Single CAR-T cell treatment controls disseminated ovarian cancer in a syngeneic mouse model

Diana Rose E Ranoa,1,2 Preeti Sharma,3 Claire P Schane,3 Amber N Lewis,3 Edward Valdez,3 Venkata V V R Marada,3 Marlies V Hager,3,4 Will Montgomery,4 Steven P Wolf,5 Karin Schreiber,5 Hans Schreiber,5 Keith Bailey,6 Timothy M Fan,1,7 Paul J Hergenrother,1,2 Edward J Roy,1,8 David M Kranz1,8

ABSTRACT

Background Treatment of some blood cancers with T cells that express a chimeric antigen receptor (CAR) against CD19 have shown remarkable results. In contrast, CAR-T cell efficacy against solid tumors has been difficult to achieve.

Methods To examine the potential of CAR-T cell treatments against ovarian cancers, we used the mouse ovarian cancer cell line ID8 in an intraperitoneal model that exhibits disseminated solid tumors in female C57BL/6J mice. The CAR contained a single-chain Fv from antibody 237 which recognizes a Tn-glycopeptide-antigen expressed by ID8 due to aberrant O-linked glycosylation in the absence of the transferase-dependent chaperone Cosmc. The efficacy of four Tn-dependent CARs with varying affinity to Tn antigen, and each containing CD28/CD3ζ cytoplasmic domains, were compared in vitro and in vivo in this study.

Results In line with many observations about the impact of aberrant O-linked glycosylation, the ID8&Cosmc knock-out (ID8&Cosmc-KO) exhibited more rapid tumor progression compared with wild-type ID8. Despite the enhanced tumor growth in vivo, 237 CAR and a mutant with 30-fold higher affinity, but not CARs with lower affinity, controlled advanced ID8&Cosmc-KO tumors. Tumor regression could be achieved with a single intravenous dose of the CARs, but intraperitoneal administration was even more effective. The CAR-T cells persisted over a period of months, allowing CAR-treated mice to delay tumor growth in a re-challenge setting. The most effective CARs exhibited the highest affinity for antigen. Antitumor effects observed in vivo were associated with increased numbers of T cells and macrophages, and higher levels of cleaved caspase-3, in the tumor microenvironment. Notably, the least therapeutically effective CAR mediated tonic signaling leading to antigen-independent cytokine expression and it had higher levels of the immunosuppressive cytokine interleukin10.

Conclusion The findings support the development of affinity-optimized CAR-T cells as a potential treatment for established ovarian cancer, with the most effective CARs mediating a distinct pattern of inflammatory cytokine release in vitro. Importantly, the most potent Tn-dependent CAR-T cells showed no evidence of toxicity in tumor-bearing mice in a syngeneic, immunocompetent system.

WHAT IS ALREADY KNOWN ON THIS TOPIC

⇒ Solid tumors have been difficult to treat with chimeric antigen receptor (CAR)-T cells because of a lack of suitable targets. Despite the tremendous promise of CAR-T cell therapy against solid tumors, a majority of preclinical studies with CARs have used human tumors in immune-deficient mouse models.

WHAT THIS STUDY ADDS

⇒ We established an immunocompetent, syngeneic murine model to characterize Tn-antigen specific CAR-T cells targeting advanced stages of ovarian solid tumors. We find evidence of direct CAR T-cell mediated cytotoxicity against tumor cells and exhibited long-term delay of tumor progression when administered to mice with disseminated tumors.

HOW THIS STUDY MIGHT AFFECT RESEARCH, PRACTICE OR POLICY

⇒ In this study, we developed guidelines for successful CAR-T cell treatment of advanced ovarian cancer. We demonstrate that a single dose of Tn-dependent CAR with sufficient affinity for a Tn-glycoprotein is able to control well-established tumors in this model, and was most effective with intraperitoneal administration. The study also provides additional evidence of the role of CAR affinity in the optimization of solid tumor targeting.

BACKGROUND

Adoptive therapies with T cells that express chimeric antigen receptors (CARs) against hematopoietic cancers have shown tremendous promise.1,2 The success of these CAR-T cell therapies has stemmed in part from the targeting of the normal B cell-specific differentiation antigen, CD19, which allows elimination of malignant B cells. In contrast, solid tumors have been difficult to treat with CAR-T cells because of a lack of suitable targets and unique immunosuppressive environments.3,4 Hematopoietic cancers offer other B cell-specific differentiation antigens,
but targeting self-antigens (eg, HER2, folate receptor-α, mesothelin, MUC16) (reviewed in study by Rodriguez-Garcia et al) expressed on non-hematopoietic cancers has been limited by toxicity to normal tissues that express the same targets.

Ovarian cancer is a particularly devastating disease, with a poor prognosis due to the lack of early diagnostics. Treatment involves surgery and chemotherapy (typically platinum-based drugs) and, more recently, inhibitors of angiogenesis and poly(ADP-ribose) polymerase-1 (PARP-1). Despite these approaches, 5-year survival statistics have seen minimal improvement. While there are CARs in development against various ovarian cancer antigens (reviewed in study by Rodriguez-Garcia et al), to date there have not been clinically effective results. Although preclinical effective CARs have been identified, these typically involved immunodeficient mouse models that did not express the human target.7 8

Recently, we focused on CARs 237 and 5E5 that recognize specific Tn-glycopeptide-antigens (GalNAcα-1- O-Ser/Thr-protein).9–13 The most studied Tn-antigens are the heavily O-glycosylated, repeat proteins of the Mucin family.14–17 MUC1 has recently been shown to induce affinity-matured antibodies,18 but these and other antibodies19 are bound only to non-glycosylated MUC1. In contrast, antibodies 237 and 5E5 are cancer-specific and require GalNAc in combination with peptide epitopes. Targeting Tn-peptide antigens has the potential to avoid antigen-loss variants as observed when targeting protein epitopes such as CD19,20 as the glycosylation defect is often critical for tumor cell function and Tn-dependent CARs that recognize multiple different protein backbones can be generated.12 13

The 237 and 5E5 antibodies were generated against the Tn-antigen linked to their cognate proteins, mouse podoplanin (also known as OTS8)9 and human MUC121 respectively. The cancer-specific expression of Tn-antigens can result from various dysregulated steps of the pathway, including the glycosyltransferase enzymes or the chaperone Cosmc.22 23 While COSMC (also known as CIGALTL1CI) mutations are found in approximately 1–5% of cancers,12 15 other mechanisms of O-linked glycosylation deregulation appear to contribute to estimates that 85% of ovarian cancers express the epitope recognized by the Tn-MUC1 antibody 5E5.17

Many previous studies have used the mouse tumor line ID8 as a model for human ovarian cancer.6 24 Various treatment approaches, including surgery, cisplatin, checkpoint inhibitors, oncolytic viruses, and PARP-1 inhibitors have been tested in the ID8 model (reviewed in study by McMullen et al), but tumor control in these studies has required that treatments be given early, before the presence of established, disseminated tumors. Our approaches here, with mouse CAR-T cells against ID8, focus on treatment when there are well-established solid tumors25 in a syngeneic system that is intended to resemble the scenario encountered in human disease.

We show that a single 237 CAR-T cell treatment of ID8Cosme knock-out (ID8Cosme-KO)-bearing mice, intravenously on day 40 after tumor transplantation, resulted in long-term delay of tumor growth. Interestingly, intraperitoneal administration of the CAR-T cells resulted in improved efficacy, with delayed progression when given as late as 60 days after tumor inoculation. At this point, mice exhibited extensive dissemination of solid tumors in the peritoneal cavity. A comparison of four Tn-dependent CARs showed that maximal efficacy required a minimal affinity for the Tn-OTS8 antigen. Among the four CARs, there was a distinct profile of inflammatory cytokine release, highlighted by the observation that the least effective CAR (5E5) exhibited tonic signaling leading to a higher basal level of cytokine expression and antigen-induced secretion of interleukin (IL)-10. We also describe long-term persistence of the CAR-T cells, and their ability to delay growth of an ID8Cosme-KO tumor re-challenge. These findings support the potential of treating ovarian cancer with affinity-optimized CAR-T cells, using intraperitoneal routes of administration for optimal effectiveness.

METHODS

Cell lines

Murine ovarian cancer cell line, ID8, and its mutant line with Cosmc gene deletion (ID8Cosme-KO) were maintained in complete Dulbecco’s Modified Eagle Medium supplemented with insulin-transferrin-sodium-selenite supplement at 1:5000 dilution. The generation of ID8Cosme-KO using clustered regularly interspaced short palindromic repeats and CRISPR-associated protein 9 (CRISPR/Cas9) gene editing technique has been described.12 13 Cell lines were submitted to IDEXX BioAnalytics for authentication, and were tested free of pathogens. All cell lines and primary cells used in this study were cultured at 37°C with 5% CO2.

Chimeric antigen receptors

The 237 single-chain Fv (scFv) was fused to CD8α hinge-CD28-CD3ζ and cloned into the pMP71 retroviral vector to generate 237 CAR plasmid.10 Structure-guided engineering of the 237 scFv fragment3 yielded CAR variants that bound with higher affinity to the cognate murine Tn-OTS8 peptide (237 scFv mutant WE, designated WE CAR here) or exhibited binding to both murine Tn-OTS8 and human Tn-MUC1 (237 scFv mutant TNGK, designated TNGK CAR here). The 5E5 scFv CAR that binds to human Tn-MUC1 was characterized in human T cells previously13 and was subcloned into the pMP71 for mouse T-cell studies.13

T cell transduction and activation

The protocol for retroviral transduction of primary T cells isolated from splenocytes of donor mice has been described.10 13 A detailed protocol is outlined (DOI: dx.doi.org/10.17504/protocols.io.6qpvr4jwpkmk/v1). Mock-transduced T cells went through the retroviral
transduction process but without recombinant virus. Transduction efficiency was measured using Tn-OTS8 tetramer for 237, WE and TNGK CARs or Tn-MUC-1 tetramer for 5E5 CARs.

**Animal studies**

Animal studies were approved by the UIUC Institutional Animal Care and Use Committee (IACUC) under protocol numbers 21041 and 20176. Female C57BL/6j mice (8–10 weeks old) were purchased from Jackson Laboratories, maintained under pathogen-free conditions in a barrier facility, ear-tagged, and assigned to treatment groups. ID8 and ID8Cosmc-KO tumor cells (10⁶) were inoculated intraperitoneally. After various days, 5×10⁶ CAR-T cells (or mock-transduced T cells) were administered intravenously or intraperitoneally. At 24 hours prior to T-cell transfer, cyclophosphamide (100 mg/kg) was administered intraperitoneally. Tumor growth and progression of disease were monitored by measuring body weight on a regular basis. Mice were euthanized when their body weights reached the endpoint criterion of 36 g or a gain of 50% of their weight from the start of treatment. For some experiments, blood samples (~70 μL) were collected by submandibular vein puncture or retro-orbital bleeding. Tissue samples were collected postmortem, fixed in 10% buffered formalin for 24 hours, and subsequently transferred to 70% ethanol prior to routine tissue sectioning at 5 micron thickness and staining with H&E.

**Antibody detection agents**

Antibodies used for flow cytometry were purchased from BioLegend or BD Bioscience: CD3 (clone 17A2), CD8α (53–6,7), CD4 (RM4–5), CD45 (30–F11), and TCR Çβ (H57–597). Fixable Viability Dye eFluor 780 was purchased from Invitrogen (P-160-66). Data were acquired on a BD LSRII and analyzed with FlowJo software. The Tn-specific monoclonal antibody 5F4 (IgM), a kind gift from Dr Henrik Clausen (University of Copenhagen, Denmark), 28 was used as culture supernatant (for flow cytometry) or purified using LigaTrap Microspin Kit (LT-145KIT) and biotinylated (Thermo Fisher/ Pierce EZ-Link NHS-PEG4 biotinylation kit 21455) (for immunohistochemistry). The 237 scFv mutant WE, cloned into pET28a with a 6X histidine and AviTag, was purified from E. coli by combining biotinylated WE scFv with streptavidin-linked Protein Ligase Reaction Kits (Avidity BIRA-500). Fluorescently-labeled WE scFv tetrantomers were prepared by combining biotinylated WE scFv with streptavidin-PE (BD Bioscience CN54061) or streptavidin-Alexa Fluor 647 (Invitrogen C32457) at a 4:1 molar ratio. Biotinylated Tn-OTS8 and biotinylated Tn-MUC1 peptides were prepared as described in study by Sharma et al. 13

**Processing and analysis of tissues and tumors**

Protocols for processing mouse tissues and staining for flow cytometry are described in DOI: dx.doi.org/10.17504/protocols.io.6qpvrljwpgmk/v1. H&E staining and immunohistochemistry of tissue sections from formalin-fixed paraffin embedded samples were performed by the University of Chicago Human Tissue Resource Center core facility. Stained slides were scanned using a Hamamatsu NanoZoomer slide scanner. Protocols for staining with biotinylated agents (anti-Tn antibody 5F4 or WE scFv), anti-CD3 (Abcam ab135372, clone SP162), anti-F4/80 (BioRad MCA497GA, clone A3-1), and anti-Ly6G (BioLegend 127602, clone 1A8), and anti-cleaved caspase 3 (Cell Signaling Technology 9661) are described in DOI: dx.doi.org/10.17504/protocols.io.6qpvrljwpgmk/v1.

**In vitro co-culture assays**

ID8 wild-type (WT) or ID8Cosmc-KO cells were co-cultured with CAR-T cells in 96-well plates at different seeding density. In one approach, the number of effector T cells was kept constant (12,000 cells/well) while the number of tumor cells was varied (tumor:effector ratios). In another, tumor cell number was constant while effector T-cell numbers were varied. After co-culturing for 24 hours at 37°C, supernatants were transferred to V-bottom 96-well plates and supernatants were stored at −20°C.

For cytokine analysis, 25 μL supernatants were used in a customized mouse cytokine array (Milliplex MCYtomag-70K) following the manufacturer’s protocol. The panels included interferon (IFN)-γ, IL-2, IL-4, IL-6, IL-10, and tumor necrosis factor (TNF)α (Milliplex MCYtomag-70K-6plex). Plates were read on Luminex 200, with quality controls and standards included.

T-cell cytotoxicity of tumor cells was measured using the sulforhodamine B colorimetric assay following a protocol described in DOI: DOI: dx.doi.org/10.17504/protocols.io.6qpvrljwpgmk/v1. IFN-γ ELispot assays were performed using splenic T cells and a mouse IFN-γ ELispot kit (CTL ImmunoSpot), following 24-hour co-culture.

T-cell proliferation assays were performed by labeling activated mock or CAR-T cells with CFSE (Invitrogen C34554) following the manufacturer’s protocol. Post-transduction (72 hours), T cells were washed, resuspended in phosphate-buffered saline, and incubated with 10 μM CFSE for 20 min at 37°C in the dark. Cells were washed with culture media, and resuspended to a final cell density of 1×10⁵ cells/mL. 100 μL cell suspension was aliquoted in a 96-well plate precoated with streptavidin followed by biotinylated peptides (described in studies by He et al and Sharma et al. 14 15). Following 72-hour incubation, T cells were harvested, washed, and analyzed in a BD Accuri flow cytometer.

**RESULTS**

**ID8Cosmc-KO as a model for CAR treatment of ovarian cancer**

The ovarian cancer cell line ID8, with a deletion of the gene for the chaperone Cosmc, expresses high levels of Tn-antigen as detected with anti-Tn antibody 5F4 (figure 1A). ID8Cosmc-KO is also recognized by Tn-dependent antibody 237 and its CAR derivatives. 13 For
detection, the high-affinity variant called 237-WE (WE from here-on) was expressed as a soluble biotinylated scFv. Tetramers of the WE scFv were used to validate the expression of the Tn-OTS8-antigen in the ID8Cosmc-KO line (figure 1B). Notably, the parental ID8 (ID8 WT) and ID8Cosmc-KO lines exhibited the same rate of growth in vitro, with doubling times of 19.2 hours and 21.8 hours, respectively (figure 1C).

To evaluate the ID8Cosmc-KO as a model for 237 CAR treatment, we inoculated $10^7$ tumor cells intraperitoneally into syngeneic C57BL/6 mice (figure 1D). We used ID8 lines without luciferase for imaging to minimize the potential for immunogenicity of foreign proteins in immunocompetent mice.36 31 Consistent with previous ID8 studies, mice with ID8 WT began to develop ascites and gained weight to the point of euthanasia criterion.

Figure 1 ID8Cosmc-KO as a model for ovarian cancer in C57BL/6J mice. Tn antigen expression on ID8Cosmc-KO was measured by flow cytometry using either (A) staining with monoclonal mouse IgM anti-Tn antibody 5F4 followed by goat anti-mouse IgM conjugated to Alexa Fluor 647 or (B) biotinylated anti-Tn-OTS8 WE scFv tetramer containing streptavidin-PE. (C) ID8 WT and ID8Cosmc-KO cell doubling times (19.2 hours and 21.8 hours, respectively) in vitro were measured over time by manual cell counting. Growth curves from two different experiments (total n=6) were combined and plotted using GraphPad Prism V.9.4.1 and cell doubling times were calculated using the exponential growth equation. (D) Timeline for ID8Cosmc-KO tumor growth in syngeneic C57BL/6J mice after intraperitoneal injection of $10^7$ tumor cells. (E) Survival curves of C57BL/6J mice inoculated intraperitoneally with $10^7$ ID8 WT or ID8Cosmc-KO. Median survival was 88 days and 73 days for ID8 WT (n=19) and ID8Cosmc-KO (n=35), respectively. For C57BL/6 Rag1-KO mice inoculated with $10^7$ ID8Cosmc-KO (n=10), the median survival was 28 days. Survival curves were plotted using GraphPad Prism V.9.4.1 and effects were calculated using the log-rank (Mantel-Cox) test.
Efficacy of different Tn-dependent CARs in treatment of ID8Cosme-KO tumors

We recently engineered a Tn-dependent CAR called TNGK, using the 237 scFv antibody backbone, that was selected with the human antigen Tn-MUC1.15 The TNGK scFv bound with higher affinity to Tn-MUC1 than 237 scFv, but it exhibited lower affinity binding to Tn-OTS8, the cognate mouse antigen recognized by 2379 (online supplemental figure S3A). Another Tn-dependent CAR called 5E5 was originally developed against Tn-MUC111,21 and has high affinity for Tn-MUC1 but low affinity for the mouse antigen Tn-OTS813 (online supplemental figure S3A,B). TNGK and 5E5 CARs have about 10-fold less sensitivity than 237 CAR using immobilized Tn-OTS8 antigen to stimulate IFN-γ.15 Both of these CARs were tested in an experiment where mice with ID8Cosme-KO were treated intravenously on day 40 (figure 4A). Although TNGK CAR delayed growth of the ID8 tumor, 5E5 showed no effect.

To extend these studies with additional Tn-dependent CARs, we also examined WE CAR-T cells that contained the scFv variant with 30-fold higher affinity than 237 (Kₐ values of 120 nM and 4 nM for 237 and WE scFv, respectively). WE, 237, TNGK, and 5E5 CARs were administered intraperitoneally at day 50 after ID8Cosme-KO inoculation (figure 4B). The 5E5 CAR showed no effect, TNGK
Figure 2  Pathological and histochemical analyses of tissue sections from ID8Cosmc-KO and ID8 WT mice at various times after tumor challenge. (A) At indicated days post-inoculation with $10^7$ tumor cells, mice were euthanized, and organs in the peritoneal cavity were examined visually for tumor nodules (black arrows). (B) Formalin-fixed paraffin embedded tissue sections from mice inoculated with ID8Cosmc-KO cells at the indicated time points were stained with H&E. Tumor foci are labeled. Scale bar=500 µm. (C, D) ID8 WT or ID8Cosmc-KO tumors at day 60 post-tumor challenge were examined by immunohistochemistry staining using biotinylated anti-Tn IgM antibody 5F4 (C) or biotinylated WE scFv (D). Scale bar=250 µm. KO, knock-out; scFv, single-chain Fv; WT, wild-type.
**Figure 3** CAR-T cell treatment of mice with established ID8Cosmc-KO tumors. (A) Schematic of 237 single-chain Fv linked by a CD8 hinge-CD28 transmembrane domain to the CD28 and CD3ζ cytoplasmic domains, cloned into the pMP71 retroviral vector. (B) Primary T cells isolated from splenocytes of naïve donor C57BL/6J mice were transduced with 237 CAR for 72 hours. Transduction efficiency was measured by flow cytometry using tetramers of biotinylated/Tn-glycosylated OTS8 peptide made with streptavidin-PE. Mock-transduced T cells were included as negative control. (C, D) Survival curves of ID8Cosmc-KO-bearing C57BL/6J mice either untreated (black), treated with mock-transduced T cells (blue) or 237 CAR-T cells (red) administered by retro-orbital intravenous injection of 5×10⁶ T cells at day 40 (C) or day 56 (D) post-tumor inoculation. Note that five of the mice in the 237 CAR group (C) were used at day 160 for a re-challenge experiment (figure 7B), such that their survival times may have been even longer without this re-challenge. (E) Comparison of intravenous (dashed red line) and intraperitoneal (solid red line) delivery of 237 CAR-T cells at day 50 post-tumor inoculation. All survival curves were plotted with GraphPad Prism V.9.4.1 using cumulative data across multiple experiments (two combined experiments for C and D, four combined experiments for E), with the indicated total n for each treatment arm. Median survival times are indicated; p values were calculated using the log-rank (Mantel-Cox) test. CAR, chimeric antigen receptor; i.p., intraperitoneal; i.v., intravenous; KO, knock-out.
CAR significantly extended median survival by 17 days (p=0.0034), and both 237 and WE CARs mediated long-term control of tumors, to median survivals of 264 and 299 days, respectively. There was no significant difference in tumor control between 237 and WE CARs, despite the 30-fold higher affinity of the latter.

To understand the limits of the CAR-T cell treatments, we performed an experiment in which mice were treated with 237 and WE CAR-T cells administered intraperitoneally at day 60, the phase at which some mice reach the weight criterion for euthanasia. Not surprisingly, there was reduced efficacy of the CAR-T cell treatment at this
advanced disease stage (figure 4C), but there was a significant delay in disease progression mediated by 237 and WE CARs (p=0.0009 and p=0.0095, respectively).

In vitro analyses comparing 237, WE, TNGK and 5E5 CARs

Given that all four CARs mediated Tn-dependent IFN-γ release in vitro when stimulated by ID8Cosmc-KO cells, the question remained what might explain their distinct differences in efficacy. To examine if there was a correlation between in vivo effects and in vitro cytokine release, we assayed in vitro-stimulated CAR-T cells for secretion of various cytokines, including IFN-γ, IL-2, TNFα, IL-6, IL-4 and IL-10 (figure 5A–F, online supplemental figure S4A–F). To present antigen, we used various ratios of ID8Cosmc-KO tumor cells. Consistent with our previous findings, all four CARs were effective in mediating release of IFN-γ, even at the lowest tumor:CAR-T cell ratio of 0:1 (online supplemental figure S4A). Of particular note, among multiple experiments, the 5E5 CAR showed a level of tonic cytokine release, especially with IFN-γ, in the absence of antigen. While all four CARs mediated release of the other cytokines, there was a distinct quantitative difference among them. The two most effective CARs in vivo, 237 and WE, exhibited greater stimulation of IL-2 compared with 5E5. Interestingly, the reciprocal was observed for the release of IL-4 and IL-10, where 5E5 CAR exhibited higher levels than 237 and WE. However, 5E5 CAR-T cells also exhibited higher tonic levels of IL-4 and IL-10 than the other CARs, and significant levels of antigen-independent TNFα release. The release of IL-10 mediated by 5E5 CAR was also observed when it was co-cultured with Jurkat, a human tumor line with a known Cosmc mutation (online supplemental figure S4F).

We also assessed in vitro tumor killing mediated by CAR-T cells. All four CARs mediated killing of ID8Cosmc-KO cells (figure 5G and online supplemental figure S5A) but not ID8 WT cells (figure 5H and online supplemental figure S5B). However, 5E5 CAR-T cells exhibited lower cytotoxic activity among the four CARs tested.

Finally, we examined antigen-dependent proliferation using CFSE-labeled T cells stimulated with various concentrations of OTS8, Tn-OTS8, MUC-1, and Tn-MUC1 peptides for 72 hours by flow cytometry (online supplemental figure S6). Gating on viable cells using forward/side scatter, we observed that 237, WE, and TNGK CAR exhibited background levels (mock T cells) of proliferation in the absence of the cognate antigen, with about 0.5% in the lymphocyte gate (online supplemental figure S6A,C). In contrast, 5E5 CAR exhibited almost 5% proliferating cells with the control peptides OTS8 and MUC1 (online supplemental figure S6B,C). All of the CARs, including 5E5, exhibited antigen-dependent proliferation, and for 237 and WE this level was highest with the mouse antigen Tn-OTS8. Similar results were observed when monitoring proliferation by flow cytometry for CFSE, where reduced mean fluorescent intensity correlates with greater proliferation (online supplemental figure S6D,E). The extent of proliferation was greatest for 237 and WE CARs with Tn-OTS8, whereas TNGK showed extensive proliferation with both Tn-OTS8 and Tn-MUC1. Again, 5E5 showed a high level of Tn-antigen-independent proliferation.

Immunophenotyping ascites and TILs from ID8Cosmc-KO mice

To assess CAR T cell distribution immediately after CAR treatment, we performed immunophenotyping by flow cytometry on samples from blood, spleen, pancreas, and ascites fluid recovered from ID8Cosmc-KO-bearing mice (online supplemental figure S7). The 237 or 5E5 CAR-T cells were administered intraperitoneally on day 54 post-tumor challenge, and mice were euthanized 3 days post-CAR treatment. Although ascites at this time point is typically not large, the volume collected from 237 CAR-treated mice was higher relative to that collected from untreated mice (online supplemental table S3). Ascites from 237 CAR-treated mice was ‘milky off-white’, compared with ‘red’ typical in untreated and 5E5 CAR-treated mice and the surface of pancreases from CAR-treated mice appeared more viscous compared with untreated controls.

We examined the distribution of CAR-T cells in circulation and within the pancreatic tissue using Tn-OTS8 tetramers (gating strategy, online supplemental figure S7A). Both 237 and 5E5 CARs (CD3+, CD8+, and CD4+ double-positive cell populations) were detectable and significantly elevated in tumor tissues and ascites samples from CAR-treated tumor-bearing groups (online supplemental figure S7B), indicating that CAR-T cells, administered to mice 3 days prior, have migrated to target sites.

To further assess immune cell infiltration in tumors 3 days after CAR treatment, immunohistochemistry was performed with biotinylated WE-scFv, anti-CD3, anti-F4/80, anti-Ly6G, and anti-cleaved caspase 3 antibodies (figure 6A and online supplemental figure S8). CAR-treated tumor sections contained prominent cellular infiltrates that outnumbered tumor cells. In the absence of CAR-treatment, few CD3+ T cells were detected in tumors. Following either 237 or 5E5 CAR administration, tumor nodules and masses were extensively infiltrated by CD3+ T cells, with 237 CAR-treated tissues showing significantly higher CD3+ T cell infiltration compared with 5E5 CAR-treated tissues (figure 6B, left panel). Based on flow analysis of TILs, it appears that only 5–10% of these CD3+ T cells represent CAR-T cells (online supplemental figure S7B). The high frequency of infiltrating immune cells could partly explain the presence of sections that appeared negative for WE immunostaining, compared with tumors from untreated mice. While most of the small infiltrating cells had lymphocyte morphology and reacted with anti-CD3, there was also an increase in F4/80+ macrophage infiltration in all tumor sections examined, with 237 CAR-treated tissues exhibiting highest levels compared with either untreated or 5E5 CAR-treated tissues (figure 6B, middle panel). Rare to small numbers of Ly6G+ cells (neutrophils) were observed in the tumor sections (online supplemental figure S8). Consistent with high levels of T cell infiltration, CD3+ T cells in circulation were detected at much lower levels than the other CARs, and significant levels of antigen-independent TNFα release. The release of IL-10 mediated by 5E5 CAR was also observed when it was co-cultured with Jurkat, a human tumor line with a known Cosmc mutation (online supplemental figure S4F).

We also assessed in vitro tumor killing mediated by CAR-T cells. All four CARs mediated killing of ID8Cosmc-KO cells (figure 5G and online supplemental figure S5A) but not ID8 WT cells (figure 5H and online supplemental figure S5B). However, 5E5 CAR-T cells exhibited lower cytotoxic activity among the four CARs tested.

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**Figure 5** In vitro cytokine and killing analyses of different CAR T cells. Mock, 237-CAR, WE-CAR, TNGK-CAR, or 5E5-CAR transduced primary T cells from C57BL/6J donor mice were co-cultured with ID8Cosmc-KO cells in a 96-well plate for 24 hours at various tumor:effector ratios. The number of effector T cells were kept constant at 12,000 cells/well across all conditions tested. After 24 hours, 25μL culture supernatants were analyzed for multiple cytokines using a Luminox plate reader to measure IFN-γ (A), IL-2 (B), TNFα (C), IL-6 (D), IL-4 (E), and IL-10 (F). Plots shown are combined cytokine array data points from five experiments. Bar graphs were plotted using GraphPad Prism V.9.4.1. Error bars are SEM. To simplify the presentation, p values are shown only for 237 versus 5E5 CAR. A complete set of tumor:effector ratio titration plots for each of the five cytokine array experiments are shown in online supplemental figure S5. (G, H) The remaining ID8Cosmc-KO (G) or ID8 WT (H) adherent tumor cells on plates were fixed in ice-cold 10% trichloroacetic acid for the SRB viability assay. The percent growth inhibition was calculated using the formula outlined in the Methods section. SRB experiments were performed twice, each with a technical replicate of n=4. Bar graphs were plotted using GraphPad Prism V9.4.1. Error bars are SEM. P values for 237, WE, TNGK were calculated against either mock T or TNGK are at <0.001 (**). CAR, chimeric antigen receptor; IFN, interferon; IL, interleukin; KO, knock-out; TNF, tumor necrosis factor; WT, wild-type.
Figure 6  Peritoneal organs were harvested from ID8Cosmc-KO inoculated mice, 3 days after intraperitoneal treatment with either 237 CAR-T or 5E5 CAR-T at day 54 post-tumor inoculation. Untreated mice were used as control. (A) Immunohistochemistry was performed to visualize Tn-OTS8-glycopeptide+ tumors, CD3+ T cells, F4/80+ macrophages, and apoptotic cells from adjacent FFPE tissue sections stained with biotinylated WE-scFv, anti-CD3, anti-F4/80, and anti-cleaved caspase 3 antibodies, respectively. H&E stained tissue sections were included as additional reference. Scale bar=250 µm. (B) ImageJ-quantified CD3+ T cells, F4/80+ macrophages, and cleaved caspase-3+ apoptotic cells from tissue sections of untreated, 237 CAR-T, 5E5 CAR-T treated mice. A total of 5–20 images at 20× magnification corresponding to tumors spotted lining the fatty tissues near stomach, small and large intestines of two mice per treatment group were analyzed and quantified using ImageJ. Dot plots and p values were generated using GraphPad Prism V.9.4.1. Error bars are SEM. Each dot represent one field of view at 20× magnification. CAR, chimeric antigen receptor; FFPE, formalin-fixed paraffin embedded; KO, knock-out; scFv, single-chain Fv.
and macrophage infiltration, 237 CAR-treated tissues also had significantly higher numbers of apoptotic cells as shown in cleaved-caspase 3 immunoreactive tissue sections (figure 6B, right panel). We also examined the tumor stroma using trichrome staining and found scant amounts of intratumoral fibrous tissue/collagen, arranged in thin, delicate strands (positive blue stain in bottom row of online supplemental figure S8), but not to an extent that we believe would interfere with CAR migration. Taken together, our results show that 237 CAR treatment recruited more CD3+ and F4/80+ macrophages to the tumor microenvironment, resulting in higher cleaved caspase 3-positive apoptotic cells compared with both 5E5 CAR treated and untreated tissues.

**Persistence of CAR-T cells in treated mice**

One of the metrics associated with efficacy of CAR-T cell treatments is persistence of CAR-T cells. To assess persistence, we tested the ability of CAR-T cell-treated mice to mount a response to re-challenge with ID8 Cosmc-KO tumor, using mice that had controlled progression of tumor for a period of 120 days post T-cell administration. Prior to re-challenge, blood cells were co-stained with anti-CD3 and Tn-OTS8 tetramer, showing that CAR-T cells persisted, compared with age-matched non-tumor bearing mice (figure 7A, left panel). Seven days after re-challenge with ID8 Cosmc-KO, the population of CD3+ 237 CAR T increased further (figure 7A, right panel). Survival data of re-challenged, CAR-treated mice (figure 7B) indicated that some mice were able to delay tumor progression, with a median survival of 108 days after re-challenge (compared with 68 days for naïve mice) (p=0.0150). In addition, analysis of blood 55 days post-re-challenge suggested that levels of CD4+CAR-T remained elevated in treated mice compared with control (online supplemental figure S9A).

To further assess T-cell persistence, we performed IFN-γ ELISPOT of spleen cells from 237 CAR-treated mice. Splenic T cells were co-cultured with ID8 WT or ID8 Cosmc-KO tumor cells at various tumor-to-effector ratios. More cells from 237 CAR-treated mice were stimulated to secrete IFN-γ by ID8 Cosmc-KO compared with ID8 WT cells (figure 7C), indicating that Tn-specific T cells persisted in treated mice. Presence of Tn-dependent CAR T cells was also shown by the stimulation of IFN-γ secreting T cells by the Cosmc-deletion line AG104A, but not other mouse tumor lines (online supplemental figure S9B).

To explore whether antigen was necessary for CAR-T cell persistence, we examined whether CAR-T cells administered 60 days prior to tumor inoculation could delay tumor progression (figure 7D). CAR-T pretreated mice survived to a median of 89 days compared with 68 days for untreated naïve mice, indicating that sufficient CAR-T cells were present, even in the absence of antigen, to delay growth (p=0.045).

**Tn-antigen requirement in CAR-T cell treated mice**

To verify that Tn-antigen was required for 237 CAR efficacy, we treated ID8 WT-bearing mice with 237 CAR (figure 7E). The 237 CAR-treated mice did not show a delay in survival compared with untreated or mock-treated controls, indicating that the in vivo potency of the 237 CAR requires Tn-antigens such as Tn-OTS8 generated by the ID8 Cosmc-KO line.

Finally, to determine if long-term, relapsed mice treated with the 237 or WE CARs showed evidence of loss of the Tn-antigen, we analyzed the Tn-expression status of tumor cells collected from ascites of relapsed mice. The frequency of Tn-expressing tumor cells in ascites collected from 237-treated and WE-CAR-treated mice that reached endpoint was reduced compared with untreated and mock T treated groups (online supplemental figure S10A). To directly address if the tumor cells in ascites from 237 CAR-treated mice contained lower level of Tn-antigen, we cultured cells from untreated mice (n=4), short-term treated mice (n=2), and long-term treated, relapsed mice (n=6). Tumor cell lines from untreated and short-term treated mice all expressed the same surface level of Tn-antigen (WE) as the parental ID8 Cosmc-KO line (online supplemental figure S10B). In contrast, all six of the tumor lines derived from relapsed long-term survivors showed either lower levels of the antigen, or complete absence of detectable Tn-antigen. Immunohistochemistry of tissue sections from two long-term relapsed 237 CAR treated mice also demonstrated heterogeneity in Tn-expression in tumors (online supplemental figure S10C).

**DISCUSSION**

The mouse ovarian cancer line ID8 has been used in many previous studies to examine the efficacy of various therapies. However, successful treatment of this tumor has been difficult, requiring early treatments (ie, often less than 2 weeks after tumor inoculation) to observe delayed tumor growth. In contrast, human ovarian cancer is rarely diagnosed at such early stages of disease development. Our most important finding is that late stage treatments with CAR-T cells, when solid tumors are detectable by gross and histologic analysis, yielded long-term control of the tumor. Our studies also contrast with studies using human tumors transplanted into immunodeficient mice, which often do not express the human target antigen on normal mouse tissues which would be needed to assess toxicity.

Successful treatment was accomplished with just a single dose of 237 or WE CAR-T cells, even when ID8 Cosmc-KO tumors were disseminated throughout the peritoneal cavity. Gross visual and histochemical analyses revealed tumor nodules and masses in adipose tissues by day 40 with progression to more extensive tumor spread by the time of CAR-T cell treatments on day 50 or 60. Several days after treatment, extensive T-cell infiltration into tumors was identified, yet there was no sign of CAR-T cell-related
**Figure 7** Evidence for in vivo persistence of 237 CAR T cells. (A) 237 CAR T-treated ID8Cosmc-KO-bearing mice were re-challenged with $10^7$ ID8Cosmc-KO cells at 120 days post-T cell transfer. Age-matched naive mice were inoculated with $10^7$ ID8Cosmc-KO cells and used as control. Blood samples were collected from some mice immediately before intraperitoneal injection of re-challenged tumors and 7 days after re-challenge; processed and stained with anti-CD3 antibody and Tn-glycosylated OTS8 peptide tetramer, and analyzed by flow cytometry. The percentage of CD3+ 237 CAR T cells were plotted using GraphPad Prism V.9.4.1. Each dot represents one mouse. Error bars are SEM. (B) Survival curve of 237 CAR T treated mice re-challenged with ID8Cosmc-KO at either 120 (n=5) or 160 days (n=5) after initial tumor injection, compared with age-matched C57BL/6 naïve mice that were inoculated with the same number of tumor cells. (C) Spleens were removed from 237 CAR T-treated ID8Cosmc-KO-bearing mice up to 5 months after CAR treatment (152 days for the data shown). Spleen cells were co-cultured with either ID8 WT or ID8Cosmc-KO tumor cells in a 96-well plate at a tumor:effector ratio of 0.3. ELISpot assays were performed to measure IFN-gamma spots 24 hours after co-culture experiment. Splenocytes from age-matched mice were used as a control, while CD3/CD28 Dynabeads were added to T cells as a positive control for T cell activation. Plates were scanned with automated ImmunoSpot analyzer. Bar graph of quantified IFN-γ spots are plotted on the right panel. Results representative of five independent experiments. (D) Non-tumor bearing mice were injected intraperitoneally with 237 CAR T (see online supplemental figure S2B) or untreated. At day 60, mice were challenged with $10^7$ ID8Cosmc-KO cells and the survival curve was compared with that of mice inoculated with tumor that did not receive prior 237 CAR T injections (untreated). (E) Survival curve of ID8 WT-bearing C57BL/6J mice either untreated (black), treated with mock-transduced T cells (blue), or treated with 237 CAR T (red), administered by intraperitoneal injection of $5 \times 10^6$ T cells at day 60. Data for D and E are from two combined experiments. All survival curves were plotted using GraphPad Prism V.9.4.1 with the indicated total n for each treatment arm. Median survival times are indicated; p values were calculated using the log-rank (Mantel-Cox) test. CAR, chimeric antigen receptor; IFN, interferon; i.p., intraperitoneal; KO, knock-out; WT, wild-type.
injury to neighboring organ systems. Although edema was observed in pancreas sections of CAR-treated mice (online supplemental table S2), there was no evidence of toxicity in either tumor-bearing or non-tumor-bearing mice in this syngeneic, immunocompetent system. The potency of the 237 CAR was associated with 237 CAR-driven immune cell infiltration which conferred an anti-tumor phenotype. This is consistent with a previous study showing that adoptively transferred T cells are capable of overcoming immune suppressive cells.\(^\text{35}\) It was also interesting that the ID8\(^{-}\)Cosme-KO tumor grew rapidly in Rag1-KO mice, indicating that endogenous B and/or T cells are capable of controlling the tumor. Whether these cells are, or could be, recruited through the activity of 237 cells are capable of controlling the tumor. Whether these CARs are in clinical development against several human ovarian cancer antigens, including folate receptor alpha (NCT03916679, NCT03799913, NCT03585764), mesothelin (NCT03916679, NCT03814417, NCT05372692, NCT03054298, NCT05057715), B7-H3 (NCT05211557, NCT04670068), TAG72 (NCT05225363), MUC16 (NCT03907527), placental alkaline phosphatase, PLAP (NCT04627740), and P-MUC1 (NCT05239143).\(^\text{3}\) Preclinical studies against other targets, such as the Mullerian inhibiting substance type II receptor, have also shown promise.\(^\text{34}\) The optimal antigen target has yet to be determined, but development of multiple CARs against different targets provides opportunities to mitigate immune-evasive strategies employed by cancer cells such as antigen loss. In the present study, the potency of the Tn-dependent CARs ultimately resulted in reduced antigen expression in long-term survivors that eventually relapsed, but we have not yet elucidated the mechanism associated with the loss or reduction of Tn-antigen.

A trial using CAR T-cells expressing the humanized 5E5 scFv against Tn-MUC1 has been initiated (NCT04025216). In this regard, it is worth comparing the binding features of the 5E5 CAR and the 237, WE, and TNGK CARs. While all of these CARs are Tn-dependent, each has a different affinity and/or peptide fine-specificity. Surface plasmon resonance (SPR) has been used to determine the affinity of the parental antibodies and their antigens: 5E5 has an affinity (K_d) of 2 nM for human Tn-MUC1\(^\text{35}\) and 237 has an affinity (K_d) of 140 nM for mouse Tn-OTS8.\(^\text{36}\) The WE scFv was derived from 237 using in vitro affinity maturation to yield an affinity increase of 30-fold, to an affinity of 4 nM for Tn-OTS8 (ie, similar to 5E5 for Tn-MUC1).\(^\text{36}\) The TNGK scFv was engineered to bind to Tn-MUC1.\(^\text{35}\) WE and TNGK CAR differ in their CDRs from 237 CAR by only two and four amino acids, respectively.

While affinities using SPR have not been determined for these scFv:Tn-glycopeptide pairs, based on antigen titrations the order of binding to the predominant mouse antigen in ID8\(^{-}\)Cosme-KO, Tn-OTS8, is WE > 237 > TNGK > 5E5 (online supplemental figure S3A). The order of binding to human Tn-MUC1 is 5E5 > TNGK > 237 > WE. Although all these CARs exhibit a degree of cross-reactivity with different Tn-peptide backbones, we show in this report that affinities of 140 nM (237 for Tn-OTS8) or higher may be equally optimal for in vivo efficacy, at least in the context of antigen densities represented with ID8\(^{-}\)Cosme-KO. The reduced in vivo efficacy of the TNGK CAR may be accounted for by its lower affinity than 237 and WE for the Tn-OTS8 antigen on ID8\(^{-}\)Cosme-KO. An affinity threshold for the scFv fragments used in CARs has been previously described.\(^\text{37}\)

Another observation of interest is that despite the ability of all four CARs to mediate in vitro release of IFN-γ, stimulation of other cytokines differed among the CARs. It was particularly notable that the most efficacious CARs in vivo showed higher IL-2 but lower IL-10 and IL-4. In contrast, 5E5 CAR-T cells secreted higher relative levels of IL-10 and IL4. This observation is consistent with the well-known immunosuppressive abilities of IL-10.\(^\text{38}\)

Perhaps most decisive with regard to the mechanism underlying the lack of efficacy of the 5E5 CAR was our observation that it yielded a low level of tonic activity, unlike the other three CARs. A head-to-head comparison of 237-CAR and 5E5-CAR (both with CD28 signaling domain) showed that the 5E5 CAR used in this study exhibited basal tonic activity and proliferation, and about a twofold lower level of surface expression compared with other CARs (online supplemental figure S3B). Tonic, antigen-independent activity of CARs has been reported to be due to aggregation of scFv. The \(V_h\) and \(V_l\) domains of 5E5 are quite different from the \(V_h\) and \(V_l\) domains of the 237, WE, and TNGK CARs, which have identical framework regions. Importantly, tonic activity has been correlated with ineffective CARs, and one way to mitigate this tonic stimulation is through CAR engineering.\(^\text{39–41}\) It is worth noting that two previous studies by Posey et al and He et al used 4-1BB instead of CD28 in their 5E5 CAR constructs, which could explain why these 5E5 CAR constructs did not yield tonic activity.\(^\text{42}\) The use of the TNGK-CD28 CAR for human translational studies, on the other hand, may be advantageous given its favorable properties of low tonic release and its recognition of human Tn-MUC1.

The question remains what surface level of Tn-protein antigen in human cancers is minimally sufficient to elicit CAR T-cell efficacy, as described here. The 5E5 CAR has been shown to mediate killing of, and IFN-γ stimulation by, a panel of diverse human cell lines \(6 \text{ out of 10 cell lines).}^{11}\) This activity was correlated with upregulation of the ST6GalNAc1 transferase, or in one case (Jurkat) the mutation of COSMC. Among the limited tumor cell lines that we have examined with 237 or its derivatives (including WE and TNGK) using transduced mouse T cells, a Cosme deletion was necessary for Tn-dependent activity.\(^\text{10–12,13}\) The high density of Tn-targets found in Cosme deletion mutants may correlate with the optimal in vitro cytokine stimulation and proliferative activity, and in vivo tumor infiltration of the 237 and WE CARs, as described here. Nevertheless, the frequency of human tumors that exhibit COSMC mutations may make treatment with Tn-dependent CARs a valuable option for some patients. Endometrial cancers are reported to have a 5% prevalence of COSMC mutations, and other cancer types including ovarian cancer...
cancer are in the range of 0.5–1%.\textsuperscript{12,13} While this may seem low, current efforts to target a specific driver mutation (eg, in p53 or KRAS), combined with the requirement for HLA restriction brings these frequencies to the same range.

Our findings that intraperitoneal administration of the CAR-T cells was more effective than intravenous administration supports such an approach in clinical trials of CAR-T cells against ovarian cancer. Furthermore, the potency of a single CAR T-cell treatment at days 40 to 50 after tumor transplant indicates that post-surgical treatment, when there might be minimal but significant disease, could be a preferred time for optimal therapeutic benefit. This is also particularly relevant to a recent study in which ID8 was implanted into the ovaries,\textsuperscript{43} despite surgical debulking at different times, all mice exhibited ID8 tumor development with pathology and dissemination much like the ID8\textsuperscript{Cosmo-KO} model described here.

A notable exception for later treatment of the ID8 model involved the use of an ‘armored’ CAR that expressed a synthetic IL-12 gene.\textsuperscript{44} It remains to be seen if CART cells engineered to secrete IL-12 or other cytokines have appropriate safety profiles, as IL-12 has shown serious toxicities clinically.\textsuperscript{45} Inducible systems have been designed to mitigate potential cytokine toxicity associated with these types of CARs.\textsuperscript{46} While the induction of some cytokines in the latter system delayed tumor growth, the exact cytokine balance that will drive optimal CAR-T cell efficacy in different solid tumors is unknown.

As with CAR or TCR-based cellular approaches and other immunotherapies, combination therapies that target other cancer antigens or pathways would be expected to reduce the chances of loss of any single tumor antigen. Combination with oncolytic viruses for induction of innate and natural killer cell responses\textsuperscript{47,48} and vaccines that elicit endogenous Tcell activity could be key combination therapies to sustain tumor control, without the emergence of CAR-antigen loss variants.\textsuperscript{49–52} There is some evidence that endogenous T cells can be recruited in the ID8 tumor model,\textsuperscript{50,53} but their existence in the ID8 CAR system described here remains to be seen. Nevertheless, our findings using a single dose of CAR T-cells are even more notable given that other strategies would likely provide greater benefit through synthetic engineering, recipient T cell selections, or combination treatments.\textsuperscript{12,54}

**Author affiliations**

1. Carl R. Woese Institute for Genomic Biology and Cancer Center at Illinois, University of Illinois at Urbana-Champaign, Urbana, Illinois, USA
2. Department of Chemistry, University of Illinois at Urbana-Champaign, Urbana, Illinois, USA
3. Department of Biochemistry, University of Illinois at Urbana-Champaign, Urbana, Illinois, USA
4. Carl R. Woese Institute for Genomic Biology, University of Illinois at Urbana-Champaign, Urbana, Illinois, USA
5. Department of Pathology and David and Etta Jonas Center for Cellular Therapy, The University of Chicago, Chicago, Illinois, USA
6. Charles River Laboratories Inc Mattawan, Mattawan, Michigan, USA
7. Department of Veterinary Clinical Medicine, University of Illinois at Urbana-Champaign, Urbana, Illinois, USA
8. Department of Pathology, University of Illinois at Urbana-Champaign, Urbana, Illinois, USA
9. Department of Microbiology and Immunology, University of Illinois at Urbana-Champaign, Urbana, Illinois, USA
10. Department of Microbiology and Immunology, University of Illinois at Urbana-Champaign, Urbana, Illinois, USA
11. Department of Pathology and David and Etta Jonas Center for Cellular Therapy, The University of Chicago, Chicago, Illinois, USA
12. Charles River Laboratories Inc Mattawan, Mattawan, Michigan, USA
13. Department of Veterinary Clinical Medicine, University of Illinois at Urbana-Champaign, Urbana, Illinois, USA

**Contributors**

Conceptualization—DRER, DMK. Funding—HS, TMF, PJH, DMK. Data curation and formal analysis—DRER. DMK. Supervision—DRER, PJH, DMK. Validation—DRER, PS, CPS, ANL, EV, HS, KB, EJR, DMK. Investigation—DRER, PS, CPS, ANL, EV, VVWM, MVH, WM, SPW, KS, HS, KB, EJR, DMK. Methodology—DRER, PS, EJR, DMK. Writing—original draft—DRER, DMK. Writing and editing—DRER, PS, CPS, ANL, EV, SPW, KS, HS, KB, TMF, PJH, DMK. Guarantor—DMK.

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**Competing interests**

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All data relevant to the study are included in the article or uploaded as supplementary information.

**Supplemental material**

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**ORCID iDs**

Diana Rose E Ranoa http://orcid.org/0000-0002-2363-6111

Amber N Lewis http://orcid.org/0009-0008-3651-9062

David M Kranz http://orcid.org/0000-0002-4951-126X

**REFERENCES**


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Not applicable.


Of course, it is important to consider the role of CAR T cells in the treatment of cancer. For instance, the use of CAR T cells in ovarian cancer has shown promise. A recent study by et al. demonstrated that CAR T cells eradicate established tumors despite cancer-immune suppression. J Immunother Cancer 2022;10:e003959.


Lavrsen K, Madsen CB, Rasch MG, et al. A sensitivity scale for targeting T cells with Chimeric antigen receptors (CAR) and Bispecific T cell Engagers (bite). Oncoimmunology 2018;375:6583.


