Bioactivity and Safety of IL13Rα2-Redirected Chimeric Antigen Receptor CD8+ T Cells in Patients with Recurrent Glioblastoma

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Abstract

Purpose: A first-in-human pilot safety and feasibility trial evaluating chimeric antigen receptor (CAR)–engineered, autologous primary human CD8+ cytotoxic T lymphocytes (CTL) targeting IL13Rα2 for the treatment of recurrent glioblastoma (GBM).

Experimental Design: Three patients with recurrent GBM were treated with IL13(E13Y)-zetakine CD8+ CTL targeting IL13Rα2. Patients received up to 12 local infusions at a maximum dose of 10^6 CAR-engineered T cells via a catheter/reservoir system.

Results: We demonstrate the feasibility of manufacturing sufficient numbers of autologous CTL clones expressing an IL13(E13Y)-zetakine CAR for redirected HLA-independent IL13Rα2-specific effector function for a cohort of patients diagnosed with GBM. Intracranial delivery of the IL13-zetakine CD8+ CTL clones into the resection cavity of 3 patients with recurrent disease was well-tolerated, with manageable temporary brain inflammation. Following infusion of IL13-zetakine CD8+ CTLs, evidence for transient anti-glioma responses was observed in 2 of the patients. Analysis of tumor tissue from 1 patient before and after T-cell therapy suggested reduced overall IL13Rα2 expression within the tumor following treatment. MRI analysis of another patient indicated an increase in tumor necrotic volume at the site of IL13-zetakine T-cell administration.

Conclusions: These findings provide promising first-in-human clinical experience for intracranial administration of IL13Rα2-specific CAR T cells for the treatment of GBM, establishing a foundation on which future refinements of adoptive CAR T-cell therapies can be applied.

Introduction

Despite aggressive standard-of-care therapies, including surgery, radiation, and chemotherapy, glioblastoma (GBM) remains one of the most universally lethal human cancers, with a 5-year survival rate of less than 10% (1). Inevitable treatment failure for most patients is largely attributable to therapy-resistant invasive malignant cells that are responsible for tumor recurrence. Adoptive cell therapy (ACT) with T cells genetically modified to express chimeric antigen receptors (CAR) is a promising therapeutic approach that may be effectively and safely applied to GBM to reduce recurrence rates (2). CAR T cells can be rapidly generated to specifically recognize antigenically distinct tumor populations independently of pre-existing antitumor immunity (3–6), and T cells can migrate through the brain parenchyma to target and kill infiltrative malignant cells (7–9). Recent clinical successes with CAR-engineered T cells mediating durable and complete responses in patients diagnosed with CD19+ malignancies, even under conditions of bulky disease, provide support for further development of this therapeutic approach (10, 11). However, broader clinical application to solid tumors, including brain tumors, has proven to be complex and is presently under intense investigation.

Our group has focused on a T-cell immunotherapy for GBM targeting IL13 receptor α2 (IL13Rα2), a monomeric high-affinity IL13 receptor that is overexpressed by more than 50% of GBM and is a prognostic indicator of poor patient survival (12–15). In the context of molecular subtypes of high-grade gliomas, IL13Rα2 expression is most closely associated with expression of mesenchymal signature genes, which may reflect its association with the proinflammatory characteristics of mesenchymal tumors (12, 16). IL13Rα2 is expressed by both stem-like and more...
Activity and Safety of IL13Rα2-Specific CAR T Cells in GBM

Translational Relevance
Standard-of-care therapies for glioblastoma (GBM), the most aggressive and common type of glioma, have resulted in an unacceptably poor median overall survival of approximately 15 months following primary diagnosis. Immunotherapeutic targeting of brain tumors offers the opportunity to redirect the potency and specificity of the immune system to improve therapeutic outcomes. However, clinical realization of this potential is still forthcoming. Here, we report a first-in-human pilot safety and feasibility study evaluating chimeric antigen receptor (CAR)–engineered IL13Rα2-specific primary human autologous CD8⁺ cytolytic T lymphocytes for the treatment of recurrent GBM. We provide evidence for feasibility of this treatment approach, and for transient anti-glioma activity in the absence of serious adverse events, thus establishing the foundation for further development of this IL13Rα2-specific CAR T-cell therapy.

differentiated malignant cells (5), as well as by tumor-infiltrating macrophages/myeloid-derived suppressor cells (17). Importantly, IL13Rα2 is not expressed at significant levels on normal brain tissue (15, 18–20), and early phase clinical trials support the safety and tolerability of targeting IL13Rα2 for the treatment of GBM with vaccine therapy (21) and IL13 immunotoxins (22), thereby qualifying IL13Rα2 as an attractive immunological therapeutic target.

We have developed an IL13Rα2-specific, MHC-independent CAR, termed IL13-zetakine. This CAR recognizes IL13Rα2 via a membrane-tethered IL13 ligand mutated at a single site (E13Y) to reduce potential binding to the more widely expressed membrane-tethered IL13 ligand mutated at a single site (E13Y) mutein, human IgG4 Fc linker, human CD4 transmembrane domain, and the cytoplasmic domain of human CD3 zeta.

Eligible patients were adults (18–70 years) with recurrent or refractory unifocal supratentorial grade III or IV glioma whose tumors did not show communication with ventricles/CSF pathways and were amenable to resection. Patients were required to have a survival expectation of greater than 3 months, a Karnofsky performance status (KPS) equal to or greater than 70, to be steroid independent, and to have completed primary therapy (≥ 2 weeks) of remission at limiting dilution to isolate single CD8⁺ T-cell clones. Patients were enrolled following initial diagnosis of high-grade glioma (WHO grade III or IV), at which time they underwent leukapheresis for collection of peripheral blood mononuclear cells (PBMC). These cells were used to engineer CD8⁺ CTLs to express the IL13-zetakine CAR and the ancillary HyTK selection/suicide fusion protein (23). Subsequently, the release-tested IL13-zetakine/HyTK cells were cryopreserved and stored for later use. At the time of first recurrence, the research participant underwent resection of tumor along with placement of a Rickham reservoir/catheter. Concurrently, the therapeutic clone was thawed, re-expanded in vitro using rapid expansion method (REM) stimulation (25). Following recovery from surgery and after baseline MR imaging, the IL13-zetakine⁺ CD8⁺ CTLs were administered directly into the resection cavity via the indwelling catheter (Supplementary Fig. S1 and Supplementary Methods). Cells were manually injected into the Rickham reservoir using a 21-gauge butterfly needle to deliver a 2 mL volume over 5 to 10 minutes, followed by 2 mL flush with preservative-free normal saline over 5 minutes.

The protocol treatment plan specified an intrapatient dose escalation schedule with a target of 12 CAR T-cell doses administered intracranially over a 5-week period comprised of weekly treatment cycles (Fig. 1A). During cycles 1, 2, 4, and 5, T-cell infusions were performed on days 1, 3, and 5 of the cycle week, and week 3 was a rest cycle. For safety, in cycle 1, we utilized an intrapatient dose escalation strategy, with CAR T-cell doses of 10⁷, 5 × 10⁷, and 10⁸ cells per infusion administered on days 1, 3, and 5, respectively, and this was followed by 9 additional CAR T-cell infusions of 10⁸ cells over 4 weeks. Imaging to assess response was performed during the week 3 rest cycle and after week 5. The guidelines provided in the NCI Common Toxicity Criteria version 2.0 (http://ctep.cancer.gov/protocolDevelopment/electronic_applications/docs/ctcmanual_v4_10_4-99.pdf) were followed for the monitoring of toxicity and adverse event reporting.

Clinical IL13-zetakine CAR vector and T-cell manufacturing
A schematic of the T-cell manufacturing process is provided in Fig. 1B. The plasmid vector, encoding the IL13-zetakine CAR and the selection/suicide HyTK transgenes, has been described previously (23). Briefly, the IL13-zetakine CAR is composed of the human GM-CSF receptor alpha chain leader peptide, human IL13 (E13Y) mutein, human IgG4 Fc linker, human CD4 transmembrane domain, and the cytoplasmic domain of human CD3 zeta.

Materials and Methods
Study design and research participants
This single-institution first-in-human pilot safety and feasibility study was conducted at the City of Hope National Medical Center. All participating patients gave written informed consent. The clinical protocol was approved by the City of Hope Institutional Review Board, conducted under an Investigational New Drug Application (IND 10109), and registered at ClinicalTrials.gov (NCT00730613).

Eligible patients were adults (18–70 years) with recurrent or refractory unifocal supratentorial grade III or IV glioma whose tumors did not show communication with ventricles/CSF pathways and were amenable to resection. Patients were required to have a survival expectation of greater than 3 months, a Karnofsky performance status (KPS) equal to or greater than 70, to be steroid independent, and to have completed primary therapy (≥ 2 weeks) of remission at limiting dilution to isolate single CD8⁺ T-cell clones. Subsequently, CAR T-cell clones were expanded using a 14-day cycle of REM consisting of OKT3, rHL-2 (Novartis Oncology), and irradiated feeders as previously described (25), the only exception...
Irradiated feeders were authenticated for viability, growth, sterility, and lack of mycoplasma prior to cryopreservation as a GMP bank, and thawed cells were cultured for less than 6 months prior to use in a REM cycle. A summary of the quality control tests performed and the requisite test results for T-cell product release are summarized in Supplementary Table S1 and Supplementary Methods.

Clinical imaging

MRI scans of the brain, including fluid attenuated inversion recovery (FLAIR) and post-gadolinium T1-weighted sequences, were acquired on a GE Signa Excite scanner (1.5 Tesla). Regions of contrast-enhancing or necrotic tumor were outlined by a radiologist, and the corresponding volumes were automatically determined using a GE AW workstation running AW version 4.3 software. Imaging with 18F-fluorodeoxyglucose (18F-FDG) was performed using a GE Advance-NXi PET scanner (15 cm axial field of view, slice separation 4.3 mm) or a GE Discovery Ste 16 PET-CT scanner (15 cm axial field of view, slice separation 3.3 mm).

Protein and mRNA analyses

IL13R2 immunohistochemistry (IHC) was performed on 5-mm sections of formalin-fixed paraffin-embedded specimens as previously described (5). The tumor regions (tumor cells >60%) were determined by a clinical neuropathologist. For Fig. 2A and B, IL13R2 expression was calculated for defined tumor areas as integrated OD (lumen x pixel²)/C2 pixel² using the DAB plug-in for ImagePro Premier v9.1 (Media Cybernetics). For Fig. 4C, IL13R2 expression levels were measured in individual pixels using the

**Figure 2.**

Tumor IL13R2 expression for each patient. A, immunohistochemical staining of IL13R2 on paraffin-embedded primary patient-derived brain tissues. Sections were scored blindly by a neuropathologist for staining intensity (0, not detected; 1−, low; 2+, moderate; 3+, high), and percentages of positive cells are indicated. B, quantification of DAB intensity (lumen x pixel²) for dense tumor regions (>60% tumor, red outline). C, gene expression of IL13R2 mRNA levels evaluated by qPCR using Taqman gene expression assay.

being the use of 0.2 mg/mL hygromycin in place of G418.
The primary objective of this study was to assess the feasibility and safety of intracranially administered autologous IL13Rzeta-kine-CD8\(^+\) T-cell clones for the treatment of recurrent high-grade glioma (WHO grade III and IV) following tumor resection. Cell products meeting all quality control tests (Supplementary Table S1) were generated for 10 of the 13 patients enrolled on this study; three T-cell products (UPN024, UPN029, UPN031) did not complete product release due to patient disease progression prior to completion of CAR T-cell manufacturing (Supplementary Table S2). Of the 10 remaining enrolled patients, three (UPN025, UPN027, UPN030) did not meet T-cell infusion eligibility criteria due to multifocal disease, death, and steroid dependence, respectively, and therefore did not receive their IL13-zetakine\(^+\) CTL product. An additional 3 patients (UPN026, UPN034, UPN038) did not develop disease progression before the protocol was closed to treatment (Supplementary Table S2), and 1 patient (UPN036) voluntarily withdrew from the study at the time of progression. The 3 remaining patients (UPN028,
UPN013, UPN031, UPN033) were treated on this study. An overview of manufactured products and patient treatment/nontreatment rationale is provided in Supplementary Table S2 and Supplementary Fig. S2. For the 3 patients that were treated on this trial, patient characteristics, including diagnosis and prior therapy, are summarized in Supplementary Table S3. For these 3 patients, analyses of excised tumors collected at the time of Rickham catheter placement revealed a range of IL13Rα2 antigen expression levels as determined by both IHC (Fig. 2A and B) and qPCR (Fig. 2C), with UPN028 exhibiting the lowest, UPN033 intermediate, and UPN031 the highest level of antigen expression. Of interest, our previous studies showed that low-passage stem-like or differentiated glioma cell lines derived from IL13Rα2-positive (by IHC) UPN033 maintained IL13Rα2 expression, and were recognized and killed by this patient’s autologous IL13-zetakine CTLs, thus suggesting that the differentiation status of malignant cells in the tumor did not impact IL13Rα2 antigen expression or targetability (5).

Safety of IL13-zetakine CAR T-cell administration

The intrapatient dose escalation schedule depicted in Fig. 1A was executed as planned for 2 of 3 patients (UPN031 and UPN033), who received a full course of 12 escalating intracavitary doses of 10^8 to 10^10 IL13-zetakine CTL. The other patient (UPN028) received all but one 10^6 cell dose at day 1 of cycle 3 due to transient worsening of a headache (for individual patient treatment regimens, see Supplementary Fig. S3). There were no Grade 3 or higher adverse events with possible correlation to administration of either 10^6 or 5 × 10^7 T cells. However, at the 10^8 T-cell dose, two cases of Grade 3 headache occurred in one subject (UPN028), which were possibly attributable to T-cell administration (Table 1). There was also one Grade 3 neurologic event (UPN031), which included shuffing gait and tongue deviation, possibly attributable to T-cell administration (Table 1). This neurologic event occurred the day after the 12 and final CAR T-cell infusion, required hospital admission, and was treated with a single infusion of 10 mg intravenous dexamethasone. At follow-up 4 days later, the patient remained stable and did not require additional dexamethasone for this adverse event. While transient neurologic worsening was anticipated in this protocol, it is possible that the high level of IL13Rα2 expression (Fig. 2) in this subject (UPN031) elicited a T-cell response that contributed to these neurological symptoms. Alternatively, tumor recurrence in UPN031, confirmed with a third surgery 14 weeks after the T-cell treatment, may also have contributed to the development of these neurologic adverse events. In contrast, UPN033, whose GBM expression of IL13Rα2 was also documented by IHC and qPCR (Fig. 2), did not exhibit any adverse events of Grade 3 or higher attributable to T-cell administration (Table 1). In summary, we conclude that repetitive dosing of up to 12 infusions of IL13Rα2-specific CAR CD8+ T-cell clones, with cumulative doses of 10.6 × 10^8 T cells administered into the resected tumor cavity, was well-tolerated and exhibited an acceptable safety profile with limited transient adverse events.

For all patients, adoptive transfer of autologous IL13-zetakine CTL clones resulted in brain inflammation at the site of T-cell infusion, as detected by increased MRI Gd-enhancement and increased signal on FLAIR images. This increased signal change is believed to be a consequence of the therapeutic cells and not indicative of disease progression, since the MRI signal change was observed immediately following the first two cycles of six total infusions for all patients, and was transient, subsiding within a few months of T-cell infusion for UPN028 and UPN033, who did not develop tumor recurrence at the treatment site (Fig. 3A and B). For UPN031, whose tumor had the highest IL13Rα2 tumor expression, the inflammatory reaction to T-cell therapy was more pronounced than in the other two subjects (Fig. 3C). Again, these early MRI enhancements in UPN031 were most consistent with T-cell–related inflammation and not tumor recurrence, since regions of enhancement showed low PET activity (Supplementary Fig. S4A). However, in contrast with the other patients, the central nervous system (CNS) inflammatory reaction for UPN031 did not substantially subside, and disease recurrence was confirmed following a third craniotomy. Within these 3 patients, the degree of brain inflammation appeared to correlate with IL13Rα2 antigen expression, since inflammation was most pronounced in the 2 patients with the highest expression of IL13Rα2 (UPN031, UPN033). While this inflammation may have resulted from other factors, such as the infusion vehicle or nonspecific T-cell activity, these factors were similar in all 3 patients.

Treatment of UPN033 was previously published in an article describing the use of 18F-FHIBG PET to detect therapeutic CTLs via the HSV tk gene included as part of the IL13-zetakine/HyTK expression construct (9). As we previously noted for this patient, following CAR T-cell administration into the resection cavity a 18F-FDG PET-avid, MR-enhancing lesion was detected contralaterally across the splenium of the corpus callosum rather than adjacent to the CAR-treated tumor cavity (Figs. 3B and 5A; and Supplementary Fig. S4B). Although the inflammatory reaction surrounding the resection cavity subsided over time, the lesion in the corpus callosum continued to progress. UPN033 was, therefore, taken off protocol 20 days after the last CAR T-cell infusion, and received additional radiation, bevacinuzumab, and BCNU over a 5-month period (Supplementary Fig. S3D). Following further

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^aOnly events of Grade 3 or higher, according to the NCI Common Toxicity Criteria, with possible or higher attribution to the T-cell administration are reported.

^bShuffling gait and tongue deviation to the left.

^cT cells administered at secondary site of recurrence.

^dSurvival after second biopsy/relapse detected on day 64 as related to timeline in Figure S3D was 11.8 months.
tumor progression, this patient was enrolled on a separate single subject protocol in which the enhancing lesion was biopsied to confirm tumor recurrence, and IL13-zetakine⁺ CTL clones were administered directly into the tumor. In this second protocol, UPN033 received a series of five daily infusions of autologous IL13-zetakine⁺ CTL with escalating intratumoral doses ranging from $2.5 \times 10^7$ to $1 \times 10^8$ cells delivered directly into the recurrent tumor site via an externalized catheter (2 mL of cells were infused over 4 hours; Supplementary Fig. S3D). Grade 3 or higher adverse events that were possibly associated with T-cell administration included transient leukopenia, headaches during the infusion period, and fatigue (Table 1), again indicating an acceptable safety profile.

Evidence for antitumor bioactivity following adoptive transfer of IL13-zetakine⁺ CTLs

Decreased tumor IL13Rα2 expression following therapy. Of the 3 patients treated on NCT00730613, the tumor excised from UPN031 prior to therapy had the highest IL13Rα2 expression (Fig. 2). As mentioned above, this patient required another craniotomy for tumor resection approximately 14 weeks after CAR T-cell therapy. In recurrent tumor tissue adjacent to the T-cell injection site, IL13Rα2 expression was significantly decreased from pre-T-cell therapy levels. This was initially observed by flow cytometry of freshly excised tumor (Fig. 4A) and confirmed by both qPCR analysis of tumor samples (Fig. 4B) and IHC staining of paraffin-embedded histologic sections (Fig. 4C). While in this single patient we cannot rule out confounding contributions of local sampling within a heterogeneous tumor, we suggest that autologous IL13-zetakine⁺ CTL successfully targeted IL13Rα2-expressing tumor cells and reduced overall IL13Rα2 expression. These data thus provide corroborating evidence for an IL13-zetakine⁺ CTL-mediated antitumor response, with treatment failure in this patient due to recurrence/progression of IL13Rα2-negative/low tumor cells. Tumor recurrence could also be impacted by lack of significant expansion and/or survival of therapeutic CAR T cells, as indicated by our ability to specifically detect only low levels of the adoptively transferred T cells in the recurrent tumor.
targeting of IL13Rα2-expressing tumor cells. A, T1-weighted postadministration of IL13-zetakine on day 314. B, MRS analysis of enhancing volume (indicative of neuroinflammation) and necrotic volume at the recurrent site following retreatment with T cells. Lack of tumor recurrence at the original treatment site (right occipital) and increase in necrosis at the recurrent site (left corpus callosum, white circles in A) are highly suggestive of therapeutic activity. C, single voxel MR spectroscopy with a pane over the lesion medial to the atrium of the left lateral ventricle on day 314. Choline (Cho), creatine (Cr), N-acetyl-l-aspartate (NAA), and lactate/lipid (Lac/Lip) peaks are indicated.

Retreatment of UPN033, comprising five daily infusions of IL13-zetakine CD8+ CTLs directly into the recurrent tumor mass within the splenium of the left corpus callosum, provided an opportunity to evaluate the therapeutic response of CAR T cells within an intact tumor. This recurrent lesion had been treated with radiation, chemotherapy (BCNU), and antiangiogenic therapy (bevacizumab) 1.5 to 3 months prior to T-cell infusion (~89 days for radiation, and ~44 days for bevacizumab/BCNU). Immediately following T-cell administration, there was a transient inflammatory response on MRI, as well as an increase in necrotic volume which persisted for several months (Fig. 5A and B). Although necrosis is a feature of GBM, the rapid onset of the inflammatory response, along with the persistent necrosis observed shortly after T-cell infusion, suggests that these responses were T-cell mediated rather than due to tumor progression or previous treatments. Further, single voxel MR spectroscopy over the lesion 88 days after the final T-cell infusion (day 314) showed a significant elevation of the lactate and lipid peaks, confirming necrosis, together with very little elevation of the choline peak relative to the creatine peak, suggesting low to negligible tumor burden (Fig. 5C). Collectively, these observations are highly suggestive of antitumor activity of IL13Rα2-specific CAR T-cell therapy.

Discussion

In this study, we present the first-in-human evaluation of local administration of CAR T cells to treat high-grade glioma. CAR T-cell therapies have demonstrated remarkable clinical success in the treatment of CD19+ hematological malignancies, with complete response rates reported for the majority of patients with acute lymphoblastic leukemias (10, 26). A major goal for this field of research is the successful application of CAR T-cell therapy to a range of solid tumors. Achieving this goal will require overcoming critical barriers to therapeutic success, including, but not limited to, tumor antigen selection, tumor heterogeneity, and the immunosuppressive tumor microenvironment. Our group is focusing on the development of CAR T-cell therapy for the treatment of high-grade glioma, a tumor class that is essentially incurable with current standard-of-care therapies. While preclinical studies had demonstrated that CAR T cells can have potent anti-tumor responses in orthotopic mouse models (5, 23), this approach had yet to be validated in a patient context. Here, we summarize our experience treating 3 patients with IL13Rα2-specific CAR T cells for recurrent GBM. We demonstrate feasibility and safety of repetitive intracranial delivery of autologous IL13-zetakine CD8+ CTL clones, and further report findings of transient anti-tumor activity for some patients in the absence of serious adverse events.

While CAR T-cell immunotherapy has the potential to improve treatment outcomes for patients with high-grade glioma, one of the major challenges for glioma immunotherapy is the identification of tumor-specific target antigens. The ideal antigen would be expressed by the majority of tumor cells, but absent from normal brain tissue, thus reducing the potential for off-tumor CNS toxicities. Promising cell surface markers that are being pursued for development of CAR-based T-cell therapy to treat...
high-grade glioma include IL13Rα2, HER2, EphA2, and EGFRvIII (3, 4, 6, 23). In this study, we focused on IL13Rα2, as previous studies have shown that IL13Rα2 is overexpressed by greater than 50% of GBMs, its expression increases with malignancy grade, and it is associated with decreased long-term survival (12, 15). Further, IL13Rα2 is expressed independently of tumor differentiation status, having been identified on stem-like malignant cells and more differentiated counterparts (5). Importantly, IL13Rα2 is not expressed at significant levels by normal brain tissue (15, 19, 20). The clinical experience treating 3 patients reported here further validates IL13Rα2 as a suitable target molecule for CAR T-cell immunotherapy by establishing patient tolerance and acceptable safety profiles for repetitive intracranial infusions of IL13Rα2-specific CAR T cells, with adverse events being transient and manageable. Further, we provide proof-of-principle that CAR T-cell targeting of a documented IL13Rα2-positive tumor can elicit tumor regression.

For this trial, autologous IL13-α2-zetakine® CD8+ CTL clones were manufactured for each patient using a process that included DNA electroperoration, followed by drug selection and ex vivo expansion using OKT3 and cell feeders. This process was feasible, with IL13-α2-zetakine® CD8+ T cell clones successfully generated for all patients (10 of 13) who did not progress within the manufacturing time-frame; however, these methodologies were cumbersome and took over 3 to 4 months to generate the final therapeutic product for each patient. To accommodate this delay, patients on this trial were enrolled following their primary diagnosis to give sufficient time for T-cell manufacturing prior to disease progression/recur. The overall time needed for clinical manufacture becomes an important issue for patients with recurrent GBM, who have a very limited therapeutic timeline. One strategy that could expedite CAR T-cell therapy is to develop an "off-the-shelf" allogeneic CAR T-cell product that would be available for administration shortly after tumor recurrence (27). Our group, in collaboration with Sangamo Biosciences, has recently shown the feasibility of infusing allogeneic IL13-α2-zetakine® CAR T cells while maintaining patients on steroids to reduce the possibility of CAR T-cell rejection (manuscript in preparation). Alternatively, newer manufacturing strategies using CD3/CD28 beads have reduced manufacturing times to less than 2 weeks, suggesting that, with these further improvements, enrollment at time of recurrence would be practical for autologous CAR T-cell therapy.

Route of delivery is an additional unresolved issue related to the application of CAR T-cell therapy for brain tumors. In this pilot safety and feasibility study, we delivered the CAR T cells intracranially via an implanted reservoir/catheter system. This enabled repetitive dosing of the CAR T cells directly into the resected tumor cavity, and, for the second treatment of UPN033, the site of recurrent tumor. Repetitive delivery of up to 1.6 × 10^9 total CAR T cells was well-tolerated by all 3 patients, as was repetitive local delivery into the site of active disease for UPN033. Further, there were no immediate or acute device-related adverse events, such as occlusion, malfunction, or infection.

For the optimal therapeutic response, it is imperative that the CAR T cells are able to migrate to distant sites of infiltrative and/or multifocal disease. The delivery strategy best able to target infiltrative and/or multifocal disease is not yet evident. Although the standard imaging modalities reported here could not evaluate T-cell trafficking using the reservoir/catheter system, our collaborators on this clinical study have previously provided evidence, using 18F-FHBG PET, that CAR/HyTK T cells were detected at the site of injection, as well as at a recurrent secondary site near the corpus callosum. This observation suggests the potential trafficking of CAR T cells to distant sites within the brain after intratumoral injection (9). In an alternative approach, both the NCI and Baylor groups, in ongoing phase I clinical trials evaluating EGFRvIII and HER2-specific CAR T cells (NCT01454596; NCT0110905), are delivering the CAR T cells intravenously (i.v.) for the treatment of GBM. Indeed, i.v. administered melanoma-specific tumor-infiltrating lymphocyte-derived T cells provide proof-of-principle for the capacity of adoptively transferred T cells to traffic to the brain and mediate regression of brain metastases (8). Thus, as future studies evaluate the treatment of brain tumors with CAR T-cell therapies, it will be important to compare differences in T-cell homing, persistence, and antitumor responses with different T-cell delivery methodologies.

While the primary objective of our pilot clinical study was assessment of safety, we observed encouraging evidence of transient antiglioma activity, including increased necrotic volume by MRI, significant loss of IL13Rα2+ tumor cell expression following CAR T-cell administration, and detection of transferred T cells at tumor microfoci removed from the site of injection (9). Although a survival benefit could not be established with such a small cohort of patients, the 3 patients treated had a mean survival of 11 months after relapse, with best survival of almost 14 months (Table 1).

This study also highlights several barriers to more durable clinical outcomes. First, we find that tumor heterogeneity may contribute to relapse by providing pools of target-deficient tumor cells that can then expand, as exemplified by the IL13Rα2-low tumor that recurred in UPN031. The recurrence of antigen-negative GBM is not unique to our study or specific for CAR T-cell immunotherapy, since it has also been observed following EGFRvIII-targeted vaccination strategies (28, 29). To address this issue, we and others are pursuing CAR T-cell approaches that target multiple antigens (30). Second, our data indicate that T-cell persistence may be another important factor (5). UPN031, whose recurrent tumor was resected 14 weeks following the last CAR T-cell infusion, displayed very low levels of CAR T-cell persistence (Supplementary Fig. S5). Although administration of dexamethasone may have influenced T-cell persistence for this patient, this lack of therapeutic T-cell persistence is also consistent with observations from our orthotopic xenograft mouse models which indicate low survival/persistence 14 days after intracranial administration of IL13-α2-zetakine® CTL clones (5). Further, previous clinical studies administering first-generation CAR T cells manufactured using a similar platform and administered i.v., also identified limited T-cell persistence as a barrier to durable clinical efficacy (31, 32). To address this limitation, we have recently optimized the IL13Rα2-specific CAR T cells, incorporating several innovations that improve both T-cell persistence and antitumor efficacy in orthotopic glioma mouse models, including engineering T cells to express a costimulatory CAR, starting with a population of central memory T cells, and utilizing CD3/CD28 beads in a manufacturing platform that limits ex vivo expansion time (manuscript in preparation). Finally, to improve the delivery methodology, our group is developing devices that allow for large infusions of therapeutic cells directly into tumors (B. Badie, unpublished data).

In summary, our clinical experience treating three research participants with recurrent GBM has demonstrated the feasibility...
of engineering, expanding, and locally delivering autologous CAR T cells without the development of serious therapy-related side effects, and has provided evidence for transient angiogenesis response. These findings support the potential of IL13Rx2-specific CAR therapy for the treatment of GBM. Future studies will focus on improving T-cell persistence, tumor targeting, and efficient delivery.

Disclosure of Potential Conflicts of Interest
M.C. Jensen has ownership interest (including patents) in, is a consultant/advisory board member for, and reports receiving commercial research grants from Juno Therapeutics, Inc. No potential conflicts of interest were disclosed by the other authors.

Authors’ Contributions
Conception and design: B. Badie, M.C. Jensen
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Acquisition of data (provided animals, acquired and managed patients, provided facilities, etc.): C.E. Brown, B. Badie, M.E. Barish, W.-C. Chang, A. Naranjo, R. Starr, J. Wagner, C. Wright, Y. Zhai, J.R. Bading, M. D’Apuzzo, M.C. Jensen
Analysis and interpretation of data (e.g., statistical analysis, biostatistics, computational analysis): C.E. Brown, B. Badie, M.E. Barish, L. Weng, J.R. Ostberg, A. Naranjo, R. Starr, Y. Zhai, J.R. Bading, M. D’Apuzzo, S.J. Forman, M.C. Jensen

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