bethesda, MD 20892, USA.

E-mail: kochendj@mail.nih.gov

4-1BB costimulatory domains have been reported to enhance persistence of CAR T cells under some circumstances; however, the relative anti-tumor activity of CD28-containing CARs versus 4-1BB-containing CARs is unclear and might be dependent on a variety of conditions.^{30–36}

In this paper, we performed comparisons between anti-SLAMF7 CARs with either CD28 or 4-1BB costimulatory domains. We also designed and tested constructs encoding both an anti-SLAMF7 CAR and the IC9 suicide gene.

RESULTS

SLAMF7 Expression Is Limited to Certain Leukocytes

We assessed SLAMF7 expression in normal tissues. SLAMF7 RNA was detected by PCR in some tissues, including lymph nodes, tonsils, stomach, bone marrow, and spleen, which is likely explained by the presence of SLAMF7⁺ leukocytes infiltrating these organs; otherwise, SLAMF7 RNA was not detected (Figure 1A). Next, we assessed SLAMF7 protein expression by immunohistochemistry using a normal human tissue microarray (Table S1). SLAMF7 protein was not expressed in major organs except in plasma cells, lymphocytes, and macrophages. SLAMF7 was not expressed on CD34⁺ hematopoietic stem cells (Figure 1B). We confirmed high and uniform expression of SLAMF7 on CD38⁺ cells from bone marrow of MM patients (Figure 1C). We also assessed SLAMF7 expression on peripheral blood mononuclear cells (PBMCs). We found SLAMF7 expression on most CD56⁺ NK cells, some CD8⁺ T cells, and small fractions of monocytes and CD4⁺ T cells (Figures 1D and 1E). We assessed SLAMF7 expression on CD4⁺ and CD8⁺ T cells before and after activation with the anti-CD3 antibody OKT3 in interleukin (IL)-2-containing medium. This is the same T cell culture process used in other experiments reported in this paper and in clinical trials conducted by our group. SLAMF7 expression increased with activation of either CD4⁺ or CD8⁺ T cells. As expected, SLAMF7 expression was higher on CD8⁺ T cells versus CD4⁺ T cells (Figures 1F–1H).

Anti-SLAMF7 CAR Expression

We constructed CARs incorporating single-chain variable fragments (scFvs) derived from either the Luc90 or Luc63 monoclonal antibodies (Figure 2A). These CARs also included hinge and transmembrane domains from CD8 α , CD28 costimulatory domains and CD3 ζ activation domains. The CARs were encoded by the mouse stem cell virus-based splice-gag vector (MSGV1) gamma-retroviral vector,³⁷ and the CARs were designated Luc90-CD828Z and Luc63-CD828Z. The Luc90 scFv is derived from a murine monoclonal antibody, and the Luc63 scFv is derived from the humanized version of the monoclonal antibody Luc63, also known as elotuzumab. After transduction, Luc90-CD828Z and Luc63-CD828Z were expressed on the surface of CD4⁺ and CD8⁺ T cells (Figures 2B and 2C); both CARs had similar transduction efficiencies (Figures 2D and 2E).

Function of Luc63- and Luc90-Containing CARs

There was no difference in SLAMF7-specific degranulation when Luc90-CD828Z and Luc63-CD828Z were compared for CD4⁺ (Figure 2F) and CD8⁺ (Figure 2G). Interferon (IFN)- γ release, when T cells were cultured with SLAMF7⁺ target cells, was slightly higher

for T cells expressing Luc90-CD828Z versus T cells expressing Luc63-CD828Z. Background IFN- γ release, when CAR T cells were cultured with SLAMF7-negative target cells and when CAR T cells were cultured alone, was slightly higher with Luc63-CD828Z versus Luc90-CD828Z (Figure 2H). Due, in part, to slightly lower nonspecific activity and slightly higher SLAMF7-specific IFN- γ release, we decided to use Luc90-CD828Z in future experiments, although functional differences between these two CARs were small.

MM patients have detectable (range, 1–80 ng/mL) serum SLAMF7.¹⁴ We sought to determine if adding soluble SLAMF7 to cultures could inhibit anti-SLAMF7 CAR T cell activity. We conducted IFN- γ ELISA by coculturing Luc90-CD828Z CAR T cells with SLAMF7⁺ target cells and adding varying concentrations of soluble SLAMF7. Soluble SLAMF7 did not block antigen-specific IFN- γ release by anti-SLAMF7 CAR T cells (Table S2).

Expression of CARs Containing CD28 versus 4-1BB Costimulatory Domains

We compared Luc90-CD828Z to Luc90-CD8BBZ, CARs that differ from each other only in their costimulatory domains, as shown in Figure 3A. Both Luc90-CD828Z and Luc90-CD8BBZ were expressed on T cells (Figures 3B and 3C). The percentages of transduced T cells that expressed the two CARs and the level of expression on each transduced T cell were not statistically different (Figures 3D and 3E). Interestingly, the CD4-to-CD8 ratio of Luc90-CD8BBZ-expressing T cells was higher than the CD4-to-CD8 ratio of Luc90-CD828Z-expressing T cells (Figure 3F). Analysis for memory phenotype showed that for both CD4⁺ and CD8⁺ T cells, the percentage of central memory T cells was statistically higher for Luc90-CD8BBZ-expressing T cells than Luc90-CD828Z-expressing T cells (Figure S1).

Proliferation of Anti-SLAMF7 CARs Containing CD28 versus 4-1BB

In proliferation assays, there was not a statistically significant difference in numbers of accumulated CD4⁺CAR⁺ T cells when Luc90-CD828Z T cells and Luc90-CD8BBZ T cells were cultured for 4 days with SLAMF7⁺ target cells (Figure 3G). However, more CD8⁺ Luc90-CD828Z T cells than CD8⁺ Luc90-CD8BBZ T cells accumulated over a 4-day culture with SLAMF7⁺ target cells (Figure 3H). There was not a statistically significant difference in SLAMF7-specific proliferation for T cells expressing Luc90-CD828Z versus Luc90-CD8BBZ for either CD4⁺ (Figure 3I) or CD8⁺ (Figure 3J) T cells. We assessed activation-induced cell death (AICD) in T cells expressing either Luc90-CD828Z or Luc90-CD8BBZ. There was not a statistically significant difference in AICD for either CD4⁺ or CD8⁺ T cells; however, there was a trend toward less AICD among CD8⁺ T cells expressing Luc90-CD828Z versus Luc90-CD8BBZ (Figure S2).

Cytokine Release and Degranulation by T Cells Expressing CARs with CD28 versus 4-1BB

Compared with T cells expressing Luc90-CD8BBZ, T cells expressing Luc90-CD828Z released higher levels of IFN- γ (Figure 3K), IL-2 (Figure 3L), and tumor necrosis factor (TNF)- α (Figure 3M) when

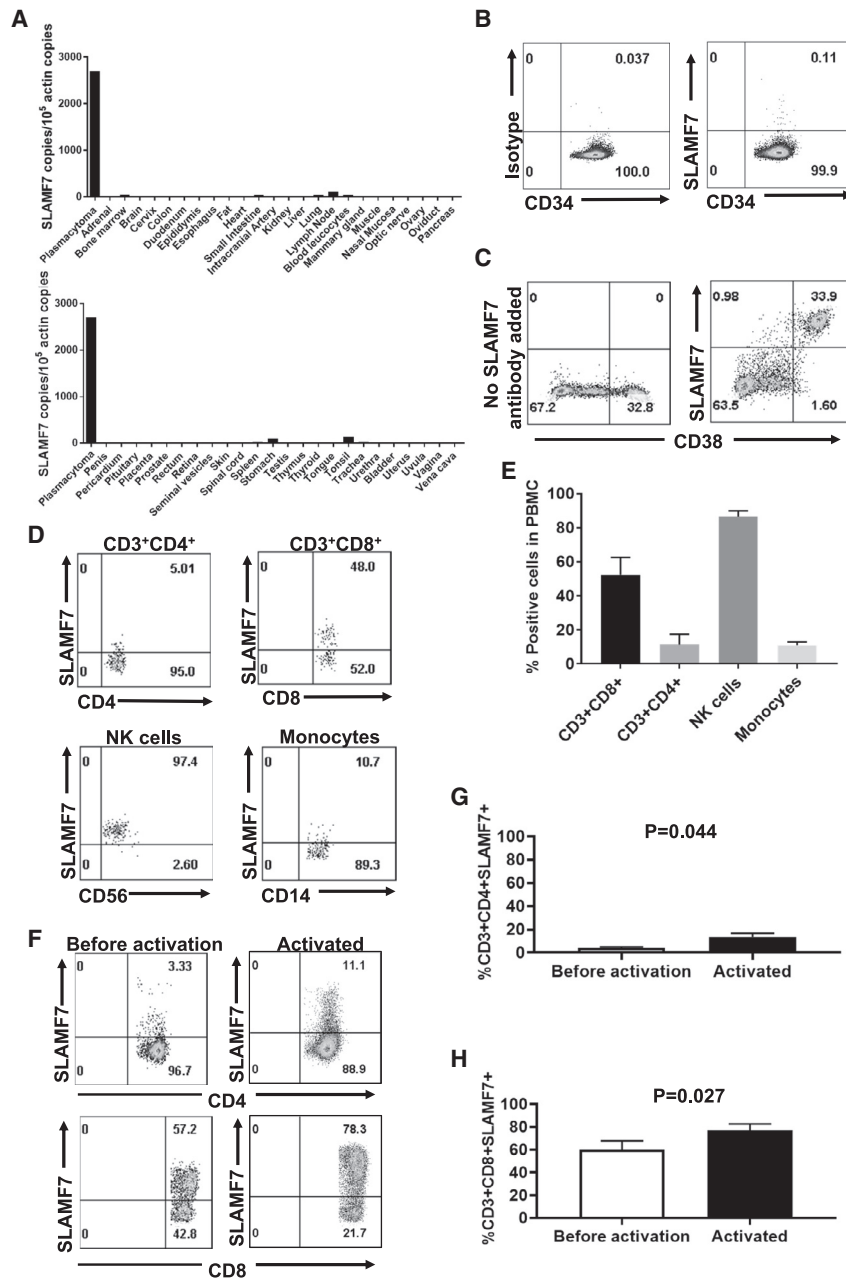


Figure 1. SLAMF7 Expression

(A) SLAMF7 RNA expression was assessed in all major normal tissues and a plasmacytoma made up of SLAMF7⁺ multiple myeloma (MM) cells. Quantitative PCR was performed on cDNA from each of the tissues. SLAMF7 cDNA copies were normalized to actin cDNA copies from each tissue. (B) Primary human CD34⁺ hematopoietic stem cells were obtained from the blood of a patient after the patient received granulocyte colony-stimulating factor to mobilize stem cells from the bone marrow. Staining with anti-CD34 and either anti-SLAMF7 or an isotype-matched control antibody was performed, followed by flow cytometry. Plots are gated on live CD34⁺ cells. One representative experiment of 2 experiments with different donors is shown. (C) Primary bone marrow cells from a patient with MM were stained with antibodies against CD38 alone or with antibodies against both CD38 and SLAMF7. Plots are gated on live CD3-negative cells. (D) Peripheral blood mononuclear cells (PBMCs) were stained for SLAMF7 along with antibodies against CD3, CD4, CD8, CD14, and CD56. The plot showing CD3⁺CD4⁺ cells is gated on CD3⁺CD4⁺ cells. The plot showing CD3⁺CD8⁺ cells is gated on CD3⁺CD8⁺ cells. The NK cells' plot is gated on CD3-negative, CD56⁺ cells. The monocyte plot is gated on CD3-negative, CD14⁺ cells. For T cells and NK cells, plots were gated on lymphocyte forward-scatter and side-scatter regions. For monocytes, a more inclusive gate that included lymphocytes as well as higher forward-scatter and side-scatter events was used. All plots had dead cells excluded. Regions were set with an isotype-control antibody for SLAMF7. (E) Blood CD3⁺CD8⁺ T cells, CD3⁺CD4⁺ T cells, NK cells, and monocytes were stained and analyzed as in (D). Percentages of each cell type that expressed SLAMF7 are shown. For monocytes, $n = 4$ different donors. For other cell types, $n = 8$ different donors. Bars represent mean plus standard error of the mean (SEM). (F) Human PBMCs were cultured overnight in culture medium alone without IL-2. The PBMCs from each donor were then divided into two portions. The first portion was stained with antibodies against CD3, CD4, CD8, and SLAMF7 and analyzed by flow cytometry. Representative example plots from one of three donors are shown. The plots shown are all gated on live CD3⁺ lymphocytes and then on either CD4⁺ or CD8⁺ cells. The second portion of PBMCs was placed in culture with IL-2 and activated with an anti-CD3 antibody for 1 day. Activated cells were stained, and flow cytometry was performed in the same manner as before activation. (G and H) PBMCs from 3 donors were cultured and analyzed before and after activation as in (F). The mean + SEM of %SLAMF7⁺ cells for (G) CD4⁺ and (H) CD8⁺ T cells is shown. Statistics were by two-tailed, paired t test; $p < 0.05$ was considered statistically significant.

stimulated with MM.1S target cells *in vitro*. Compared with Luc90-CD828Z T cells, Luc90-CD8BBZ T cells exhibited higher levels of nonspecific IFN- γ release against SLAMF7-negative target cells *in vitro* (Figure S3). We did not find a statistically significant difference in SLAMF7-specific degranulation by CD107a expression when Luc90-CD828Z T cells and Luc90-CD8BBZ T cells were compared (Figure S4).

Anti-tumor Activity of CD28 CARs Was Superior to Anti-tumor Activity of 4-1BB CARs

We evaluated anti-tumor efficacy of Luc90-CD8BBZ and Luc90-CD828Z CAR T cells in NOD.Cg-Prkdc^{scid} Il2rg^{tm1Wjl}/SzJ (NSG) mice bearing solid tumors of the MM.1S human MM cell line. Mice were left untreated or received a single intravenous infusion of 2×10^6

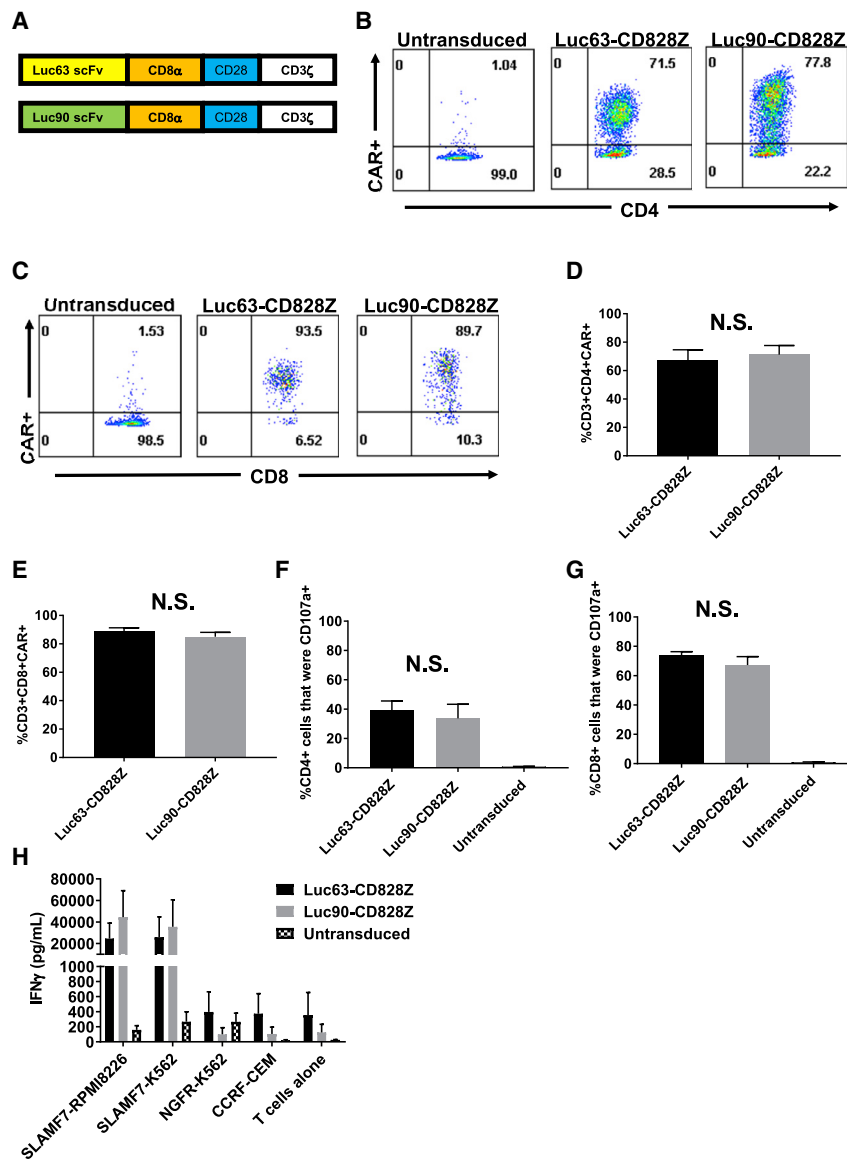


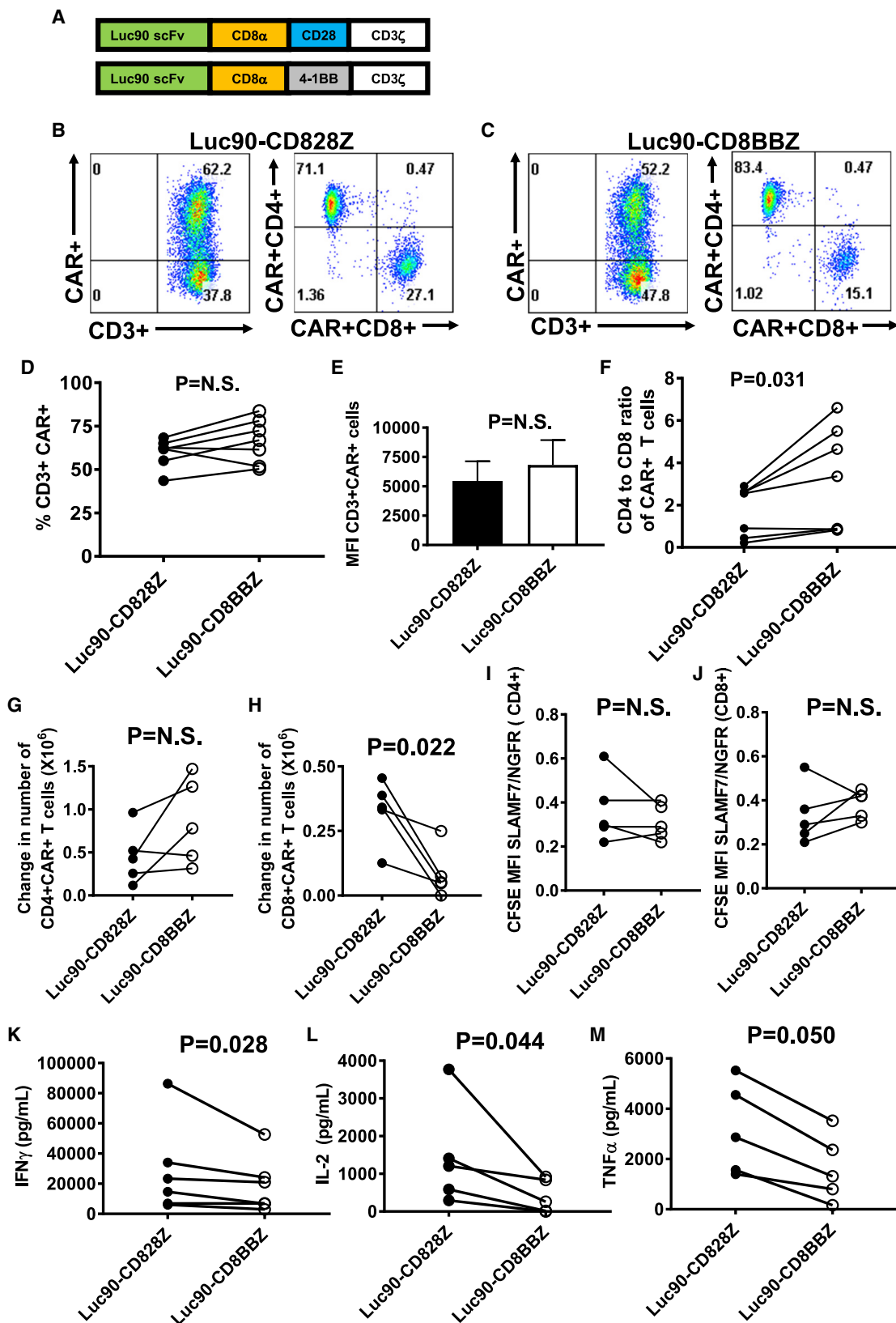
Figure 2. Comparison of Different Anti-SLAMF7 CAR Designs

(A) Luc63-CD828Z contained the Luc63 scFv, and Luc90-CD828Z contained the Luc90 scFv. Both CARs contained CD8 α hinge and transmembrane domains, CD28 costimulatory domains, and CD3 ζ activation domains. (B) An example of CAR expression on CD4⁺ T cells is shown. Whole PBMCs were stimulated on day 0 of culture with an anti-CD3 antibody and cultured in IL-2-containing medium. On day 2 of culture, transductions were carried out with MSGV1 gamma-retroviral vectors encoding either Luc63-CD828Z or Luc90-CD828Z. 5 days after transduction, flow cytometry was performed with protein L to detect CARs, and staining was also performed against CD3, CD4, and CD8. Plots are gated on live CD3⁺CD4⁺ lymphocytes. (C) CAR expression on CD8⁺ T cells is shown. Experiment was performed as in (B). Plots are gated on live CD3⁺CD8⁺ lymphocytes. Mean \pm SEM percentages of (D) CD4⁺ T cells or (E) CD8⁺ T cells expressing Luc63-CD828Z or Luc90-CD828Z is shown; n = 5 different donors for all groups. T cells were cultured, transduced, and analyzed as in (B). For both CD4⁺ and CD8⁺ T cells, there was no significant difference (N.S.) in the percentage of T cells expressing Luc63-CD828Z versus Luc90-CD828Z. (F) CD4⁺ T cells and (G) CD8⁺ T cells were assessed for SLAMF7-specific CD107a upregulation. T cells were transduced with either Luc63-CD828Z or Luc90-CD828Z or left untransduced. T cells were cultured and transduced as in (B). Assays were conducted 5 days after transduction. T cells were cultured with either SLAMF7⁺ SLAMF7-RPMI8226 cells or SLAMF7-negative NGFR-K562 cells for 4 h in the presence of anti-CD107a antibodies. After the incubation, T cells were stained for CD3, CD4, and CD8. Data are presented after normalization for CAR expression, and %SLAMF7-specific CD107a⁺ events were calculated as %CD107a⁺ events with SLAMF7-RPMI8226 stimulation minus %CD107a⁺ events with NGFR-K562 stimulation. Bars represent mean \pm SEM. For both CD4⁺ and CD8⁺ T cells, there was no statistically significant difference in CD107a upregulation when Luc63-CD828Z and Luc90-CD828Z were compared; n = 5. (D–G) Statistical testing was by paired, two-tailed t tests, and statistical significance was defined as p < 0.05. (H) T cells were cultured

and transduced with Luc63-CD828Z or Luc90-CD828Z or left untransduced. 5 days after transduction, T cells were cultured overnight with target cells. SLAMF7-RPMI8226 and SLAMF7-K562 were SLAMF7⁺. NGFR-K562 and CCRF-CEM were SLAMF7 negative. After the overnight culture, supernatant was assayed by ELISA for IFN- γ . Bars show mean \pm SEM of 3 experiments with cells from 3 different donors.

T cells expressing Luc90-CD828Z, Luc90-CD8BBZ, or Hu19-CD828Z. Only T cells expressing Luc90-CD828Z eradicated all MM.1S solid tumors. The anti-tumor activity of T cells expressing Luc90-CD8BBZ was less than that of T cells expressing Luc90-CD828Z (Figures 4A and 4B). When we doubled the dose of anti-SLAMF7 CAR T cells to 4×10^6 CAR T cells per mouse, both Luc90-CD828Z and Luc90-CD8BBZ CAR T cells consistently eliminated MM.1S solid tumors (Figure S5). At the 4×10^6 CAR T cell dose, tumor elimination was superior at early time points by Luc90-CD828Z CAR T cells compared with Luc90-CD8BBZ CAR T cells.

Similar to solid tumors, clearance of disseminated MM.1S cells was more complete with Luc90-CD828Z T cells versus Luc90-CD8BBZ T cells (Figure 4C). On day 12 after CAR T cell infusion, when all mice in both groups were still alive, the tumor burdens of mice that received Luc90-CD828Z CAR T cells were statistically lower than the tumor burdens of mice that received Luc90-CD8BBZ CAR T cells (Figure 4D). Survival of mice with disseminated MM.1S burdens that received Luc90-CD828Z CAR T cells was significantly higher than survival of mice that received Luc90-CD8BBZ CAR T cells (Figure 4E).



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We assessed CAR T cells infiltrating MM.1S tumors 7 days after infusion of CAR T cells intravenously. To avoid the issues of CAR down-regulation on CAR T cells, which can be a problem with flow cytometry detection of tumor-infiltrating CAR T cells, we performed quantitative PCR (qPCR) to detect CAR⁺ cells. We found a statistically higher number of tumor-infiltrating CAR⁺ cells in mice receiving Luc90-CD828Z CAR T cells versus mice receiving Luc90-CD8BBZ CAR T cells. (Figure 4F). We recognize the limitations of only assessing tumor-infiltrating CAR⁺ cells at one time point, but rapid elimination of tumors in mice treated with Luc90-CD828Z T cells precluded comparison at later time points.

Anti-tumor Activity of Anti-CD19 CARs with CD28 or 4-1BB Domains

To assess whether superior anti-tumor efficacy of T cells expressing CARs with CD28 versus 4-1BB occurred with CARs targeting a different antigen, we utilized anti-CD19 CARs. We compared the CD28-containing anti-CD19 CAR Hu19-CD828Z to the 4-1BB-containing CAR Hu19-CD8BBZ. Hu19-CD8BBZ was identical to Hu19-CD828Z except for the difference in costimulatory domains. We used a model in which solid tumors of CD19⁺ NALM6 cells were established in NSG mice. In agreement with anti-SLAMF7 CAR results, we found that tumor treatment with Hu19-CD828Z CAR T cells was superior to treatment with Hu19-CD8BBZ CAR T cells (Figure S6). Taken together, the anti-SLAMF7 and anti-CD19 data showed that CAR T cells with CD28 domains were superior to CAR T cells with 4-1BB domains at eliminating solid tumors in the NSG mouse models used. Hu19-CD828Z T cells and Hu19-CD8BBZ T cells were specifically activated by CD19⁺ target cells (Table S3). As with anti-SLAMF7 CARs (Figure S3), nonspecific IFN- γ release was higher with 4-1BB-containing anti-CD19 CARs than with CD28-containing anti-CD19 CARs (Table S3). The CD28 and 4-1BB sequences in the CARs used in this work are shown in Figure S7.

T Cells Expressing an Anti-SLAMF7 CAR and a Suicide Gene Can Be Eliminated *In Vitro*

We designed two constructs encoding the Luc90-CD828Z CAR and the IC9 suicide gene. Diagrams of these constructs, which were designated Luc90-CD828Z-IC9 and IC9-Luc90-CD828Z are shown in Figure 5A. The Luc90-CD828Z-IC9 and the IC9-Luc90-CD828Z constructs differ only in the order of the CAR and suicide genes. The CAR and the IC9 genes were separated by a *Thoseaasigna* virus (T)2A ribosomal skip sequence. The CARs were encoded by MSGV1. Luc90-CD828Z was expressed on T cells that were transduced with either MSGV1-Luc90-CD828Z-IC9 or MSGV1-IC9-Luc90-CD828Z (Figure 5B). Next, we evaluated the efficiency of CAR T cell elimination by adding AP1903 (rimiducid) to the cultures of T cells expressing Luc90-CD828Z-IC9 or IC9-Luc90-CD828Z (Figures 5C and 5D). Concentrations of AP1903 used in the assay were equal to concentrations of AP1903 achieved in clinical trials.^{26,27} We found that T cells transduced with either Luc90-CD828Z-IC9 or IC9-Luc90-CD828Z could be rapidly eliminated when AP1903 was added to the T cell cultures. Elimination of CAR⁺ T cells after AP1903 treatment was statistically superior with T cells expressing IC9-Luc90-CD828Z compared with T cells expressing Luc90-CD828Z-IC9 (Figure 5D). Almost all residual IC9-Luc90-CD828Z CAR⁺ T cells were apoptotic 6 h after AP1903 treatment by Annexin V staining (Figure 5E).

Elimination of Anti-SLAMF7 CAR T Cells by AP1903 Prevents NK Cell Depletion by Anti-SLAMF7 CAR T Cells

To demonstrate elimination of NK cells by anti-SLAMF7 CAR T cells, we cocultured T cells expressing either the anti-CD19 CAR Hu19-CD828Z or IC9-Luc90-CD828Z or untransduced control T cells with autologous PBMC. In one coculture containing IC9-Luc90-CD828Z and PBMC, AP1903 was added to eliminate the CAR T cells. We then assessed NK cells by flow cytometry. NK cells were defined as CD56⁺CD3⁻ cells (Figure 5F). NK cells were depleted from cultures containing PBMCs plus IC9-Luc90-CD828Z without AP1903 (Figure 5G). Addition of AP1903 to cultures containing IC9-Luc90-

Figure 3. Comparison of Anti-SLAMF7 CARs with Different Costimulatory Domains

(A) Luc90-CD8BBZ was identical to Luc90-CD828Z except for substitution of 4-1BB for CD28. (B and C) T cells were stimulated with anti-CD3 on day 0 and transduced on day 2 of culture. Flow cytometry with protein L staining was performed on day 7 of culture to detect CARs. (B) CAR expression on MSGV1-Luc90-CD828Z-transduced T cells. (C) CAR expression on MSGV1-Luc90-CD8BBZ-transduced T cells. (B and C) The left plot is gated on live CD3⁺ lymphocytes. The right plot is gated on the CD3⁺CAR⁺ population from the left plot. (D) The percentages of live CD3⁺ T cells expressing cell-surface Luc90-CD828Z or Luc90-CD8BBZ from cultures of transduced T cells are shown; lines connect values for each CAR from cultures of the same patient. Differences in percentages of T cells expressing Luc90-CD828Z versus Luc90-CD8BBZ were not statistically significant. (E) There was not a statistically significant difference in median fluorescence intensities (MFIs) of CD3⁺CAR⁺ cells transduced with either Luc90-CD828Z or Luc90-CD8BBZ. (F) CD4-to-CD8 ratios of T cells expressing either Luc90-CD828Z or Luc90-CD8BBZ are shown. Lines connect results from the same patient. (D–F) T cells were cultured and plots gated as in (B) and (C). (D–F) n = 7 different donors; statistics were by paired, two-tailed t test. (G and H) T cells were transduced with either Luc90-CD828Z or Luc90-CD8BBZ. T cells were labeled with CFSE and cultured with either SLAMF7-K562 cells or NGFR-K562 cells for 4 days. The changes in numbers of (G) CD4⁺CAR⁺ T cells or (H) CD8⁺CAR⁺ T cells between day 0 and day 4 of culture are shown. The change in cell numbers for Luc90-CD828Z and Luc90-CD8BBZ T cells from the same patient is connected by a line. Statistical testing was by paired, two-tailed t tests, and statistical significance was defined as p < 0.05. (I and J) At the end of the 4-day culture period, the same T cell cultures in (G) and (H) were assessed for the ratio of CFSE MFI with SLAMF7-K562 stimulation versus NGFR-K562 stimulation (SLAMF7/NGFR ratio). A decrease in CFSE MFI indicates proliferation. Lower SLAMF7/NGFR ratios indicate more SLAMF7-specific proliferation. Cultures from the same patient are connected by a line. Differences were not statistically different. (G–J) n = 5 different donors. (K–M) SLAMF7-specific cytokine production was compared between paired cultures of T cells from the same donor expressing Luc90-CD828Z or Luc90-CD8BBZ; the paired cultures are connected by lines. T cells were cultured overnight with either MM.1S cells or NGFR-K562 cells, and ELISAs were performed on supernatants. Results were normalized for CAR expression. MM.1S-specific cytokine release was calculated by subtracting cytokine levels with NGFR-K562 stimulation from cytokine levels with MM.1S stimulation. (K) IFN- γ , (L) IL-2, and (M) TNF- α . n = 6 different donors for IFN- γ ; n = 5 different donors for IL-2 and TNF- α . Statistical testing was by two-tailed ratio, paired t tests; statistical significance was p < 0.05. There was a statistically significant difference for IFN- γ and IL-2. The difference for TNF- α was of borderline statistical significance.

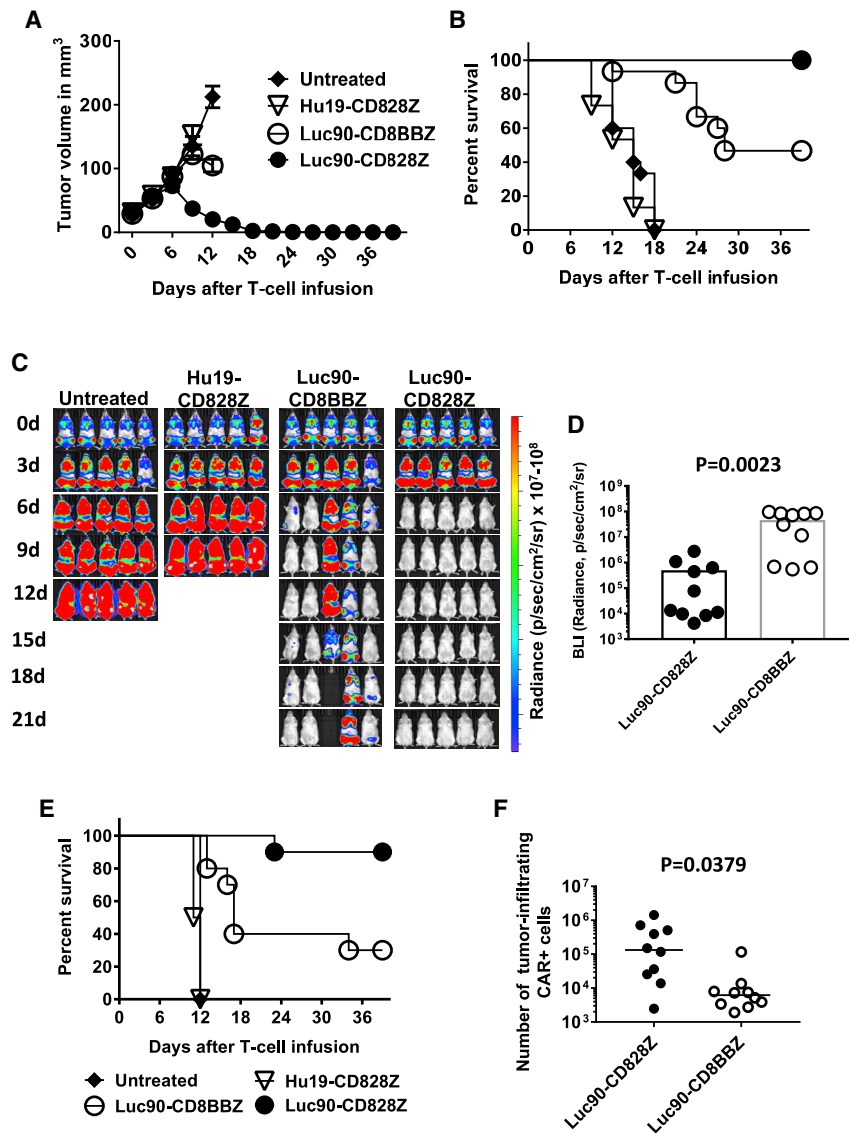


Figure 4. Luc90-CD828Z CAR T Cells Are Superior to Luc90-CD8BBZ in Eliminating Tumors

(A) Solid tumors were established in NSG mice by intradermal injection of 4×10^6 MM.1S cells. 7 days after tumor implantation, mice were either left untreated or infused intravenously with 2×10^6 T cells transduced with Luc90-CD828Z, Luc90-CD8BBZ, or the negative-control CAR Hu19-CD828Z. Graphs show mean tumor volume \pm SEM for each time point. The graph shows combined data from three independent experiments with different donor T cells. There were five mice in each group per experiment for a total of 15 mice for each group shown on the graph. Lines end when the first mouse of the group, represented by the line, was sacrificed or when the experiment ended at day 39. (B) Kaplan-Meier survival graph of the same mice as in (A). By the log rank test, there were statistically significant differences in survival between mice receiving Hu19-CD828Z T cells and Luc90-CD828Z T cells ($p < 0.0001$) and mice receiving Hu19-CD828Z T cells and Luc90-CD8BBZ T cells ($p < 0.0001$). There was also a statistically significant difference in survival between mice receiving Luc90-CD828Z T cells and Luc90-CD8BBZ T cells ($p = 0.0011$). (C) Disseminated tumor burdens were established in NSG mice by injecting 8×10^6 MM.1S-luciferase cells intravenously. 13 days after tumor cell injection, mice received 4×10^6 T cells transduced with either Luc90-CD828Z or Luc90-CD8BBZ or the negative-control CAR Hu19-CD828Z. Another group of mice was left untreated. Tumor burden was evaluated by bioluminescent imaging (BLI) every 3 days. The figure is representative of one experiment out of two independent experiments with T cells from different donors. (D) Average BLI intensity was compared for mice with disseminated MM.1S-luciferase burdens receiving 4×10^6 T cells transduced with either Luc90-CD828Z or Luc90-CD8BBZ. The graph includes BLI from 10 mice from each treatment group, 12 days after CAR T cell infusion. There was a statistically significant difference in BLI between Luc90-CD828Z and Luc90-CD8BBZ ($p = 0.0023$; unpaired, two-tailed t test). The graph shows one point for each mouse. Bars represent mean \pm SEM. Five mice in each group are from (C); an additional 5 mice from a second experiment are also included in the graph. (E) Kaplan-Meier survival plot of mice in the experiments reported in (D). The

graph includes combined data from two independent experiments with different donor T cells. By a log rank test, there was a statistically significant difference in survival between mice receiving T cells expressing Hu19-CD828Z and mice receiving T cells expressing either Luc90-CD828Z or Luc90-CD8BBZ ($p < 0.0001$ for both comparisons). There was also a statistically significant difference in survival between mice receiving T cells that expressed Luc90-CD828Z and Luc90-CD8BBZ ($p = 0.0056$). There were 10 mice in each group. (F) Solid MM.1S tumors were established in NSG mice. Mice were infused with 4×10^6 CAR⁺ T cells. 7 days after CAR T cell infusion, tumors were excised and processed into single-cell suspensions. Quantitative PCR was performed with a PCR reaction capable of detecting both Luc90-CD828Z and Luc90-CD8BBZ to quantify the number of tumor-infiltrating T cells in the tumors of each mouse. Mice that received Luc90-CD828Z T cells had more tumor-infiltrating T cells ($p = 0.0379$; unpaired, two-tailed t test; $n = 10$ mice per group). The horizontal bars represent the medians.

CD828Z CAR T cells prevented elimination of NK cells from cultures presumably by killing most CAR T cells (Figure 5G).

Function of T Cells Expressing IC9-Luc90-CD828Z or Luc90-CD828Z-IC9

For both CD4⁺ and CD8⁺ T cells, cell-surface CAR expression of IC9-Luc90-CD828Z and Luc90-CD828Z-IC9 was not different on day 7 of culture; however, cell-surface CAR expression of IC9-Luc90-CD828Z-transduced T cells was lower than cell-surface expression of Luc90-

CD828Z-IC9-transduced T cells on day 14 of culture (Figures 6A and 6B). CAR expression on the T cell surface was lower for T cells transduced with either MSGV1-IC9-Luc90-CD828Z or MSGV1-Luc90-CD828Z-IC9 when compared with T cells transduced with MSGV1-Luc90-CD828Z that encodes the CAR without a suicide gene (Figure S8). Both CD4⁺ (Figure 6C) and CD8⁺ (Figure 6D) IC9-Luc90-CD828Z and Luc90-CD828Z-IC9 CAR T cells degranulated in a SLAMF7-specific manner; levels of degranulation were higher for CD8⁺ T cells than CD4⁺ T cells. T cells expressing either IC9-Luc90-CD828Z or

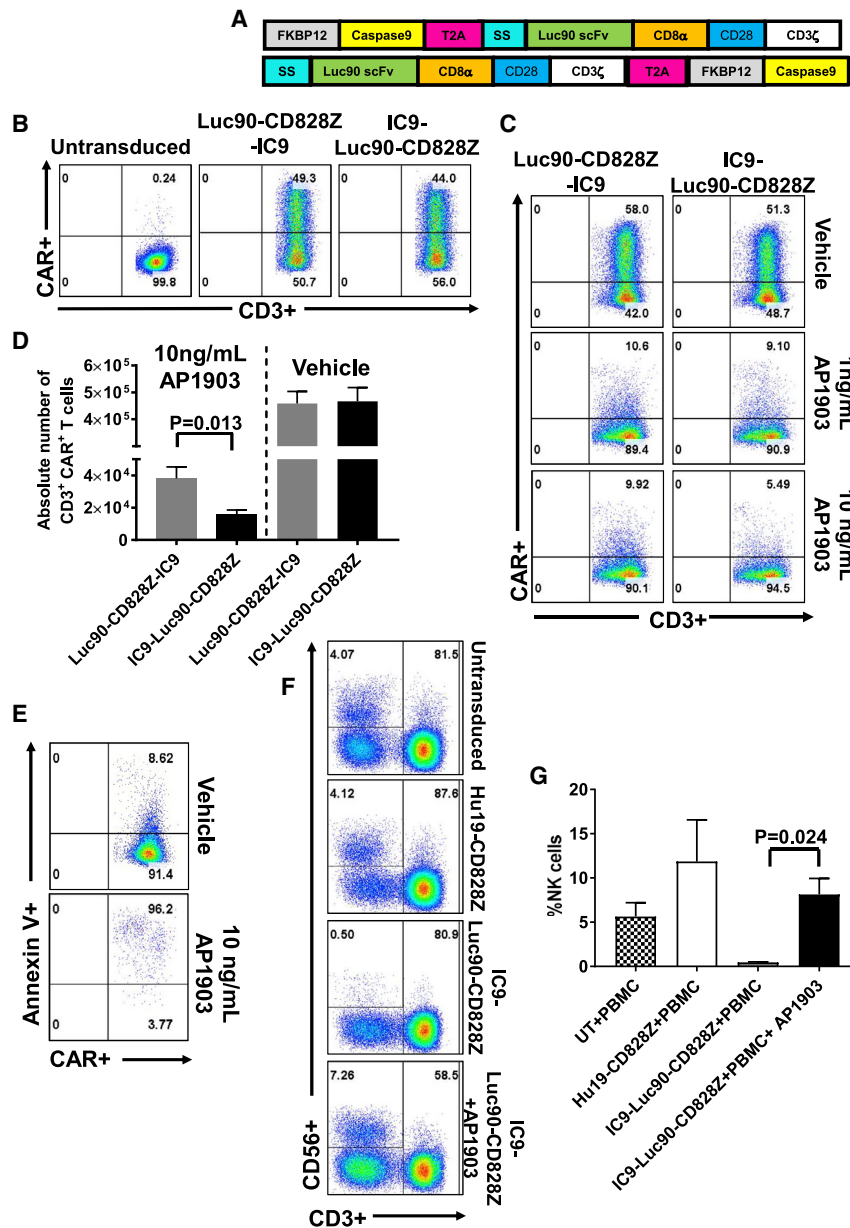


Figure 5. Constructs Containing an Anti-SLAMF7 CAR and the IC9 Suicide Switch

(A) Schematics of the Luc90-CD828Z-IC9 and IC9-Luc90-CD828Z constructs that both encode the Luc90-CD828Z CAR and the IC9 suicide switch. IC9 was made up of a modified FKBP12 domain followed by a modified caspase-9 sequence. In each construct, CAR sequences and IC9 sequences were separated by T2A sequences. Luc90-CD828Z-IC9 and IC9-Luc90-CD828Z only differ in the order of the CAR and IC9. Both CARs were encoded by the MSGV1 gamma-retroviral vector. SS, signal sequence. (B) Representative examples of CAR expression on CD3⁺ T cells transduced with MSGV1-Luc90-CD828Z-IC9 or MSGV1-IC9-Luc90-CD828Z are shown. Plots are gated on live CD3⁺ lymphocytes. (C) 5 days after transduction, T cells transduced with either MSGV1-Luc90-CD828Z-IC9 or MSGV1-IC9-Luc90-CD828Z were exposed to the indicated concentrations of AP1903 or vehicle (DMSO) for 6 h. The plots shown are gated on CAR⁺, CD3⁺ live lymphocytes and are representative of 5 independent experiments with lymphocytes from different donors. (D) Absolute numbers of CAR⁺CD3⁺ live lymphocytes were quantified after treatment with 10 ng/mL of AP1903 or vehicle (DMSO) for 6 h. AP1903 treatment eliminated significantly more T cells expressing IC9-Luc90-CD828Z than T cells expressing Luc90-CD828Z-IC9 ($p = 0.013$; two-tailed, paired t test). Each bar represents mean + SEM; $n = 5$ different donors for all groups. (E) The plots show that most residual IC9-Luc90-CD828Z-expressing T cells from the culture treated with 10 ng/mL of AP1903 shown in (C) were Annexin V⁺, which indicates that the cells were apoptotic. Plots are gated on CAR⁺CD3⁺ lymphocytes. This is one of 5 experiments with similar results. (F) 6 days after transduction, T cells transduced with the indicated constructs were cultured with autologous PBMC at a 1:1 ratio for 24 h. One of the cultures containing IC9-Luc90-CD828Z CAR T cells was treated with 10 ng/mL AP1903. Plots show percent live CD3⁻negative, CD56⁺ NK cells in the upper left. (G) The graph shows percentages of NK cells from assays conducted as in (F). Results show eradication of NK cells in the presence of IC9-Luc90-CD828Z CAR T cells and prevention of NK cell eradication by eliminating IC9-Luc90-CD828Z CAR T cells with AP1903. $N = 4$ different donors for all groups. Bars represent mean + SEM; statistical testing was by paired, two-tailed t tests; statistical significance was $p < 0.05$.

Luc90-CD828Z-IC9 proliferated and accumulated during a 4-day culture with SLAMF7⁺ target cells (Figure 6E). There was no statistically significant difference in the degree of SLAMF7-specific proliferation between IC9-Luc90-CD828Z and Luc90-CD828Z-IC9 CAR T cells in (Figure 6F).

Cytokine Release and Cytotoxicity of T Cells Expressing IC9-Luc90-CD828Z

IC9-Luc90-CD828Z and Luc90-CD828Z-IC9 CAR T cells released large amounts of IFN- γ in the presence of SLAMF7⁺ target cells and very low levels of IFN- γ in the presence of SLAMF7-negative target cells (Figure 6G). T cells expressing

IC9-Luc90-CD828Z released TNF- α (Figure 6H) and IL-2 (Figure 6I) in a SLAMF7-specific manner. T cells expressing IC9-Luc90-CD828Z specifically killed MM.1S cells (Figure 6J). In addition, IC9-Luc90-CD828Z cells did not recognize a variety of SLAMF7-negative target cells when assessed by IFN- γ ELISA (Table S4).

T cells expressing Luc90-CD828Z-IC9 and IC9-Luc90-CD828Z exhibited similar *in vitro* function. However, IC9-Luc90-CD828Z T cells can be eliminated slightly more effectively than Luc90-CD828Z-IC9 T cells after AP1903 treatment (Figure 5D), so IC9-Luc90-CD828Z was selected for further development

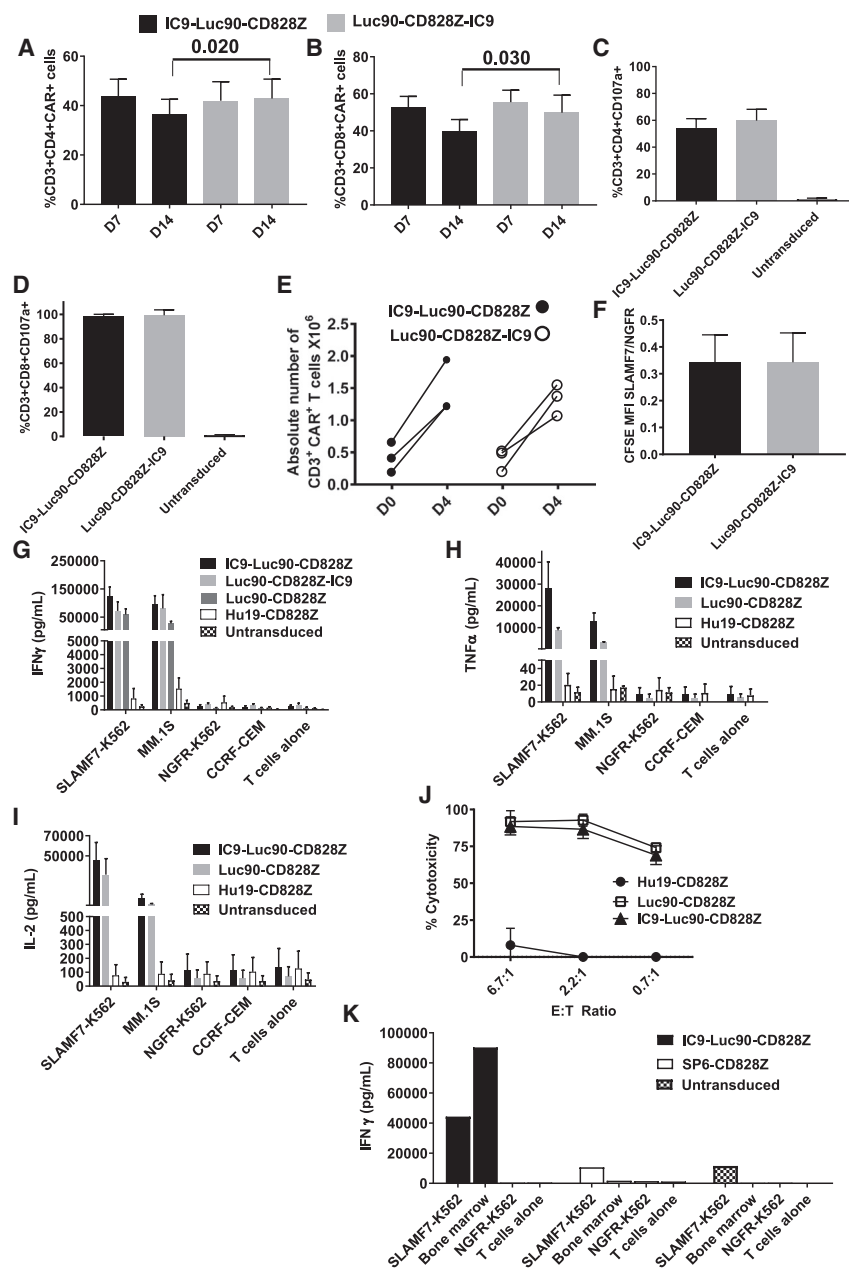


Figure 6. T Cells Expressing IC9-Luc90-CD828Z Exhibit SLAMF7-Specific Reactivity *In Vitro*

T cells were transduced with MSGV1-IC9-Luc90-CD828Z or MSGV1-Luc90-CD828Z-IC9. On day 7 (D7) and day 14 (D14) of culture, flow cytometry was performed to detect (A) CAR⁺CD3⁺CD4⁺ lymphocytes or (B) CAR⁺CD3⁺CD8⁺ lymphocytes. Mean \pm SEM percentages of T cells expressing IC9-Luc90-CD828Z or Luc90-CD828Z-IC9 are shown; $n = 8$ for day 7; $n = 6$ for day 14. For CD4⁺ and CD8⁺ T cells, there was no significant difference in the percentages of T cells expressing IC9-Luc90-CD828Z versus Luc90-CD828Z-IC9 at day 7, but there was a higher level of expression for Luc90-CD828Z-IC9 versus IC9-Luc90-CD828Z at day 14. (C and D) T cells expressing IC9-Luc90-CD828Z or Luc90-CD828Z-IC9 or left untransduced were cultured with either SLAMF7⁺ MM.1S cells or SLAMF7-negative NGFR-K562 along with antibodies against CD107a for 4 h. Mean \pm SEM of %MM.1S-specific CD107a⁺ events are shown. %MM.1S-specific CD107a⁺ events were calculated as %CD107a⁺ events with MM.1S stimulation minus %CD107a⁺ events with NGFR-K562 stimulation. Data are normalized for CAR expression. For both (C) CD4⁺ and (D) CD8⁺ T cells, there was no statistical difference in CD107a upregulation between IC9-Luc90-CD828Z and Luc90-CD828Z-IC9; $n = 4$ different donors for all groups. (A–D) Statistical testing was by paired, two-tailed t tests; statistical significance was $p < 0.05$. (E) T cells were labeled with CFSE and cultured with either SLAMF7-K562 cells or NGFR-K562 cells for 4 days. Changes in numbers of CAR⁺ T cells during the 4-day culture are shown for T cells expressing each CAR from 3 different donors. CAR⁺ T cells increased from day 0 to day 4 ($p = 0.017$ for IC9-Luc90-CD828Z; $p = 0.035$ for Luc90-CD828Z-IC9 by two-tailed, paired t test). (F) Proliferation was assessed by CFSE dilution. The degree of SLAMF7-specific proliferation is presented as the ratio of MFI of CFSE-labeled T cells cultured with SLAMF7-K562 cells (SLAMF7) divided by the MFI of CFSE-labeled T cells cultured with NGFR-K562 cells (NGFR). A SLAMF7/NGFR ratio < 1 indicates SLAMF7-specific proliferation. There was no statistical difference in CFSE dilution between IC9-Luc90-CD828Z and Luc90-CD828Z-IC9. (G) T cells expressing Hu19-CD828Z, Luc90-CD828Z, IC9-Luc90-CD828Z, or Luc90-CD828Z-IC9 or untransduced control T cells were cultured overnight with target cells. SLAMF7-K562 and MM.1S were SLAMF7⁺. NGFR-K562 and CCRF-CEM were SLAMF7-negative. Supernatant was assayed by ELISA for IFN- γ . Bars show mean \pm SEM. The number of experiments with cells from different donors was $n = 7$ for untransduced, IC9-Luc90-CD828Z, and Luc90-CD828Z; $n = 4$ for Luc90-CD828Z-IC9; and $n = 3$ for Hu19-CD828Z. (H and I) Bars show mean \pm SEM of 3 experiments with cells from 3 donors. After overnight culture, ELISAs were performed on culture supernatants for (H) TNF- α or (I) IL-2; data were normalized for CAR expression. (J) T cells were transduced with the indicated CARs, and a 4-h cytotoxicity assay was performed with SLAMF7⁺ MM.1S target cells. Mean \pm SEM of duplicate wells is graphed. The experiment was conducted 2 times with cells from different donors with similar results. (K) T cells were transduced with IC9-Luc90-CD828Z, SP6-CD828Z, or left untransduced. The T cells were cocultured with primary bone marrow MM cells, SLAMF7-K562, or NGFR-K562 overnight, and an IFN- γ ELISA was performed. Bars represent average IFN- γ in duplicate wells. Two experiments that used cells from 2 different donors were conducted with similar results.

IC9-Luc90-CD828Z T Cells Specifically Recognize Primary MM Cells

T cells expressing IC9-Luc90-CD828Z released IFN- γ specifically in response to primary bone marrow cells from a patient with extensive bone marrow MM that was SLAMF7⁺ (Figure 6K). Similarly, IC9-Luc90-CD828Z T cells degranulated specifically in response to SLAMF7⁺ primary MM bone marrow cells (Figure S9).

Anti-tumor Activity of IC9-Luc90-CD828Z

T cells expressing IC9-Luc90-CD828Z have dose-dependent activity against established tumors of the MM.1S MM cell line. A single intravenous infusion of 2×10^6 or more of IC9-Luc90-CD828Z-expressing T cells/mouse eradicated MM.1S solid tumors (Figures 7A and 7B). A single intravenous dose of 2×10^6 CAR T cells/mouse of

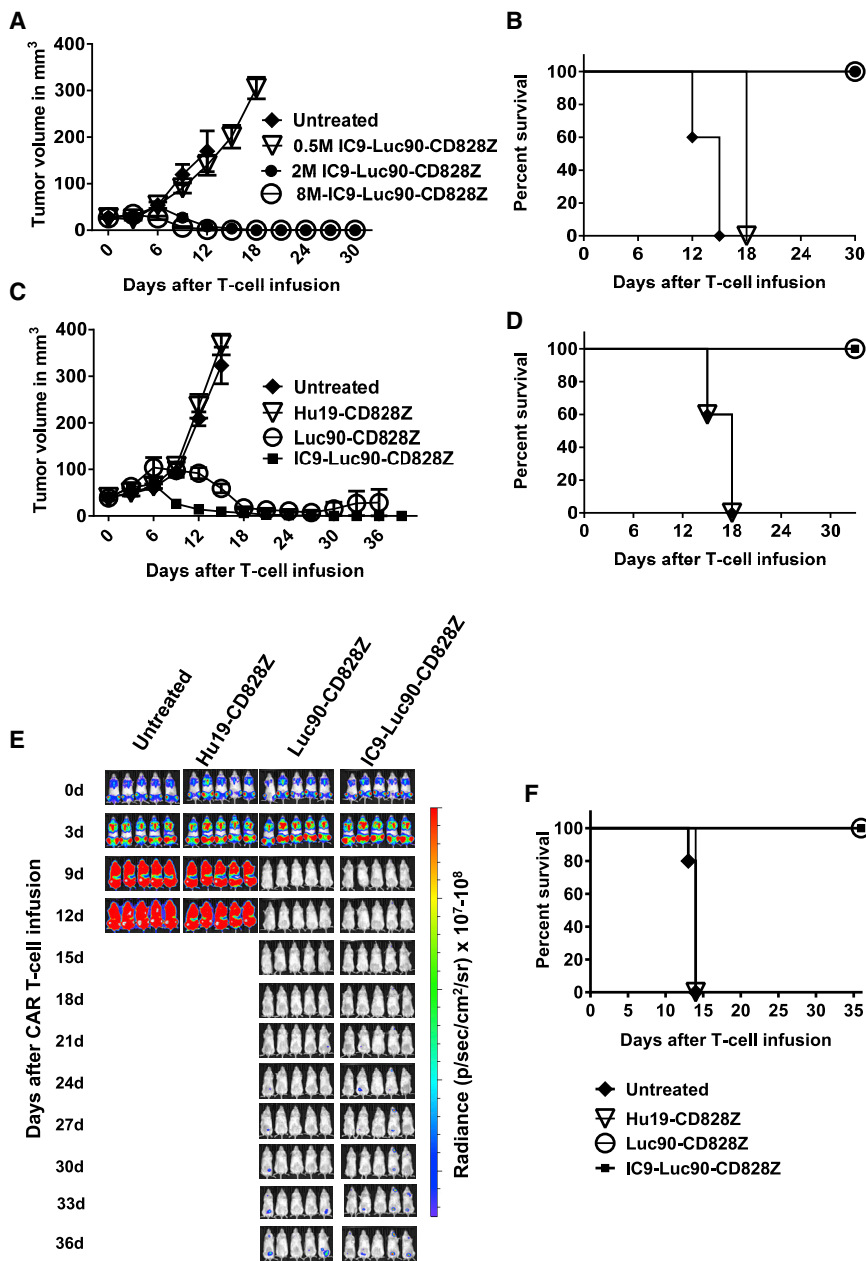


Figure 7. Anti-SLAMF7 CAR T Cells with IC9 Eradicated Solid and Disseminated Tumor Burdens

(A) 7 days after injection of MM.1S cells into NSG mice when palpable tumors were present, the indicated doses of CAR⁺ T cells were infused intravenously into the mice, and tumors were measured every 3 days in a blinded manner. The graph shows mean tumor volume \pm SEM for each time point. There were five mice in each group. (B) Kaplan-Meier survival plot of the same mice as in (A). (C) MM.1S solid tumors were established in NSG mice. 7 days after MM.1S cell injection, mice were either left untreated or received infusions of 2×10^6 T cells expressing Luc90-CD828Z, IC9-Luc90-CD828Z, or the negative-control CAR Hu19-CD828Z. The graph shows mean tumor volume \pm SEM for each time point. There were five mice in each group. The graph is representative of one of two independent experiments that used T cells from different donors. (D) Kaplan-Meier survival plot of the same mice as in (C). $p = 0.0031$ for the comparisons of either IC9-Luc90-CD828Z or Luc90-CD828Z versus Hu19-CD828Z by log rank test. (E) Disseminated tumor burdens were established in NSG mice by intravenous injection of 8×10^6 MM.1S-luciferase cells. The mice were left untreated or received infusions of 4×10^6 T cells expressing Luc90-CD828Z, IC9-Luc90-CD828Z, or the negative-control CAR Hu19-828Z. Tumor burden was evaluated by BLI every 3 days. The figure shows BLI intensity of mice for each time point. The figure is representative of one experiment of two experiments with T cells from different donors. (F) Kaplan-Meier survival plots of the same mice as in (E). $p = 0.0027$ for the comparison of either IC9-Luc90-CD828Z or Luc90-CD828Z versus Hu19-CD828Z by log rank test.

T cells expressing IC9-Luc90-CD828Z eradicated solid MM.1S tumors, whereas T cells expressing the anti-CD19 CAR Hu19-CD828Z did not have an anti-tumor effect (Figures 7C and 7D). IC9-Luc90-CD828Z T cells had activity against disseminated malignancy burdens of MM.1S cells in NSG mice. MM.1S cells reached undetectable levels within 9 days of infusion of 4×10^6 IC9-Luc90-CD828Z or Luc90-CD828Z CAR T cells/mouse (Figures 7E and 7F).

In Vivo Assessment of Suicide Switch Function

We established MM.1S solid tumors in NSG mice and injected IC9-Luc90-CD828Z T cells intravenously with or without administration

of AP1903 that activates the IC9 suicide switch. Mice were treated with AP1903 or vehicle control 3 days and 6 days after the CAR T cell infusion. Mice that received IC9-Luc90-CD828Z T cells plus AP1903 had progressive tumors; in contrast, tumors were eliminated from mice that received IC9-Luc90-CD828Z T cells plus the vehicle control (Figures 8A and 8B).

We examined AP1903-induced CAR T cell elimination *in vivo*. Disseminated MM.1S tumor burdens were established in NSG mice. The mice received IC9-Luc90-CD828Z T cells intravenously and were treated with two doses of AP1903 or a vehicle control. Mice were sacrificed 9 days after CAR T cell infusion. We found significantly higher numbers of CD4⁺ and CD8⁺CAR⁺ T cells among splenocytes of mice that received vehicle control compared with mice that received AP1903 (Figures 8C–8E).

CD4-to-CD8 Ratio of Anti-SLAMF7 CAR T Cells in Culture

The CD4:CD8 ratio of *in vitro* cultures of human anti-SLAMF7 CAR T cells is shown in Figures 8F and 8G. Compared to T cell cultures

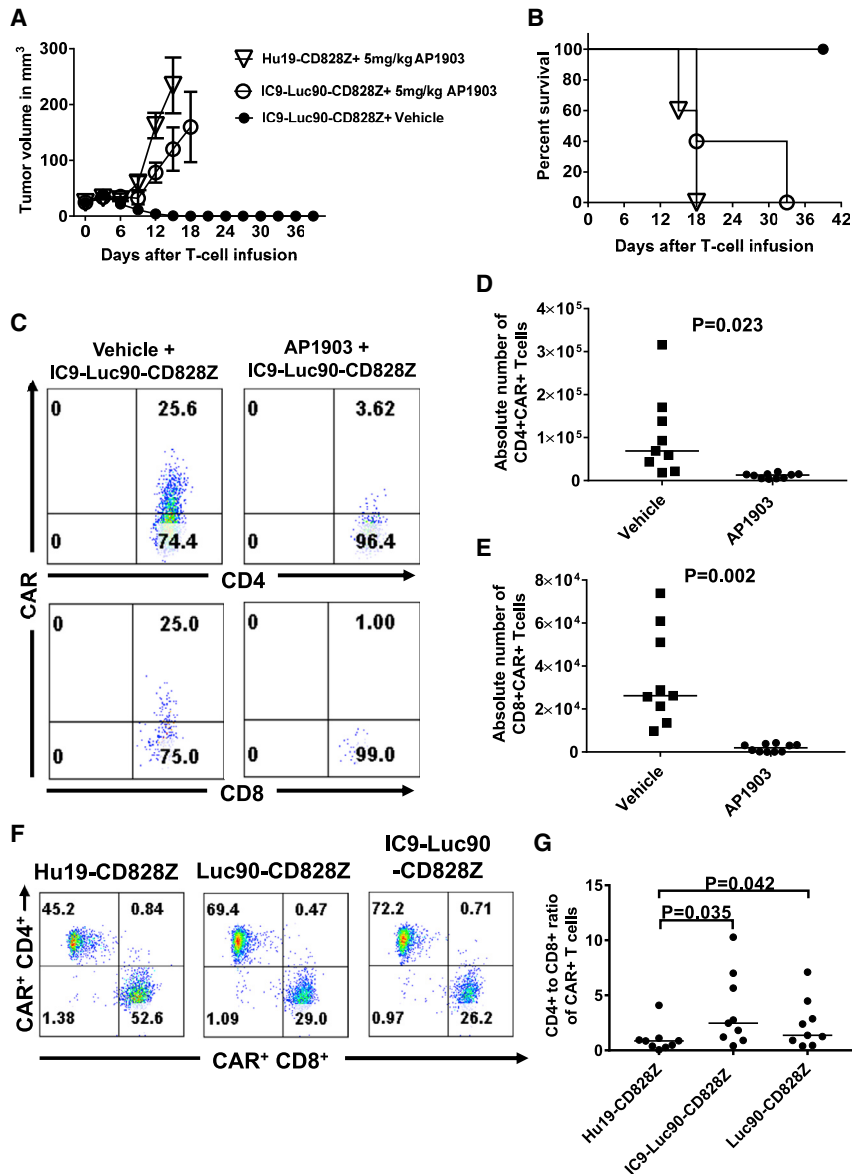


Figure 8. AP1903 Treatment Eliminated IC9-Luc90-CD828Z T Cells

(A) MM.1S solid tumors were established in NSG mice, and then mice received 2×10^6 CAR T cells intravenously. 3 days and 6 days after CAR T cell infusions, mice were injected intraperitoneally with 5 mg/kg of AP1903 or vehicle. Tumors were measured every 3 days in a blinded manner. The mean tumor volumes \pm SEM of each time point are shown. There were five mice in each group. (B) Kaplan-Meier survival plot of the same mice as shown in (A). There was a statistical difference in survival between the IC9-Luc90-CD828Z T cells plus AP1903 and IC9-Luc90-CD828Z plus vehicle; $p = 0.0023$ by log rank test. (C) Disseminated MM.1S cells were established in NSG mice, and the mice then received 4×10^5 T cells transduced with IC9-Luc90-CD828Z. 3 and 6 days after infusion of CAR T cells, mice were injected intraperitoneally with 5 mg/kg of AP1903 or vehicle. On day 9, mice were sacrificed. Splenocytes were isolated and stained for CAR expression on human T cells. The plots show human CD4⁺ (top) and CD8⁺ (bottom) T cells. Plots were first gated on live human CD3⁺ lymphocytes. The plots are from cells of one mouse and are representative results from two independent experiments performed with cells from different donors. (D) CD4⁺ and (E) CD8⁺CAR⁺ T cells present in spleens of mice treated with AP1903 or vehicle control as in (C) were quantified. Compared to the vehicle-control treatment group, the AP1903 treatment group had significantly lower numbers of splenic IC9-Luc90-CD828Z T cells. Each data point indicates the splenic CAR⁺ T cell number of an individual mouse. Horizontal bars indicate medians; $n = 9$ for vehicle-treated group, and $n = 10$ for AP1903-treated group. Statistical testing was by unpaired, two-tailed t tests. $p < 0.05$ was statistically significant. (F) T cells transduced with Hu19-CD828Z, Luc90-CD828Z, or IC9-Luc90-CD828Z were stained for CAR T cell expression on day 7 of culture (5 days after transduction). Plots are gated on CD3⁺CAR⁺ cells. These plots show a higher CD4-to-CD8 ratio with T cells expressing anti-SLAMF7 CARs versus T cells expressing the Hu19-CD828Z anti-CD19 CAR. Plots are representative of 9 experiments. (G) The graph shows CD4-to-CD8 ratios of T cells expressing different CARs. Experiments were performed as in (F). Each data point is the CD4-to-CD8 ratio in one culture of T cells expressing

the indicated CARs. Medians are indicated on the graph by black horizontal bars; $n = 9$ for all groups. Statistical testing was by 1-way ANOVA, followed by Tukey test for multiple comparisons. p values for comparisons of pairs of CAR T cell cultures are shown. Statistical significance was defined as $p < 0.05$.

that were transduced with the Hu19-CD828Z anti-CD19 CAR, there was a statistically higher CD4:CD8 T cell ratio in cultures of T cells expressing anti-SLAMF7 CARs (Figure 8G). Despite possible elimination of some CD8⁺ T cells from cultures transduced with anti-SLAMF7 CARs, T cells in the cultures accumulated steadily (Figure S10).

DISCUSSION

Our work focused on two important aspects of anti-SLAMF7 CAR development: first, selection of either CD28 or 4-1BB as the costimulatory molecule for the CAR; second, development of a construct en-

coding both an effective anti-SLAMF7 CAR and a suicide switch capable of eliminating CAR T cells on demand.

Previous articles have reported anti-SLAMF7 CARs with CD28 costimulatory domains,^{13,20–22} and two articles have reported 4-1BB-containing CAR constructs that targeted both SLAMF7 and BCMA.^{23,38} None of these prior reports compared anti-SLAMF7 CARs with CD28 versus 4-1BB costimulatory domains. We compared the CD28-containing CAR Luc90-CD828Z to Luc90-CD8BBZ, a CAR that was identical to Luc90-CD828Z except for replacement of CD28 with 4-1BB. Compared with Luc90-CD8BBZ, T cells expressing Luc90-CD828Z cleared solid tumors and

disseminated tumor cells more consistently and more quickly (Figures 4A and 4C). We also compared the anti-tumor activity of T cells expressing anti-CD19 CARs with either CD28 or 4-1BB domains (Figure S6), and we found that the CD28-containing CAR had superior anti-tumor activity when compared to the 4-1BB-containing CAR against solid tumors. Our findings are in agreement with prior reports of murine experiments that showed superior anti-tumor activity of CD28-containing CARs versus 4-1BB-containing CARs,^{32–34} however, some prior comparisons of CD28-containing CARs and 4-1BB-containing CARs showed equivalent anti-tumor activity under some conditions and superiority of 4-1BB under other conditions.^{35,36} Finally, some reports have shown superior anti-tumor activity of T cells expressing 4-1BB-containing CARs versus T cells expressing CD28-containing CARs.^{30,31}

7 days after CAR T cell infusion, when tumor regressions were starting, we found higher numbers of tumor-infiltrating T cells when the T cells expressed CD28-containing CARs versus 4-1BB-containing CARs (Figure 4F). The higher numbers of tumor-infiltrating T cells expressing the CD28-containing CAR could explain the superior clearance of tumors with Luc90-CD828Z CAR T cells versus Luc90-CD8BBZ CAR T cells. One mechanism that could be important in the greater anti-tumor activity and greater number of T cells at tumor sites could be the increased cytokine release by T cells expressing CD28-containing CARs versus 4-1BB-containing CARs (Figures 3K–3M). Cytokine production by CAR-expressing human T cells might be especially important in models, such as those used in this work, in which human T cells are infused into mice that lack human cytokines.

As a way to explain the greater numbers of tumor-infiltrating T cells when T cells expressed a CD28-containing versus a 4-1BB-containing CAR, we hypothesized that Luc90-CD828Z CAR T cells might be less prone than Luc90-CD8BBZ T cells to undergo AICD. We performed AICD assays *in vitro*. Although there was no statistically significant difference in AICD between Luc90-CD828Z and Luc90-CD8BBZ CAR T cells for either CD4⁺ or CD8⁺ T cells (Figure S2), there was a trend in CD8⁺ T cells toward more AICD in T cells expressing Luc90-CD8BBZ versus T cells expressing Luc90-CD828Z. A prior study has reported higher levels of apoptosis among T cells expressing 4-1BB-containing CARs versus CD28-containing CARs, especially when, as in the current report, a gamma-retroviral vector was used.³⁹

We found a higher percentage of T cells with a central memory phenotype among T cells expressing a 4-1BB-containing CAR versus T cells expressing a CD28-containing CAR (Figure S1). The reason for this difference is not clear and was not assessed by our experiments. It is possible that CAR signaling caused by SLAMF7⁺ leukocytes in the cultures contributed to this. Other investigators have previously reported a higher percentage of central memory T cells among T cells expressing a 4-1BB-containing CAR versus a CD28-containing CAR, and these investigators also associated differences in memory phenotype with differences in cellular metabolism.⁴⁰

When compared to cultures of anti-CD19 CAR T cells, the CD4:CD8 ratio was higher for cultures of anti-SLAMF7 CAR T cells (Figures 8F and 8G). This increase in CD4:CD8 ratio in cultures of T cells transduced with anti-SLAMF7 CARs suggests that SLAMF7⁺CD8⁺ T cells could have been eliminated by fratricide conducted by the anti-SLAMF7 CAR T cells against SLAMF7⁺CD8⁺ T cells. Note that SLAMF7 expression is substantially higher on CD8⁺ versus CD4⁺ T cells (Figure 1), so fratricide could preferentially eliminate CD8⁺ T cells.

Changes in any part of a CAR sequence might affect CAR T cell function. For example, hinge and transmembrane domains have been shown to strongly affect CAR T cell function.^{41,42} Anti-CD19 CARs commonly used clinically have had either CD28 or 4-1BB costimulatory domains, and 4-1BB has been thought to be an important factor promoting persistence of 4-1BB-containing CAR T cells in clinical trials;⁴³ however, persistence of T cells expressing 4-1BB-containing CARs might not be completely attributable to 4-1BB because the CD28-containing CARs and 4-1BB-containing CARs used in clinical trials contained different hinge and transmembrane domains and differed in other parameters.^{43–46} It is possible that increased T cell persistence attributed to 4-1BB was due, in part, to the difference in hinge and transmembrane domains.

We selected the murine Luc90 scFv rather than the humanized Luc63 scFv for three reasons. First, our results showed slightly superior recognition of SLAMF7⁺ target cells for Luc90-CD828Z versus Luc63-CD828Z when assessed by IFN- γ release (Figure 2H). Second, the humanized Luc63 antibody that the Luc63 scFv is derived from is the commonly used commercial monoclonal antibody elotuzumab.⁴⁷ We reasoned that even though the Luc63 scFv is humanized, there is still a chance of a recipient immune response against the Luc63 scFv, particularly in patients with many prior exposures to elotuzumab. Finally, elotuzumab has modest anti-myeloma activity when used as a single agent,⁴⁸ which also encouraged us to use Luc90.

Anti-SLAMF7 CAR T cells can eliminate NK cells (Figure 5F). NK cell deficiencies are associated with life-threatening viral infections, especially infections caused by members of the herpes virus family, such as cytomegalovirus.²⁴ The risk of elimination of normal NK cells by anti-SLAMF7 CAR T cells makes inclusion of a suicide gene with an anti-SLAMF7 CAR prudent. T cells expressing IC9-Luc90-CD828Z were able to eradicate normal human NK cells from cultures, and we were able to abrogate NK cell depletion by eliminating CAR T cells with AP1903 *in vitro* (Figure 5F). During potential clinical use, IC9-Luc90-CD828Z T cells could be eliminated from patients experiencing severe adverse events by administration of AP1903 (rimiducid).

Previous studies reported anti-SLAMF7 CARs plus depletion systems that utilize the monoclonal antibody cetuximab.^{13,21} One of the main mechanisms for monoclonal antibodies to eliminate target cells is antibody-dependent cellular cytotoxicity (ADCC); NK cells are one of the main cell types participating in ADCC.⁴⁹ Because anti-SLAMF7

CAR T cells and the lymphocyte-depleting chemotherapy that is normally given before clinical CAR T cell infusions will likely eliminate most NK cells in patients, use of a suicide-gene system, such as IC9, which does not depend on ADCC, is preferable.

Despite expression of SLAMF7 on some CAR-transduced CD8⁺ T cells, anti-SLAMF7 CAR T cells accumulated in cultures and sufficient CAR T cells for use in a clinical trial can be obtained. The pre-clinical results with IC9-Luc90-CD828Z have led us to initiate testing of T cells expressing this construct in a clinical trial for MM.

MATERIALS AND METHODS

Patient Samples and Mice

All cells used were human cells. Use of samples from patients enrolled in National Cancer Institute (NCI) clinical trials was approved by the NCI Institutional Review Board. Informed consent was obtained from all patients. Mouse studies were carried out on protocols approved by the NCI Animal Care and Use Committee.

Cell Lines

Cell lines are described in [Supplemental Materials and Methods](#).

PCR to Quantify SLAMF7 Expression

Detection of SLAMF7 RNA in tissues by qPCR is described in [Supplemental Materials and Methods](#).

Tissue Staining of SLAMF7

A paraffin-fixed normal tissue microarray (number MN0661; Pan-tomics, Fairfield, CA) was stained with anti-SLAMF7 clone OTI3B3 (number LS-C340266; LSBio, Seattle, WA).

Design and Construction of Plasmids Encoding CARs

The sequences of Luc90-CD828Z and Luc63-CD828Z followed this pattern from N terminus to C terminus: CD8 α signal sequence, anti-SLAMF7 scFv, human CD8 α hinge and transmembrane domains, the cytoplasmic portion of human CD28, and the cytoplasmic portion of human CD3 ζ . One CAR designated Luc90-CD828Z had a scFv derived from the murine Luc90 antibody. The Luc90 scFv had a sequence from the N terminus to C terminus of the light-chain variable domain, linker, and heavy-chain variable domain. Luc63-CD828Z had an identical sequence as Luc90-CD828Z, except it had a scFv derived from the humanized Luc63 antibody. Sequences of Luc90 and Luc63 variable regions were obtained from a patent.⁵⁰ Luc90-CD8BBZ was identical to Luc90-CD828Z, except a 4-1BB sequence replaced the CD28 sequence of Luc90-CD828Z.

Combination Luc90-CD828Z plus suicide-gene constructs were designed based on prior work.^{26,51,52} The IC9 suicide gene used in this work includes a modified FKBP12 domain linked to a modified caspase-9 domain. The FKBP12 sequence in IC9 was based on the natural FKBP12 sequence (GenBank: AH002818). The caspase-9 sequence used in IC9 was based on the natural caspase-9 sequence (GenBank: NM_001229). The linker sequence connecting the

FKBP12 and caspase-9 components was serine-glycine-glycine-glycine-serine. One CAR plus suicide-gene construct has the IC9 suicide gene 5' to the CAR and was designated IC9-Lu90-CD828Z. The other CAR plus suicide-gene construct has IC9 3' to the CAR and was designated Luc90-CD828Z-IC9. For both Luc90-CD828Z-IC9 and IC9-Luc90-CD828Z, the CAR and IC9 sequences were separated by a T2A sequence.⁵³ We also utilized 3 previously reported CARs: the anti-CD19 CARs Hu19-828Z and Hu19-CD8BBZ^{41,42} and the SP6-CD828Z negative-control CAR.⁷ All CAR DNA sequences were synthesized by GeneArt/Thermo Fisher Scientific (Regensburg, Germany) and ligated into the MSGV1 gamma-retroviral vector backbone³⁷ by standard methods.

T Cell Culture and Gamma-Retroviral Transductions

Gamma-retroviral vector supernatant was prepared as previously described.⁵⁴ In brief, cultures were initiated with whole PBMC by stimulating with the anti-CD3 antibody OKT3, and transduction was performed 2 days after initiation of T cell cultures, as previously described.³⁶ AIM V media refers to AIM V T cell culture media (Thermo Fisher Scientific) plus 5% human serum (Valley Biomedical, Winchester, VA) plus penicillin and streptomycin.

CAR Detection and T Cell Phenotyping

SLAMF7-fragment crystallizable (Fc) protein labeled with phycoerythrin (PE; Creative Biomart, Shirley, NY) or biotin-labeled protein L (GenScript, Piscataway, NJ) was used as a CAR-detection reagent in a manner similar to previous work.^{36,41} The percentage of CAR⁺ T cells was calculated as the percentage of transduced T cells stained with a CAR-detection reagent minus the percentage of untransduced T cells from the same donor stained with the CAR-detection reagent. Antibodies used are in Supplemental Methods. Dead cells were excluded with 7-amino-actinomycin D (7-AAD; BD Biosciences, San Jose, CA) in all experiments. Flow cytometry analysis was performed with FlowJo (Tree Star, Ashland, OR).

In Vitro T Cell Function Assays

CD107a degranulation assays, cytotoxicity assays, ELISA, and proliferation assays were performed as previously described.³⁶ For normalization, ELISA (pg/mL) and CD107a (% CD107a⁺ events) results were divided by the fraction of CAR⁺ T cells for each T cell population used.

AP1903 T Cell Death Analysis

T cell cultures that were either untransduced or CAR transduced were suspended at 1×10^6 T cells/mL in AIM V media + 300 IU of IL-2. The cells were distributed to wells and treated with 1, 10, or 100 ng/mL of AP1903 (rimiducid; MedChemExpress, Monmouth Junction, NJ) or dimethyl sulfoxide (Sigma, St. Louis, MO) as a vehicle control. 6 h after these cultures were set up, the cells were counted and stained with protein L as described above. Next, the cells were incubated with Annexin V (BD Biosciences, San Diego, CA) and 7-AAD, and flow cytometry was performed immediately.

NK Cell Depletion Assay

Untransduced T cells or T cells expressing either IC9-Luc90-CD828Z or Hu19-CD828Z were cultured with autologous PBMCs in a 1:1 ratio. The cocultures were treated with either 10 ng/mL of AP1903 or vehicle control. 24 h after the cultures were set up, the cells were counted, washed, and assessed for NK cells by flow cytometry.

Solid Tumor and Disseminated Tumor Cell Mouse Experiments

Mouse tumor experiments are described in figure legends and were conducted with MM.1S cells as previously described.³⁶ Tumor experiments with NALM6 cells were carried out as previously described.⁴¹

Real-Time qPCR for Quantifying CAR⁺ Cells in Tumors

Similar to previous work, we made serial 1:5 dilutions of DNA from the infused T cells of each experimental group into untransduced T cell DNA, and we made standard curves by performing qPCR on this DNA.^{42,55,56} We used the standard curves to determine the percentage of CAR⁺ cells in tumors as described in [Supplemental Materials and Methods](#). The absolute number of CAR⁺ cells in each tumor was calculated by multiplying the percentage of CAR⁺ cells by the total number of cells in the tumor.

Solid Myeloma Tumor with AP1903 Treatment

Solid tumors of MM.1S cells were established in NSG mice. The mice were infused with 2×10^6 CAR T cells. 3 and 6 days after CAR T cell infusion, mice received intraperitoneal injections of 5 mg/kg AP1903 or vehicle. The dose of AP1903 was determined from a previous study.⁵²

T Cell Persistence in Mice after AP1903 Treatment

Mice were injected with 8×10^6 MM.1S-luciferase cells intravenously. After 13 days, 4×10^6 CD3⁺CAR⁺ T cells/mouse were infused intravenously. Mice received an intraperitoneal injection of either 5 mg/kg of AP1903 or vehicle on days 3 and 6 after T cell infusion. Mice were sacrificed 9 days after T cell infusion. Splenocytes were counted, and flow cytometry was performed for SLAMF7-Fc-PE for CAR detection, anti-CD3, anti-CD4, and anti-CD8. To calculate the number of CAR T cells/spleen, the total number of splenocytes per mouse was multiplied by the product of the %CD3⁺, %CD4⁺, or %CD8⁺ and the %CAR⁺.

AICD Assay

T cells expressing Luc90-CD828Z or Luc90-CD8BBZ were flow sorted using a BD FACSAria. T cells were then cocultured with either MM.1S or NGFR-K562 in a 1.5:1 target-to-effector ratio for 18 h. T cells were then stained for CAR and Annexin V as described above.

Statistical Analyses

Statistical comparisons were performed as described in the figure legends with GraphPad Prism version 7.

SUPPLEMENTAL INFORMATION

Supplemental Information can be found online at <https://doi.org/10.1016/j.ymthe.2020.10.008>.

AUTHOR CONTRIBUTIONS

C.A. conducted experiments, planned experiments, analyzed data, and wrote the manuscript. M.A.P., N.L., C.G., D.V., S.C., and S.M.H. conducted experiments, analyzed data, and edited the manuscript. S.A.F. designed CAR constructs and edited the manuscript. J.N.K. designed CAR constructs, conducted experiments, planned experiments, analyzed data, and wrote the manuscript.

CONFLICTS OF INTEREST

This work was supported by NIH, NCI, and Center for Cancer Research intramural funding. J.N.K. and S.A.F. are inventors on a patent application for the anti-SLAMF7 plus suicide switch constructs presented in this work. The content of this publication does not necessarily reflect the views or policies of the Department of Health and Human Services nor does mention of trade names, commercial products, or organizations imply endorsement by the US government.

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Supplemental Information

Development of CAR T Cells Expressing a Suicide Gene Plus a Chimeric Antigen Receptor Targeting Signaling Lymphocytic-Activation Molecule F7

Christina Amatya, Melissa A. Pegues, Norris Lam, Danielle Vanasse, Claudia Geldres, Stephanie Choi, Stephen M. Hewitt, Steven A. Feldman, and James N. Kochenderfer

Supplemental Materials

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Supplemental Figure S1: Memory Phenotype

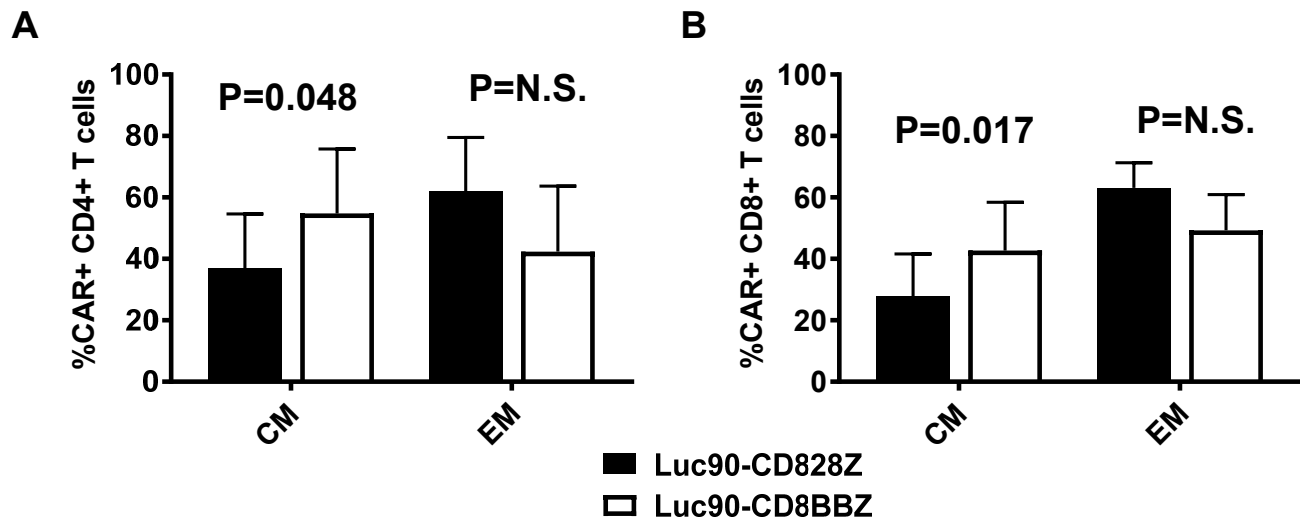


Figure S1. Memory phenotype of T cells expressing different anti-SLAMF7 CARs. T cells were transduced with vectors encoding either Luc90-CD828Z or Luc90-CD8BBZ. On day 9 of T-cell culture, cells were stained for detection of CAR, CD45RA, and CCR7. The graphs show percentages of (A) CD4⁺ and (B) CD8⁺ CAR T cells that were either CD45RA-negative, CCR7⁺ central memory cells (CM) or CD45RA-negative, CCR7-negative effector memory cells (EM). N=3 different donors for all groups. For both CD4⁺ and CD8⁺ CAR-expressing T cells, the percentage of CM cells was higher for Luc90-CD8BBZ T cells compared with Luc90-CD828Z T cells. There was not a statistically-significant difference in the percentage of EM cells between the two CARs. Bars represent mean+SEM. Statistical testing was done by paired, 2-tailed T tests, and statistical significance was defined as P<0.05. N.S. stands for not statistically significant.

Supplemental Figure S2: Activation-induced Cell Death of CARs with CD28 Versus 4-1BB

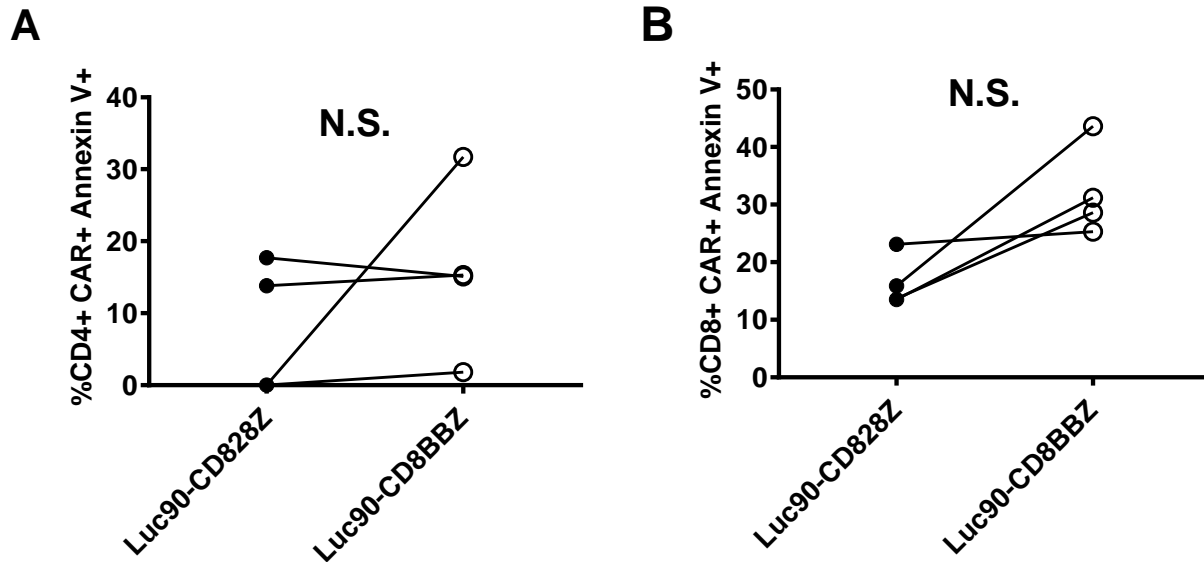


Figure S2. Apoptosis of CAR-transduced T cells was assessed by Annexin V staining. CD4⁺ or CD8⁺ T cells expressing either Luc90-CD828Z or Luc90-CD8BBZ were cultured overnight with either MM.1S cells (SLAMF7⁺) or NGFR-K562 cells (SLAMF7-negative). Flow cytometry was then performed to assess Annexin V staining of live (A) CD4⁺ or (B) CD8⁺ CAR-expressing T cells. The graphed results show the %Annexin V⁺ cells with NGFR-K562 stimulation subtracted from the %Annexin V⁺ cells with MM.1S stimulation. N=4 different donors. Statistical significance was assessed with a 2-tailed, paired T test. P<0.05 was considered statistically significant. Lines connect paired results with cells from the same donors. There was no statistically significant difference between Luc90-CD828Z T cells and Luc90-CD8BBZ for either CD4⁺ or CD8⁺ T cells, but there was a trend toward lower %Annexin V⁺ cells in CD8⁺ T cells expressing Luc90-CD828Z compared with T cells expressing Luc90-CD8BBZ (P=0.059). N.S. stands for not statistically significant.

Supplemental Figure S3: IFN γ Release

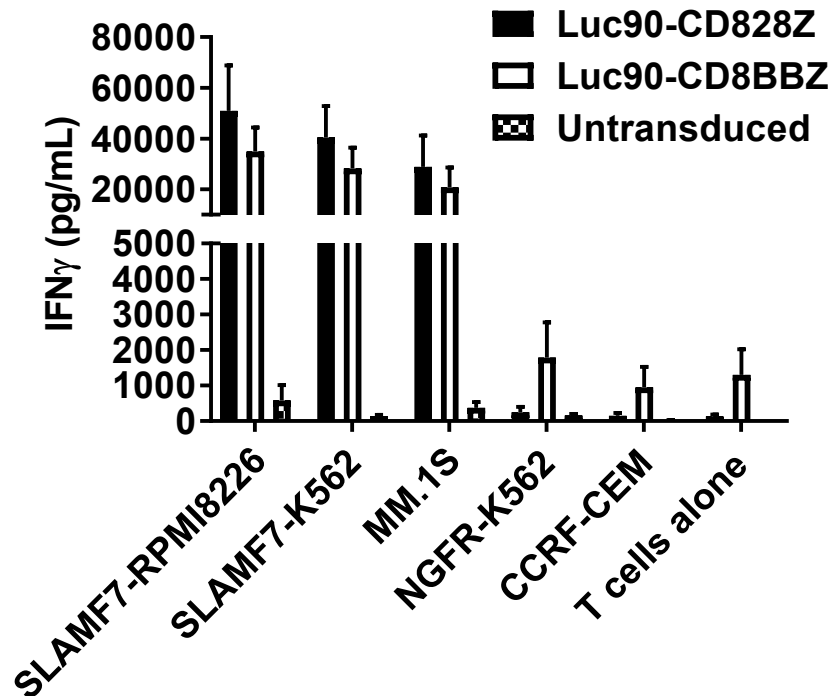


Figure S3 IFN γ release by Luc90-CD828Z and Luc90-CD8BBZ T cells.

T cells were transduced with either Luc90-CD828Z or Luc90-CD8BBZ and cultured overnight with the indicated target cells. Culture supernatants were then assayed for IFN γ by ELISA. Units are pg/mL. Bars show mean+SEM. SLAMF7-RPMI8226, SLAMF7-K562, and MM.1S were SLAMF7⁺. NGFR-K562, and CCRF-CEM were SLAMF7-negative. N=3 replicates with T cells from different patients for SLAMF7-RPMI8226. N=6 replicates with T cells from different patients for other target cells. Note relatively high nonspecific IFN γ release with SLAMF7-negative targets and T cells alone for Luc90-CD8BBZ T cells.

Supplemental Figure S4: Luc90-CD828Z Versus Luc90-CD8BBZ Degranulation

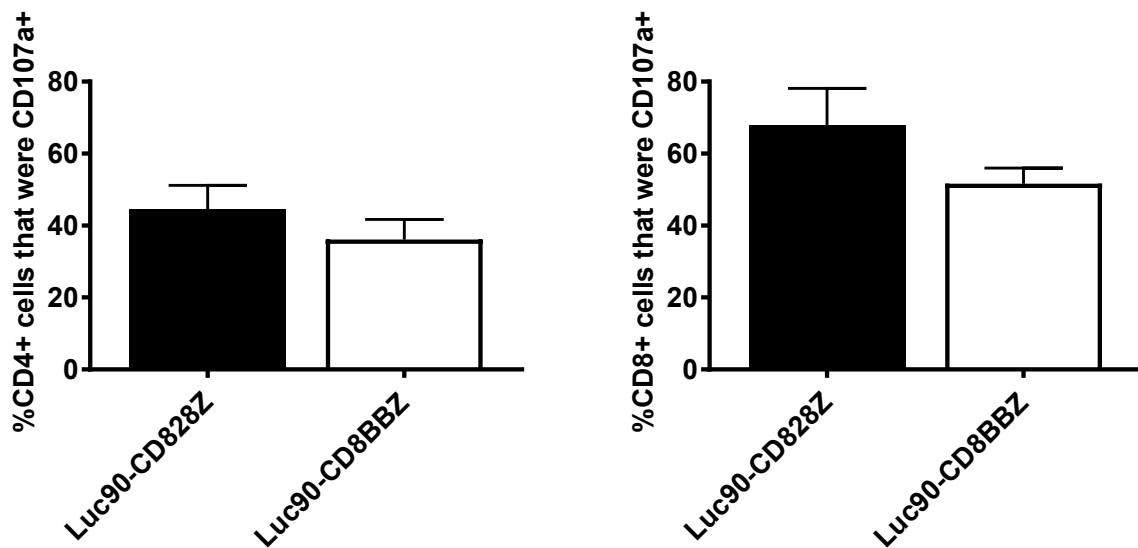


Figure S4. Degranulation of T cells expressing either Luc90-CD828Z or Luc90-CD8BBZ. T cells expressing the indicated CARs were cultured 4 hours with either MM.1S or NGFR-K562 cells in the presence of an antibody against CD107a. Cells were then stained with antibodies against CD3, CD4, and CD8; n=3 different donors for both the CD4 and CD8 analyses. Values are CD107a⁺ events with MM.1S stimulation minus CD107a⁺ events in a parallel culture with NGFR-K562. The percentage of T cells that degranulated was not statistically different between Luc90-CD828Z and Luc90-CD8BBZ for either CD4⁺ or CD8⁺ T cells. The %CD107a⁺ events were normalized for CAR expression. Statistical comparison was by 2-tailed paired T test; P<0.05 was considered statistically significant.

Supplemental Figure S5: Tumor Treatment with 4×10^6 CAR T-cell Dose

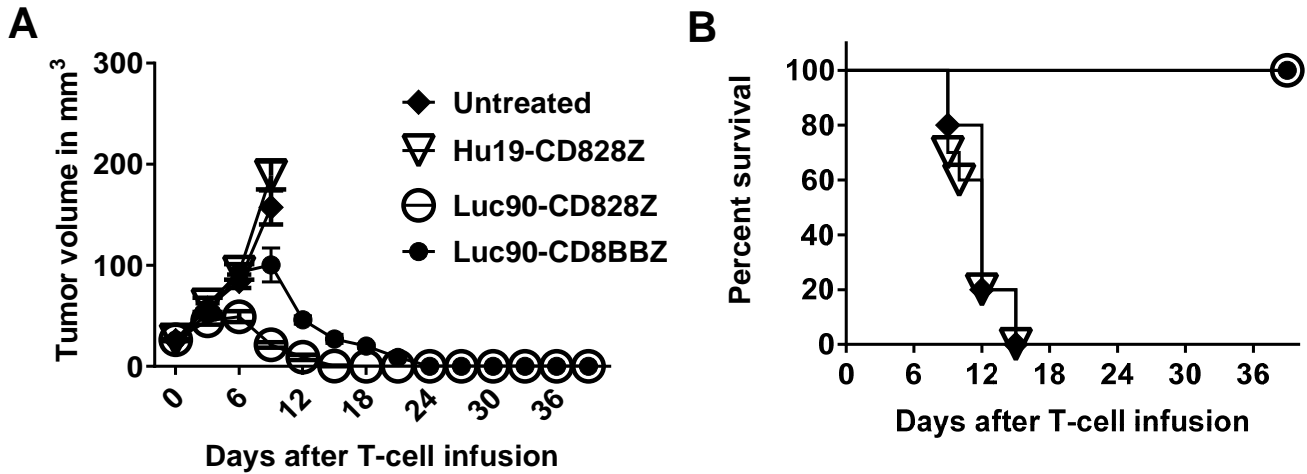


Figure S5. Both Luc90-CD828Z and Luc90-CD8BBZ CAR T cells eradicated solid tumors at a dose of 4×10^6 CAR T cells/mouse. (A) MM.1S tumors were established in NSG mice. Mice were either left untreated or received infusions of 4×10^6 T cells expressing Luc90-CD828Z, Luc90-CD8BBZ, or, as a control, the anti-CD19 CAR Hu19-CD828Z. The graph shows mean tumor volume \pm SEM for each time point. The graph includes data from two independent experiments with different donor T cells. There were five mice in each group per experiment, so $n=10$ total mice for each treatment group. (B) Kaplan-Meier survival plot of the same mice as in A. By log-rank test, there was a statistically-significant survival difference between mice that received T cells expressing Hu19-CD828Z versus Luc90-CD828Z ($P < 0.0001$). There was also a statistically-significant survival difference between mice that received T cells expressing Hu19-CD828Z versus Luc90-CD8BBZ ($P < 0.0001$). Early tumor control was superior for Luc90-CD828Z T cells versus Luc90-CD8BBZ T cells. For example, on day 9 after CAR T-cell infusion, the mean tumor volume was 21.3 mm^3 for Luc90-CD828Z T cells and 100.3 mm^3 for Luc90-CD8BBZ T cells ($P = 0.0002$ by 2-tailed unpaired T test).

Supplemental Figure S6: Anti-tumor Activity of Anti-CD19 CARs Containing CD28 Versus 4-1BB Domains

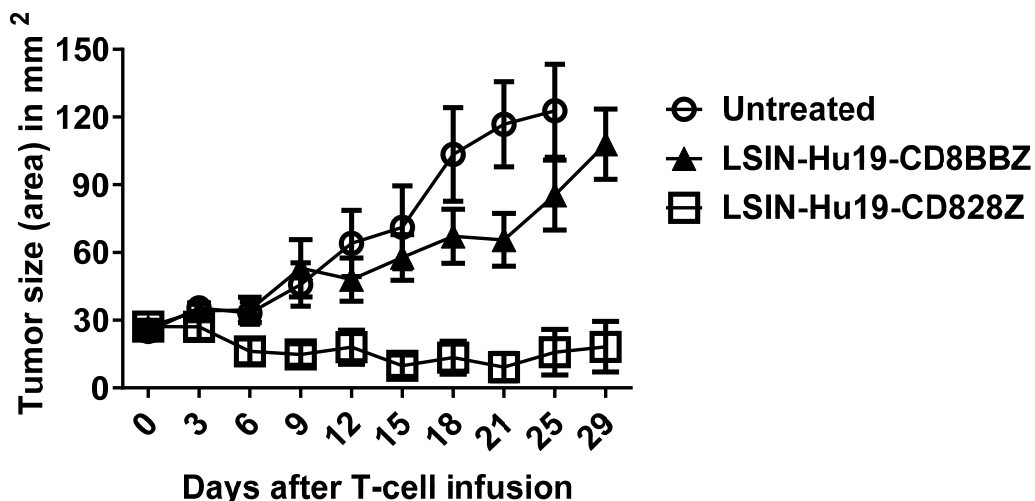


Figure S6. Comparison of anti-tumor activity of T cells expressing anti-CD19 CARs with 4-1BB versus CD28 costimulatory domains. NSG mice were injected with 4 million NALM6 cells to form solid tumors. Human donor T cells were transduced with either LSIN-Hu19-CD8BBZ or LSIN-Hu19-CD828Z, and mice received intravenous infusion of 10 million CAR T cells after solid tumors were established. A third group of tumor-bearing mice was left untreated. The day of CAR T-cell infusion was day 0 on the graph. Tumor sizes were monitored by measuring with calipers. The graph shows the mean \pm SEM of tumor sizes of 5 mice per group at each time point. These results are from one of two experiments with similar results. In both of the experiments, tumor growth in mice that received 4-1BB-containing CAR T cells was more similar to tumor growth in untreated mice than to tumor growth in mice that received CD28-containing CAR T cells.

Supplemental Figure S7: Costimulatory Sequences

CD28 sequence in all CD28-containing CARs

RSKRSRLLHSD YMNMPRRPG PTRKHYPYA
PPRDFAAAYS

4-1BB sequence in Luc90-CD8BBZ

KRGRKKL LYIFKQPFMR PVQTTQEEDG
CSCRFPEEEE GGCEL

4-1BB sequence in Hu19-CD8BBZ

RFSVVKRGRKKL LYIFKQPFMR PVQTTQEEDG
CSCRFPEEEE GGCEL

Figure S7. Sequences of costimulatory domains in CARs.

The same CD28 sequence was used in all CARs in this report. The anti-SLAMF7 CAR Luc90-CD8BBZ and the anti-CD19 CAR Hu19-CD8BBZ contained the indicated 4-1BB sequences that differed by 5 amino acids at the N-terminus.

Supplemental Figure S8: Comparison of expression

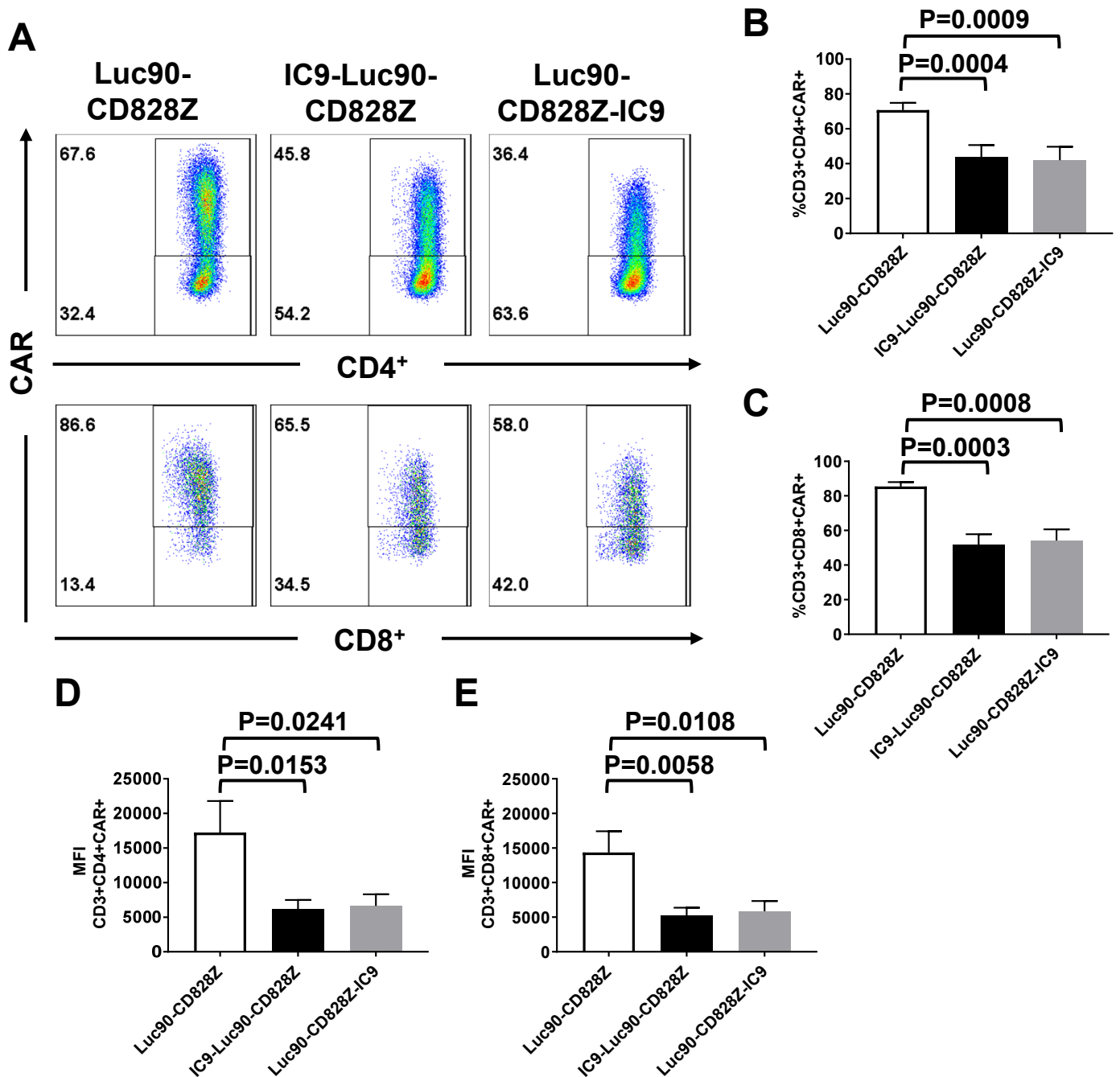


Figure S8. Comparison of percentage and intensity of CAR expression among T cells transduced with Luc90-CD828Z and IC9-bearing Luc90-CD828Z CAR constructs. T cells were transduced with vectors encoding either Luc90-CD828Z or IC9-Luc90-CD828Z or Luc90-CD828Z-IC9. On day 7 of T-cell culture, T cells were stained for detection of CAR. (A) The plots are representative examples of the percentages and intensities of CAR expression on CD4⁺ and CD8⁺ T cells. (B-C) The graphs show percentages of (B) CD4⁺ and (C) CD8⁺ T cells that were CAR⁺. (D-E) The graphs show median fluorescence intensities (MFI) of CAR expression on (D) CD4⁺ and (E) CD8⁺ T cells. For B-E, n=8 different donors for all groups. For both CD4⁺ and CD8⁺ CAR-expressing T cells, the percentages of both CAR expression and MFI were statistically higher for Luc90-CD828Z T cells compared with IC9-Luc90-CD828Z and Luc90-CD828Z-IC9 T cells. Bars represent mean+SEM. There was no statistical difference between IC9-Luc90-CD828Z and Luc90-CD828Z-IC9 in either percent CAR expression or MFI. Statistical testing was done by paired, 2-tailed T tests. After correction for multiple comparisons with the Bonferroni method, statistical significance was defined as P<0.017.

Supplemental Figure S9: Degranulation in Response to Primary Multiple Myeloma Cells

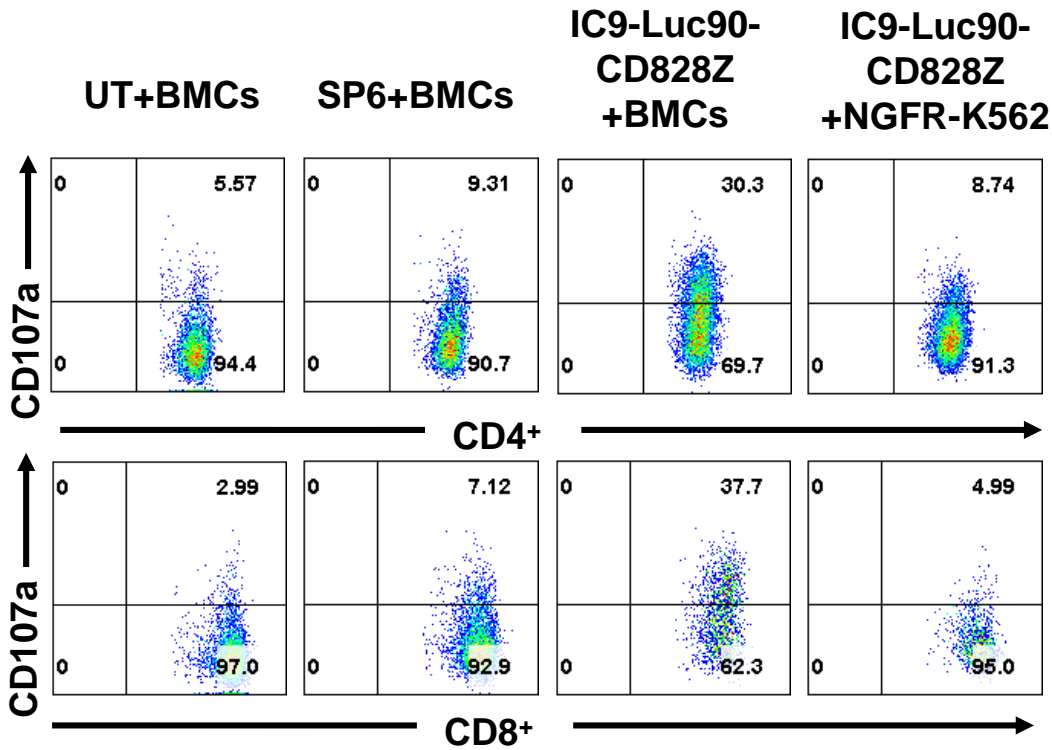
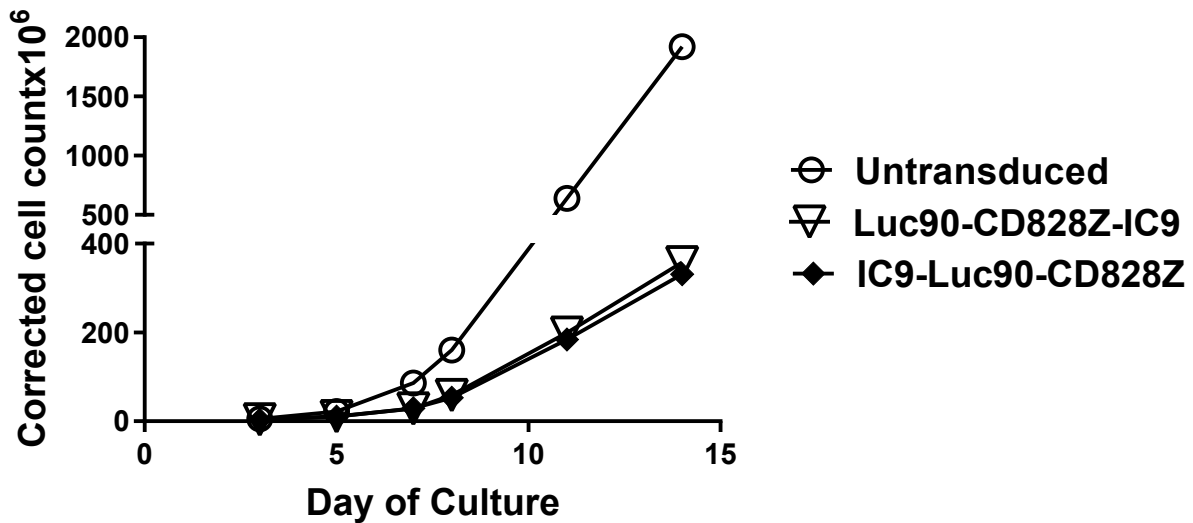


Figure S9

IC9-Luc90-CD828Z CAR T cells specifically degranulate in the presence of primary MM cells. T cells were transduced with vectors encoding SP6-CD828Z or IC9-Luc90-CD828Z or left untransduced. On day 8 of T cell culture, T cells were cultured with either primary bone marrow cells (BMCs) from a multiple myeloma patient or NGFR-K562 (negative control) for 4-hours in the presence of an antibody against CD107a. Flow cytometry was performed. Plots are gated on live CD4⁺ or CD8⁺ T cells. The figure shows one of 2 experiments with similar results performed with cells from different donors.

Supplemental Figure S10: T-cell Accumulation in Culture



Supplemental Figure S10. The cumulative increases in numbers of T cells left untransduced, transduced with the gene for Luc90-CD828Z-IC9, or transduced with the gene for IC9-Luc90-CD828Z are shown. Untransduced T cells accumulated more rapidly than the CAR-transduced T cells, but CAR-transduced T cells did accumulate steadily. One representative example of 4 experiments with cells from 4 different donors is shown.

Supplemental Table S1: SLAMF7 Immunohistochemistry

Organs stained for SLAMF7 by immunohistochemistry and found to not have SLAMF7 expression	
adrenal	lung
bladder	ovary
bone	pancreas
breast	parathyroid
cerebellum	pituitary
cerebral cortex	placenta
eye	prostate
fallopian tube	skin
esophagus	spinal cord
stomach	spleen
small intestine	skeletal muscle
colon	testis
rectum	thymus
heart	thyroid
kidney	tonsil
liver	uterine cervix
	uterine endometrium

Table S1

SLAMF7 is not expressed on non-hematopoietic tissues from various organs. A paraffin-fixed normal tissue microarray (Pantomics, number MN0661) was stained with anti-SLAMF7 (clone OT13B3, LSBio number LS-C340266). The table includes tissues where SLAMF7 protein expression was not detected by IHC except on infiltrating plasma cells, some macrophages and some lymphocytes.

Supplemental Table S2: Soluble SLAMF7

SLAMF7 (ng/mL)	SLAMF7-positive targets				SLAMF7-negative targets				T cells only			
	0	100	200	400	0	100	200	400	0	100	200	400
	SLAMF7-K562	SLAMF7-K562	SLAMF7-K562	SLAMF7-K562	NGFR-K562	NGFR-K562	NGFR-K562	NGFR-K562	T cells	T cells	T cells	T cells
UT	38.6	42.5	39.7	37.3	51.4	46.9	49.9	44.7	24.4	23.0	28.0	25.5
MSGV-Hu19-CD828Z	35.2	35.0	34.5	38.3	40.1	35.6	38.3	40.5	31.1	32.1	30.6	31.6
MSGV-Luc90-CD828Z	12553.6	10969.2	9486.7	13422.8	17.2	16.2	12.8	13.1	19.5	16.1	13.6	13.3

Table S2. Soluble SLAMF7 protein does not interfere with anti-SLAMF7 CAR T-cell target recognition. T cells were transduced with vectors encoding Hu19-CD828Z or Luc90-CD828Z or left untransduced. Seven days after transduction, T cells were co-cultured with SLAMF7-K562 or NGFR-K562 target cells or were cultured alone. The co-cultures included either vehicle or 100ng/mL, 200ng/mL or 400ng/mL of soluble SLAMF7 protein. After overnight culture, supernatant was harvested and IFN γ was evaluated by ELISA.

Supplemental Table S3: IFN γ Release by Anti-CD19 CAR T Cells

	CD19-K562	CLL	CCRF-CEM	T Cells alone	% CAR+
LSIN-Hu19-CD828Z	32645	2003	64	61	60.3
LSIN-Hu19-CD8BBZ	25240	3548	495	504	51.2
Untransduced	112	<12	239	<12	0

Table S3. T cells expressing either Hu19-CD828Z or Hu19-CD8BBZ produced IFN γ in an antigen-specific manner in vitro. Human T cells from the same donor were left untransduced or transduced with LSIN-Hu19-CD828Z or LSIN-Hu19-CD8BBZ. The T cells were cultured overnight with the target cells indicated on the top row. CD19-K562 and primary CLL cells were CD19⁺. CCRF-CEM was CD19-negative. After the overnight culture, an IFN γ ELISA was performed. The percentage of the T cells that were CAR⁺ is indicated. All values are the mean of duplicate wells. Units are pg/mL of IFN γ .

Table S4: Specificity of IC9-Luc90-CD828Z T cells

	SLAMF7 -K562	A549	SAOS2	COLO205	U251	624	TC71	A431-H9	Panc 10.5	CCRF- CEM	T cells only
Untransduced	<12	30.3	33.6	19.2	28.8	30.2	<12	<12	<12	22.6	<12
Hu19-CD828Z	<12	27.6	37.7	37.7	28.1	18.7	13.4	0.4	<12	32.7	<12
IC9-Luc90- CD828Z	45231.4	15.6	29.5	34.2	19.0	121.9	12.3	31.8	12.3	24.0	30.3

Table S4. IC9-Luc90-CD828Z does not recognize SLAMF7-negative cell lines. T cells from the same donor were transduced with vectors encoding the anti-CD19 CAR Hu19-CD828Z or IC9-Luc90-CD828Z or left untransduced. The T cells were cultured overnight with the target cells listed on the top row. All targets were SLAMF7-negative except for SLAMF7-K562. After the overnight culture, an ELISA for IFN γ was performed. All values are pg/mL of IFN γ .

Supplemental Methods

Real-time qPCR to Quantify SLAMF7 Expression

SLAMF7 cDNA copies were quantitated in samples of cDNA from human tissues included in the Human Major Tissue qPCR Panel II (Origene, Rockville, MD) by performing qPCR with a SLAMF7-specific primer and probe set (Applied Biosystems, Foster City, CA). Plasmacytoma cells of a patient with advanced multiple myeloma served as a positive control for SLAMF7 expression. A standard curve for the SLAMF7 qPCR was created by amplifying dilutions of a plasmid that encoded the full-length cDNA of SLAMF7 (Origene, Rockville, MD). β -actin cDNA copy numbers were quantified in the same tissues with a Taqman β -actin primer and probe kit (Applied Biosystems, Foster City, CA) and used to normalize SLAMF7 copy numbers. PCR was conducted on a Roche LightCycler480.

Cell Lines

MM.1S is an SLAMF7⁺ MM cell line from ATCC (Manassas, VA). SLAMF7-K562 and SLAMF7-RPMI8226 are cell lines transduced with the SLAMF7 gene in our laboratory. The following CD19-negative human cell lines from ATCC were used: A549 (lung carcinoma), CCRF-CEM (T cell leukemia), Saos2 (osteosarcoma), Panc10.05 (pancreatic carcinoma), COLO205 (colon carcinoma). U251 (glioblastoma) and H431-H9 (epidermoid carcinoma) cell lines were gifts from Dr. Robert Somerville (NIH). TC71 (Ewing's sarcoma) were a gift of Dr. M. Tsokos, (NIH). The 624 melanoma cell line was a gift from Dr. Steven A. Rosenberg, NIH. SLAMF7-negative, NGFR-K562 were transduced with the gene for low-affinity nerve growth factor in our laboratory. CD19⁺

NALM6 cells (acute lymphoid leukemia from DSMZ, Braunschweig, Germany) were used.

Antibodies Used in Flow Cytometry

For T-cell phenotyping, cells were stained with the following antibodies: CD3 APC-Cy7 (Clone UCHT1, BD Biosciences, San Jose, CA), CD4 FITC or BV510 (Clone RPA-T4, BD Biosciences, or Biolegend, San Diego, CA), CD8 PE-Cy7 or eFluor450 (Clone RPA-T8, BD Biosciences or Thermo Scientific). For memory T-cell phenotyping, the following antibodies were included: CD45RA FITC (Clone HI100, BD Biosciences) and CCR7 APC (Clone 150503, BD Biosciences).

Real-time qPCR for quantifying CAR⁺ Cell in Tumors

In murine experiments, the percentages of infused T cells that were CAR⁺ were determined by flow cytometry. These percentages were used in making CAR⁺ T cell standard curves. Tumor cells were mechanically disaggregated and treated with a 1:1 mix of ACK lysing buffer to PBS. The cells were counted, and DNA was extracted from the lysates using a Qiagen (Valencia, CA) DNeasy kit. DNA from the tumor was amplified in duplicate with a primer and probe set (IDT, Coralville, IA) that was specific for the MSGV1 retroviral vector. Real-time PCR was carried out with a Roche (Basel, Switzerland) Light Cycler 96 real-time PCR system. Similar to an approach used previously, we made serial 1:5 dilutions of DNA from the infused T cells of each experimental group into untransduced T-cell DNA, and we made standard curves by performing qPCR on this DNA¹⁻³ To determine the percentage of total tumor cells that contained the CAR gene, the qPCR results obtained with each tumor's DNA were compared to the qPCR results obtained from each experimental group's CAR⁺ T-cell

standard curve. Note that total tumor cells included MM.1S cells and infiltrating human T cells.

PCR results were normalized to tumor β -actin. For the actin PCR, an actin plasmid (Origene plasmid ACTB NM_001101) was amplified in duplicate with a primer and probe set (Bio-rad ACTB qHsaCEP0036280, Hercules, CA) to generate a standard curve. This standard curve was used to determine actin copy numbers in tumor DNA samples amplified with the same actin primer and probe set.

After the percentages of CAR⁺ cells were determined by qPCR, the absolute number of CAR⁺ T cells in each tumor was calculated by multiplying the percentage of CAR⁺ T cells by the total number of cells in the tumor. PCR results showed very similar efficiencies in reactions with DNA from Luc90-CD828Z-transduced T cells versus Luc90-CD8BBZ-transduced T cells. All tested tumors had levels of CAR DNA that were above the lower limit of detection for the PCR reactions.

References

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2. Morgan, RA, Yang, JC, Kitano, M, Dudley, ME, Laurencot, CM, and Rosenberg, SA (2010). Case report of a serious adverse event following the administration of t cells transduced with a chimeric antigen receptor recognizing ERBB2. *Molecular Therapy* **18**: 843-851.
3. Brudno, JN, Lam, N, Vanasse, D, Shen, YW, Rose, JJ, Rossi, J, *et al.* (2020). Safety and feasibility of anti-CD19 CAR T cells with fully human binding domains in patients with B-cell lymphoma. *Nat Med* **26**: 270-280.