Visualizing Spatial and Stoichiometric Barriers to Bispecific T-Cell Engager Efficacy

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ABSTRACT

Bispecific T-cell engager (BiTE) molecules are biologic T-cell–directing immunotherapies. Blinatumomab is approved for treatment of B-cell malignancies, but BiTE molecule development in solid tumors has been more challenging. Here, we employed intravital imaging to characterize exposure and pharmacodynamic response of an anti-muCD3/anti-huEGFRvIII mouse surrogate BiTE molecule in EGFR variant III (EGFRvIII)-positive breast tumors implanted within immunocompetent mice. Our study revealed heterogeneous temporal and spatial dynamics of BiTE molecule extravasation into solid tumors, highlighting physical barriers to BiTE molecule function. We also discovered that high, homogeneous EGFRvIII expression on cancer cells was necessary for a BiTE molecule to efficiently clear tumors. In addition, we found that resident tumor-infiltrating lymphocytes (TIL) were sufficient for optimal tumor killing only at high BiTE molecule dosage, whereas inclusion of peripheral T-cell recruitment was synergistic at moderate to low dosages. We report that deletion of stimulatory conventional type I DCs (cDC1) diminished BiTE molecule–induced T-cell activation and tumor clearance, suggesting that in situ antigen-presenting cell (APC) engagements modulate the extent of BiTE molecule efficacy. In summary, our work identified multiple requirements for optimal BiTE molecule efficacy in solid tumors, providing insights that could be harnessed for solid cancer immunotherapy development.

Introduction

In the past decade, dramatic improvements for the treatment of cancers have come from immunotherapy, particularly in the field of T cell–targeting therapies, where immune checkpoint inhibitors (ICI) and chimeric antigen receptor (CAR)-expressing T cells are currently approved for clinical use (1–3). Another promising strategy is the bispecific T-cell engager (BiTE; refs. 4, 5), a recombinant bispecific protein that has two linked single-chain variable fragments (scFv), one targeted to the CD3 coreceptor on T cells (e.g., CD3ε) and the other targeting a tumor-associated antigen (TAA; ref. 6). BiTE molecules have been shown to redirect and stimulate both CD4+ and CD8+ T-cell populations to kill tumor cells (7), independent of intrinsic T-cell receptor (TCR) and MHC-restricted recognition, and this mechanism-of-action has been validated clinically, most notably via the approval of CD19-targeting blinatumomab (Blinicyte) for the treatment of B-cell acute lymphoblastic leukemia (5, 8, 9).

One potential limitation of a BiTE molecule when targeting solid tumors is drug accessibility to the intended site of action. The molecular weights of BiTE molecules span a large range [~54 kDa for first-generation molecules through ~110 kDa for half-life extended (HLE) BiTE molecules], and solid tumor exposures could potentially be limiting (16), given the often narrow therapeutic index of this modality (4). Furthermore, the tumor vasculature is often heterogeneous, leading to additional physical barriers to drug extravasation and sufficient T-cell infiltration (17). There have been few reports that characterize BiTE molecule biodistribution to solid tumors (18), and it is anticipated that a combination of poor drug exposure within the often highly immunosuppressive TME remain as hurdles to BiTE efficacy (5). Ideally, real-time observation within tumors to define anatomic, cellular, and/or biochemical barriers to BiTE efficacy in the immunocompetent state could be beneficial for optimizing BiTE-based immunotherapy, including dissecting the potential role of ancillary immune components in the solid TME.

A second limitation of BiTE molecules is the selection of tumor-associated antigens (TAA), which have minimal background tissue expression to avoid off-tumor effects in nonmalignant cells and yet are of sufficiently high expression to allow high valency cross-linking of T cells. One putative TAA is EGFR with activating mutations specific to cancers, including the extensively studied...
EGFRvIII (deletion of exons 2–7) that promotes cell proliferation, angiogenesis, and invasion in different model systems (19). The EGFRvIII mutation occurs most commonly in glioblastoma (GBM) with inconsistent expression in breast, head and neck, lung, and prostate cancers reported (20, 21). Despite early evidence of clinical activity with the EGFRvIII BiTE AMG 596 in patients with GBM, the relationship between EGFRvIII expression and BiTE molecule activity remain unclear (12), especially given its known heterogeneity and possible downregulation in tumors (22).

Monitoring intratumoral T-cell responses has become important to understand the tumor response to immune manipulation and address questions related to effective immunotherapies (23). Intravital imaging (IVM) in tumors has revealed previously unknown immune processes inside the TME, including the link between T-cell behavior and dysfunctional differentiation (24), as well as the interaction of stimulatory, conventional type I dendritic cells (cDC1) and cytotoxic T cells (CTL; ref. 25) via real-time analysis of immune processes inside the tumor. IVM has also uncovered unexpected modes of action of ICIs and CAR T cells (26). In this study, we examined a mouse surrogate EGFRvIII-targeting BiTE molecule in a similarity adapted (syngeneic) breast cancer tumor model. IVM has also uncovered unexpected modes of action of ICIs and CAR T cells (26).

In this study, we examined a mouse surrogate EGFRvIII-targeting BiTE molecule in a similarly adapted (syngeneic) breast cancer tumor model to visualize BiTE molecule diffusion into solid tumors, with the goal of understanding the tissue pharmacokinetics in relation to the pharmacodynamic exposure–immune response in a fully immunocompetent mouse model. We visualized tumor exposure to a fluorescently labeled BiTE molecule and found it to be characterized by multiple distinct diffusion profiles from heterogeneous intratumoral vasculatures. By engineering variable target expression on tumor cells and by modulating tumor-infiltrating CTL and DC populations, we discovered critical, and sometimes unexpected, requirements for maximum BiTE molecule efficacy that included: (i) high, uniform target expression; (ii) a dose-dependent peripheral lymphocyte recruitment sensitivity that could be overcome at a high dosage when resident T-cell populations proved sufficient; and (iii) a role for ancillary CD103+ DC populations that enhanced T-cell expansion and recruitment.

Materials and Methods

Study design

The main objective of this study was to visualize and characterize human(anti)-EGFRvIII/murine(mu)CD3ε-muFc BiTE molecule tissue extravasation into EGFRvIII-expressing breast tumors and understand the key requirements for BiTE molecule efficacy in solid tumors. An immunocompetent mouse tumor model, as described below, was used in this study to better understand the complex interactions among the immune cells in both the peripheral immune system and the local TME of the solid tumor. Data for tumor intravital imaging was collected from three regions of interest (ROI) of a tumor, as indicated below, and was representative of >5 tumors from different mice. In the orthotopic tumor growth experiments, mice were randomly assigned to groups based on the tumor size so that control and experimental groups had the same average size before the start of treatment. The number of mice used for each group is indicated in the figure or figure legends. Animal numbers were chosen to ensure adequate statistical power and were based on the consistency of tumor growth observed in preliminary studies.

Mouse strains

All mice were housed and bred at University of California, San Francisco (UCSF), and were maintained under specific pathogen-free conditions in accordance with the regulatory standards of the NIH and American Association of Laboratory Animal Care standards and were with the approval of UCSF institution of Animal Care and Use Committee (IACUC approval no. AN170208-01B). C57BL/6 and Batf3−/− mice were purchased from The Jackson Laboratory. PyMT-ChOVA transgenic mice were generated as previously described (25). CD2-DsRed mice were obtained from Mark Coles (University of Oxford, Oxford, United Kingdom; refs. 24, 27). XCR1DTR mice were a gift from Tsuneyasu Kaisho (Osaka University, Osaka, Japan). All mice used in experiments were female age 8 to 20 weeks and bred onto a C57BL/6 background.

Cell lines

A PyMT-ChOVA cell line (PyMT-Control) was derived from PyMT-ChOVA spontaneous mammary tumors as previously described and verified by sequencing for the transgene cassettes (28). Cells tested negative for Mycoplasma on October 19, 2018, by PCR. Cell lines were used at less than five passages after thawing for the described experiments. To generate the PyMT-EGFRvIII cell line, a pLenti-humanEGFRvIII construct was generated by inserting the human EGFRvIII gene (NCBI Reference Sequence: NP_001333870.1) into the pLenti-puro vector (Addgene; #39481) and sequenced for accuracy. Lentivirus was produced in HEK293 cells (ATCC) using TransIT-Lenti Transfection Reagent (Mirus Bio). 500 μL of lentivirus containing supernatant were collected from HEK293 T cells after 48 hours and were applied to 5 × 10⁵ PyMT-ChOVA cells. Transduced cells were sorted for EGFRvIII-positive cells on day 2 after infection on a BD FACSAria III using a chimeric EGFRvIII human variable/murine IgG1 constant domain antibody generated as previously described (29). Following an additional week of culture, cells were sorted for a second time based on the expression EGFRvIII to establish PyMT-EGFRvIII-Hi (high expression of EGFRvIII) and PyMT-EGFR-Lo (low expression) cell lines. These two cell lines were stained for EGFRvIII and sorted into high- and low-expressing cells to maintain the proper EGFRvIII expression when needed. All the cell lines were cultured at 37°C in 5% CO₂ in DMEM (Invitrogen) with 10% FCS (Benchmark) and 1% penicillin–streptomycin–glutamine (Thermo Fisher Scientific).

BiTE molecule construction

The mouse surrogate BiTE molecule consisted of a human anti-huEGFRvIII scFv, anti-muCD3 (KE3) scFv linked via a SG4S linker, and a single-chain murine IgG1 Fc domain for half-life extension (HLE). The mouse surrogate CD3ε engager moiety, KE3, was based on a rat anti-mouse CD3ε scFv derived from the KT3 mb sequence (30). KT3 engagement on CD3ε has been demonstrated by competition for T-cell binding with the monoclonal CD3ε antibody, 145-2C11 (31). An analogous human-CD3ε–targeted HLE BiTE construct was used as a control in human breast cancer cell coculture assays that harbored an identical huEGFRvIII scFv, but with an anti-huCD3ε scFv sequence and single-chain human IgG1 Fc domain. The BiTE molecules were constructed by recombinant DNA technology, produced in stably transfected CHO cells, and purified by chromatography as described previously (32).

Characterization of CD3ε and EGFRvIII target binding affinity by SPR

The binding kinetics and affinity of the mouse surrogate anti-EGFRvIII/anti-muCD3ε BiTE-Fc (mulgG1) to either EGFRvIII or muCD3ε target was characterized by surface plasmon resonance
You et al.

(1) pH surface was regenerated between cycles using 10 mmol/L glycine, during association, and a 400-second interval of dissociation. The conducted at a in running buffer (HBS-Pþ (R&D Systems, #9565-ER-050) at 2 for the Meso Scale Discovery Analyzer (MSD) QuickPlex platform. an electrochemiluminescent (ECL) immunoassay was developed an du s to a l o a dr e c o m b i n a n tE G F R v I I - H i st a gp r o t e i n( S i n o Biological, #29662-H08B) to a maximum density of 180 RU. A serial a n du s et ol o a dr e c o m b i n a n tE G F R v I I - H i st a gp r o t e i n( S i n o Biological, #29662-H08B) to a maximum density of 180 RU. A serial

Pharmaceutical preparation and characterization of anti-EGFRVIII-Fc (muIgG1) was conducted in naive female C57BL/6 mice. The BiTE molecule was administered by intravenous (IV) bolus injection at either 0.05 mg/kg or 0.5 mg/kg using a formulat-

drug concentration–time profiles over 240 hours postdose (time points: 0.083, 1, 4, 8, 24, 48, 96, 168, and 240 hours). At each time point, approximately 0.06 mL of whole blood was collected via subman- dibular or tail vein puncture in SARSTEDT Micrucer serum separator tubes (Thermo Fisher Scientific) for processing serum. After whole blood was collected into the SARSTEDT tubes, the tubes were inverted several times and remained at room temperature for approximately 20 minutes or until fully clotted. CLOT formation was verified by manual inversion. The specimens were centrifuged using a calibrated Eppendorf 5417R Centrifuge (Brinkmann Instruments, Inc.). Blood specimens were centrifuged at 2 to 8 °C at approximately 11,500 rpm for 15 minutes. To quantify the concentration of anti-EGFRVIII BiTE molecule, an electrochemiluminescent (ECL) immunoassay was developed for the Meso Scale Discovery Analyzer (MSD) QuickPlex platform. Briefly, a streptavidin functionalized MSD plate (Meso Scale Diagnostics, LLC) was treated with biotinylated-huEGFRVIII pro-

A single-dose pharmacokinetics study of anti-huEGFRVIII mouse surrogate BiTE–Fc (muhlG1) was conducted in naive female C57BL/6 mice. The BiTE molecule was administered by intravenous (IV) bolus injection at either 0.05 mg/kg or 0.5 mg/kg using a formulat-
ed dosing solution of 0.0125 mg/mL and 0.125 mg/mL, respectively. Each study arm included n = 6 animals that were split into two staggered subgroups (n = 3) for sparse sampling to obtain concentration–time profiles over 240 hours postdose: (time points: 0.083, 1, 4, 8, 24, 48, 96, 168, and 240 hours). At each time point, approximately 0.06 mL of whole blood was collected via submandibular or tail vein puncture in SARSTEDT Microvette serum separator tubes (Thermo Fisher Scientific) for processing serum. After whole blood was collected into the SARSTEDT tubes, the tubes were inverted several times and remained at room temperature for approximately 20 minutes or until fully clotted. CLOT formation was verified by manual inversion. The specimens were centrifuged using a calibrated Eppendorf 5417R Centrifuge (Brinkmann Instruments, Inc.). Blood specimens were centrifuged at 2 to 8 °C at approximately 11,500 rpm for 15 minutes. To quantify the concentration of anti-EGFRVIII BiTE molecule, an electrochemiluminescent (ECL) immunoassay was developed for the Meso Scale Discovery Analyzer (MSD) QuickPlex platform. Briefly, a streptavidin functionalized MSD plate (Meso Scale Diagnostics, LLC) was treated with biotinylated-huEGFRVIII protein (R&D Systems, #9565-ER-050) at 2 μg/mL concentration to facilitate analyte capture. Serum samples were diluted up to 1:10 in Blotto solution (Thermo Fisher Scientific) prior to dispensation on the plate for room temperature incubation/capture (2 hours), along with a standard anti-EGFRVIII BiTE analyte curve (range 10,000–0.61 ng/mL) and quality control samples consisting of serum-spiked EGFRVIII BiTE analyte at variable nominal concentra-
tion to assess accuracy throughout the full concentration range of the prepared standard curve. Postincubation, the MSD plates were washed with KPL buffer (LGC Clinical Diagnostics, Inc.), followed by addition of polyclonal anti-mouse IgG1 (Jackson ImmunoResearch, #115-005-205) in Blotto (1 μg/mL) that had been previously conjugated to a MSD SULFO-TAG (Meso Scale Diagnostics, LLC). After incubation and multiple plate wash steps in KPL buffer, tripropylamine read buffer (Meso Scale Diagnostics, LLC) was added to enable detection of electrically stimulated light emission at 620 nm wavelength, where the instrument responses are proportional to the amount of anti-EGFRVIII BiTE molecule contained in a given serum sample. For quantitation, the ECL assay response versus anti-EGFRVIII BiTE molecule concentra-
tion relationship was regressed using a four-parameter logistic (Marquardt) regression model to estimate lower and upper curve plateaus, the Hill slope, and E50 using a weighting factor of 1/Y2. Noncompartmental analysis (NCA) was performed on serum concentration–nominal time data using Phoenix WinNonlin (Version 6.4.0.768, Pharsight Corporation). The pharmacokinetic parameters estimates for AUC, clearance (CL), volume of distribu-
tion at steady state (Vss), mean residence time (MRT), and terminal phase elimination half-life (t1/2,z) are reported.

Tumor model

A total of 2 × 105 PyMT-Control (Ctrl) cells, or PyMT-EGFRVIII-Hi, or PyMT-EGFRVIII-Lo tumor cells were suspended in PBS and mixed 1:1 with Matrigel GFR (Corning) for a final injection volume of 50 μL. For the mixed tumor experiment, PyMT-Ctrl and PyMT-EGFRVIII-Hi cells were mixed at ratio of 3:1 or 1:3 for a total number of 2 × 107. Cells were injected in the fat pad of C57BL/6 mice anesthetized with 3% isofluorane (Henry Schein). Tumors were then measured three times per week using electronic calipers. Tumor volume was calculated through the formula V = 0.5(w2 × l). Mice were euthanized when tumors exceeded a volume of 1,000 mm3. To block lymphocyte egress from lymph nodes (LN), 50 μL FTY729 (Cayman Chemical Company) at 1 mg/kg was injected intraperitoneally everyday beginning on day 21 post tumor injection and continued through the end of the experiment. 50 μL DMSO was injected as mock treatment in control groups. To deplete cDC1s in XCR1 DTR mice, diphtheria toxin (DT; List Biological Laboratories) at 500 ng/mouse was injected intraperitoneally every three days beginning on day 10 to day 25 post tumor injection. The same dosage of DT was injected into wild-
type C57BL/6 mice as the control groups.

Intravitral imaging of mouse tumor and analysis

PyMT-EGFRVIII-Hi tumor-bearing mice were kept under anesthesia using 2% isoflurane, and mammary tumors were surgically exposed when palpable. Confocal imaging was performed using a Nikon A1R laser scanning confocal microscope with NIS-Elements software (Nikon Instruments) and a 16X LWD water-dipping objective. Before surgery, 75 μg DyLight 649–labeled Griffonia Simplicifolia Lectin I isoelectric B4 (Vector Laboratories) was injected by retro-orbital injection for visualization of blood vessels. During imaging, 20 μg AF488–labeled BiTE molecule was injected by retro-orbital injection as

700 Cancer Immunol Res; 10(6) June 2022 CANCER IMMUNOLOGY RESEARCH
described in figure legends. Time-lapse videos were acquired every 90 seconds with a 2-step of 3 μm. Images were taken at 100-μm deep inside the tumor and in the center areas with obvious blood vessels, where tumor cells were also abundant. Data were analyzed using Imaris (Bitplane), including drift correction, video generation, cell surface detection, and generation of cell speed and track displacement data.

**Analysis of in vivo BiTE molecule dispersal and target binding**

The fluorescence intensity from AF488-labeled HLE-BiTE was used to calculate the spatiotemporal kinetics of intratumoral BiTE diffusion. Using Distance Transformation on Imaris, the distance of each voxel from the nearest isolectin-labeled blood vessel was calculated and binned into 5-μm blocks. The average BiTE intensity of all voxels within each bin was then calculated and centered spatially at the midpoint of the bin (e.g., the average intensity of the bin from 10-15 μm would be centered at 12.5 μm). Next, we used the BiTE channel intensity prior to BiTE injection (usually a 15- to 20-minute time-lapse was obtained before BiTE injection) to subtract the background autofluorescence. Finally, to dissociate the absolute amount of dispersal from its spatiotemporal dynamics, the intensities were normalized to the minimum and maximum intensities observed at that field of view (Fig. 3B-G), as follows:

\[ I_{\text{norm}} = \frac{I - I_{\text{min}}}{I_{\text{max}} - I_{\text{min}}} \]

To quantify the essence of the patterns of BiTE dissemination, an effective diffusion coefficient (\( D_{\text{eff}} \)) was calculated (Fig. 3H) using the relationship

\[ R^2 = D_{\text{eff}} \cdot t \]

where \( R \) is distance traversed by the molecule from its source in time \( t \). Although this relationship is only approximate for complex mass transport scenarios, such as drug dispersal in a tumor, it offers a quick and reliable comparison of transport kinetics across fields of view. The time of entry (t) of the BiTE at a particular spatial location x-μm away from the nearest labeled vasculature was taken to be the first time point when the intensity threshold at that spatial bin reached a value >10. The theoretical \( D_{\text{eff}} \) of AF488-labeled HLE-BiTE (MW ~ 120 kDa) in the interstitial fluid at 37°C was calculated using Stokes–Einstein equation \( D_{\text{eff}} = k \rho T / 6\pi \eta r \), where \( k = 1.38 \times 10^{-23} \text{J/K} \), \( T = 310 \text{K} \), \( \eta = 7 \times 10^{-3} \text{kg m}^{-1} \text{s}^{-1} \), \( r \sim 4 \text{nm} \).

The frequency of the specific diffusion type was calculated using the volume of the ROI in which the specific type of diffusion appeared divided by the total volume of the tissue that we imaged from that tumor. For the contained type, the percentage was calculated using 15 ROIs from 5 mice using the same method. To calculate BiTE binding kinetics on target cells, the target cells were first delineated by surface rendering on Imaris, and the average BiTE fluorescence intensity on these surfaces was calculated. Using similar distance transformations as described above, the minimum distance of these target cells from the vasculature were also concurrently obtained. These were then used to plot traces of T-cell-associate BiTE accumulation over time on individual cells proximal to the vasculature (Fig. 2C) or an aggregate of cells at different distances from the vasculature (Fig. 3I).

Similarly, larger images obtained from PyMT-Ctrl or PyMT-EGFR-Hi or PyMT-EGFRvIII-Lo tumor tissue sections fixed by 2% paraformaldehyde (Electron Microscopy Sciences) for overnight were used to quantify BiTE distribution and retention in the tumor microenvironment. Average voxel-by-voxel intensity and distance transformations were used, as described above. A sum of all BiTE intensity across same-sized fields of view were used as a surrogate to measure total amount of BiTE dispersed a given time after injection, as shown in Supplementary Fig. S5C. Analysis of Imaris-processed images was done on Matlab (Mathworks Inc.) using custom-Matlab scripts and either Matlab or GraphPad Prism was used to plot the analyzed data. Matlab scripts are available upon request.

**Breast cancer samples, digestion, and coculture assays**

Three breast tumor samples were acquired from the NCI’s Cooperative Human Tissue Network (CHTN) and from the Department of Surgery, Feinberg School of Medicine, Northwestern University (Chicago, IL). Specimens were anonymized and then collected and distributed by CHTN. Specimens are covered under collection/distribution of tissues under consent or waiver of consent. Samples were minced with surgical scissors and transferred to a GentleMACs C Tube (Miltenyi Biotec) with 100 μg/mL Liberase TL (Roche) and 200 μg/mL DNase I (Roche) at 3 mL per gram of tissue. C tubes were incubated in the GentleMACS octo dissociator (Miltenyi Biotec) with heaters, following the manufacturer’s disassociation protocol (Miltenyi Biotec Tumor Dissociation Kit). 10 mL of FACS buffer (PBS + 2% FCS) was added to samples and filtered through a 100-μm filter and centrifuged at 350 × g for 5 minutes at 4°C, and red blood cells were lysed with 175 mM ammonium chloride. CD45+ live cells from the breast cancer tumor were then sorted on a FACS Aria II (BD) and cocultured with CD8+ T cells at a 1 (12,500 cells) to 8 (1,250,000 cells) ratio for 72 hours. For both tumor and LNs, cells were washed with PBS plus 2% FCS and filtered through a 100-μm filter and centrifuged at 350 × g for 5 minutes at 4°C, and red blood cells were lysed with 175 mM ammonium chloride. CD45+ live cells from the breast cancer tumor were then sorted on a FACS Aria II (BD) and cocultured with CD8+ T cells at a 1 (12,500 cells) to 8 (1,250,000 cells) ratio for 72 hours. The huCD3/huEGFrVIII BiTE molecule (obtained from Amgen Inc.) was added at the indicated concentrations. CD8+ T cells were isolated from PBMCs of a healthy donor using EasySep Human CD8+ T-cell Isolation Kit (StemCell Technologies). T cells stimulated with Dynabeads Human T-Activator CD3/CD28 (Thermo Fisher Scientific) were used as controls.

**Mouse tumor and lymph node digestion**

For tumor digests, tumors from mice were isolated, minced, and incubated in digestion buffer containing 100 U/mL collagenase type I (Roche), 500 U/mL collagenase type IV (Roche), and 200 μg/mL DNase I (Roche) in RPMI1640 (Gibco) for 30 minutes on a shaker at 37°C. For LN digests, tumor-draining LNs (the inguinal, axillary, and brachial) were dissected, pierced, and torn with sharp forceps, and incubated in the same digestion buffer as tumors for 15 minutes at 37°C. Digestion mixtures were then pipetted repeatedly, followed by another 15-minute incubation. For both tumor and LNs, cells were washed with PBS plus 2% FCS and filtered through a 100-μm cell strainer before staining for flow cytometry. The number of BiTE molecules bound to T cells or tumor cells was calculated using quantitative fluorescence cytometry as described below.

**Quantitative fluorescence cytometry**

The number of BiTE molecules bound to T cells or tumor cells was calculated based on the degree of labeling (1.5 fluorochrome per BiTE) using quantitative fluorescence cytometry of BiTE molecules. The Quantum Alexa Fluor 488 MEF kit (Bangs Laboratories, Inc) was used per the manufacturer’s instructions. The numbers of molecules of the fluorochrome were measured to be 9,000 for T cells and 4,500 for tumor cells, so the numbers of BiTEs bound to T cells and tumor cells were 6,000 and 3,000, respectively.
Flow cytometry antibodies

The following mouse antibodies were purchased from BioLegend and were used for surface staining: anti-CD45–Bright Violet (BV)510, –BV605, or –Alexa Fluor (AF)488 (clone 30-F11); anti-CD90.2–AF700 (clone 30-H12); anti–CD8a–PerCP/Cy5.5 (clone 53-6-7); anti–CD4–BV421 (clone RM4-5); anti–CD11b–BV785 or –BV510 (clone M1/70); anti–CD44–BV711 (clone IM7); anti–CD69–BV650 or –PE (clone H1.2F3); anti–CD279 (PD-1)–BV605 (clone 29F.1A12); anti–I–A/I–E–AF711 (clone M5/114.15.2); anti–CD11c–BV650 (clone N418); anti–CD19–BV785 (clone 6D5); anti–B220–BV785 (clone RA3–82B); anti–Ly6G–BV785 (clone 1A8); anti–CD24–PE/Cy7 (clone M1/69); anti–CD103–APC (clone 2E7); anti–NK1.1–BV785 (clone PK136); anti–F4/80–AF488 (clone BM8); anti–Ly6C–AF700 (clone H1K1.4); anti–CD62L–APC (clone MEL-14); and anti–CD25–APC (clone 3C7) Anti–IFNγ–APC (clone XMG1.2); anti–TNFα–PE (clone MP6–XT22), and anti–granzyme B–PE (clone Cy Janeiro7) (clone QA16A02) were purchased from BioLegend; anti–Ki–67–PE–eFluor 610 (clone SolA15) was purchased from Thermo Fisher Scientific. These antibodies were used for intracellular staining. Fluorochrome minus one control was used to identify positive populations.

The following human antibodies were used for surface staining: anti–CD45–APC/Cy7 (clone 30-F11); anti–CD25–APC (clone BC96); anti–CD45R0–AF700 (clone UC7H1), anti–CD69–PE (clone FN50), and anti–CD8a–BV605 (clone RPA–T8). All were purchased from BioLegend.

Flow cytometry

Cells from tumor digests or cell lines were stained with fixable Live/Dead Zombie NIR (BioLegend) in PBS per the manufacturer’s instructions, followed by staining with anti-Fc receptor (clone 2.4G2; BioLegend) and antibodies to surface antigens in FACS buffer for 30 minutes on ice. Intracellular staining was subsequently performed using the eBioscience Foxp3/Transcription Factor Staining Buffer Set (Thermo Fisher Scientific) when needed (all antibodies listed in the section above). For intracellular cytokine staining, cells were incubated in RPMI1640 plus 10% FCS supplemented with 10 ng/mL PMA (Sigma–Aldrich), 0.5 μg/mL ionomycin (Thermo Fisher Scientific), and 10 μg/mL Brefeldin A (Sigma–Aldrich) for 4 hours before staining. For sorting, cells were stained with anti-Fc receptor (clone 2.4G2) and antibodies to surface antigens, followed by DAPI staining (BioLegend) to exclude dead cells. Flow cytometry was performed on BD Fortessa or LSR II cytometer. Cell sorting was performed using BD FACS Aria 2 or Aria Fusion. Analysis of flow cytometry data was done using FlowJo (Treestar) software.

Coupling assay and T cell–tumor cell in vitro cocultures

Mouse CD8+ T cells from naïve C57BL/6j mice were isolated from LNs using the CD8 EasySep enrichment kit (STEMCELL Technologies) and were stained with Violet Proliferation Dye 450 (VPD; BD Biosciences). For the coupling assay, 1 × 10^6 PyMT–Control or PyMT–EGFRVIII cells were labeled with CellTracker Orange CMTMR Dye (Thermo Fisher Scientific) per the manufacturer’s instructions. Labeled CD8+ T cells and PyMT cells at a 1 to 1 ratio were cocultured in 96-well V bottom plate for 1 hour at 37°C in 100 μL RPMI1640 plus 10% FCS with 1% penicillin–streptomycin–glutamine (Thermo Fisher Scientific) with a gentle spin-down before the incubation. After incubation, 100 μL 4% paraformaldehyde (PFA) per well was added into each well for 10 minutes, and cells were washed with FACS buffer. T cell–tumor doublets were measured by flow cytometry as described in the Flow cytometry section. For T cell–tumor cell long-term cocultures, cells were collected at the indicated time points, and flow cytometry was performed to determine T-cell activation states and tumor killing effect.

Calcium influx imaging

Mouse CD8+ T cells from naïve C57BL/6 mice were isolated as described previously and were stimulated in vitro with Dynabeads Mouse T-activator CD3/CD28 for T-Cell Expansion and Activation (Thermo Fisher Scientific) plus IL2 (50 IU, Roche) for 2 to 3 days in RPMI-1640 plus 10% FCS with 1% penicillin–streptomycin–glutamine (Thermo Fisher Scientific). Activated T cells were harvested and labeled with Fura-2 AM (Thermo Fisher Scientific) per manufacturer’s instructions. PyMT–EGFRVIII cells were labeled with CellTracker Orange CMTMR Dye (Thermo Fisher Scientific). T cells and PyMT–EGFRVIII cells were mixed at 1:1 ratio and incubated in 0.5% low-melting agarose in RPMI1640 without Phenol red ( Gibco; T-cell imaging medium) in Nunc Lab–Tek II Chambered Coverglass (Thermo Fisher Scientific) for 15 minutes before imaging. Time-lapse videos were acquired every 90 seconds using a widefield inverted Zeiss AxioScope 200M microscope equipped with a 20× long working distance lens, an Andor Clara CCD Camera, and 340 and 380 excitation filters paired with a Lambda XL light source (Sutter Instruments). Samples were kept at 37°C using a heated stage top incubator with 5% CO2. 95% O2. BiTEs were added after 270 seconds of imaging.

Cell death imaging

Mouse CD8+ T cells from naïve C57BL/6 mice were stained with Violet Proliferation Dye 450 (VPD; BD Biosciences). PyMT–EGFRVIII cells were labeled with the Live/Dead Cell Imaging Kit (Thermo Fisher Scientific) per manufacturer’s instructions. T cells and tumor cells were cocultured in T-cell imaging medium, as described in the previous section, plus the Live/Dead staining reagent and imaged as indicated in the section above.

Statistical analysis

Statistical analyses were performed using GraphPad Prism software. Data are shown as mean ± SD. Specifically, an unpaired Student t test or one-way ANOVA with multiple comparison test was used as indicated in the figure legends, with P values described in detail.

Data and material availability

All data are available in the main text or the Supplementary Materials. Matlab code for imaging analysis is available to share upon request.

Results

HuEGFRVIII/muCD3: surrogate BITE molecule and PyMT–EGFRVIII model

A previously developed anti–huEGFRVIII/muCD3 BiTE construct (AMG 596) was investigated in patients with glioblastoma in clinical trial NCT03296696 (12). In this study, we constructed an analogous EGFRVIII-targeted, mouse surrogate molecule with a similar half-life extended (HLE–BITE) design based on murine IgG1 Fc. For this, an anti–EGFRVIII scFv moiety was fused to the anti-murine CD3ε engagement domain (KE3), along with a murine IgG1 Fc domain harboring a N297G mutation to minimize Fc effector function (huEGFRVIII/muCD3ε–muFc BITE) and to impart favorable pharmacokinetics (Fig. 1A). The single-dose serum pharmacokinetics was characterized in C57BL/6

Cancer Immunol Res; 10(6) June 2022

CANCER IMMUNOLOGY RESEARCH
between mouse CD8\(^{\text{Fc}}\) domain harboring a N297G mutation to minimize Fc effector function. MW, molecular weight; kDa: kilodaltons.

Live/Dead Zombie dye.

EGFRvIII-Hi cells for 48 hours in the presence of different concentrations of BiTEs as indicated. Measurement of tumor cell death was done by staining with VPD dilution), activation (indicated by surface CD25 and CD69 expression), and tumor killing induced by BiTEs when T cells were cocultured with PyMT-ChOVA cells expressing fluorescent mCherry protein to aid visualization of tumors (25). Given that altered T-cell function has been shown to correlate with changes in dynamic behavior inside PyMT tumors (24, 33), this model provides an ideal imaging platform to study the relationship between BiTE molecule exposure within the TME and pharmacodynamic response, including target engagement and cell killing. Moreover, normalization of the tumor vasculature in this mouse model enhances drug delivery and delays tumor growth (24), supporting it as a valid mouse model to study drug diffusion properties. We overexpressed huEGFRvIII on a PyMT-ChOVA cell line and sorted the cells with high expression (EGFRvIII-Hi) to access BiTE diffusion and efficacy. To understand if target expression influenced BiTE function, for some studies we also sorted cells with low EGFRvIII expression (PyMT-EGFRvIII-Lo), which was roughly 6 to 7 times lower than its expression on EGFRvIII-Hi cells (Supplementary Fig. S2A). The use of PyMT tumors engineered to express EGFRvIII antigen was further validated as a relevant model of human disease by screening several primary breast cancer samples, where we detected variable expression of EGFRvIII on CD45+ immune cells (28). PyMT-ChOVA cells express fluorescent mCherry protein to aid visualization of tumors (25). Given that altered T-cell function has been shown to correlate with changes in dynamic behavior inside PyMT tumors (24, 33), this model provides an ideal imaging platform to study the relationship between BiTE molecule exposure within the TME and pharmacodynamic response, including target engagement and cell killing. Moreover, normalization of the tumor vasculature in this mouse model enhances drug delivery and delays tumor growth (34), supporting it as a valid mouse model to study drug diffusion properties. We overexpressed huEGFRvIII on a PyMT-ChOVA cell line and sorted the cells with high expression (EGFRvIII-Hi) to access BiTE diffusion and efficacy. To understand if target expression influenced BiTE function, for some studies we also sorted cells with low EGFRvIII expression (PyMT-EGFRvIII-Lo), which was roughly 6 to 7 times lower than its expression on EGFRvIII-Hi cells (Supplementary Fig. S2A). The use of PyMT tumors engineered to express EGFRvIII antigen was further validated as a relevant model of human disease by screening several primary breast cancer samples, where we detected variable expression of EGFRvIII on CD45+ immune cells (28). PyMT-ChOVA cells express fluorescent mCherry protein to aid visualization of tumors (25). Given that altered T-cell function has been shown to correlate with changes in dynamic behavior inside PyMT tumors (24, 33), this model provides an ideal imaging platform to study the relationship between BiTE molecule exposure within the TME and pharmacodynamic response, including target engagement and cell killing. Moreover, normalization of the tumor vasculature in this mouse model enhances drug delivery and delays tumor growth (34), supporting it as a valid mouse model to study drug diffusion properties. We overexpressed huEGFRvIII on a PyMT-ChOVA cell line and sorted the cells with high expression (EGFRvIII-Hi) to access BiTE diffusion and efficacy. To understand if target expression influenced BiTE function, for some studies we also sorted cells with low EGFRvIII expression (PyMT-EGFRvIII-Lo), which was roughly 6 to 7 times lower than its expression on EGFRvIII-Hi cells (Supplementary Fig. S2A). The use of PyMT tumors engineered to express EGFRvIII antigen was further validated as a relevant model of human disease by screening several primary breast cancer samples, where we detected variable expression of EGFRvIII on CD45+ immune cells (Supplementary Fig. S2B–S2D), with the highest expression present in a triple-negative breast cancer patient, which was about 50% lower compared with the PyMT-EGFRvIII-Hi cell line, whereas the EGFRvIII expression on other breast cancer samples was similar to that of PyMT-EGFRvIII-Lo cells. Furthermore, adding the analogous huCD3ε/huEGFRvIII BiTE molecule in an in vitro coculture assay consisting of the isolated CD45+ populations from the triple-negative breast cancer sample in combination with allogenic human CD8+...
You et al.

T cells from a healthy donor induced T-cell activation similar to that observed with the surrogate mouse BiTE molecule (Supplementary Fig. S3), suggesting this to be a high-fidelity mouse surrogate molecule.

CD8+ T-cell activation and tumor killing by huEGFRvIII/muCD3ε BiTE in vitro

To further validate the efficacy of the huEGFRvIII/muCD3ε BiTE molecule, we assessed its ability to simultaneously engage both effector and target cells using a flow cytometry-based trans-coupling assay, where T cell (CD8+) - PyMT-EGFRvIII-Hi cell doublet formation was monitored as a function of BiTE molecule concentration. The BiTE was observed to efficiently mediate the formation of T cell-PyMT cell doublets in an EGFRvIII-dependent manner (Fig. 1B), yet in line with theoretical predictions, cell–cell coupling was observed with increasing BiTE molecule concentration to a maximum doublet frequency, followed by a decrease in coupling efficiency above the 100 nmol/L concentration, consistent with the classic Hook effect anticipated for three-body binding equilibria (35). The huEGFRvIII/muCD3 BiTE molecule also induced a dose-dependent calcium influx, but only when T cells were in close contact with PyMT-EGFRvIII-Hi cells (Fig. 1C; Supplementary Video S1). In addition, more T cells were observed in close proximity to tumors with BiTE molecule treatment (Supplementary Fig. S4A).

We next examined the tumor killing efficiency of the mouse surrogate BiTE molecule in vitro and observed PyMT-EGFRvIII-Hi tumor cell death within 2 hours at ≥1 nmol/L BiTE molecule concentration in cocultures with CD8+ T cells (Fig. 1D; Supplementary Video S2). Additional immunophenotyping conducted at 48 hours demonstrated dose-dependent expression of the activation markers, CD25 and CD45RO, expansion of T-cell numbers, and tumor killing (Fig. 1E). We validated a similar upregulation of CD25 and CD45RO of allogenic human T cells when mixed with a EGFRvIII-expressing human breast cancer sample and the human BiTE molecule (Supplementary Fig. S3). No effect was seen if T cells were cocultured with PyMT-control cells (PyMT-Ctrl), demonstrating that BiTE molecule–TCR binding alone is not sufficient for T-cell activation, consistent with the intended mechanism of action for this modality. In total, we successfully validated an in vitro tumor–T-cell system and demonstrated that huEGFRvIII/muCD3ε BiTE molecule-activated T cells could kill tumors expressing EGFRvIII in a dose-dependent manner, with the anticipated potency observed with primary human cell types exposed to the human-targeted BiTE molecule.

Characterization of huEGFRvIII/muCD3ε BiTE molecule in solid tumors

To extend our characterization of the BiTE molecule dose–response to the added complexity inherent in an immunocompetent mouse, we examined the effects of target engagement on T cells in vivo with intravitreal imaging (IVM) of PyMT-EGFRvIII-Hi tumor and combined multicolor flow analysis of disaggregated tumor harvests to characterize the TME (Fig. 2A). To this end, an AF488-labeled BiTE molecule (average degree of labeling/DoL ~1.5±1) was administered at 20 μg/animal (~0.8 mg/kg) retro-orbitally, where fluorescence indicated accumulation on T cells or tumor cells predominantly near the intratumoral vasculature (Fig. 2B and C). Surface punctalike structures were visible at T cell–tumor cell interfaces by 2.5 hours postdose, suggesting that clusters of CD3-BiTE-EGFRvIII bridged cellular complexes (Fig. 2B). We also visualized real-time T-cell recruitment to target tumor cells in regions near the vasculature (Supplementary Fig. S4B; Supplementary Video S3).

To better understand the pharmacokinetics in tumor tissues and to dissect BiTE cellular partitioning within the TME, we quantified the fluorescent BiTE molecule bound to PyMT-EGFRvIII tumor cells or CD8+ tumor-infiltrating lymphocytes (TIL) over 72 hours. At a low BiTE dosage (2 μg), preferential accumulation on CD8+ T cells was observed (Fig. 2D and E), with up to ~75% of CD8+ T cells harboring a bound BiTE molecule, but only ~2% of isolated PyMT-EGFRvIII cells produced a detectable BiTE molecule signal. In contrast, at the higher 20 μg dose, both cell types were equivalently saturated. This phenomenon would not be anticipated for a monovalent binding event, based on the target affinity indicated by Kd measurement of EGFRvIII and CD3ε (Supplementary Fig. S1A), nor given that target expression (and cellularity) within the TME would intuitively favor the higher affinity, higher abundance EGFRvIII target. However, the preferential accumulation on T cells could be due to the better accessibility of the BiTE molecule to T cells in blood and peripheral lymphoid organs (Supplementary Fig. SSD), followed by the migration of the BiTE molecule–bound T cells to the tumor. This effect would be more obvious with a low BiTE dosage because BiTE T-cell binding in blood could be significant, contributing to a sufficient “sink” effect. We also noted that maximum BiTE molecule exposure occurred within the tumor at ~12 hours (tmax) following a single dose, inferred by the absence of a further increase in BiTE-AF488 intensity on target cells at subsequent times (Fig. 2E; Supplementary Fig. SSA–SSC).

Using bead-based calibration, we determined that at tmax (~12 hours post 20 μg dose), CD8+ T cells and mCherry+ tumor cells bound to ~6,000 and ~3,000 BiTE-AF488 molecules per cell, respectively (Fig. 2E; see methods). A dose dependence of CD8+ target cell coverage was also observed in 2-cell–rich lymph node nodules (Supplementary Fig. SSD), although a more dramatic difference in T-cell coverage at 20 μg versus 2 μg was observed, possibly indicating that abundant EGFRvIII expression in tumor tissue may stabilize, or otherwise enhance, binding or retention of the drug to T cells. These drug concentrations were efficacious based on the induction of extensive tumor killing 72 hours postinjection, particularly by high-dose BiTE therapy (Fig. 2F and G).

Heterogeneous tissue extravasation of BiTE molecule within solid tumor

We next visualized the impact of the heterogeneous tumor vasculature on early BiTE molecule extravasation into the tumor space in PyMT-EGFRvIII-Hi tumors (Fig. 3A; Supplementary Video S4). Following injection of BiTE-AF488, we observed three distinct patterns of diffusion (Fig. 3B, D, and F). By plotting normalized pixel intensities against the distance from vasculature over time, we described these as normally diffusive, irregular diffusive, and contained. (Fig. 3C, E, and G). Diffusive behavior was observed in 39.2% of fields as a gradual and steady decrease in BiTE molecule intensity away from vessels (Fig. 3B and C). This was best modeled by a constant slope, when time of BiTE molecule entry was plotted against the square of the distance (Fig. 3H), consistent with minimal physical barriers and/or indicating that binding sites for BiTE molecules were homogeneous throughout the region of interest. In stark contrast, in 28.2% of fields, we also often classified BiTE molecule extravasation as irregular (Fig. 3D), which was characterized by a sharp slope of BiTE molecule signal gain at less than 25 μm from the vessel boundary, followed by a blunt increase at 30 to 50 μm away from the vasculatures (Fig. 3E). BiTE molecule extravasation kinetics in this regime were found to be biphasic (Fig. 3H), with the calculated Doff being almost five times higher when BiTE molecules were closer to the vessels.
than when they were away. The theoretical $D_{eff}$ of this AF488-labeled, HLE-BiTE molecule in the interstitial fluid is about 450 $\mu m^2/minute$, calculated using Stokes–Einstein equation for the diffusion coefficient of particles undergoing Brownian motion (see Materials and Methods), which is an order or two magnitudes higher than the $D_{eff}$ estimated from IVM, indicating binding site barriers and/or physical barriers to BiTE molecule diffusion. Finally, a third, tightly restricted BiTE-AF488 ("Contained", Fig. 3F and G) profile was also observed in 33% of fields, with insignificant signal for BiTE molecule above the limit of detection emanating beyond the vessel boundary, even at 2.5 hours. This BiTE molecule diffusion heterogeneity was observed both across separate tumor-bearing animals, and within a given tumor for an individual animal. Such variation has predicted consequences of accumulation of BiTE molecules on CD3$^+$ cells (Fig. 3I; Supplementary Fig. S6). Under the diffusive condition, BiTE molecule binding to T cells accumulated at a steady rate, regardless of where T cells were located (Fig. 3I). However, in the irregular/biphasic case, appreciable BiTE molecule binding to T cells was restricted to the regions

Figure 2.
BITE-mediated target-binding and tumor killing in breast tumors. A, Schematic of the experimental design. PyMT-EGFRvIII-Hi cells and CD2-DsRed mice were used. Panels with corresponding data are indicated. B, Representative images showing BiTEs (20 $\mu g$) binding to T cells and tumor cells near vessels (<10 $\mu m$). Blue outlines, tumor cells; red outlines, T cells. Scale bar, 15 $\mu m$. C, Quantification of individual T cell- or tumor cell-associated BiTE intensity over time. Each colored line represents one individual cell. Histogram (D) and quantification (E) of BiTE-AF488 binding on CD8$^+$ T cells or PyMT-EGFRvIII-Hi mCherry$^+$ tumor cells at two dosages and different time points as indicated. NT, no treatment. Representative flow plots (F) and quantification (G) of live tumor cells out of all CD45$^-$ live cells or all live cells in the tumors at 72 hours with high or low dosage of BiTEs. Live cells were gated using Zombie Aqua fixable viability dye and then mCherry$^+$ cells were gated as live tumor cells. $N=3$ per group (D–G). Data are representative of two independent experiments; ***$, P<0.0001$, **$, P<0.001$, $^*, P<0.01$, $^!, P<0.05$ as determined by one-way ANOVA and multiple comparison test.

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Cancer Immunol Res; 10(6) June 2022 705
of rapid “leaky” dissemination (<25 μm from vessels, Fig. 3D and E). Such pronounced heterogeneity in HLE-BiTE molecule tumor extravasation could limit efficacy in a clinical setting.

High and homogenous target expression maximized BiTE efficacy

To characterize the significance of these findings, we varied the requisite target expression for BiTE molecule efficacy in our immunocompetent mouse model, using the high- and low-expressing lines, spanning the range of human tumors examined (Supplementary Fig. S2). The BiTE diffused farther away from the vasculature in PyMT-EGFRvIII-Hi tumors compared with low-expressing tumors at 4 hours post BiTE injection, with a higher binding to tumor cells. This difference was more pronounced when measured at 72 hours (Supplementary Fig. S7). A single 10-μg dosage failed to clear PyMT-EGFRvIII-Lo tumors (Fig. 4A and B) and did not significantly increase overall CD8⁺ T cells compared with PyMT-Ctrl tumors (Fig. 4C). In contrast, the same dose given to mice bearing PyMT-EGFRvIII-Hi tumors resulted in complete tumor clearance (Fig. 4A and B) and significant CD8⁺ T-cell expansion (Fig. 4C), suggesting that high
target expression either enhanced intratumoral T-cell proliferation or recruited more T cells from lymphoid tissues, or both. Immunophenotyping demonstrated increased CD44 and IFNγ expression in EGFRvIII-Hi tumors but not in EGFRvIII-Lo tumors, as well as increased proliferative capability indicated by Ki67 (Fig. 4D and E).

Given that the expression of a specific tumor antigen is often variable in solid tumors, we mixed EGFRvIII-Hi tumor cells with control tumor cells at two different ratios of 3:1 or 1:3 to determine whether the BiTE molecule could lead to clearance of nontargeted tumors. Despite that a higher percentage of EGFRvIII-expressed cells resulted in smaller overall tumor size at every time point, full tumor regression was not observed in either condition. This indicated that, although effective against EGFRvIII+ tumors, BiTE molecule-induced killing of tumors was incomplete when not every cell had high expression (Supplementary Fig. S8).

Peripheral T-cell recruitment for BiTE efficacy is dose dependent

We next sought to understand whether the BiTE molecule-induced tumor clearance required peripheral T cells to be recruited
from lymphoid tissues. T-cell egress from lymph nodes was blocked by treatment with FTY720, an S1P agonist (36), and this was evidenced by a reduction in total and naive (CD44−CD62L+) circulating blood CD8+ T cells compared with mock-treated animals (Fig. 5A and B). At a high BiTE molecule intravenous dosage (20 μg), FTY720 treatment did not have a significant effect and permitted complete clearance of PyMT-EGFRvIII-Hi tumors, similarly to controls. In contrast, low BiTE molecule dosage (2 mg) was sufficient for complete regression of PyMT-EGFRvIII-Hi tumors, but FTY720 treatment significantly decreased BiTE molecule antitumor activity (Fig. 5C). IFNγ and granzyme B production from tumoral CD8+ T cells for the indicated groups of mice was not differentially upregulated by high-dose BiTE under FTY720 treatment compared with control groups. However, it was lower in FTY720-treated groups with low-dose BiTE therapy (Fig. 5D and E). The data taken together suggest a significant contribution of cell recruitment from the lymphatics that regulates T-cell function in the TME with BiTE treatment, but that limitations produced by this can be overcome with a high BiTE dosage due to sufficient activity from resident TILs.

**Stimulatory DCs boosted local T-cell pool to enhance BiTE efficacy**

We extended our investigation of the mechanisms of BiTE molecule–mediated T-cell activation in solid tumor to focus on the role of other immune cells potentially involved. Specifically, we focused on CD103+ intratumoral cDC1 because they can potently stimulate T cells to drive antitumor responses (37). Batf3 is the master transcriptional factor for cDC1 development (38), and we used Batf3 knockout mice (Batf3−/−) to understand if cDC1s were required for CD8-mediated tumor killing in response to BiTE treatment. In PyMT-EGFRvIII-Hi tumors in the absence of BiTE molecule, the loss of CD103+ DCs did not change the already-robust tumor outgrowth nor affect the CD4+ T-cell population but did result in reduced numbers of CD8+ T cells in tumors, with those...
present having similar expression of effector molecules (e.g., granzyme B and IFNγ; Supplementary Fig. S9A–S9C). In contrast to controls, in Batf3−/− mice, the BiTE molecule was unable to elicit clearance of PyMT-EGFRvIII-Hi tumors (Fig. 6A and B). As expected, CD103+ DCs were largely absent in the tumors of the Batf3−/− group, whereas other DC populations remained unchanged (Supplementary Fig. S9D; Fig. 6C). Concordant with failed tumor clearance, we also observed significantly lower IFNγ and granzyme B production in the global absence of CD103+ DCs in treated mice (Fig. 6D). We also found that DCs were critical in a genetic model in which DCs were present throughout development but were depleted just days prior to BiTE treatment using XCR1 DTR mice (Supplementary Fig. S10A and S10B; refs. 39, 40). Taken together, the data suggest that an established ongoing endogenous APC, is necessary to permit optimal BiTE function.

**Discussion**

Here, we identified key requirements of perfusion, target expression, lymph node involvement, and a DC-driven endogenous response for promoting BiTE efficacy. We found these requirements using a preclinical solid tumor mouse model with endogenous immunocompetency, whose features closely mimicked human disease, including target expression and heterogeneity, and the model also allowed a clinical drug surrogate to be studied to determine pharmacologic barriers to its efficacy in the context of a solid tumor.

Prior to this work, other studies have described the efficacy of various types of T-cell bispecific molecules in diverse solid tumor models (41), including glioblastoma (42), prostate (43), colorectal (44), breast (45), and lung (46) cancers. In those models, immuno-deficient mice were typically transplanted with human cancer cell lines or primary tumor tissue and supplemented with exogenous human T cells. Although these studies provide strong evidence of target-dependent T-cell activation, proliferation, and infiltration into the tumor and tumor regressions induced by BiTE immunotherapy, they are incapable of mimicking the complex interactions of immune cells in both the peripheral immune system and the TME of solid tumors or depicting BiTE molecule diffusion patterns into solid tumors.

The pharmacokinetics of immunotherapeutic reagents defining the exposure of biologic immunotherapies within solid tumors has remained an underappreciated area of study to rationalize efficacy, or lack thereof, in patients. Numerous studies have shown how vascular transport of many drugs into tissue are limited by poorly perfused tumor vessels due to vessel compression, hyperpermeability, abnormal morphology, and organization (47). Lessons learned from immune checkpoint therapy development include strategies to normalize vascularization in solid tumors, highlighted by promising improvements in anti–PD-1/PD-L1 efficacy in lung and kidney cancers (48). Prior to our study, the actual biodistribution of BiTE molecules within the heterogeneous TME was poorly defined and remains a significant barrier to our understanding of the tumor accessibility of half-life–extended, T cell–directing therapies. Our study suggests that some tumors may impose additional diffusion barriers distal to the vessels themselves. At present, we are unable to...
You et al.

directly measure how significantly this limits overall tumor exposure but surmise that such a barrier could potentially complicate BiTE therapy.

Our results provide a first and direct picture of BiTE molecule extravasation in a solid tumor model through intravital imaging. We found that within a tumor, the vascular permeability to the BiTE molecule exhibited a high degree of variability, with several distinct diffusion topographies, ranging from facile, or widespread “leakage”, to vessels of low apparent permeability that limited tissue/target exposure to therapy. Such variable diffusion patterns suggest that BiTE molecule extravasation within solid tumors may be inconsistent in the clinical setting, and that vascularization is just one of the criteria determining tumor accessibility. Given enough time and a high dosage of BiTE therapy, we observed ample coverage of the tumor by the drug in our model. However, the tumor vasculature usually differs between mouse and human tumors (49, 50) and such differences affect the efficacy of anti-tumor drugs (51). Therefore, an even more restricted diffusion pattern could theoretically exist in the clinical setting that potentially limits drug exposure. The heterogeneous diffusion patterns of the BiTE molecule revealed in our study highlighted that in some cases, better delivery of the drug may be needed to improve BiTE efficacy in solid tumors. One important caveat to our analysis is that our observations were conducted after single dose BiTE treatment, whereas typical clinical cycles involve multiple doses over several weeks. Thus, it is possible that sufficient tumor exposures are achieved when dosing to steady state under the variety of tumor architectures probed here. The relationships between total vascular surface area, vessel size, and receptor expression and their effects on both active and passive (paracellular) BiTE molecule transport to the tumor interstitium warrant further study.

The abundance of TILs is a positive prognostic indicator for some tumor types and is also a potential predictor of the responsiveness to immune checkpoint inhibitors in patients with cancer (52, 53). The number of TILs is determined by both T-cell expansion in situ, and the recruitment of peripheral T cells. BiTEs have been shown to increase TILs in many xenograft models (54, 55), as well as in our immunocompetent PyMT-EGFRvIII model presented here; however, the additional requirement for T-cell recruitment from the periphery, most notably from the lymphoid organs, for efficient activity leading to tumor clearance was not possible to assess prior to this analysis. One possible explanation for this requirement is that suboptimal BiTE dosages preferentially bind to T cells in the periphery and not the tumor and thus their trafficking becomes a requirement to carry the agent into the tumor and affect clearance. Our study demonstrated that a high TIL count may be prognostic of the potential for maximal BiTE efficacy, and that at lower tumor exposures, we clearly delineated the role of peripheral T-cell populations in maintaining antitumor responses. Our numerous observations also suggested not only variable TIL content within a given tumor region, but also unpredictable dynamic behaviors for TILs, ranging from active swarm like phenotypes to T-cells apparently at rest prior to BiTE molecule dosing (Supplementary Video S3 vs. Video S4, respectively). The clinical relevance of baseline T-cell dynamic cellular phenotypes in relation to BiTE molecule responses in solid tumor is unknown and is an area in need of further study.

In addition, high, homogenous EGFRvIII target abundance was determined to be a critical component for maximal efficacy. In this investigation, we mimicked the potential for heterogeneous target expression within the TME by grafting tumor cell lines with varying EGFRvIII expression, which in turn was found to impact BiTE molecule potency significantly. Our study indicates that low target expressers may require a higher BiTE molecule exposure to be completely cleared. Nonuniform target expression is a potential limiting factor for efficacy of any T-cell directing therapy, and the impact of target heterogeneity is likely context-dependent.

Among the abundant immunosuppressive myeloid cells residing in the TME, CD103⁺ cDC1 in mice (defined rather by expression of BDCA-3 in humans), demonstrate potent cross-presentation capability and the ability to stimulate CD8⁺ T cells to drive antitumor immune responses (37). The abundance of this type of stimulatory DC in solid tumors positively correlates with increased patient survival and predicts responsiveness to anti-PD-1 immunotherapy (56). Although BiTE molecules stimulate T cells regardless of MHC restriction, the demonstration that BiTE efficacy required DCs for optimal outcomes was not previously known. Consistent with previous findings about the role of Batf3⁺ DCs in effector T-cell trafficking (57), it is most likely that CD103⁺ DC deficiency diminished BiTE efficacy in solid tumors due to a deficiency in the number of available T cells to be acted upon by BiTE molecules. Although clinical data is currently lacking, we propose that the abundance of CD103⁺/BDCA-3⁺ DCs, a rare myeloid population in solid tumors, may be a key predictive factor for responsiveness to BiTE immunotherapy in human solid tumors.

Authors’ Disclosures

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Authors’ Contributions

R. You: Conceptualization, data curation, formal analysis, methodology, writing—original draft, writing—review and editing. J. Artighoker: Conceptualization, data curation, formal analysis, methodology. A. Ray: Data curation, formal analysis, methodology. H. Gonzalez Velozo: Resources, methodology. D.A. Rock: Conceptualization, resources, supervision, funding acquisition. K.P. Conner: Conceptualization, resources, data curation, formal analysis, supervision, funding acquisition, methodology, writing—original draft, writing—review and editing. M.F. Krummel: Conceptualization, resources, supervision, funding acquisition, writing—original draft, writing—review and editing.

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