Checkpoinnt Blockade Reverses Anergy in IL-13Rα2 Humanized scFv-Based CAR T Cells to Treat Murine and Canine Gliomas

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We generated two humanized interleukin-13 receptor α2 (IL-13Rα2) chimeric antigen receptors (CARs), Hu08BBz and Hu08BBz, that recognized human IL-13Rα2, but not IL-13Rα1. Hu08BBz also recognized canine IL-13Rα2. Both of these CAR T cell constructs demonstrated superior tumor inhibitory effects in a subcutaneous xenograft model of human glioma compared with a humanized EGFRvIII CAR T construct used in a recent phase 1 clinical trial (ClinicalTrials.gov: NCT02209376). The Hu08BBz demonstrated a 75% reduction in orthotopic tumor growth using low-dose CAR T cell infusion. Using combination therapy with immune checkpoint blockade, humanized IL-13Rα2 CAR T cells performed significantly better when combined with CTLA-4 blockade, and humanized EGFRvIII CAR T cells’ efficacy was improved by PD-1 and TIM-3 blockade in the same mouse model, which was correlated with the levels of checkpoint molecule expression in co-cultures with the same tumor in vitro. Humanized IL-13Rα2 CAR T cells also demonstrated benefit from a self-secreted anti-CTLA-4 minibody in the same mouse model. In addition to a canine glioma cell line (J3T), canine osteosarcoma lung cancer and leukemia cell lines also express IL-13Rα2 and were recognized by Hu08BBz. Canine IL-13Rα2 CAR T cell was also generated and tested in vitro by co-culture with canine tumor cells and in vivo in an orthotopic model of canine glioma. Based on these results, we are designing a pre-clinical trial to evaluate the safety of canine IL-13Rα2 CAR T cells in dog with spontaneous IL-13Rα2-positive glioma, which will help to inform a human clinical trial design for glioblastoma using humanized scFv-based IL-13Rα2 targeting CAR T cells.

INTRODUCTION

Malignant gliomas, including grade IV gliomas, also called glioblastomas (GBMs), are the most common primary malignant brain tumors and are associated with high morbidity and mortality.1,2 The aggressive nature of glioma cell infiltrative growth in the CNS makes total resection impossible to achieve. Despite best available therapy, including surgical resection, radiotherapy, chemotherapy, and tumor-treating field, the median survival is only 12–17 months for patients with GBMs and 2–5 years for patients with grade III gliomas.2,3

Adaptive immunotherapy with redirected T cells is a feasible strategy to treat these malignant tumors. Long-term disease-free survival was
achieved in a patient with refractory chronic lymphocytic leukemia after treatment with CD19 targeting chimeric antigen receptor modified autologous T (CAR T) cells, and complete remission was achieved in 90% of patients with relapsed acute lymphoblastic leukemia (ALL) with this strategy. However, to date, the anti-tumor activity of CAR T cells in solid tumors has been much more modest. Our group previously utilized humanized anti-epidermal growth factor receptor (EGFR) variant III (EGFRvIII) CAR T cells (2173BBz) in a phase 1 clinical trial (ClinicalTrials.gov: NCT02209376) of 10 patients with recurrent GBMs. There were obvious changes in the tumor microenvironment after CAR T cell infusion, including reduction of the EGFRvIII target antigen associated with CAR T cell trafficking and in situ functional activation. However, the study was not powered to determine clinical response (median overall survival was 251 days). A recent report described the use of repeated intratumoral and intrathecal infusions of redirected T cells expressing an interleukin-13 (IL-13) zetakine, a mutated IL-13 cytokine, fused with a T cell-signaling domain in a single patient with recurrent multifocal GBM, which led to complete tumor regression for 7.5 months. Taken together, these studies raise hope for the treatment of GBMs and other solid tumors with redirected T cells.

Interleukin-13 receptor α2 (IL-13Rα2) is expressed in different human tumor types, but no expression is seen on normal human tissues, except adult testes (Figure S1B). IL-13 signaling through IL-13Rα2 plays a critical role in cell migration and invasion. A previous study found 82% of GBM cases expressed IL-13Rα2, making it a promising target for immunotherapy. Neutralizing antibody and drug-conjugated antibody targeting IL-13Rα2 inhibiting tumor growth in xenograft mouse models. IL-13Rα2-based tumor vaccine also benefitted pediatric glioma patients. Although IL-13 zetakine redirected T cells bind IL-13Rα2 and induced a limited clinical response, they also bind IL-13Rα1 (Figure S1A), which is expressed in some normal human tissues and has demonstrated adverse, off-target effects. To circumvent these effects, a single-chain variable fragment (scFv)-based IL-13Rα2-targeting CAR T construct without reactivity against IL-13Rα1 was previously made using a murine scFv (clone 47), but this raised the possibility of inducing a human anti-mouse antibody (HAMA) response and anaphylaxis, which would limit the function of the CAR T cells and potentially induce severe adverse effects. Therefore, to improve the performance of this promising strategy in the clinical treatment of GBMs, it is necessary to generate a fully humanized, highly specific, scFv-based IL-13Rα2 CAR T cell and demonstrate its function both in vitro and in vivo in clinically relevant pre-clinical models of GBM.

The tumor microenvironment of malignant gliomas is immunosuppressive, and this has been shown after CAR T cell infusion. Immune checkpoint receptors (e.g., PD-1, CTLA-4, TIM-3, and LAG-3) are a series of molecules that downregulate the stimulation of activated T cells with different temporal and spatial profiles to regulate T cell functions. Checkpoint inhibitors have been applied in cancer therapy to overcome T cell inhibition within the immunosuppressive tumor microenvironment and recruit the T cell repertoire to target tumor cells. To date, most combinatorial studies have used anti-PD-1 checkpoint blockade together with endogenous T cell response to tumor antigens and a few selected reports on engineered T cells. Determining the most effective combination of different checkpoint inhibitors with different CAR T cells is critical for optimal clinical effect. Checkpoint inhibitors also can be directly delivered by the adoptively transferred CAR T cells via gene modification, which aims to reduce the adverse effects that can be caused by systemic delivery of checkpoint inhibitors. This is an additional strategy we will utilize to further liberate the function of IL-13Rα2 CAR T cells in immunotherapy of malignant gliomas.

Although clinical trials are often based on numerous pre-clinical animal studies, most of these studies have been performed on rodent small animal models. While more physiologic than in vitro models, there remains a gap between rodent models and human clinical medicine. Only 10.4% of new therapeutic compounds entering phase 1 clinical trials between 2003 and 2011 achieved FDA approval. A high percentage of clinical trial failures are associated with loss of patients’ treatment opportunities and limited financial resources. The contribution of naturally occurring diseases in larger animals to tumor models has historically attracted the attention of scientists. In particular, employment of immunologically intact, client-owned dogs, with spontaneous tumors that present comparable barriers to effective immunotherapies, can provide additional, clinically relevant safety and efficacy data that will help to inform human clinical trial design. While non-human primate models more closely resemble human physiology, there are no oncologic models as well as challenges and ethical concerns. A series of therapeutic interventions were evaluated in tumor-bearing domestic canine patients, based on the disease similarity to humans, including bone marrow transplantation and cancer vaccines, which contribute to early-phase human studies as well as later stages of development. Furthermore, CAR T cell therapies are currently being generated and tested in dogs with spontaneous tumors, e.g., a CD20-targeting CAR T cell approach in relapsed B cell lymphoma. Therefore, based on the encouraging result of the treatment efficiency of IL-13 zetakine redirected T cells in human GBM and our access to canine patients with GBMs, we aimed to establish highly specific scFv-based IL-13Rα2-targeting canine CAR T cells that may be used in vivo to facilitate the study of IL-13Rα2-targeting CAR T cell therapy on solid tumors in both human and canine cancer patients.

Here we established two humanized IL-13Rα2-targeting CAR T cells, and we explored the feasibility of combinational therapy with different checkpoint blockades (anti-PD-1, anti-CTLA-4, and anti-TIM-3) compared with humanized EGFRvIII CAR T cells. We further incorporated endogenous checkpoint blockade secretion in our IL-13Rα2-targeting CAR T cells, to test the feasibility of local secretion of checkpoint inhibitors in combination with highly specific CAR T cell function. Finally, we developed and evaluated the function
of canine IL-13Rα2-targeting CAR T cells as the penultimate step required for a companion animal clinical trial in GBM.

RESULTS

Humanized IL-13Rα2-Targeting CAR T Cells

The Human Protein Atlas illustrates that IL-13Rα1 is widely expressed in normal human tissues (Figure S1A), while IL-13Rα2 is restricted to expression in testes (Figure S1B). In contrast, The Cancer Genome Atlas demonstrates IL-13Rα2 was expressed in multiple different tumor samples with different tissue origins. Very high expression of IL-13Rα2 was found in GBM (Figure S1C). To make IL-13Rα2-targeting CAR T cells, a second-generation CAR construct with human CD8 hinge and transmembrane domains linked with human 4-1BB and CD3ζ intracellular signaling domains was used. Human IL-13Rα2-targeting murine scFv sequences, Mu07 and Mu08 generated by Dr. Ma and colleagues, were cloned into the pGEM vector (Figure S2A). mRNA encoding the IL-13Rα2 CAR was made in vitro with the pGEM template. After mRNA electroporation into human T cells, the two CAR structures, Mu07BBz and Mu08BBz, were detected on the T cell surface (Figure S2B). Three glioma cell lines (U87, U251, and D270) and two T cell cancer lines, Sup-T1 and Jurkat, were chosen as target cells for testing the specificity and function of the murine scFv-based IL-13Rα2 CAR T cells in vitro. IL-13Rα2 was detected on all three glioma cell lines, but not on the Sup-T1 and Jurkat T cell cancer lines, confirming their negative control status (Figure S2C). To determine antigen-specific CAR T cell activation, we co-cultured electroporated murine scFv-based IL-13Rα2 CAR T cells with target tumor cells. Interferon (IFN)γ production was only detected within CAR T cells co-cultured with IL-13Rα2-positive tumor cell lines (Figure S2D), and it was not detected within CAR T cells co-cultured with the negative control cell lines. The production of interleukin-2 (IL-2) and tumor necrosis factor alpha (TNF-α) also demonstrated the same pattern as IFNγ production (Figure S4A).

To avoid HAMA responses and anaphylaxis, we utilized humanized 07 (Hu07) and 08 (Hu08) scFvs to generate the humanized IL-13Rα2-targeting CAR T cells. Hu07 and Hu08 scFvs were prepared by complementarity-determining region (CDR) grafting with frame back mutations. DP-54 and DPK9 were utilized as the human acceptor framework. Based on the binding activity and thermal stability of the humanized scFvs described previously, Hu07 and Hu08 sequences were cloned into the second-generation CAR construct in the pGEM vector (Figure 1B). After human T cell mRNA electroporation, the Mu07/08 and Hu07/08 scFvs were detected on the T cell surface with anti-murine or anti-human immunoglobulin G (IgG) antibodies (Figure 1A). All four structures were detected by the anti-murine IgG, but only the humanized CARs were recognized by anti-human IgG (Figure 1A). To stably express the IL-13Rα2 CARs on the human T cell surface, Hu07BBz and Hu08BBz CAR constructs were cloned into the pTRPE vector, which is a transfer plasmid used in lentivirus production (Figure 1B). CAR expression was detected on the cell surface of transduced T cells (Figure 1C).

To determine the specificity of both IL-13Rα2 CARs, we co-cultured transduced IL-13Rα2 CAR T cells with target cell lines that expressed neither IL-13Rα1 nor IL-13Rα2 (supT1 and Jurkat cells), IL-13Rα1 only (the lung cancer cell line A549), IL-13Rα2 only (D270), or both IL-13Rα1 and IL-13Rα2 (U87 and U251) (Figure 1D). Both humanized 07/08BBz CAR constructs produced IFNγ when co-cultured with IL-13Rα2-positive target cells (Figure 1E). Additionally, the humanized IL-13Rα2-targeting CAR T cells did not cross-react with IL-13Rα1, as evidenced by a lack of IFNγ production when co-cultured with A549. IL-2 and TNF-α production also corresponded with the production of IFNγ (Figure S4B). These co-culture results are consistent with those of murine scFv-based CARs (Figure S2D).

To determine the ability of the humanized IL-13Rα2-targeting CAR T cells to mediate antigen-specific cytotoxicity, we performed chromium release assays at different effector/target (E:T) ratios (1:1, 3:1, 10:1, and 30:1) of humanized IL-13Rα2-targeting CAR T cells to target tumor cells. The humanized CAR T cells specifically killed IL-13Rα2-positive target cells (U87, U251, and D270) during 4 hr of co-culture, even at the lower E:T ratios (Figure 1F). No killing activity above background was detected in the negative control groups. Hu07 and Hu08BBz CAR T cells (Figure S3A) were also co-cultured with normal human primary cells. We detected the different levels of IL-13Rα1 expression on several types of human primary cell (Figure S3B), specifically human small airway epithelial cells, human renal epithelial cells, and human keratinocytes. No stimulation was found in the co-cultured humanized IL-13Rα2-targeting CAR T cells with either of these targets by intracellular cytokine (IFNγ, IL-2, and TNF-α) staining (Figures S3C and S4C). IL-13Rα2 expression was also detected on the co-cultured human aortic smooth muscle cells and pulmonary artery smooth muscle cells with IL-13Rα2 antibody (clone 47) (Figure S3B), which also induced stimulation of both CAR T cells (Hu07BBz and Hu08BBz) illustrated as the type I cytokine production (IFNγ, IL-2, and TNF-α) (Figures S3C and S4C). Taken together, IL-13Rα2 represented a viable target in GBM, and Hu07 and Hu08BBz CAR T cells target this receptor with a high degree of specificity.

IL-13Rα2 CAR T Cells Control Tumor Growth In Vivo

To further test the function of the two humanized IL-13Rα2 CAR T cells in vivo, we developed subcutaneous and orthotopic glioma xenograft models in NSG mice. The D270 glioma cell line was chosen for the in vivo work, based on pathophysiologic characteristics that closely match human primary glioma invasive and aggressive growth patterns. The status of the orthotopic implanted D270 glioma cells was monitored in the NSG mouse model (Video S1), using a two-photon microscope, after skull thinning, as described previously. The D270 cell line not only expressed IL-13Rα2 endogenously but also EGFRvIII (Figure 2A). The expression of both targets was detected on days 0, 1, 2, 3, 5, and 7 of D270 culture in vitro (Figure S4D). This allowed us to include the previously described 2173BBz CAR T cells in this experiment. 2173BBz is a humanized,
EGFRvIII-targeting CAR with the same CAR backbone as Hu07/08BBz. We co-cultured EGFRvIII-targeting (2173BBz) and IL-13Rα2-targeting (Hu08BBz) CAR T cells with D270 glioma cell line. The stimulation of T cells was illustrated by FITC-conjugated anti-CD69 antibody staining, the median fluorescence intensity (MFI) was quantified on CD4 and CD8 CAR-positive T cells after 24- or 48-hr co-culture, controlled with un-transduced T cells. Statistically significant differences were calculated by one-way ANOVA with post hoc Tukey test. (C) Human T cells enumerated in the spleens of D270-injected NSG mice (n = 3), 11 days after i.v. transferring equal numbers of un-transduced T cells, EGFRvIII-targeting (2173BBz), or IL-13Rα2-targeting (Hu08BBz) CAR T cells. (D) Five million CAR-positive EGFRvIII-targeting (2173BBz) or IL-13Rα2-targeting (Hu08BBz) CAR T cells or the same number of un-transduced T cells after i.v. infusion in a D270 subcutaneously implanted NSG mouse model (n = 10 per group), 7 days after tumor implantation. Tumor volume measurements (left panel) and bioluminescence imaging (right panel) were performed to evaluate the tumor growth. Linear regression was used to test for significant differences between the experimental groups. Endpoint was predefined by the mouse hunch, inability to ambulate, or tumor reaching 2 cm in any direction, as predetermined IACUC-approved morbidity endpoint. Survival based on time to endpoint was plotted using a Kaplan-Meier curve (Prism software). Statistically significant differences were determined using log-rank test. (E) 800,000 IL-13Rα2-targeting CAR-positive (Hu08BBz) CAR T cells or the same number of un-transduced T cells were given by i.v. infusion in NSG mice (n = 8 per group) orthotopically implanted with the D270 tumor, 8 days after tumor injection. Bioluminescence imaging was repeated every 3–4 days to evaluate the tumor growth. Endpoint was predefined and statistically significant differences were determined as described in (D). *p < 0.05, **p < 0.01, ***p < 0.001, ****p < 0.0001. Data are presented as means ± SEM.

Figure 1. Humanized IL-13Rα2-Targeting CAR T Cells
(A) Flow cytometric detection of CAR expression by human T cells, after mRNA electroporation of murine and humanized scFv- (07 and 08) based CAR constructs using rabbit anti-mouse or rabbit anti-human IgG antibodies. (B) Vector maps of tested anti-IL-13Rα2 CAR design based on the size of each components. (C) CAR expression staining of the humanized IL-13Rα2 CAR transduced T cells used in the co-culture experiments. (D) IL-13Rα1 and IL-13Rα2 expression analysis on the human tumor cell lines (Sup-T1, Jurkat, A549, U87, U251, and D270) with isotype antibodies staining control in blue. (E) Flow-based intracellular cytokine (IFNγ) staining of the humanized IL-13Rα2 CAR T cells co-cultured with human tumor cell lines in (D) controlled with un-transduced T cells (UTD). Human CD8 was stained to distinguish the CD4-positive and CD8-positive subgroups of T cells along the x axis. (F) Chromium release assay of humanized IL-13Rα2 CAR T cells co-cultured with tumor cell lines in (D) was analyzed at different effector/target (E:T) ratios (1:1, 3:1, 10:1, and 30:1) compared with the UTD T cells with one-way ANOVA post hoc Tukey test. **p < 0.01, ***p < 0.001, ****p < 0.0001. Data are presented as means ± SEM.

Figure 2. IL-13Rα2 CAR T Cells Control Tumor Growth In Vivo
(A) Flow-based EGFRvIII and IL-13Rα2 expression on the D270 tumor cell line controlled with control antibodies. (B) EGFRvIII-targeting (2173BBz) and IL-13Rα2-targeting (Hu08BBz) CAR T cells co-cultured with D270 tumor cell line. The stimulation of T cells was illustrated by FITC-conjugated anti-CD69 antibody staining, the median fluorescence intensity (MFI) was quantified on CD4 and CD8 CAR-positive T cells after 24- or 48-hr co-culture, controlled with un-transduced T cells. Statistically significant differences were calculated by one-way ANOVA with post hoc Tukey test. (C) Human T cells enumerated in the spleens of D270-injected NSG mice (n = 3), 11 days after i.v. transferring equal numbers of un-transduced T cells, EGFRvIII-targeting (2173BBz), or IL-13Rα2-targeting (Hu08BBz) CAR T cells. (D) Five million CAR-positive EGFRvIII-targeting (2173BBz) or IL-13Rα2-targeting (Hu08BBz) CAR T cells or the same number of un-transduced T cells after i.v. infusion in a D270 subcutaneously implanted NSG mouse model (n = 10 per group), 7 days after tumor implantation. Tumor volume measurements (left panel) and bioluminescence imaging (middle panel) were performed to evaluate the tumor growth. Linear regression was used to test for significant differences between the experimental groups. Endpoint was predefined by the mouse hunch, inability to ambulate, or tumor reaching 2 cm in any direction, as predetermined IACUC-approved morbidity endpoint. Survival based on time to endpoint was plotted using a Kaplan-Meier curve (Prism software). Statistically significant differences were determined using log-rank test. (E) 800,000 IL-13Rα2-targeting CAR-positive (Hu08BBz) CAR T cells or the same number of un-transduced T cells were given by i.v. infusion in NSG mice (n = 8 per group) orthotopically implanted with the D270 tumor, 8 days after tumor injection. Bioluminescence imaging was repeated every 3–4 days to evaluate the tumor growth. Endpoint was predefined and statistically significant differences were determined as described in (D). *p < 0.05, **p < 0.01, ***p < 0.001, ****p < 0.0001. Data are presented as means ± SEM.

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subgroups of CAR T cells after 24 hr (p < 0.0001 and p = 0.0021) and 48 hr of co-culture (p = 0.0008 and p = 0.0038) (Figure 2B), suggesting that the Hu08BBz CAR T cells were more activated in response to target cells compared to the 2173BBz CAR T cells.

To determine antigen-specific proliferation, we co-cultured carboxy-fluorescein succinimidyl ester (CFSE)-labeled UTD T cells, 2173BBz CAR T cells, and Hu08BBz CAR T cells with the D270 cell line, as well as the target negative cell line A549. The intensity of CFSE signaling on UTD T cells and CAR-positive T cells (2173BBz and Hu08BBz) was determined by flow cytometry (Figure S4E). The MFI of both CAR T populations was progressively lower than the UTD T cells during 3, 5, and 8 days co-culture with the D270 cell line (p < 0.0001), indicating increased proliferation compared with the UTD T cells. The spleen, as an important lymphoid organ, is a reservoir of large amounts of lymphocytes. The status of CAR T cells in the spleen has been demonstrated to correspond with their function in vivo.\textsuperscript{15} We transplanted CellTrace Violet- (blue) and tetramethylrhodamine (TRITC)- (red) labeled CAR T cells intravenously into an orthotopically implanted glioma NSG mouse model, where they were visualized in the mouse spleen with 2-photon microscopy (Video S2). To determine CAR-mediated T cell expansion in vivo, we also intravenously infused 2 x 10^6 human CAR T cells (2173BBz and Hu08BBz) or UTD T cells into mice, 7 days after D270 subcutaneous implantation in NSG mice. At 11 days after T cell transfer, human T cells were counted in the spleen of three mice per group. There were 7 times more human T cells in mice treated with 2173BBz (n = 7.3 x 10^5) than in those treated with UTD T cells (n = 1 x 10^5), while there were 30 times more human T cells in the Hu08BBz (n = 3 x 10^6) group than the UTD group (Figure 2C); but, no statistical differences were detected between each group with one-way ANOVA.

To determine whether CAR T cells could control tumor growth, we implanted D270 cells subcutaneously, and 7 days later we administered 5 x 10^6 CAR T cells (2173BBz/Hu07BBz/Hu08BBz) or the same number of UTD T cells via the intravenous route. Compared with UTD cells, all three CAR T cells tested (2173BBz/Hu07BBz/Hu08BBz) significantly inhibited tumor growth, as determined by caliper measurement (p < 0.0001) and decreased bioluminescent signal (p < 0.0001), as detected by the in vivo imaging system (IVIS) and representative tumor size (Figure 2D). For mice treated with the humanized IL-13Rz2 CAR T cells (Hu07BBz and Hu08BBz), no tumor was palpable 16 days after intravenous (i.v.) T cell implantation. Only background signal (2 x 10^4p/s/cm^2/sr) was captured in the humanized IL-13Rz2 CAR T cell co-cultures. Furthermore, no tumor recurrence was observed in either of the two groups (Hu07BBz and Hu08BBz) over 43 days, based on repeated flank palpation and bioluminescence imaging (BLI), significantly better tumor eradication (p < 0.0001), and overall survival (p = 0.0012) than the EGFRvIII (2173BBz) CAR T cells group in this mouse model (Figure 2D).

Next we evaluated the effects of the humanized IL-13Rz2 CAR T cells (Hu08BBz) in an orthotopic glioma model. D270 glioma cells were implanted intracranially, and 8 days later 8 x 10^5 CAR T cells (Hu08BBz) or control UTD T cells were administered intravenously. All mice in the UTD group became hunched and symptomatic by day 17–20 after tumor implantation, and they were euthanized based on the predetermined IACUC-approved endpoint. Although 25% of the mice in the Hu08BBz group were euthanized during the same period (day 20), bioluminescent signals from the D270 tumor cells were not detected in any other mice in that treatment group (p < 0.0001), and mice treated with Hu08BBz showed a clear survival advantage over mice treated with UTD T cells (control group) (p = 0.0027) (Figure 2E). These results suggest Hu07 and Hu08BBz have potent anti-tumor activity in vivo.

**Immune Checkpoint Blockade Selectively Enhances the Function of CAR T Cells**

Immune checkpoint receptors are expressed on the surface of T cells. We next decided to test the effects of the three most frequently studied immune checkpoint receptors (PD-1, CTLA-4, and TIM-3) on CAR T cell function to determine whether immune checkpoint blockade could augment CAR T cell function.

First, we assessed the expression of PD-1, CTLA-4, and TIM-3 on human T cells during in vitro expansion on days 0, 3, 7, and 13 (Figure S6A). T cell activation was illustrated by the expression of CD69, which peaked on day 3 of in vitro culture. The percentage of checkpoint receptor (PD-1, CTLA-4, and TIM-3)-positive T cells also increased during early stimulation, and then it decreased in both the CD4 and CD8 T cells subgroups. We also detected the ligands of PD-1 (PD-L1), CTLA-4 (CD80 and CD86), and TIM-3 (galecin-9) on the surface of the T cells. The percentage of ligand-positive T cells similarly fluctuated with time after T cell stimulation (Figure S6A). Investigation of the D270 glioma cell line revealed expression of PD-L1 and galecin-9 (Figure S6B), making it an appropriate tumor target to study the effects of checkpoint blockade on CAR T cells.

To determine the effects of CAR target engagement on checkpoint molecule expression over time, after 12 days of T cell expansion, we co-cultured the humanized EGFRvIII-targeting CAR T cells (2173BBz) and the humanized IL-13Rz2-targeting CAR T cells (Hu08BBz) with either D270 tumor cells (EGFRvIII and IL-13Rz2+) or A549 tumor cells (EGFRvIII and IL-13Rz2+, IL-13Rz1-; negative control), in vitro for 24 or 48 hr (Figures S5A and S5B). Compared with A549 cell co-culture groups or the group of UTD T cells, the expression of PD-1, CTLA-4, and TIM-3 on D270 cell co-cultured 2173BBz and Hu08BBz CAR T cells was increased at both time points (Figure 3A). Interestingly, the expression level of these checkpoint receptors differed between the 2173BBz and Hu08BBz CAR T cells (Figure 3A). CTLA-4 expression was higher on the Hu08BBz CAR T cells than on the 2173BBz CAR T cells during 24 and 48 hr of co-culture, in both CD4 (p = 0.0003 and p = 0.0010) and CD8 (p = 0.0006 and p = 0.0050) T cell subsets, which corresponded to the level of CD69 expression when co-cultured with the D270 cell line (Figure 2B). Although PD-1 expression was
not statistically different between 2173BBz and Hu08BBz CAR T cells after 24 hr of co-culture, it was significantly higher on the CD4- and CD8-positive 2173BBz CAR T cells than on the Hu08BBz CAR T cells after 48 hr of co-culture (p = 0.0021 and p = 0.0456). Finally, the expression of TIM-3 was higher with 2173BBz than Hu08BBz CAR T cells, independent of the duration of co-culture or the CD4 and CD8 subsets (p = 0.0371 and p = 0.0026 for 24-hr co-culture; p = 0.0002 and p = 0.0004 for 48-hr co-culture).

To further study if blocking the immune checkpoint receptors enhanced the tumor-killing activity of the CAR T cell, we combined 2173BBz and Hu08BBz CAR T cell treatment with intraperitoneal (i.p.) administration of checkpoint inhibitor (anti-PD-1, anti-CTLA-4, and anti-TIM-3) in NSG mice with intracranially implanted D270 tumors. For the majority of mice, only background signal was detectable at later time points, making it difficult to show any benefit of combined checkpoint blockade. Therefore, to enable us to determine whether checkpoint blockade enhanced the anti-tumor effects of CAR T cell therapy, we decreased the number of CAR T cells administered and studied the effects of combination CAR T cells and checkpoint blockade on mice with subcutaneously implanted D270 glioma cells. In this mouse model, intraperitoneal delivery of checkpoint inhibitor (anti-PD-1, anti-CTLA-4, and anti-TIM-3) did not have any effect on reducing tumor size, because the tumor grew at the same rate when mice were injected with UTD T cells (p = 0.1600, 0.1194, and 0.4565) (Figure 3B). Significant inhibition of tumor growth was seen when either 2173BBz or Hu08BBz CAR T cells were combined with anti-PD-1 and anti-TIM-3 (p < 0.0001 for 2173BBz groups and p = 0.0325 and p = 0.0032 for Hu08BBz groups). In addition, Hu08BBz CAR T cells also showed enhanced anti-tumor effects...
when used in combination with anti-CTLA-4, whereas 2173BBz CAR T cells did not benefit from CTLA-4 checkpoint blockade (p = 0.5817 for 2173BBz groups and p < 0.0001 for Hu08BBz groups) (Figure 3C).

Next, we compared the effects of the different checkpoint inhibitors on tumor growth in mice treated with either the 2173BBz CAR T cells or Hu08BBz CAR T cells (Figure 3D). Combination therapy with either anti-PD-1 or anti-TIM-3 produced greater anti-tumor effects with 2173BBz CAR T cells than did anti-CTLA-4 (p < 0.0001), with combination anti-PD-1 having the greatest effect (p = 0.0185 compared with anti-TIM-3 group). For the Hu08BBz CAR T cells, CTLA-4 blockade presented the best combinational therapy (p = 0.0010 and p < 0.0001). These results correspond with the expression levels of checkpoint receptors on 2173BBz and Hu08BBz CAR T cells during co-culture with D270 tumor cells in vitro (Figure 3A). Only combination therapy with anti-PD-1 prolonged survival in the 2173BBz CAR T cell treatment group (p = 0.00135), whereas anti-CTLA-4 prolonged survival in the Hu08BBz CAR T cell treatment group (p = 0.0135). We also compared the number and activation status of human T cells in mouse spleens between the 2173BBz CAR T cell group and the 2173BBz CAR T cell plus anti-PD-1 group (Figure S6C). PD-1 expression was efficiently blocked in the combination anti-PD-1 group (p = 0.0034 and 0.0037, respectively), and there was a higher percentage of CD69+ human T cells (p = 0.0006 and 0.0340) and a larger percentage of CD8+ human T cells in this group compared to those treated with 2173BBz CAR T cells alone (p = 0.0177 and 0.0022). Thus, checkpoint blockades enhanced the function of CAR T cells with specific efficacy on different CARs.

IL-13Rα2 CAR T Cells Are Selectively Enhanced by In Situ-Secreting Anti-CTLA-4 Checkpoint Blockade

Given the finding that systemic checkpoint blockade enhanced the anti-tumor effect of both 2173BBz and Hu08BBz CAR T cells, we sought to evaluate the effects of in situ checkpoint blockade by modifying CAR T cells to secrete checkpoint inhibitors. We rationalized that local checkpoint inhibition would enhance CAR T cell activity and would reduce adverse effects induced by systemic checkpoint blockade. We transduced T cells with the pTRPE vector containing the Hu08BBz CAR construct linked to anti-PD-1, anti-CTLA-4, and anti-TIM-3 constructs via the ribosomal skipping sequence (P2A), which enables simultaneous expression of the Hu08BBz CAR and the checkpoint inhibitor molecules. To decrease T cell burden of molecule secretion, we reduced the size of the checkpoint inhibitors by directly linking their scFvs with the CH3 domain of the human IgG1 molecule to generate minibodies (Figure 4A). We refer to these as minibody-secreting T cells (MiSTs).

We confirmed by flow cytometry that the surface expression of the Hu08BBz CAR was detected on the Hu08BBz CAR T cells and on the Hu08BBz CAR T cells secreting the minibodies (Figure 4B). Conditioned media from the CAR T cells were collected, concentrated, and used in a standard direct ELISA to confirm the secretion and binding of anti-PD-1 and anti-CTLA-4 minibodies from CAR T cells to recombinant hPD-1 and hCTLA-4 (p = 0.0017 and 0.0075) (Figure 4C). To evaluate the specificity of the anti-TIM-3 minibody secreted from Hu08BBz CAR T cells, we co-cultured Hu08BBz CAR T cells with or without minibodies with the D270 tumor cell line for 24 or 48 hr, and a competitive inhibition experiment was performed with fluorochrome-conjugated anti-TIM-3 antibody. The MFI-determined binding of fluorochrome-conjugated anti-TIM-3 antibody was significantly lower in the anti-TIM-3 minibody-secreting CAR T cell group, suggesting effective secretion and blockade by the TIM-3 minibody (Figure 4D).

Furthermore, with the exception of CD4-positive anti-TIM-3-secreting CAR T cells at 48 hr, there was no statistical difference between TIM-3 expression on anti-TIM-3-secreting CAR T cells and UTD T cells in these co-cultures (CD4 p = 0.6642 and CD8 p = 0.8771 for 24-hr co-culture; CD4 p = 0.0014 and CD8 p = 0.4578 for 48-hr co-culture). The MFI-determined binding of fluorochrome-conjugated anti-PD-1 and anti-CTLA-4 antibodies was also lower on anti-PD-1/anti-CTLA-4 MiSTs than non-minibody-secreting CAR T cells (Hu08BBz) when co-cultured with D270 cells, demonstrating competitive binding by the anti-PD-1/anti-CTLA-4 minibodies secreted by MiSTs (anti-PD-1 MiST and Hu08BBz CD4 p = 0.0678 and CD8 p = 0.0140 for 24-hr co-culture; CD4 p < 0.0001 and CD8 p = 0.0012 for 48-hr co-culture; anti-CTLA-4 MiST and Hu08BBz CD4 p < 0.0001 and CD8 p < 0.0001 for 24-hr co-culture; CD4 p = 0.0002 and CD8 p = 0.0006 for 48-hr co-culture) (Figure S7A). Taken together, these data suggest that Hu08BBz MiSTs secreting anti-PD-1, anti-CTLA-4, and anti-TIM-3 minibodies were successfully generated.

We next evaluated the effects of minibody secretion on Hu08BBz CAR T cell activation by D270 target cells in vitro for 48 hr, using CD69 expression as a marker of activation. The stimulation of Hu08BBz CAR T cells without minibody secretion was higher than the minibody-secreting cells in the 24 or 48 hr of co-culture, but no statistical difference was seen with anti-CTLA-4 minibody-secreting Hu08BBz CAR T cells in the CD8-positive T cell subgroup (p = 0.0614 and 0.4561) (Figure S7B). Among the minibody-secreting groups, the stimulation of anti-PD-1 Hu08BBz CAR T cells was significantly lower than the other two groups in the 24-hr co-culture (p < 0.0001) and in the 48-hr co-culture (p = 0.0008 and 0.0010) of CD4-positive T cells. For the anti-CTLA-4- and anti-TIM-3-secreting Hu08BBz CAR T cells, the stimulation of anti-CTLA-4 secretion was higher than anti-TIM-3 secretion in the CD4+ CAR T cells at 24 hr (p < 0.0001), while no difference was seen in the other subgroups (Figure S7B). We also compared the cytokine secretion (IFNγ, IL-2, and TNF-α) of these CAR T cells when co-cultured with the D270 tumor cell line. Compared with the other Hu08BBz CAR T cell groups, a significantly lower percentage of the anti-PD-1 minibody-secreting group secreted cytokines in each subgroup (Figure S7B). A greater percentage of anti-CTLA-4- and anti-TIM-3-secreting groups produced IFNγ and TNF-α than the no minibody-secreting group.
The production of IL-2 was 1.5-fold more frequent in the anti-CTLA-4-secreting group than in the anti-TIM-3-secreting group, and it was significantly higher in the CD4+ subgroup (p = 0.0001) (Figure S7C).

To determine the effects of checkpoint blockade via minibody secretion from CAR T cells, we intravenously administered a sub-therapeutic dose of Hu08BBz CAR T cells (8 x 10^5 cells per mouse), and the same amount of MiSTs or UTD T cells into NSG mice, 8 days after subcutaneous implantation of D270 cells in one set of experiments. Despite this low dose, Hu08BBz CAR T cells mediated transient tumor regression until day 22, but the tumors progressed after this time point (Figure 4E). Among the minibody-secreting CAR T cell groups, only the anti-CTLA-4 minibody-secreting CAR T cells prolonged the Hu08BBz CAR T cell function and further inhibited the tumor growth (p = 0.0195) (Figure 4E), which was consistent with our in vitro results and in vivo results using systemic checkpoint blockade. These results not only demonstrated the feasibility of MiST but also confirmed the specificity of benefits from checkpoint blockades on CAR T cells.

IL-13Rα2 CAR T Cells Show Potent Anti-tumor Activity against Canine IL-13Rα2+ Tumors

Besides generating the D270 glioma cell line, we also harvested glioma tissues from surgical excision to generate glioma stem cell lines. We detected IL-13Rα2 expression on many of these lines (Figure 5A). The expression was heterogeneous as demonstrated by the percentage of target-positive cells and target expression level. Further considering about the potential on target off tumor toxicity and specific benefits from checkpoint blockades, prior to their use in human GBM patients, IL-13Rα2 CAR T cells should be evaluated.
in a clinically relevant, spontaneous, large animal model of human disease.

To this end, the domestic dog develops high-grade GBM that mimics the biology and clinical course of the human disease and has been suggested as a preclinical mode for GBM. Therefore, we sought to determine whether Hu07BBz and Hu08BBz CAR T cells recognize epitopes on canine IL-13R\(\alpha_2\). First, we compared the amino acid sequence of human IL-13R\(\alpha_2\) and canine IL-13R\(\alpha_2\), and we found them to share 71.6% sequence homology (Figure S8A). Next, we electroporated human T cells with Hu07BBz and Hu08BBz mRNA and confirmed the expression of both CARs on the T cell surface (Figure 5B). Electroporated Hu07BBz and Hu08BBz human CAR T cells were then co-cultured with human and canine IL-13R\(\alpha_2\) protein controlled with BSA. CD8 staining was used to distinguish CD4- and CD8-positive T cell groups on the x axis.

Intracellular cytokine (IFN\(\gamma\)) staining was performed after these CAR T cells were co-cultured with human and canine IL-13R\(\alpha_2\) protein. Surprisingly, only Hu08BBz CAR T cells were activated by canine IL-13R\(\alpha_2\). Neither Hu07BBz nor Hu08BBz CAR T cells were activated by BSA (negative control) (Figure 5B). We next investigated the expression of canine IL-13R\(\alpha_1\) and IL-13R\(\alpha_2\) mRNA level in a variety of canine tumor cell lines. All four of the canine osteosarcoma cell lines (BW-KOSA, CS-KOSA, MC-KOSA, and SK-KOSA) was detected with RT-PCR, controlled with canine GAPDH. The percentage of cytokine-positive (IFN\(\gamma\), IL-2, and TNF-\(\alpha\)) T cells in CD4- and CD8-positive T cell subgroups was analyzed for mRNA-electroporated IL-13R\(\alpha_2\)-targeting (Hu07BBz and Hu08BBz) human CAR T cells and un-transduced T cells after co-culture with canine tumor cell lines mentioned before. (D) Two million Hu08BBz-transduced human CAR-positive T cells were injected i.v. after 7 days of five million MC-KOSA subcutaneous implantation (n = 5 per group). Tumor size was caliperd and compared with the same amount of un-transduced T cell control group. Statistically significant difference of tumor growth was determined by linear regression. ***p < 0.0001. Data are presented as means ± SEM.
MC-KOSA, and SK-KOSA), while a lower percentage of CD4 and CD8 Hu08BBz CAR T cells produced these cytokines in response to GL-1, Cacal3, and Cacal5 tumor cell lines (Figure 5C). Cytokine production corresponded with the level of canine IL-13Ra2 expression in these tumor cell lines. Although Camac2 expressed canine IL-13Ra1, none of the CAR T cells co-cultured with these cells produced cytokines, demonstrating a lack of cross-reactivity of the IL-13Ra2 CAR T cells with canine IL-13Ra1. Furthermore, no cytokine-positive T cells were detected in the UTD T cell group and Hu07BBz CAR T cell group (Figure 5C).

Prior to testing the anti-tumor activity of Hu08BBz CAR T cells against IL-13Ra2+ tumors, we established a canine osteosarcoma model in NSG mice. Three different doses of canine osteosarcoma tumor cells were implanted subcutaneously into the right flank of NSG mice, and bioluminescence imaging was used to evaluate the tumor growth. The average radiation of the canine osteosarcoma mouse model with the MC-KOSA tumor cell line reached $1 \times 10^7$ p/s/cm²/sr and increased with time. The average radiation in the other canine osteosarcoma cell lines was much lower and did not consistently increase (Figure S8B). NSG mice implanted with $5 \times 10^5$ MC-KOSA tumor cells showed a significant difference in radiation compared with the other tumor cell groups (p < 0.0001). We chose the highest tolerated dose ($5 \times 10^6$) of MC-KOSA to establish IL-13Ra2+ osteosarcoma tumors in the NSG mice, and, 7 days after implantation, we intravenously administered $2 \times 10^6$ Hu08BBz CAR-transduced human T cells. Tumor growth was significantly inhibited in the CAR T cell treatment group compared to the UTD T cell treatment group (Figure 5D). These results highlighted the potential in generating canine CAR T cells to target canine IL-13Ra2-positive tumors.

**Canine IL-13Ra2 CAR T Cells Control Canine Tumor Growth**

In the next step toward evaluating IL-13Ra2 CAR T cells in dogs with spontaneous GBM, we generated canine IL-13Ra2 CAR T cells, and we evaluated their antigen-specific function in vitro and in vivo. We electroporated primary canine T cells with Hu08BBz CAR mRNA and co-cultured them with different canine tumor cell lines for testing their activation. Canine IFNγ was secreted by Hu08BBz canine CAR T cells, but not by UTD canine T cells, when co-cultured with IL-13Ra2-expressing tumor cells (all osteosarcoma cell lines plus Cacal5). Canine IFNγ was not secreted in response to IL-13Ra2-negative canine cell lines (Camac2 and CLBL-1). Interestingly, the canine glioma cell line, J3T, induced the greatest amount of IFNγ by canine Hu08BBz CAR T cells, reaching the maximum detectable limit in this assay ($1.26 \times 10^4$ pg/mL) (Figure 6A).

To mimic the physiological expression and stimulation status of canine T cells, we established a second-generation canine IL-13Ra2 CAR by switching the human CD8z domain and human 4-1BB and CD3ζ intracellular signaling domains with canine CD8z and canine 4-1BB and CD3ζ (Figure 6B). Canine IFNγ secretion was compared between Hu08-human-BBz (Hu08HuBBz) and Hu08-canine-BBz (Hu08CaBBz) canine T cells when they were co-cultured with canine tumor cell lines (CLBL-1 and J3T). Hu07-human-BBz (Hu07HuBBz) CAR T cells were included as a negative control. Canine T cells expressing either the Hu08HuBBz or Hu08CaBBz CAR produced IFNγ in response to the J3T tumor cell line, but not the CLBL-1 cell line, and no significant difference in IFNγ production was detected between the two CAR constructs (p = 0.2736) (Figure 6C).

Next, we orthotopically injected J3T glioma cells into mice to further evaluate the function of these CAR T cells in vivo. The J3T tumor cell line was transduced with the click beetle green luciferase gene for visualizing in the IVIS. After J3T implantation, $1.2 \times 10^7$ mRNA-electroporated Hu08HuBBz, Hu08CaBBz canine CAR T cells, or UTD canine T cells were injected intravenously by mouse tail vein on days 7, 10, and 13. Bioluminescence imaging was performed until day 41 after tumor implantation. Both Hu08HuBBz and Hu08CaBBz CAR T cells mediated prolonged inhibition of tumor growth when compared to UTD T cells (p < 0.0001; p = 0.0015) (Figure 6D). Canine T cells used in the second implantation were analyzed in vitro. Hu08HuBBz and Hu08CaBBz CAR constructs were detected on the surface of canine T cells (Figure 6E, left panels), although the expression of the canine CAR construct was less. Canine IFNγ production was also detected in the J3T-co-cultured canine CAR T cells (Figure 6E, right panels). Thus, we successfully generated canine IL-13Ra2-targeting CAR T cells for translational studies.

**DISCUSSION**

CAR T cells are a promising treatment method for cancer patients. CD19-targeting CAR T cell is the first gene therapy drug approved by the U.S. Food and Drug Administration (FDA).\textsuperscript{44} The data on treatment efficacy of CAR T cells are still limited for solid tumors. IL-13 zetakine-redirected T cells reversed the tumor growth and led to a clinical remission of 7.5 months in a recurrent multifocal GBM patient, demonstrating the potential of CAR T cell therapy for the solid tumors.\textsuperscript{11} However, IL-13 zetakine-redirected T cells are not as specific for IL-13Ra2 as scFv-based CAR T cells. Although the IL-13 molecule on the zetakine CAR structure was mutated to increase the affinity with IL-13Ra2, IL-13 zetakine-redirected T cells still displayed off-target effects, as they were stimulated by IL-13Ra1,\textsuperscript{18} which is extensively expressed in normal tissues (Figure S1A). Balyasnikova and colleagues\textsuperscript{49} established IL-13Ra2-targeting CAR T cells with a murine IL-13Ra2-targeting scFv (clone 47) to avoid cross-reactivity, but they raised the possibility of rejection effect from the murine scFv sequence, which comprised the majority of the extracellular domain of the CAR structure. In several clinical trials of murine scFv-based CAR T cell therapy, HAMA development was observed after CAR T infusion,\textsuperscript{8,20,21} resulting in a decrease in both persistence and function of CAR T cells in patients. Of note, a mesothelioma patient suffered a serious anaphylactic reaction leading to cardiac arrest in their third infusion of murine-based scFv CAR T cells.\textsuperscript{21} Humanizing the scFv of the CAR structure is important for the efficiency and safety of CAR T therapy.\textsuperscript{9,10} In an ALL CAR T retreatment clinical trial, 64% of murine CD19 CAR T cell-resistant patients had disease remission induced by humanized CD19 CAR T cells.\textsuperscript{45}
In our work, after establishing two murine scFv-based IL-13Rα2 CAR T cells (Mu07BBz/Mu08BBz), we tested the specificity and efficiency of Hu07 and Hu08 scFvs as CAR molecules. The humanized IL-13Rα2 CAR T cells did not cross-react with IL-13Rα1 on either normal human cells or tumor cells. They also demonstrated responses to the target cells. In mouse models, IL-13Rα2 CAR T cells not only inhibited tumor growth in all of the mice in the subcutaneous model but also reduced growth in 75% of the intracranially implanted, highly invasive D270 tumor model. Although IL-13Rα2-targeting antibody (clone 47) inhibited IL-13 binding with IL-13Rα2 and improved the survival of glioma xenograft mouse model, superior performance of the humanized IL-13Rα2 CAR T cells might be due to the high affinity of their scFvs. The K_D for the Mu07 and Mu08 are 0.648 and 0.964 nM, respectively, for IL-13Rα2. Humanized scFvs (Hu07 and Hu08) demonstrated comparable affinity with corresponding murine scFvs. 2173 was humanized from murine EGFRvIII-targeting scFv (3C10), the affinity of which is decreased during humanization to prohibit the cross-reaction with EGFR. The K_D for 3C10 and 2173 are 25.8 and 101 nM for EGFRvIII, which is lower than IL-13Rα2-targeting scFv in this experiment.

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Immune checkpoints are a series of molecules downregulating the function of T cells. They induce T cell tolerance of self-normal tissues.

Figure 6. Canine IL-13Rα2 CAR T Cells Control Canine Tumor Growth
(A) mRNA-electroporated Hu08BBz canine CAR T cells were co-cultured with canine tumor cell lines (Camac2, CLBL-1, GL-1, Cacal3, Cacal5, BW-KOSA, CS-KOSA, MC-KOSA, SK-KPSA, and J3T). Canine IFNγ secretion was detected with ELISA and compared the stimulation with un-transduced canine T cells. (B) Vector maps of anti-IL-13Rα2 human Hu08BBz CAR structure (Hu08HuBBz) and canine Hu08BBz CAR structure (Hu08CaBBz). (C) mRNA-electroporated Hu07HuBBz, Hu08HuBBz, and Hu08CaBBz canine T cells co-cultured with CLBL1 and J3T tumor cell lines. Canine IFNγ secretion was detected with ELISA. Unpaired t test was used to determine the statistically significant difference of IFNγ secretion between Hu08HuBBz and Hu08CaBBz co-cultured with J3T glioma cells. (D) J3T canine glioma cell line orthotopically implanted into the NSG mouse brain. 12 million electroporated Hu08HuBBz, Hu08CaBBz, or un-transduced canine T cells were i.v. injected into the mouse model (n = 4 per group) on days 7, 10, and 13 after tumor implantation. Tumor growth was evaluated by bioluminescence imaging every 3–4 days. Statistically significant differences of tumor growth were determined by linear regression. (E) The canine T cells used on the second injection on day 10 were analyzed for CAR expression and canine IFNγ secretion after co-culture with the J3T tumor cell line. Canine CD4 was stained to distinguish the canine CD4- and CD8-positive subgroups along the x axis. ns, not significant; **p < 0.01, ****p < 0.0001. Data are presented as means ± SEM.
and limit auto-inflammatory reactions by binding with their ligands, but also they raise the possibility of tumor tolerance, especially when tumors express the respective checkpoint ligands. The treatment effect of CAR T cell therapy is based on the function of cytotoxic T cells, which is also modulated by immune checkpoints. Compared to normal T cells, the stimulation of CAR T cells is more specific, mainly based on the CAR structure recognizing the specific epitope of the target protein, without dependence on the variability of major histocompatibility complex molecules. In a previous study, two CD19-targeting CARs with different hinges performed different levels of anti-tumor efficacy, accompanied with different levels of PD-1 and LAG-3 expression. Specific framework of scFv (e.g., IgG2) in the CAR was also prone to induce the expression of T cell exhaustion receptors (PD-1, TIM-3, and LAG-3). In this study, we included EGFRvIII- (2173) and IL-13Ra2- (Hu08) targeting CAR T cells with different scFvs, which displayed different levels of stimulation when co-cultured with the D270 tumor cell line. The expression levels of immune checkpoints (PD-1, CTLA-4, and TIM-3) were also different between the two CAR T treatments, consistent with the different stimulation levels. CTLA-4 was more prominently upregulated on Hu08BBz CAR T cells, while expression of PD-1 and TIM-3 was more prominent on 2173BBz CAR T cells.

Checkpoint blockade molecules were utilized to recruit T cell reservoir and release inhibitory T cell functions. For TCR gene modified specific targeting T cells, Moon and colleagues demonstrated intraperitoneal injection of PD-1 blockade enhanced the function of NY-ESO-1-targeting T cells to control tumor growth. In our study, we combined PD-1, CTLA-4, and TIM-3 checkpoint blockades with two different specificity CAR T cells (2173BBz and Hu08BBz) to explore the feasibility of this strategy. The treatment effect of CAR T cells was enhanced by checkpoint blockade in all groups but at different levels, with the exception of the 2173BBz combined with anti-CTLA-4, which demonstrated no increase in activity. Interestingly, anti-PD-1 and anti-TIM-3 treatments worked significantly better than the anti-CTLA-4 for the 2173BBz CAR T cells, whereas anti-CTLA-4 worked best for the Hu08BBz CAR T cells, which corresponded to the expression level of the checkpoint receptors when the CAR T cells were co-cultured with their target cells.

Although the checkpoints are known to downregulate T cell functions, they demonstrate varying effects with different ligands. CTLA-4 has a high degree of homology with the T cell costimulatory molecular CD28 and can outcompete CD28 for ligand binding (CD80 and CD86), which predominantly regulate ICOS+ CD4+ T cells and exhausted-like CD8+ T cells at the initiation of T cell activation. PD-1 is induced later during T cell activation and works on the CD8-positive T cells. We also detected an increasing frequency of human CD8+ T cells in the mouse spleen with anti-PD-1 intraperitoneal injection in this study. When PD-1 engages with its ligand (PD-L1 and PD-L2), the signaling mediated by the T cell receptor is attenuated by recruitment of tyrosine phosphatases. Hui and colleagues demonstrate that dephosphorylation of CD28 by PD-1-recruited Shp2 phosphatase primarily contributes to this function. TIM-3 is usually co-expressed with PD-1, which could explain both anti-PD-1 and anti-TIM-3 treatment demonstrating increased activity with the 2173BBz CAR T cells in this study. But unlike PD-1, there are no tyrosine-based inhibition motifs or tyrosine-based switch motifs on the TIM-3 molecule; human leukocyte antigen B-associated transcript 3 (Bat3) might be involved in its function when binding with galectin-9.

In this study, 2173BBz and Hu08BBz CAR T cells preferred different checkpoint blockades in the same tumor model. Although the same signaling domain (4-1BB and CD3ζ) should induce the same stimulation signal, the stimulation levels were significantly different in these two CAR T cells, which could explain the different responses observed with checkpoint blockades. This may also indicate which checkpoint blockades should be combined with specific CAR T cells. In addition, the expression levels of checkpoint molecules predicted the response level of these CAR T cells when combining with corresponding checkpoint blockades. In a phase 1 clinical trial performed in our group, after infusion of 2173BBz CAR T cells, PD-1 and PD-L1 were significantly upregulated in the tumor microenvironment, which also suggests that PD-1 checkpoint blockade may benefit 2173BBz CAR T cell therapy in vivo.

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While checkpoint blockades enhance the immune response in tumors, they also raise the possibility of auto-immune-related adverse events. There were 10%–15% of patients in a clinical study of anti-CTLA-4 treatment that had grade 3 or 4 immune-related adverse events, which led to patients dropping out of the study and requiring administration of high-dose systemic corticosteroids. Checkpoint blockade-secreting CAR T cells, not only avoid the adverse events triggered by systemic delivery but also increase the bioavailability of checkpoint blockades localized in the tumor. This is of particular importance al GBMs, which require drugs to pass through the blood-brain barrier to reach the tumor.

Overexpression of checkpoint blockades as intact antibodies (around 150 kDa) might increase the T cell burden, due to the increased synthesis of mRNA and protein as well as the competition of complexes and other molecules during these procedures. Li’s group showed that local delivery of an anti-PD-1 scFv by CD19-targeting CAR T cells enhanced the function of CAR T cells. Although an scFv is much smaller (around 28 kDa) and retains the binding affinity of an antibody, the clearance is also fast after infusion into the body with a half-life less than an hour. scFvs fused to human IgG1 CH3 (around 40 kDa) promote and stabilize dimerization and significantly prolong plasma half-life.

We used a similar structure to locally produce checkpoint blockades (anti-PD-1, anti-CTLA-4, and anti-TIM-3) in Hu08BBz CAR T cells in this study. Checkpoint blockade-secreting CAR T cells do not show a significant advantage of stimulation or cytokine secretion over non-checkpoint blockade-secreting CAR T cells in a 1- or 2-day co-culture experiment with target cells, which may be due to the cell burden of checkpoint blockade secretion. Regulating the
secretion of checkpoint blockades via the stimulation of CAR T cells might be a feasible strategy to decrease the cell burden. Among the checkpoint blockade-secreting CAR T cells, anti-CTLA-4–secreting Hu08BBz CAR T cells performed better than the other two CAR T cells, although in some groups there was no statistically significant difference. Anti-CTLA-4–secreting Hu08BBz CAR T cells were the only checkpoint blockade-secreting CAR T cells that demonstrated enhancement of the function of the Hu08BBz CAR T cells, which highlights the feasibility of this strategy and further confirms the beneficial effect of anti-CTLA-4 minibody with Hu08BBz CAR T cells.

Pet dogs develop spontaneous gliomas at an estimated incidence of 7 cases/100,000 dogs/year or approximately 4,000 spontaneous canine gliomas per year in the United States, which is similar to the incidence of human glioma. Canine gliomas exhibit comparable clinical, biological, histopathological, and genetic similarities to human gliomas, including invasive tumor growth with gross hemorrhage, necrosis and cyst formation, neovascularization, pleomorphism, endothelial cell proliferation, and T cell and macrophage infiltration. Furthermore, similar to human gliomas, canine gliomas frequently harbor p53 mutations and EGFR and IL-13Rα2 overexpression. Based on these similarities and the immunocompetent nature of pet dogs with spontaneous tumors, there is escalating interest in the use of dogs with spontaneous brain tumors to inform human clinical trial design. Given the comparable brain size between dogs and humans, ready access to state-of-the-art imaging modalities, such as computed tomography (CT) and MRI in specialized veterinary centers, and the high expression of IL-13Rα2 in canine gliomas, the development of canine IL-13Rα2-targeting CARs described here now sets the stage for the performance of a pilot clinical trial in canine glioma patients that aims to provide information on the safety and early effectiveness of this approach.

In conclusion, our study illustrated two humanized scFv-based IL-13Rα2-targeting CARs effectively controlled tumor growth in vitro and in vivo without cross-reaction with IL-13Rα1. Checkpoint blockades benefitted the function of IL-13Rα2 CAR T cells (Hu08BBz) as well as the humanized EGFRvIII CAR T cells (2173BBz), but these two types of CAR T cells required different checkpoint blockades, which can be predicted by the expression level of the corresponding checkpoint receptors and indicates the checkpoint blockades should be selectively chosen to combine with specific CAR T cells. We also in situ delivered checkpoint blockades as minibodies secreted by CAR T cells (MiSTs) to further confirm IL-13Rα2 CAR T cells (Hu08BBz) were selectively enhanced by CTLA-4 blockade. The secretion of minibodies could be further optimized by an inducible system rather than a constitutive expression system to decrease the burden of CAR T cells. We also generated canine IL-13Rα2-targeting CAR T cells and tested the function in vitro and in vivo. The canine IL-13Rα2-targeting CAR T cells will be used in the treatment of canine spontaneous GBM patients and in translational studies.

MATERIALS AND METHODS

Study Design

The aims of this study were to design fully humanized IL-13Rα2-specific targeting CAR T cells and test the possibility of combinational therapy with different checkpoint blockades by systemic and local delivery for potential use as a therapeutic agent in patients with IL-13Rα2-expressing tumors, such as malignant glioma. Canine IL-13Rα2-targeting CAR T cells are generated to treat canine malignancies expressing IL-13Rα2. In this study, two murine IL-13Rα2-targeting scFv and their humanized scFv CAR T cells were cloned and tested by co-culturing with tumor cell lines or human normal cell lines in vitro. The function of humanized IL-13Rα2-targeting CAR T cells are also tested by intravenous infusion into subcutaneous or orthotopic xenograft glioma mouse models. Expression of checkpoint receptors was detected after in vitro T cell stimulation. Tumor sizes were measured through caliper and compared between groups of IL-13Rα2-targeting (Hu08BBz) or EGFRvIII-targeting (2173BBz) CAR T cells combined with checkpoint blockade (anti-PD-1, anti-CTLA-4, and anti-TIM-3) in the glioma mouse model. IL-13Rα2-targeting (Hu08BBz) CAR T cells were modified to express these different checkpoint blockade minibodies to further explore the feasibility of combinational therapy by this strategy in vitro and in vivo. Canine IL-13Rα2-targeting CAR T cells were sorted out by co-culturing with canine IL-13Rα2 protein and confirmed by co-culturing with different canine tumor cell lines. Human and canine protein component-based canine
IL-13RA2-targeting CAR T cells were established and tested in a canine glioma orthotopic xenograft mouse model. Each experiment was performed multiple times with T cells derived from various normal donors.

**Cell Lines and Culture**

The human tumor cell lines (Sup-T1, Jurkat clone E6-1, A549, and 293T cells) and canine tumor cell lines (CLBL-1 and GL-1) were maintained in RPMI-1640 plus GlutaMAX-1, HEPES, pyruvate, and penicillin and streptomycin (Thermo Fisher Scientific) supplemented with 10% fetal bovine serum (FBS) (R10 media). U87 was purchased from the American Type Culture Collection (ATCC) and maintained in MEM (Richter & Co.) supplemented with penicillin and streptomycin (Thermo Fisher Scientific). Canine T cells were cultured with aAPCs at a 2:1 ratio to a final concentration of 1 × 10^6 canine T cells and 5 × 10^5 aAPCs per mL with 0.5 μg/mL mouse anti-canine CD3 (Bio-Rad) in RPMI-1640 plus GlutaMAX-1, HEPES, pyruvate and penicillin and streptomycin (Thermo Fisher Scientific) supplemented with 10% fetal bovine serum (FBS) (R10 media). Canine T cells were cultured with aAPCs at a 2:1 ratio to a final concentration of 1 × 10^6 canine T cells and 5 × 10^5 aAPCs per mL with 0.5 μg/mL mouse anti-canine CD3 (Bio-Rad) in R10 media with 30 IU/mL recombinant human IL-2 (rhIL-2, Proleukin, Chiron). Stably transduced human CAR T cells used in the in vivo study were normalized to 30% CAR+ before transplantation.

**Human T Cell Transduction and Culture In Vitro**

Human T cell transduction and culture were performed as previously described. Briefly, isolated T cells were derived from leukapheresis products obtained from the Human Immunology Core at the University of Pennsylvania using de-identified healthy donors under an institutional review board-approved protocol. T cells were stimulated with Dynabeads Human T-Activator CD3/CD28 (Life Technologies) at a bead-to-cell ratio of 3:1. After 24-hr stimulation, lentivirus was added into the culture media and thoroughly mixed to produce stably transduced CAR T cells. The concentration of the expanding human T cells was calculated on a Coulter Multisizer (Beckman Coulter) and maintained at 1.0–2.0 × 10^6 per mL in R10 media supplemented with 30 IU/mL recombinant human IL-2 (rhIL-2, Proleukin, Chiron). Stably transduced human CAR T cells used in the in vivo study were normalized to 30% CAR+ before transplantation.

**Canine T Cell Culture and Expansion In Vitro**

Canine T cells were collected from leukapheresis products obtained from peripheral blood of healthy research dogs at the University of Pennsylvania. Canine T cells were cultured and expanded with cell-based antigen-presenting cells (aAPCs) as described before. In brief, the human erythroleukemic cell line K562 transduced with lentiviral vector to stably express human FcγRII (CD32) and canine CD86 was used as artificial APCs, which were provided by Nicola Mason (School of Veterinary Medicine, University of Pennsylvania). Before expanding canine T cells, aAPCs were irradiated with 10,000 rad and washed with R10 media. Canine T cells were cultured with aAPCs at a 2:1 ratio to a final concentration of 1 × 10^6 canine T cells and 5 × 10^5 aAPCs per mL with 0.5 μg/mL mouse anti-canine CD3 (Bio-Rad) in R10 media with 30 IU/mL rhIL-2. The concentration of the expanding canine T cells was calculated on a Coulter Multisizer (Beckman Coulter) and maintained at 1.0–2.0 × 10^6 per mL R10 media with rhIL-2.

**mRNA In Vitro Transcription and Electroporation**

RNA was synthesized and electroporated as previously described. Briefly, pGEM plasmids were linearized by digestion with SpeI.
mRNA in vitro transcription was performed using the T7 mScript Standard mRNA production system (CellScript) as per the manufacturer’s instructions to obtain capped and tailed mRNA. Production was aliquoted and stored at −80°C until use. Expanded T cells were washed three times with Opti-MEM media (Gibco) and resuspended at 1 × 10⁸ cells/mL. 10 mg mRNA was mixed with 1 × 10⁷ T cells and moved into cuvettes for electroporation. After electroporation with 500 V for 700 μs, T cells were recovered in the R10 media with rhIL-2.

**Flow Cytometry**

For CAR detection, cells were stained with biotinylated protein L (GenScript), goat anti-mouse IgG, and rabbit anti-mouse and anti-human IgG (Jackson ImmunoResearch Laboratories), and secondary detection was carried out by the addition of streptavidin-coupled phycoerythrin (PE) or fluorescein isothiocyanate (FITC) (BD Biosciences). Before and after each detection, cells were washed three times with PBS containing 2% fetal bovine serum (FBS) twice and resuspended at 10⁷ cells/mL. 10 mg mRNA was mixed with 10⁶ target cells after tumor implantation, based on the dosage applied in the other experiments. Cells were added in a volume of 100 μL at different E:T ratios (1:1, 3:1, 10:1, and 30:1). Effector and targets were incubated together for 4 h at 37°C. Supernatant from each well was collected and transferred onto the filter of a LumaPlate. The filter was allowed to dry overnight. Radioactivity released in the culture medium was measured using a β-emission reading liquid scintillation counter. Percentage specific lysis was calculated as follows: (sample counts – spontaneous counts)/(maximum counts – spontaneous counts) × 100.

**Chromium Release Assays**

Cytotoxicity of the CAR-expressing T cells was tested in a 4-hr ⁵¹Cr release assay, which was described. 1 × 10⁶ target cells were labeled with radioactive ⁵¹Cr (50 μCi) for 1 h at 37°C. After labeling, cells were washed with 10 mL non-phenol red RPMI medium plus 5% FBS twice and resuspended at 1 × 10⁶ cells/mL. 5,000 (100 μL) labeled target cells was plated in each well of a 96-well plate. Effector cells were added in a volume of 100 μL at different E:T ratios (1:1, 3:1, 10:1, and 30:1). Effector and targets were incubated together for 4 h at 37°C. Supernatant from each well was collected and transferred onto the filter of a LumaPlate. The filter was allowed to dry overnight. Radioactivity released in the culture medium was measured using a β-emission reading liquid scintillation counter. Percentage specific lysis was calculated as follows: (sample counts – spontaneous counts)/(maximum counts – spontaneous counts) × 100.

**Mouse Models**

All mouse experiments were conducted according to IACUC-approved protocols and described. For orthotopic models, 2 × 10⁶ D270 cells or J3T cells were implanted intracranially into 6- to 8-week-old female NSG mice (JAX). Tumor progression was also evaluated by luminescence emission on a Xenogen IVIS spectrum after intraperitoneal D-luciferin injection according to the manufacturer’s directions (GoldBio). Anti-PD-1, anti-CTLA-4, and anti-TIM-3 checkpoint blockades (BioLegend) were intraperitoneally injected 200 μg per mouse every 4 days from day 6 after tumor implantation, based on the dosage applied in the other studies. Survival was followed over time until the predetermined IACUC-approved endpoint was reached.

**RT-PCR**

cDNA of canine tumor cell lines was synthesized with reverse transcription kit from the extracted RNA. Phusion polymerase
(New England Biolabs) was used to amplify DNA fragments. Reaction was set up as indicated in the PCR protocol for Phusion polymerase. Primers designed for the experiments are as follows: IL-13Rα1 forward 5'-CAATGTGACGCTCAAGCCTTC-3', reverse 5'-GCTGTACGTAGCTGAACTAC-3'; IL-13Rα2 forward 5'-CTATGCCAGAGCTTATTAGC-3', reverse 5'-GATTCGCTGAATAGCCCTTTC-3'; and GAPDH forward 5'-GCCATCAATGACCCCTTCATTGATC-3', reverse 5'-CTATGCCACCAGACTACCTTAGTC-3'. After 35 cycles of reaction, PCR products were run on a 1% agarose gel and visualized in a gel documentation system (GDS touch, ENDURO).

ELISA
For detecting anti-PD-1 and anti-CTLA-4 minibodies, T cells were transduced and maintained as described above, between 1.0–2.0 × 10^6 cells/mL. 70 mL supernatant from day 11 of T cell expansion in vitro was collected and concentrated with Centricon Plus-70, as per the manufacturer's instructions. A standard direct ELISA was performed with DuoSet Ancillary Reagent Kit 2 (R&D systems). After coating with recombinant human PD-1 and CTLA-4 protein (Abcam), a 96-well plate was loaded with the concentrated supernatants followed by peroxidase goat anti-human IgG (Jackson ImmunoResearch Laboratories) detection antibody. For detecting canine IFNγ, supernatant was collected from canine T cell and target cell 16-hr co-culture at 1:1 ratio. The detection was performed with canine IFN-gamma DuoSet ELISA kit (R&D Systems) as the introduction indicated.

2-Photon Microscopy
Mice were anaesthetized and maintained at a core temperature of 37°C. Thinned-skull surgery was performed as described previously.44 For ex vivo imaging, as described before,63 CellTrace Violet (Life Technologies) and TRITC- (Thermo Fisher Scientific) labeled CAR T cells were intravenously transplanted; 4 hr later, the mice were euthanized, and the spleen was removed immediately and placed in a heated chamber where specimens were constantly perfused with warmed (37°C), oxygenated medium (phenol red-free RPMI 1640 supplemented with 10% FBS, Gibco). The temperature in the imaging chamber was maintained at 37°C using heating elements, and it was monitored using a temperature-control probe (Fine Science Tools). Imaging was performed with a Leica SP5 two-photon microscope system (Leica Microsystems) equipped with a picosecond or femtosecond laser (Coherent). Images were obtained using a 20× water-dipping lens. The resulting videos were analyzed with Volocity software (PerkinElmer).

Statistical Analysis
Data are presented as means ± SEM. Cytotoxicity assays, intracellular cytokine analysis, and MFI results of flow cytometry; and bioluminescence imaging. Survival curves were analyzed with Kaplan-Meier (log-rank test). All statistical analyses were performed with Prism software version 7.0 (GraphPad).

SUPPLEMENTAL INFORMATION
Supplemental Information includes eight figures and two videos and can be found with this article online at https://doi.org/10.1016/j.omto.2018.08.002.

AUTHOR CONTRIBUTIONS

CONFLICTS OF INTEREST

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