Combining bempegaldesleukin (CD122-preferential IL-2 pathway agonist) and NKTR-262 (TLR7/8 agonist) improves systemic antitumor CD8\(^+\) T cell cytotoxicity over BEMPEG+RT

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ABSTRACT

Background Tumor cell death caused by radiation therapy (RT) triggers antitumor immunity in part because dying cells release adjuvant factors that amplify and sustain dendritic cell and T cell responses. We previously demonstrated that bempegaldesleukin (BEMPEG: NKTR-214, an immunostimulatory IL-2 cytokine prodrug) significantly enhanced the antitumor efficacy of RT through a T cell-dependent mechanism. Because RT can induce either immunogenic or tolerogenic cell death, depending on various factors (radiation dose, cell cycle phase), we hypothesized that providing a specific immunogenic adjuvant, like intratumoral therapy with a novel toll-like receptor (TLR) 7/8 agonist, NKTR-262, would improve systemic tumor-specific responses through the activation of local innate immunity. Therefore, we evaluated whether intratumoral NKTR-262 combined with systemic BEMPEG treatment would elicit improved tumor-specific immunity and survival compared with RT combined with BEMPEG.

Methods Tumor-bearing mice (CT26; EMT6) received BEMPEG (0.8 mg/kg; intravenously), RT (12 Gy \(\times\) 1), and/or intratumoral NKTR-262 (0.5 mg/kg). Flow cytometry was used to evaluate CD4\(^+\) and CD8\(^+\) T cell responses in the blood and tumor 7 days post-treatment. The contribution of specific immune subsets was determined by depletion of CD4\(^+\), CD8\(^+\), or NK cells. CD8\(^+\) T cell cytolytic activity was determined by an in vitro CTL assay. Data are representative of 1–2 independent experiments (n=5–14/ group) and statistical significance was determined by 1-way analysis of variance (ANOVA) or repeated measures ANOVA (p value cut-off of 0.05).

Results BEMPEG+NKTR-262 significantly improved survival compared with BEMPEG+RT in a CD8\(^+\) T cell-dependent manner. Response to BEMPEG+NKTR-262 was characterized by a significant expansion of activated CD8\(^+\) T cells (GzmA\(^+\); Ki-67\(^+\); ICOS\(^+\); PD-1\(^-\)) in the blood, which correlated with reduced tumor size (p<0.05). In the tumor, BEMPEG+NKTR-262 induced higher frequencies of GzmA\(^+\) CD8\(^+\) T cells exhibiting reduced expression of suppressive molecules (PD-1\(^-\)), compared with BEMPEG+RT (p<0.05). Further, BEMPEG+NKTR-262 treatment induced greater tumor-specific CD8\(^+\) T cell cytolytic function than BEMPEG+RT.

Key messages

What is already known on this topic

- Combining systemic and local therapies can induce highly effective antitumor immunity. For example, combining systemic IL-2 therapy with local radiation therapy (RT) improved responses over IL-2 alone, but produced limited abscopal effects. Here, we sought to evaluate the extent to which combining BEMPEG (immunostimulatory IL-2 cytokine prodrug) with intratumoral toll-like receptor 7/8 agonist (NKTR-262) therapy would augment tumor-specific responses and increase survival as compared with BEMPEG+RT.

How this study might affect research, practice or policy

- BEMPEG+NKTR-262 induced superior systemic adaptive immunity and survival as compared with BEMPEG+RT. These data demonstrate the potential therapeutic potential of BEMPEG+NKTR-262 and suggest that clinical evaluation of this combination is warranted.

Conclusions BEMPEG+NKTR-262 therapy elicited more robust expansion of activated CD8\(^+\) T cells compared with BEMPEG+RT, suggesting that intratumoral TLR stimulation provides superior antigen presentation and costimulatory activity compared with RT. A clinical trial of BEMPEG+NKTR-262 for patients with metastatic solid tumors is in progress (NCT03435640).

INTRODUCTION

While applications of immunotherapies, either individually or in combination with other treatments, continue to improve patient outcomes,1,2 determining which immunotherapy combinations achieve the best possible outcome is a critical and unresolved issue. Identifying highly efficacious combination therapies will require elucidating the
molecular mechanisms by which they induce antitumor responses and an in-depth, side-by-side characterization of different therapies. Here, we compare efficacy of two combination therapies, bempegalsleukin (BEMPEG, immunostimulatory IL-2 cytokine prodrug)\(^3\)\(^-\)\(^5\) combined either with radiation therapy (RT) or with NKTR-262, a toll-like receptor (TLR) 7/8 agonist.

Cytokine-based immunotherapies aim to increase the proliferation and survival of pre-existing antitumor effector T cells. For example, high-dose IL-2 (HD IL-2) is an FDA-approved cytokine therapy with a 15%–20% objective response rate in metastatic renal cell carcinoma and melanoma.\(^6\)\(^-\)\(^8\) However, HD IL-2 efficacy is limited by the expansion of suppressive CD25\(^+\) FoxP3\(^+\) T regulatory (Treg) cells\(^9\) and by treatment toxicity, which includes vascular leak syndrome, hypotension, and liver toxicity.\(^10\) One prodrug modified to reduce those toxicities is BEMPEG, an engineered IL-2R agonist with six releasable polyethylene glycol (PEG) units attached to the IL-2R\(\alpha\) binding region. These PEG units increase the duration of IL-2 receptor agonism and preferentially reduce binding to IL-2R\(\alpha\) (CD25)\(^5\) compared with IL-2R\(\beta\)\(\delta\), supporting effector T cell expansion over Treg expansion, and thereby increasing efficacy and reducing toxicity. Indeed, BEMPEG-induced T cell activation and expansion activity in vivo increases Teff:Treg ratios in tumor tissue compared with IL-2.\(^4\)

Because BEMPEG supports adaptive immunity, we sought to evaluate BEMPEG in combination with different innate immune agonists capable of boosting antitumor immunity, aiming to enhance overall therapeutic potential. Both RT and TLR targeting have different innate immunostimulatory effects that can unleash antitumor CD8\(^+\) T cell responses.\(^11\)\(^-\)\(^12\) RT-induced tumor cell death results in increased cross-presentation of tumor antigen,\(^13\) increased numbers of IFN-\(\gamma\)-secreting tumor-specific tumor-infiltrating lymphocytes (TIL),\(^14\) and increased expression of chemokines that promote T and NK cell trafficking to the tumor.\(^15\)\(^-\)\(^16\) TLR7/8 stimulation can induce functional APC differentiation,\(^17\) induce Th1-biased responses,\(^18\) and enhance CD8\(^+\) T cell effector functions.\(^11\) Furthermore, TLR7/8 agonists have demonstrated antitumor activity in preclinical models.\(^19\) BEMPEG synergizes with RT and provides the greatest benefit to immunologically ‘hot’ (well infiltrated) tumors,\(^16\) but only a modest benefit to immunologically ‘cold’ (poorly infiltrated) tumors.\(^20\) The modest benefit observed may be a result of the immunosuppressive effects elicited by RT, which include inactivating NK cells, recruiting myeloid-derived suppressor cells (MDSCs), and altering macrophage polarization towards an M2 tumor-promoting phenotype.\(^21\)

Whether the innate stimulation provided by the TLR7/8 agonist NKTR-262 combined with BEMPEG will improve response rates over BEMPEG +RT is unknown. We hypothesized that the proinflammatory signals provided via intratumoral NKTR-262 would enhance the priming of tumor-reactive T cells that would then be supported by systemic BEMPEG treatment, resulting in BEMPEG +NKTR-262 eliciting more robust tumor regression than BEMPEG +RT. To address these hypotheses, we comprehensively characterized functional and phenotypic immune responses induced by these two combination therapies, which provided insight into the mechanisms associated with efficacious combination therapies.

**MATERIALS AND METHODS**

**Mice**

Wild-type 6–8 week-old C57BL/6 and BALB/c mice (Jackson Labs; Bar Harbor, ME), and Nur77-GFP transgenic mice (Dr. Andrew Weinberg; EACRI, Portland, OR)\(^25\) were bred in the EACRI facility. Mice were maintained under specific pathogen-free conditions in the Providence Portland Medical Center animal facility.

**Tumor cell lines**

CT26 (colon carcinoma, BALB/c), EMT6 (mammary carcinoma, BALB/c), and MCA-205 (fibrosarcoma, C57BL/6) tumor cell lines (all from ATCC) were maintained in complete RPMI-1640 (cRPMI; 10% FBS, 10 mmol/L HEPES, 1% non-essential amino acids, sodium pyruvate (Lonza), and penicillin–streptomycin–glutamine (Invitrogen)). Cell line identity was verified through monthly assessment of morphology and growth kinetics. Cell lines were tested annually using the MycoAlert mycoplasma detection kit (Lonza, Basel, Switzerland).

**In vivo tumor studies**

Mice were inoculated with \(1\times10^6\) (CT26 and MCA-205) and \(1\times10^5\) (EMT6) tumor cells in dual-flank subcutaneous injections. Tumor growth was monitored using two-dimensional (length \(\times\) width) caliper measurements 2–3 \(\times\)/week. Treatments began 10 days following implant, when tumors reached 50–80 mm\(^2\). Mice received control (diluent; 10 mM citric acid, 7% trehalose, pH 4, iv or HBSS it), RT (12 Gy), BEMPEG (Nektar Therapeutics, San Francisco, CA) (10 mg/kg, intravenously), NKTR-262 (Nektar Therapeutics, San Francisco, CA) (0.8 mg/kg, intravenously), NKTR-262 +RT, or BEMPEG +RT, or BEMPEG +NKTR-262 concurrently. CT-guided photon RT with a beam energy of 220 kV was delivered using a Small Animal Radiation Research Platform (XStrahl, Gulmay Medical, Suwanee, GA) to an isocenter in the center of the tumor. Dosimetry was performed using Murislice software (XStrahl), and irradiated lesions received 12 Gy in a single fraction using opposed tangential fields. For survival experiments, tumor size was monitored until all animals reached a primary endpoint (tumor-free or total tumor burden >250 mm\(^3\)). In BALB/c animals, CD4\(^+\) and CD8\(^+\) T cell depletion experiments were performed using 200 \(\mu\)g anti-CD4 (clone GK1.5) given 1 \(\times\)/week (ip) for 6 weeks, or 200 \(\mu\)g anti-CD8 (clone 53–6.7) delivered once (ip) on day 9. In C57BL/6 animals, NK depletions experiments were performed using 200 \(\mu\)g aNK1.1 (PK136, BioXcell).
delivered 1× ip starting the day before therapy. For IFNAR blocking experiments, 200 µg aIFNAR (MAR1-5A3, BioXcell) was delivered on days 9, 13, 17, and 21 (ip) post-tumor implants. For statistical analyses, endpoints were defined as the first time point that tumor area exceeded 250 mm² or was non-palpable and did not recur.

**Blood, tumor, and lymph node collection and processing**

Peripheral blood (PBL) samples were drawn on day 7 post-treatment; 25 µL of fresh heparinized blood was incubated with fluorescence-conjugated antibodies (online supplemental table 2) for 30 min at 4°C in the dark. Tumors were harvested 3 or 7 days post-treatment, cut into small fragments, and digested in 1 mg/mL collagenase and 20 mg/mL DNase (Sigma) in serum-incubated with fluorescence-conjugated antibodies (online supplemental table 2) for 30 min at room temperature (RT). TIL were filtered through 70 µm nylon mesh (Cell Treat), washed with 10 mL cRPMI, and collected by centrifugation (1500 rpm, 4 min). Pelleted cells were resuspended for staining and analysis by flow cytometry (see below). Lymph nodes (LNs) were harvested 7 days post-treatment and processed to obtain single-cell suspensions. Red blood cells were lysed with ACK buffer (Lonza) for 2 min and processed to obtain single-cell suspensions and 20 mg/mL DNase (Sigma) in serum-incubated with fluorescence-conjugated antibodies (online supplemental table 2) for 30 min at 4°C in the dark. Tumors were harvested 3 or 7 days post-treatment, cut into small fragments, and digested in 1 mg/mL collagenase and 20 mg/mL DNase (Sigma) in serum and incubated with fluorescence-conjugated antibodies (online supplemental table 2) for 30 min at 4°C in the dark.

**Flow cytometry**

For PBL, TIL, and LN phenotyping, single cell suspensions were stained with intracellular targets (online supplemental table 2). Cells were fixed and permeabilized following manufacturer’s instructions (FoxP3/Transcription Factor Staining Buffer Set, ThermoFisher, San Diego, CA) and stained with intracellular targets (online supplemental table 2). Flow cytometry data were acquired on an LSR II flow cytometer running FACSDiva software (BD Biosciences), and data were processed and analyzed with FlowJo (BD Biosciences).

**Cytokine bead array**

PBL was incubated in a 96-well round bottom plate coated with agonistic aCD3/aCD28 (100 µL solution per well of aCD3 (5 µg/mL, 145-2 C11, BD Biosciences) and aCD28 (2 µg/mL, 37.5.1, BD Biosciences)) prior to staining. Plates were incubated for 44 hours in cell culture conditions (37°C, 5% CO₂, 95% humidity) for a cytokine bead array using a ProcartaPlex Mouse Cytokine/Chemokine Panel 1A 36-Plex kit (EPX360-26012-901; Invitrogen). Data were acquired on a Luminex 200 (R&D Systems).

**In vitro T cell coculture assays**

Cancer cells were plated at a density of 2000 cells/well in a 96-well plate. After 24 hours, adherent cells were rinsed twice with cRPMI supplemented with β-ME. CD8⁺ T cells were isolated and sorted (BD FACSAria) from CT26 tumors 7 days post-BEMPEG+RT or BEMPEG+NKTR-262 or vehicle control therapy. Isolated CD8⁺ T cells were added to the cancer cell cultures at a 25:1 effector:target ratio. Cells settled for 15 min at RT prior to hourly tracking of cellular confluence and death (Caspase 3/7 Green Dye, Sartorius) in the Incucyte (Sartorius, Goettingen, Germany), housed at 36.5 C and 5% CO₂ until untreated cells reached confluence. Cancer cell growth and death were analyzed using Zoom software (Incucyte, Sartorius).

**Statistical analysis**

Data presented in box and whisker plots: the line within the box indicates median, the box spans the IQR, and the whiskers extend to the highest and lowest observations. We used one-way analysis of variance (ANOVA) or repeated measures ANOVA as appropriate along with Šídás or Dunnett’s multiple comparisons tests. Spearman’s correlation was used to correlate PBL and TIL populations with tumor size, and Kaplan-Meier plots and log-rank tests were used for tumor survival analysis. All statistical analyses were performed using GraphPad Prism software (GraphPad, San Diego, CA) or R. A p <0.05 was considered significant.

**RESULTS**

**BEMPEG+NKTR-262 improves survival over BEMPEG+RT in a CD8⁺ T cell-dependent manner**

To compare efficacy of BEMPEG +RT against BEMPEG +NKTR-262, we chose two tumor models: CT26 (colon carcinoma) and EMT6 (mammary carcinoma). CT26 allowed us to track tumor-specific (AH1-A5 tetramer⁺) responses and based on our historical data indicating a 46% survival rate following BEMPEG +RT, there was room to statistically distinguish either an improved or worsened response to BEMPEG +NKTR-262. EMT6 is less responsive to RT and therefore was selected as a secondary tumor model. Tumors were established subcutaneously in the bilateral flanks of BALB/c mice; 10 days later, animals received monotherapies of either 12 Gy RT or 10 µg NKTR-262 in the treated tumor (intratumorally, it) or 0.8 mg/kg BEMPEG (intravenously). For combination therapies, mice received BEMPEG +RT or BEMPEG +NKTR-262. Tumor growth was assessed over time. PBL and, in following experiments, TIL were assessed for phenotype and function 7 days post-treatment (figure 1), which is the peak of the adaptive immune response and before tumor regression impairs our ability to evaluate TIL.

In the treated tumor (right side), we observed a significantly reduced tumor size by day 17 post-treatment following all therapies compared with control (figure 2A–B, p<0.05, online supplemental table 1). In the non-treated tumor (left side), we saw a significantly reduced tumor size by day 17 post-treatment only following BEMPEG (p<0.05), BEMPEG+RT (p<0.01), or BEMPEG+NKTR-262 (p<0.01) compared with control (figure 2A–B, online supplemental table 1), demonstrating the systemic influence of BEMPEG on these local therapies. In the non-treated tumor, we observed a significantly larger tumor size after RT, NKTR-262, and BEMPEG in comparison to BEMPEG+NKTR-262.
n=0/5 for BEMPEG+R 100% of control mice (n=0/7 for BEMPEG+NKTR-protected against tumor growth while tumors grew in following rechallenge with CT26 tumors, and all were surviving mice were tested for tumor-EMT6 tumor model (online supplemental figure 1). All efficacy depends on CD8+ T cells, with some contribution of TR-
tumors after BEMPEG+NKTR-62, we depleted NK cells.16 To address this for BEMPEG+NKTR-262, we
monotherapies to BEMPEG+RT (figure 2A–B, online supplemental table 1). The difference between BEMPEG+NKTR-262 and BEMPEG+RT therapy is seen when comparing the treated and non-treated tumors within each group. There was a significant difference in tumor size between the non-treated and treated tumors after BEMPEG+RT (p<0.05) but not after BEMPEG+NKTR-262, where both non-treated and treated tumors reduced in size at a similar rate (figure 2C). Importantly, most non-treated tumors following BEMPEG+RT grew out (13/15; 86.6%; figure 2A,D), similar to our previous observations.16 In contrast, many of the non-treated tumors after BEMPEG+NKTR-262 cured. This resulted in significantly increased survival after BEMPEG+NKTR-262 (9/15; 60%) versus BEMPEG+RT (2/15; 13.3%) (figure 2D, p<0.05). We confirmed these results in the EMT6 tumor model (online supplemental figure 1). All surviving mice were tested for tumor-specific memory following rechallenge with CT26 tumors, and all were protected against tumor growth while tumors grew in 100% of control mice (n=0/7 for BEMPEG+NKTR-262; n=0/5 for BEMPEG+RT; n=5/5 untreated control mice).

We previously demonstrated that BEMPEG+RT efficacy depends on CD8+ T cells, with some contribution of NK cells.16 To address this for BEMPEG+NKTR-RT, we performed depletion studies (online supplemental figure 2A) and found that efficacy requires CD8+ T cells (0% survivors) but not CD4+ T cells (37% survivorship vs 36% for BEMPEG+NKTR-262) (figure 2E–F, online supplemental figure 2B). Notably, even without CD8+ or CD4+ T cells, BEMPEG+NKTR-262 therapy induced a significant delay of both treated and non-treated tumor growth in comparison to control (p<0.05 at d14). Because TLR7/8 agonists can activate NK cells directly,23 we depleted NK cells and determined that NK cells neither influenced overall survival nor eliminated BEMPEG+NKTR-262-delayed tumor growth, as tumor size was still significantly smaller than control by day 13 (p<0.05; online supplemental figure 2C), suggesting BEMPEG+NKTR-262 induced an NK-independent early response (day 1–5) sufficient to delay tumor growth.

**BEMPEG+NKTR-262 therapy increases the frequency of activated CD8+ T cells in the PBL in comparison to BEMPEG+RT**

Our data suggested that BEMPEG+NKTR-262 induced a more potent systemic CD8+ T cell-dependent antitumor response than BEMPEG+RT; thus, to understand the efficacious systemic response to BEMPEG+NKTR-262, we performed LN (online supplemental figure 3) and PBL (figure 3) immunophenotyping 7 days post-therapy. Focusing on the CT26 model, we examined frequencies of CD8+, NK, CD4+FoxP3+ T regulatory (Treg), and CD4+FoxP3+ T effector (Teff) cells, and markers including PD-1, TIM-3, LAG-3, CD62L, ICOS, Ki-67, granzyme A (GzmA), CD25 (IL-2Rα), CD122 (IL-2Rβ), and AH1-A5. In the LN, we found patterns of CD8+ T cell activation, with average frequencies of CD8+ AH1-A5, GzmA+, IFN-γ, and TNF-α all increased 4–5-fold over control after BEMPEG+NKTR-262 but not after BEMPEG+RT (online supplemental figure 3A). Although statistically significant, the overall frequencies of these populations were small. For example, GzmA+ CD8+ T cells ranged from only 1.5%–4.5% after BEMPEG+NKTR-262 (online supplemental figure 3B). Therefore, we focused on PBL for further analysis of peripheral immune responses. For PBL samples, we correlated cell types, phenotypes, and functions with tumor size across treatment groups (figure 3A). We found no significant correlations between PBL phenotypes and treated tumor size; however, we found many significant negative correlations (p<0.05) between CD8+ T cell markers and non-treated tumor size, including ICOS+, AH1-A5+, PD-1+, Ki-67+, and GzmA+ (figure 3A), which highlights the value of the peripheral response in monitoring the non-treated tumor response. Comparing CD8+ T cell phenotypes between treatment groups revealed that BEMPEG+NKTR-262 significantly expanded CD8+ T cells over BEMPEG+RT (figure 3B,D; p<0.0001). The CD8+ T cell phenotypes that significantly distinguished BEMPEG+NKTR-262 from BEMPEG+RT were PD-1, CD122, GzmA, ICOS, and Ki-67 (for all: p<0.0001), indicating that BEMPEG+NKTR-262 is superior to BEMPEG+RT at expanding active functional CD8+ T cells (figure 3B–D).
Teff and Treg populations did not have many differences between combination treatment groups because many phenotypes were driven by BEMPEG and not altered by additional therapy (online supplemental figure 4A). However, CD25+ and ICOS+ frequencies were significantly increased in Teff and Treg after both BEMPEG monotherapy and BEMPEG+NKTR-262 combination therapy in comparison to BEMPEG+RT, suggesting a negative effect of RT on these phenotypes (online supplemental figure 4A).

To determine whether these therapies altered Th1/Th2 polarization, we analyzed serum cytokine levels via multiplex ELISA. We found an increase in IFN-γ (Th1) and a decrease in IL-4 (Th2) and IL-3 (immune regulating cytokine), suggesting a more inflammatory helper T cell response in the periphery after BEMPEG+NKTR-262 in comparison to BEMPEG+RT (online supplemental figure 4B). Other proinflammatory cytokines induced by BEMPEG and significantly elevated after BEMPEG+NKTR-262 compared with BEMPEG+RT included IL-1β,
Figure 3  BEMPEG+NKTR-262 induces a greater expansion of active CD8+ T cells than BEMPEG+RT. (A) We determined immune phenotypes from PBL 7 days post-therapy. These data are presented as Spearman correlations of PBL phenotypes with tumor size, across individuals and treatment groups. Significant correlations after FDR correction are indicated. (B) Representative flow cytometry plots of CD45+, CD4+, and CD8+ gates. (C) Representative flow cytometry plots demonstrating CD8+ cell expression of CD62L, PD-1, GzmA, ICOS, Ki-67, and AH1-A5 after BEMPEG+RT (blue) or BEMPEG+NKTR-262 (red). (D) PBL immune phenotypes determined by flow cytometry. Box and whisker plots represent the min and max (whiskers), the quartiles (box) and median (line). Each point represents an individual mouse. N=10–20, from two independent experiments, except AH1-A5, which is one of two representative experiments N=4–8 per experiment. One-way ANOVA with Šidák’s multiple comparisons test. (E) (Left) Representative flow cytometry plot showing AH1-A5+ and GzmA+ CD8+ T cells. (right) Granzyme A (GzmA), proliferation (Ki-67), and CD62L expression on tumor specific (AH1-A5+, filled circle) or not (AH1-A5-, open circle). Only mice that had more than 70 AH1-A5+ cells were analyzed for AH1-A5+ phenotypes. Data from one of two representative experiments. For comparisons between AH1-A5+ within one treatment group, Student’s t-test. For comparisons among treatment groups, one-way ANOVA with Šidák’s multiple comparisons test. *P<0.05, **p<0.01, ***p<0.001, ****p<0.0001. ANOVA, analysis of variance; FDR, false discovery rate; PBL, peripheral blood; RT, radiation therapy.
IL-1α, GM-CSF, and RANTES (CCL5) (online supplemental figure 4B), which are associated with the inflammatory process and leukocyte recruitment, further supporting that BEMPEG+NKTR-262 induces an immune response that supports antitumor immunity.

AH1-A5+ CD8 T cells were expanded over control in 50% of the mice that received BEMPEG+NKTR-262 therapy, but not expanded over control in any mice after BEMPEG+RT; however, statistically there was no difference between combination therapies (figure 3D). Due to the lack of difference in AH1-A5+ CD8 T cell frequency between the two combination therapies, we thought there may have been a broadening of the tumor-specific TCR repertoire after BEMPEG+NKTR-262 therapy due to NKTR-262-induced antigen presenting cell (APC) differentiation. Therefore, we examined AH1-A5+ and AH1-A5+ CD8 T cells in depth, hypothesizing that tumor-specific AH1-A5 cells would express similar levels of activation markers as AH1-A5+. Indeed, we found similar frequencies of proliferating (Ki-67+) AH1-A5 cells after BEMPEG+NKTR-262 as of AH1-A5+ after BEMPEG+RT and significantly more proliferating AH1-A5+ CD8 T cells after BEMPEG+NKTR-262 than BEMPEG+RT (p<0.0001; figure 3E). Furthermore, GzmA MFI was statistically indistinguishable comparing AH1-A5+ to AH1-A5+ CD8 T cells from BEMPEG+NKTR-262-treated tumors, and GzmA MFI from AH1-A5+ CD8+ T cells after BEMPEG+NKTR-262 was significantly increased in comparison to AH1-A5+ CD8+ T cells after BEMPEG+RT (p<0.01) (figure 3E). Similarly, CD62L MFI (figure 3E), was significantly reduced in AH1-A5+ CD8+ T cells after BEMPEG+NKTR-262 compared with BEMPEG+RT (p<0.05). This data could suggest that the BEMPEG+NKTR-262 induced AH1-A5+ CD8+ T cells were activated ‘bystander’ CD8+ T cells driven by proinflammatory cytokines such as type I interferons (IFN) induced by TLR agonists, or that they were activated tumor-specific CD8+ T cells that targeted a different tumor antigen, perhaps an indication of increased priming after BEMPEG+NKTR-262 therapy.

IFN-α/β signaling contributes to BEMPEG+NKTR-262 efficacy

Type I IFN signaling supports bystander CD8+ T cell activation and drives increased cross-presentation capacity of dendritic cells (DCs) after RT. Therefore, we blocked the IFN alpha/beta receptor (IFNAR-1) to ask what role IFN signaling plays in the context of BEMPEG+NKTR-262 therapy. IFNAR signaling contributed to BEMPEG+NKTR-262 efficacy, as IFNAR blockade reduced survival from 80% to 50% (figure 4) primarily due to increased growth of non-treated tumors (online supplemental figure 5). Examination of PBL immune phenotypes 7 days post-treatment revealed that IFNAR signaling contributed to CD8+ T cell expansion (p<0.0001) and effector function (GzmA, ICOS, PD-1, CD62L) (figure 4B). In all cases the influence of type I IFN signaling was significant for BEMPEG+NKTR-262 and not for BEMPEG+RT (figure 4).

We observed reduced AH1-A5+ CD8+ T cell frequency after BEMPEG+RT plus IFNAR blockade that did not reach statistical significance (figure 4B). However, AH1-A5+ CD8 T cell frequency was significantly reduced after BEMPEG+NKTR-262 plus IFNAR blockade (figure 4B, p<0.01), which may reflect the contribution of IFN-α/β signaling to cross-presentation of tumor antigens. Interestingly, BEMPEG+NKTR-262 induced significantly more AH1-A5+ cells than BEMPEG+RT in the absence of type I IFN signaling (p<0.01), yet AH1-A5+ CD8+ T cells had similar frequencies of CD62L, GzmA, and Ki-67 in the presence or absence of IFNAR signaling across combination therapies (figure 4C), suggesting type I IFNs do not influence AH1-A5+ CD8+ T cell function. However, for AH1-A5+ CD8 T cells, frequencies of GzmA, Ki-67, and PD-1 of CD8+ T cells were significantly decreased after BEMPEG+NKTR-262 plus IFNAR blockade (figure 4D), indicating type I IFNs influenced the function of AH1-A5+ CD8+ T cells after BEMPEG+NKTR-262. This pattern did not occur after BEMPEG+RT. Collectively, these data suggest two results: first, unlike BEMPEG+RT, BEMPEG+NKTR-262-induced type I IFN signaling likely leads to increased priming/cross-priming; and second, BEMPEG+NKTR-262-induced type I IFN signaling supports a highly functional AH1-A5+ CD8+ T cell population that may be comprised of activated bystander cells or non-AH1-specific tumor-reactive cells.

Both TLR signaling and type I IFNs can induce DC maturation and regulate priming. Because BEMPEG+NKTR-262 efficacy relies, in part, on type I IFNs to induce tumor-specific CD8+ T cells (figure 4B), we asked whether we could detect differences in the DC compartment comparing BEMPEG+NKTR-262 and BEMPEG+RT. We harvested CT26 tumors 3 days post-treatment and analyzed DC (CD11c+MHCIICD24+F4/80) phenotypes (CD103, PD-L1, Arg1, iNOS) by flow cytometry. The UMAP distribution of DCs (online supplemental figure 6A) and an unbiased clustering algorithm (FlowSOM) (online supplemental figure 6B) revealed that BEMPEG drove a significantly different distribution of DCs, with a majority of DCs in cluster 0 for the three treatment groups that received BEMPEG (online supplemental figure 6B). Within cluster 0, BEMPEG+NKTR-262 treatment induced higher MHCIIC and CD103 expression than BEMPEG+RT (online supplemental figure 6C), both markers of conventional type 1 DC maturation (cDC1). CD103+ DC frequency was driven by both BEMPEG and NKTR-262 on the treated side, but not by RT (online supplemental figure 6D). Thus, BEMPEG+NKTR-262 resulted in a synergistic increase in CD103+ DC frequency over BEMPEG+RT.
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These data suggest BEMPEG + NKTR-262 induces a greater cDC1 frequency than BEMPEG + RT, which may support increased T cell priming.

On further assessment of NKTR-262’s impact on innate immune responses in the tumor microenvironment (TME), we found a pattern of changes driven by NKTR-262 on day 1 post-treatment that were maintained by BEMPEG on day 3. These changes included an increased ratio of M1-like (iNOS+)-like to M2-like (Arginase+) macrophages and an increased frequency of PD-L1+ macrophages (online supplemental figure 7A). Other NKTR-262-driven changes to the TME included a reduced frequency of monocytic MDSCs...
and increased frequency of polymorphonuclear MDSCs (online supplemental figure 7B). Taken with the observed increase in cDC1s, these results suggest an ‘antitumor’ TME that may better support an active adaptive immune response.

**BEMPEG+NKTR-262 induces CD8+ T cells with reduced checkpoint receptor expression and increased functional marker expression as compared with BEMPEG+RT**

We next looked at TIL phenotypes and functions from the treated and non-treated tumors 7 days post-therapy by flow cytometry (figure 5A). There was no significant difference in CD8+ T cell frequency or density (cells/mm^2) in the treated or non-treated tumors at this timepoint after BEMPEG+NKTR-262 compared with BEMPEG+RT (figure 5B), though there was a significant increase in Teff and concomitant decrease in Treg frequency in the treated tumor after BEMPEG+NKTR-262 compared with BEMPEG+RT (p<0.05, online supplemental figure 8A,B). Those changes in frequency were not reflected

**Figure 5**  BEMPEG+NKTR-262 treatment induces CD8+ T cells in the tumor with reduced checkpoint receptor expression and increased functional marker expression than BEMPEG+RT. (A) Representative flow cytometry gating strategy for CT26 tumors harvested 7 days post-therapy. (B) Percent (left) and density (right) of CD8+ T cells in the tumor. N=8–18, from three or four independent experiments. (C) Checkpoint receptors expressed on CD8+ T cells in the tumor. N=15 for PD-1 (from three experiments), 5 for Tim-3 and LAG-3 (from one experiment). (D) Activation markers expressed on CD8+ T cells in the tumor. For ICOS, N=5–9 from two experiments; for GzmA, N=13–23 from four experiments; for Ki-67, N=9–13 from three experiments. (E) Frequency of AH1-A5+ CD8+ T cells. N=15–20 from four experiments. For comparisons among treatment groups, one-way ANOVA with Šídák’s multiple comparisons test. *P<0.05, **P<0.01, ***P<0.001, ****P<0.0001. NT, non-treated tumor; RT, radiation therapy; T, treated tumor.
CD8+ TIL had reduced checkpoint receptor expression and increased functional marker expression after BEMPEG+NKTR-262 compared with BEMPEG+RT (figure 5C). For example, PD-1+ CD8+ T cell frequency was significantly decreased after BEMPEG+NKTR-262 compared with BEMPEG+RT in the treated and non-treated tumors (p<0.01), and exhaustion markers LAG-3 and TIM-3 were slightly decreased (figure 5C). GzmA+ CD8+ T cells were significantly increased after BEMPEG+NKTR-262 compared with BEMPEG+RT in the treated and non-treated tumors (p<0.001, figure 5D), while the activation/proliferation markers ICOS+ and Ki-67 tended higher in CD8+ T cells after BEMPEG+NKTR-262 compared with BEMPEG+RT. These changes were specific to CD8+ T cells and not observed in T effector or T regulatory cells (online supplemental figure 8C,D). Consistent with the minimal difference observed between the combination therapies in peripheral AH1-A5 CD8+ T cells, we found no difference in AH1-A5 CD8+ T cell frequency in the tumor (figure 5E). The increase in GzmA+ CD8+ T cells coupled with the decrease in PD-1+ CD8+ T cells in the TME suggests that BEMPEG+NKTR-262 T cells supports increased cell activation and function as compared with BEMPEG+RT.

**BEMPEG+NKTR-262 induces CD8+ T cells with greater cytotoxic capacity than BEMPEG+RT**

Because BEMPEG+NKTR-262 induced more GzmA+ CD8+ T cells than BEMPEG+RT, regardless of AH1-A5 status, we hypothesized that increased priming after BEMPEG+NKTR-262 led to an increased proportion of tumor-specific CD8+ T cells. To test this hypothesis, we used transgenic Nur77 reporter mice in which the strength of TCR stimulation correlates with GFP expression.22 Surprisingly, Nur77+ CD8+ T cell frequency after BEMPEG+NKTR-262 was similar to BEMPEG+RT (figure 6A, left), which suggests that there was not an increased proportion of TCR-stimulated CD8+ T cells at this timepoint. However, after BEMPEG+NKTR-262, a significantly greater proportion of Nur77+ cells expressed GzmA in the treated tumor than their BEMPEG+RT-treated counterparts (figure 6A, right). Therefore, we...
hypothesized that BEMPEG+NKTR-262 induced more potent cytotoxic function on a per cell basis as compared with BEMPEG+RT.

To test this, we asked whether Nur77⁺ CD8⁺ T cells correlated with tumor area (figure 6B). On examining treated and non-treated tumors from RT or NKTR-262-treated mice separately, we observed that Nur77⁺ T cell frequency did not correlate with tumor size after RT. In contrast, increased Nur77⁺ T cell frequency correlated with reduced tumor size in both treated and non-treated tumors after NKTR-262 (online supplemental figure 9). Therefore, we combined the treated and non-treated tumors for further analysis. We saw no correlation between Nur77⁺ CD8⁺ T cell frequency and tumor size after BEMPEG+RT (figure 6B, left); however, Nur77⁺ CD8⁺ T cell frequency significantly correlated with tumor size after BEMPEG+NKTR-262 (figure 6B, right). Because higher Nur77⁺ frequency correlated with smaller tumor size, we hypothesized that BEMPEG+NKTR-262 induced more potent CD8⁺ T cell cytotoxic function. To test this hypothesis, we sorted CD8⁺ T cells from treated and non-treated tumors and measured their cytolytic activity against autologous CT26 cells in vitro. BEMPEG+NKTR-262 elicited significantly increased CD8⁺ T cell cytolytic capacity compared with BEMPEG+RT (figure 6C). Taken together, these data demonstrate that BEMPEG+NKTR-262 induced and recruited to the tumor activated CD8⁺ T cells that were characterized by increased GzmA and greater cytolytic capacity than those generated by BEMPEG+RT.

**DISCUSSION**

Combining drugs with complementary mechanisms of action, such as chemotherapy regimens in hematological malignancies and breast cancer, can improve therapeutic outcomes. However, determining the combinations that will achieve optimal clinical responses is an ongoing and unresolved question. Herein, we compared the IL-2 agonist prodru BEMPEG in combination with either RT or the TLR agonist NKTR-262 and found that in multiple mouse models of multifocal disease, BEMPEG+NKTR-262 resulted in significantly slower tumor growth, specifically in non-treated tumors, and increased survival in comparison to BEMPEG+RT. Both combinations rely on CD8⁺ T cells for efficacy; however, BEMPEG+NKTR-262 induces more potent tumor-specific cytotoxic CD8⁺ T cells than BEMPEG+RT.

One feature that defines the immune response to BEMPEG is the preferential expansion of CD8⁺ and CD4⁺ Teff cells over Tregs in tumor tissue. This attribute of BEMPEG has the potential to mechanistically synergize with other therapies capable of alleviating T cell exhaustion, like immune checkpoint blockade (ICB), and/or with therapies that evoke new antigen responses, like RT or TLR agonists. Indeed, BEMPEG has shown synergy in combination with ICB preclinically and clinically (NCT02983045, NCT03138889, NCT03635983). BEMPEG/ICB-mediated synergy occurs through BEMPEG-driven Treg reduction and tumor-specific T cell expansion in the tumor, which are prevented from reaching exhaustion by ICB. By comparison, RT increases TIL Treg frequency, and as a result, BEMPEG+RT reduces the favorable Teff:Treg ratio driven by BEMPEG in comparison to BEMPEG monotherapy. Interestingly, we found BEMPEG+NKTR-262 significantly increased TIL CD4⁺ Teff frequency as compared with BEMPEG+RT. While CD4⁺ T cell depletion did not influence BEMPEG+RT or BEMPEG+NKTR-262 efficacy, given the favorable Teff:Treg ratio and shift towards Th1 polarization after BEMPEG+NKTR-262, it is likely that these CD4⁺ T cell phenotypes contribute to a TME that supports CD8⁺ T cell differentiation and function. Further, maintenance of the BEMPEG-induced Teff:Treg ratio following the inclusion of NKTR-262 suggests compatible mechanistic synergy between these two modalities.

One challenge for improving immunotherapy efficacy is inducing new antitumor immune responses. A key step in the initiation of new antitumor immunity is cross-presentation of tumor antigens by DCs and other professional APCs. Both RT and TLR ligands can induce DC maturation through pro-inflammatory signals. RT primarily provokes local proinflammatory signals through tumor cell death, which can be immunogenic or tolerogenic depending on a myriad of factors including cell cycle phase, cell type, and microenvironmental factors like hypoxia. Further, RT-induced DC maturation depends on the TME: radioimmunogenic tumors support cDC1 activation, while non-radioimmunogenic tumors do not. Additionally, preexisting immunity is required for RT-induced immune responses. Thus, BEMPEG+RT therapy likely expands and activates an existing population of tumor-reactive T cells and, at least in preclinical models, does not induce new antitumor responses.

In contrast to RT, intratumoral delivery of TLR agonists may generate new antitumor responses through APC activation. TLR7/8 agonists are particularly interesting as plasmacytoid DCs, B cells, monocytes, and myeloid DCs all express TLR7/8, allowing agonists to activate a wide-ranging group of APCs. Further, TLR7/8 agonists can induce tumor cell death through activated tumoricidal DCs. Thus, we hypothesized that NKTR-262 provides more potent immunogenic proinflammatory signals required for DC activation than RT. Indeed, 5 days post-treatment, BEMPEG changed the dominant DC phenotype, like a more mature phenotype, as IL-2 signaling indirectly drives DC expansion. Combination with NKTR-262 further boosted this DC population, as after BEMPEG+NKTR-262 therapy DCs exhibited an expanded CD103⁺ population in comparison to BEMPEG+RT. We also detected increased CCL5 (RANTES) and GM-CSF in the serum after BEMPEG+NKTR-262, both cytokines that promote leukocyte migration; given that migration follows a gradient, we speculate increased levels of these cytokines in the tumor. Further, our data suggest...
BEMPEG+NKTR-262 preferentially expands and/or increases cDC1 recruitment and influences macrophage polarization. While we focused on comparing different innate immune stimulators paired with BEMPEG, combining RT with a TLR7/8 agonist has shown DC-based synergistic effects in mouse models.38 Thus, the triple combination of RT, NKTR-262, and BEMPEG would be interesting to pursue in the future.

Type I IFNs can stimulate immune responses by activating DCs through promoting cross-presentation and by activating CD8α T cell effector function.39 In addition to BEMPEG+NKTR-262 more effectively inducing cDC1 maturation in the CT26 model than BEMPEG+RT, we found an increased dependence on IFNα/β signaling for BEMPEG+NKTR-262 over BEMPEG+RT. We note that RT dependence on IFNα/β signaling depend on the tumor model and RT dose applied,39 with fractionated doses inducing a greater abscopal effect.38 BEMPEG+NKTR-262 induced IFNα/β-signaling contributed to priming, as the proportion of AH1-A5+ cells was reduced in its absence. We also observed a significant reduction in GzmA+ and Ki-67+ AH1-A5+ CD8α T cell frequency in the absence of IFNα/β signaling, which may be bystander CD8α T cells. Highly cytolytic activated bystander CD8α T cells induced by anti-CD40 and IL-2 can drive antitumor effects39 and, in viral infections, the bystander response is rapidly induced by type I IFNs and TLR agonists to control infection before infections, the bystander response is rapidly induced by CD40 and IL-12

More activated (GzmA+) Nur77+ CD8α T cells over CD8α T cell cytotoxic capacity and induced significantly cell trafficking to the tumor and frequencies of AH1-A5+ cells was reduced in its absence. We also observed a significant reduction in GzmA+ and Ki-67+ AH1-A5+ CD8α T cell frequency in the absence of IFNα/β signaling, which may be bystander CD8α T cells. Highly cytolytic activated bystander CD8α T cells induced by anti-CD40 and IL-2 can drive antitumor effects39 and, in viral infections, the bystander response is rapidly induced by type I IFNs and TLR agonists to control infection before infections, the bystander response is rapidly induced by CD40 and IL-12

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