Bispecific antibody-mediated redirection of NKG2D-CAR natural killer cells facilitates dual targeting and enhances antitumor activity

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ABSTRACT

Background Natural killer group 2D (NKG2D) is an activating receptor of natural killer (NK) cells and other lymphocytes that mediates lysis of malignant cells through recognition of stress-induced ligands such as MICA and MICB. Such ligands are broadly expressed by cancer cells of various origins and serve as targets for adoptive immunotherapy with effector cells endogenously expressing NKG2D or carrying an NKG2D-based chimeric antigen receptor (CAR). However, shedding or downregulation of NKG2D ligands (NKG2DL) can prevent NKG2D activation, resulting in escape of cancer cells from NKG2D-dependent immune surveillance.

Methods To enable tumor-specific targeting of NKG2D-expressing effector cells independent of membrane-anchored NKG2DLs, we generated a homodimeric recombinant antibody which harbors a N-terminal single-chain fragment variable (scFv) antibody domain for binding to NKG2D, linked via a human IgG4 Fc region to a second C-terminal scFv antibody domain for recognition of the tumor-associated antigen ErbB2 (HER2). The ability of this molecule, termed NKAB-ErbB2, to redirect NKG2D-expressing effector cells to ErbB2-positive tumor cells of different origins was investigated using peripheral blood mononuclear cells, ex vivo expanded NK cells, and NK and T cells engineered with an NKG2D-based chimeric receptor.

Results On its own, bispecific NKAB-ErbB2 increased lysis of ErbB2-positive breast carcinoma cells by peripheral blood-derived NK cells endogenously expressing NKG2D more effectively than an ErbB2-specific IgG, mini-antibody able to induce antibody-dependent cell-mediated cytolysis via activation of CD16. Furthermore, NKAB-ErbB2 synergized with NK-92 cells or primary T cells engineered to express an NKG2D-CD3ζ chimeric antigen receptor (NKAR), leading to targeted cell killing and greatly enhanced antitumor activity, which remained unaffected by soluble MICA known as an inhibitor of NKG2D-mediated natural cytolysis. In an immunocompetent mouse glioblastoma model mimicking low or absent NKG2D expression, the combination of NKAR-NK-92 cells and NKAB-ErbB2 effectively suppressed outgrowth of ErbB2-positive tumors, resulting in treatment-induced endogenous antitumor immunity and cures in the majority of animals.

Conclusions Our results demonstrate that combining an NKAB antibody with effector cells expressing an activating NKAR receptor represents a powerful and versatile approach to simultaneously enhance tumor antigen-specific as well as NKG2D-CAR and natural NKG2D-mediated cytotoxicity, which may be particularly useful to target tumors with heterogeneous target antigen expression.

BACKGROUND

Immunotherapy with chimeric antigen receptor (CAR)-engineered T cells has demonstrated remarkable clinical efficacy in patients with B-cell malignancies. However, successful targeting of more prevalent solid tumors with CAR-T cells still remains a challenge, owing in part to an immunosuppressive tumor microenvironment, limited persistence of infused effector cells, and heterogeneity of target antigen expression. With respect to the latter, current CAR effector cells typically recognize a single tumor-associated surface antigen, resulting in only limited cell killing in tumors with uneven distribution of the target, and favoring outgrowth of subclones with low or absent antigen expression. Hence, different strategies are being followed to enable recognition of multiple target antigens, which include a combination of different monospecific CAR-T cells and coexpression of two CARs in the same T cells, or the more complex generation of CARs harboring more than one cell-binding domain.

Another approach harnesses the broad tumor specificity of the natural killer group 2D (NKG2D) molecule in a CAR design, linking the extracellular ligand-binding domain of NKG2D to a heterologous signaling molecule such as CD3ζ. NKG2D is an activating receptor expressed by all natural killer
(NK) cells, CD8⁺ T cells, and most natural killer T cells, as well as subpopulations of CD4⁺ T cells and γδ T cells.11–13 NKG2D has multiple membrane-anchored ligands, which in humans include the MHC class I-related molecules MICA and MICB and six UL16 binding proteins (ULBP1-6).14–16 These ligands are selectively upregulated by transformed and stressed cells and are widely expressed in almost all cancer types, including lung, breast, kidney, pancreatic, ovarian and prostate cancer, melanoma, leukemia, and glioblastoma.16,17 Hence, NKG2D ligands (NKG2DLs) are promising targets for adoptive immunotherapy with cytotoxic lymphocytes that naturally express NKG2D or are engineered to express NKG2D-based CARs.9–11 16–18 General safety of NKG2D-CAR T cells has recently been demonstrated in a phase I trial in patients with acute myeloid leukemia/myelodysplastic syndrome and seven other solid tumors.19 Nevertheless, activity of such NKG2D-CAR cells may be reduced against leukemic stem cells with low or absent expression of NKG2DLs.20 Natural NKG2D-mediated immune surveillance can be counteracted through a variety of mechanisms, including downregulation of NKG2DLs on exposure of cancer cells to interferon gamma (IFN-γ) or transforming growth factor beta (TGF-β).21 22 Tumor cells can also decrease NKG2DL density by proteolytic shedding.23 Accordingly, in the sera of patients with cancer, elevated levels of soluble NKG2DLs have been found, in many cases correlating with disease stage and metastasis.12,17 High concentrations of shed NKG2DLs can act as a competitor and block the interaction of NKG2D with tumor cells, induce NKG2D internalization and degradation, and desensitize immune effector cells.12–25

To limit interference by soluble NKG2DLs and redirect NKG2D expressing immune cells specifically to cancer cells of solid tumor origins, here, we designed a bispecific antibody combining two NKG2D-binding and two ErbB2 (HER2)-specific single-chain fragment variable (scFv) domains, linked by an IgG1 Fc region in a single tetravalent molecule. Specific targeting by this NKAB-ErbB2 antibody to antigen-positive tumor cells and enhancement of NKG2D-mediated cytotoxicity was evaluated in vitro cell killing experiments with primary lymphocytes endogenously expressing NKG2D and established NK-92 and primary T cells which carry an NKG2D-based chimeric antigen receptor (NKAR) encompassing the extracellular domain of NKG2D fused to transmembrane and intracellular domains of CD3ζ. Combined antitumor effects of the NKAB-ErbB2 molecule and the NKAR-NK-92 cells in vivo were investigated in a syngeneic tumor model in immunocompetent mice with ErbB2-positive murine glioblastoma cells, mimicking low or absent NKG2DL expression.

Expression and purification of bispecific NKAB-ErbB2 antibodies

The IgG₁-based NKAB-ErbB2 sequence was designed by in silico assembly of an immunoglobulin heavy-chain signal peptide, an scFv of NKG2D-specific antibody KYK-2.0,28 hinge, CH2 and CH3 domains of human IgG1 (UniProtKB-P01861, amino acid residues 104–327), a (G,S)₃ linker, and the ErbB2-specific scFv (FRP5) antibody fragment.29,30 For the similar NKAB-ErbB2 (IgG₃) molecule, the hinge, CH2 and CH3 domains of human IgG1 were used (UniProtKB-P01857, amino acid residues 99–330). In an alternative protein design, the scFv antibody fragments within the IgG₁-based NKAB-ErbB2 sequence were exchanged, resulting in NKAB-ErbB2 (rev) with reverse orientation of the binding domains. Codon-optimized fusion genes were de novo synthesized (GeneArt; Thermo Fisher Scientific, Darmstadt, Germany) and inserted into mammalian expression vector pcDNA3, resulting in plasmids pcDNA3-NKAB-ErbB2, pcDNA3-NKAB-ErbB2 (IgG₁), and pcDNA3-NKAB-ErbB2 (rev). As a control, monospecific mini-antibody FRP5-Fc was generated, which encompassed an immunoglobulin heavy-chain signal peptide, the ErbB2-specific scFv (FRP5) antibody fragment, and hinge, CH2 and CH3 domains of human IgG₁. Recombinant antibodies were expressed in...
transiently transfected HEK 293 T cells and purified from culture supernatant by affinity chromatography using a HiTrap Protein-G column on an ÄKTA FPLC system (GE Healthcare Europe, Freiburg, Germany). Purity and integrity of NKAB antibodies were determined by sodium dodecyl sulphate-polyacrylamide gel electrophoresis (SDS-PAGE) and Coomassie staining, or immunoblotting with horseradish peroxidase (HRP)-conjugated anti-human IgG antibody (Sigma-Aldrich, Munich, Germany). Protein concentrations were determined using a Nanodrop 1000 spectrophotometer (Thermo Fisher Scientific).

Expression of lineage markers, NKG2D and natural cytotoxicity receptors

Expression of NKG2D and lineage markers by peripheral blood mononuclear cells (PBMCs) from healthy donors was assessed using PE-conjugated anti-NKG2D (Miltenyi Biotec, Bergisch Gladbach, Germany), BV421-conjugated anti-CD56, and allophycocyanin (APC)-conjugated anti-CD3 (BD Biosciences) antibodies. Phenotypical characterization of ex vivo expanded primary NK cells was performed with BV421-conjugated anti-CD56, PE-conjugated anti-CD3, Alexa Fluor 647-conjugated anti-CD16, PE-conjugated anti-Nkp30, Alexa Fluor 647-conjugated anti-Nkp44 (all BD Biosciences), PE-conjugated anti-NKG2D, and APC-conjugated anti-Nkp46 (both Miltenyi Biotec) antibodies. In all staining procedures, a human Fc receptor blocking agent was included (BD Biosciences). Expression of NK-cell activating receptors by NK-92 and NKAR-NK-92 cells was determined using PE-conjugated anti-NKG2D, PE-conjugated anti-Nkp30, APC-conjugated anti-Nkp46 (all Miltenyi Biotec), and APC-conjugated anti-Nkp44 (R&D Systems, Wiesbaden-Nordenstadt, Germany) antibodies. Flow cytometric analysis was performed with FACS Canto II or BD LSRFortessa flow cytometers (BD Biosciences), and data were analyzed using FACSDiva or FlowJo software V.10.0.7 (FlowJo, Ashland, Oregon, USA). Binding of NKAB-ErbB2 to lymphocytes was either investigated using purified NKAB-ErbB2 followed by APC-conjugated secondary antibody (Dianova, Hamburg, Germany) or purified NKAB-ErbB2 covalently labeled with APC using a conjugation kit (Abcam, Cambridge, UK) according to the manufacturer’s recommendations.

Time-lapse imaging

MDA-MB453 breast carcinoma cells as targets were transduced with a lentiviral vector encoding luciferase and superfolder green fluorescent protein (GFP). As effector cells, PBMCs were isolated from buffy coats as described previously and either cultured for 3 days in X-VIVO 10 medium supplemented with 5% heat-inactivated human plasma, 500 IU/mL IL-2 and 50 ng/mL IL-15, or on anti-CD3 antibody-coated plates in RPMI 1640 medium supplemented with 10% heat-inactivated human plasma, 100 IU/mL IL-2 and 1 pg/mL anti-CD28 antibody (BioLegend, Koblenz, Germany) to preferentially stimulate NK or T cells, respectively. GFP-positive target cells were seeded on poly-L-lysine-coated glass bottom dishes (MatTek, Ashland, Massachusetts, USA) and allowed to adhere for 2–8 hours. Effector cells were treated with human Fc block (BD Biosciences) and then stained with PE-conjugated anti-CD3 and BV421-conjugated anti-CD56 antibodies (BD Biosciences) (IL-2/IL-15-stimulated PBMCs), or PE-conjugated anti-CD3 and BV421-conjugated anti-CD8 (BioLegend) antibodies (anti-CD3/anti-CD28-stimulated PBMCs). Then stained PBMCs were added to the GFP-positive target cells in the presence of 0.64 nM (100 ng/mL) APC-conjugated NKAB-ErbB2 protein. Time-lapse imaging was performed using a CQ1 Confocal Quantitative Image Cytometer (Yokogawa, Tokyo, Japan). Phase-contrast and fluorescent images were taken every 5 min at ×40 magnification by sample excitation with lasers at 405, 488, 561 and 640 nm. Images were captured at four z-stacks and processed to maximum intensity projections (MIPs).

Generation of NKAR-expressing effector cells

The NKG2D-based chimeric antigen receptor NKAR consists of an immunoglobulin heavy-chain signal peptide, the NKG2D extracellular domain (UniProtKB-PAI0B), and a modified CD8α hinge region, followed by CD3ζ transmembrane and intracellular domains. The codon-optimized NKAR sequence was de novo synthesized (GeneArt, Thermo Fisher Scientific) and inserted into lentiviral transfer plasmid pHRSIN-cPPT-SIEW upstream of IRES and enhanced green fluorescent protein (EGFP) sequences, yielding vector pS-NKAR-IIEW. VSV-G pseudotyped vector particles were produced and NK-92 cells were transduced as described. NKAR-positive cells were enriched by sorting with a FACS Aria fluorescence-activated cell sorter (BD Biosciences), with selection based on EGFP expression and enhanced NKG2D signals detected with anti-NKG2D antibody (Clone 149810, R&D Systems) followed by APC-coupled secondary antibody. NKAR expression by sorted cells was confirmed by SDS-PAGE of cell lysates and immunoblotting with anti-CD3ζ (6B10.2) or anti-CD8α antibodies (H-160; both Santa Cruz Biotechnology, Heidelberg, Germany), followed by HRP-conjugated secondary antibody and chemiluminescent detection. Similarly, CAR-engineered primary T cells were derived by lentiviral transduction with the NKAR construct. Interaction of NKAR with soluble MICA (sMICA) was investigated by flow cytometry with recombinant His-tagged human MICA (Biozol, Eching, Germany) followed by APC-conjugated anti-His-tag antibody (BioLegend).

Cytotoxicity assays

Cytotoxicity of effector cells was analyzed in flow cytometry-based assays as described. Briefly, tumor cells were labeled with calcein violet AM (CV) (Invitrogen, Thermo Fisher Scientific) and incubated with effector cells at various effector to target (E/T) ratios for 3 hours
at 37°C in the presence or absence of bispecific antibodies. Then 150 µL of a 1 µg/mL propidium iodide (PI) solution was added to each sample before flow cytometric analysis in a FACSCanto II flow cytometer. Dead target cells were identified as CV and PI double positive. Spontaneous target cell lysis was subtracted to calculate specific cytotoxicity. Data were analyzed using FACSDiva software. Unless stated otherwise, experiments were conducted at least twice with technical triplicates in each case. For competition experiments, sMICA-Fc fusion protein (R&D Systems) or recombinant human IgG4 protein (Biozol) was added to the cultures.

In vivo tumor model
Female C57BL/6 N mice (Charles River, Sulzfeld, Germany) 6–8 weeks old were used for a syngeneic GL261/ErbB2 murine glioblastoma model. Mice were inoculated with 1×10⁷ tumor cells at the right flank. Seven days later, animals with palpable tumors were randomized into groups of eight to nine animals and treated by peritumoral injection of 1×10⁷ NKAR-NK-92 or parental NK-92 cells in 200 µL of injection medium, with or without addition of 5 µg of NKAB-ErbB2 antibody. Treatment was repeated two times per week for 3 weeks. Tumor growth was followed by caliper measurements at preset intervals, and tumor volumes were calculated using the formula: volume = length × (width)² × 0.5. Symptom-free survival was assessed by daily inspection. Group sizes were chosen to allow statistical analysis of differences in symptom-free survival. All treated animals were included in the analysis. Animal experiments were approved by the responsible government committee (Regierungspräsidium Darmstadt, Darmstadt, Germany) and were conducted according to the applicable guidelines and regulations in the animal facility of Georg-Speyer-Haus.

To assess programmed cell death protein 1 (PD-1) expression by tumor-infiltrating lymphocytes, tumor-bearing mice from the different treatment groups were sacrificed. Tumor tissues were collected and single cell suspensions were prepared by incubating the minced tissues with collagenase (Sigma-Aldrich) at 37°C for 1 hour. Likewise, peripheral blood was collected and single cell suspensions of splenocytes were prepared. Erythrocytes in blood and splenocyte samples were lysed using BD Pharm Lyse buffer (BD Biosciences). The prepared cells were stained with PE-conjugated anti-mouse PD-1 (BioLegend), APC-conjugated anti-mouse CD8α (eBioscience, Thermo Fisher Scientific), Pacific Blue-conjugated anti-mouse CD3ε and PerCP-Cy5.5-conjugated anti-mouse CD4 (both BD Biosciences) antibodies in the presence of mouse Fc block (BD Biosciences). Dead cells were excluded by staining with eFluor780 (eBioscience, Thermo Fisher Scientific). A PE-conjugated isotype control antibody (BD Biosciences) was used to differentiate non-specific background signals. Flow cytometric analysis was performed with a BD LSRFortessa flow cytometer and data were analyzed using FlowJo software.

Statistical analysis
Data were analyzed by two-tailed unpaired or paired Student’s t-test and two-way analysis of variance. Symptom-free survival was analyzed by Kaplan-Meier plot and log-rank (Mantel-Cox) test. P values of <0.05 were considered statistically significant. Prism V.9 software (GraphPad Software, La Jolla, California, USA) was used for all statistical calculations.

RESULTS
Generation of a bispecific antibody binding to NKG2D and ErbB2
To target NKG2D-expressing lymphocytes to the tumor-associated antigen ErbB2, we designed a bispecific antibody similar in structure and molecular mass to an IgG molecule. This fusion protein (termed NKAB-ErbB2) carries an N-terminal scFv antibody domain derived from an NKG2D-specific antibody,28 and a second C-terminal scFv domain derived from ErbB2-specific antibody FRP5.29 To enable dimerization and provide flexibility between the two scFv domains, they were linked via the hinge, CH2 and CH3 domains of human IgG, and a (G,S)₆ peptide sequence (figure 1A). For expression as a secreted protein, the codon-optimized antibody sequence was fused to an immunoglobulin heavy-chain signal peptide sequence in a pcDNA3 expression plasmid. The recombinant NKAB-ErbB2 molecule was then expressed in transiently transfected HEK 293T cells and purified from culture supernatant by protein G affinity chromatography. SDS-PAGE and immunoblot analysis under reducing and non-reducing conditions confirmed purity and identity of the protein and revealed expression of the molecule as a tetravalent disulfide-linked homodimer, with only a minor fraction present in monomeric form under non-reducing conditions (figure 1B). Flow cytometric analysis demonstrated specific binding of the NKAB antibody to ErbB2-expressing but NKG2D-negative MDA-MB453 breast carcinoma and ErbB2-negative but NKG2D-positive NK-92 cells, but not to MDA-MB468 breast carcinoma cells which lack expression of ErbB2 and NKG2D (figure 1C and online supplemental figure S1). This demonstrates that the NKG2D-specific and ErbB2-specific binding domains in the molecule were both functionally active.

Effect of bispecific NKAB-ErbB2 antibody on the lytic activity of NKG2D-expressing peripheral blood lymphocytes and NK cells
Next, we investigated whether NKAB-ErbB2 influences the antitumor activity of NKG2D-positive peripheral blood lymphocytes in in vitro cytotoxicity assays using freshly isolated PBMCs from three healthy donors. Proportions of effector lymphocytes varied, depending on the individual donor, with NK cells (CD56⁺ CD3⁻) ranging from 3.5% to 6.1%, CD56-positive T cells (CD56⁺ CD3⁻) ranging from 3.6% to 13.9%, and CD56-negative T cells (CD56⁻ CD3⁺) ranging from 55.4% to 63.2% (figure 2A).
All lymphocyte subpopulations tested showed NKG2D surface expression (figure 2B). Thereby, purified NKAB-ErbB2 displayed binding to NKG2D-positive NK cells, and CD56-positive and CD56-negative T cells in a manner very similar to that of a commercial anti-NKG2D antibody (online supplemental figure S2). In the absence of bispecific antibody, unstimulated PBMCs displayed little to moderate cytotoxicity against MDA-MB453 breast cancer cells ranging from 3.4 (D1) to 15.3% (D3) specific cell killing after 3 hours of coincubation at an E/T ratio of 10:1 (figure 2C), likely due to NKG2D-mediated activation of the effector cells by NKG2DLs endogenously expressed by the target cells (online supplemental figure S1). In the presence of NKAB-ErbB2 antibody, cytotoxicity against the ErbB2-overexpressing cancer cells increased in a dose-dependent manner, with maximum cell killing of 2.0-fold to 2.6-fold over baseline reached at an antibody concentration of 0.64 nM (100 ng/mL; D1 and D2) or 3.2 nM (500 ng/mL, D3) (figure 2D). Cytotoxic activity decreased again at NKAB-ErbB2 concentrations above saturation of bispecific binding, indicative of competition by free antibody molecules. To investigate whether different NKG2D-positive cell types can contribute to target cell killing, the interaction of PBMCs prestimulated to preferentially activate NK or T cells with MDA-MB453 breast cancer cells was investigated in time-lapse imaging experiments. Thereby both, NK cells and CD8+ T cells induced lysis of tumor cells opsonized by NKAB-ErbB2 as indicated by membrane blebbing and formation of apoptotic bodies, with some of the NK cells exhibiting serial target cell killing (figure 3).
In a next set of experiments, we evaluated the activity of the bispecific NKAB-ErbB2 molecule with purified and ex vivo expanded pNK cells. To allow direct comparison between NKG2D- and CD16-mediated cytotoxicity, we included a FRP5-Fc mini-antibody consisting of the same ErbB2-specific scFv domain used for NKAB-ErbB2 but linked to the CD16-binding Fc portion of human IgG1 (online supplemental figure S3). pNK cells from three different donors were expanded for 2–3 weeks in medium containing IL-2 and IL-15, with around 75%–85% of cells in the resulting populations coexpressing NKG2D and CD16 (figure 4A). On coincubation with MDA-MB453 tumor cells for 3 hours at an E/T ratio of 5:1, pNK cells from all donors demonstrated moderate baseline cytotoxicity in the absence of antibody, which depending on the donor was enhanced in a concentration-dependent manner to up to 3.9-fold by NKAB-ErbB2, with maximum cell killing again reached at 0.64 nM (100 ng/mL; D4 and D6) or 3.2 nM (500 ng/mL, D5) (figure 4B). Also antibody-dependent cell-mediated cytotoxicity triggered by FRP5-Fc through activation of CD16 resulted in increased antitumor activity of pNK cells, although for all donors to a lower degree than NKAB-ErbB2 (figure 4C). To clearly distinguish NKG2D- and CD16-mediated effects, we based the NKAB-ErbB2 molecule on IgG4, which does not bind CD16 with high affinity. To assess potential additive effects by ligating both NKG2D and CD16 to ErbB2-positive tumor cells, we also generated an NKAB-ErbB2 (IgG1) molecule (online supplemental figure S3) and compared its activity to that of the original IgG4-based NKAB-ErbB2 antibody with pNK cells from another three donors. Thereby a very similar increase in cytotoxic activity of pNK cells against ErbB2-positive breast cancer cells was found for NKAB-ErbB2 and NKAB-ErbB2 (IgG4).
Enhancement of NK-cell activity by an NKG2D-based CAR

CARs which employ NKG2D for recognition of stress-induced NKG2DLs can enhance cytotoxic activity of CAR-engineered T and NK cells towards tumor cells of various origins. Since NKG2D is a type II transmembrane glycoprotein with an extracellular C-terminus, in NKG2D-based CARs heterologous signaling domains derived from type I transmembrane proteins such as CD3ζ are either added in a reverse orientation with regard to the plasma membrane in front of the cytoplasmic N-terminus of NKG2D, or an N-terminally truncated NKG2D ectodomain is artificially extended at the C-terminus with heterologous hinge and transmembrane regions followed by the respective cytoplasmic signaling motif, with both types of CARs being functional. Here we followed the latter type I transmembrane protein design and generated a CAR molecule (termed NKAR) that harbors an immunoglobulin heavy-chain signal peptide and the extracellular domain of human NKG2D at the N-terminus, fused to transmembrane and intracellular domains of CD3ζ via a flexible linker, a Myc tag and an optimized CD8α hinge region (figure 5A). For coexpression of the CAR sequence together with EGFP as a marker, VSV-G pseudotyped lentiviral vector particles were generated and used for transduction of human NK-92 cells as a clinically relevant model. Resulting NKAR-NK-92 cells were enriched by flow cytometric cell sorting. NKAR expression was examined by SDS-PAGE under non-reducing conditions and immunoblot analysis with CD3ζ and CD8α-specific antibodies, revealing the presence of NKAR monomers, disulfide-linked NKAR-NKAR homodimers and NKAR-CD3ζ heterodimers (figure 5B). Surface expression of the NKAR molecule was confirmed by flow cytometry, identified by a markedly increased NKG2D signal in NKAR-NK-92 cells when compared with parental NK-92 (figure 5C). NKAR expression also led to increased levels of Nkp30, while Nkp44 and Nkp46 were not or only marginally affected. This may be due to a stabilizing effect of the CD3ζ-containing NKAR on overall CD3ζ levels and Nkp30, which associates with CD3ζ for signaling. NKAR expression resulted in strongly enhanced cytotoxicity of NKAR-NK-92 cells against K562 leukemia cells which express different NKG2DLs (figure 5D and online supplemental figure S1), indicating that the CAR molecule was functional.

Synergistic effects of NKAR-ErbB2 and NKAR

Next, we investigated the activity of NKAR-NK-92 cells combined with NKAR-ErbB2 against MDA-MB453 breast cancer cells, which express high levels of ErbB2 and different NKG2DLs (online supplemental figure S1) but are largely resistant to parental NK-92 cells. As seen with K562 target cells, NKAR-NK-92 cells already displayed increased lysis of MDA-MB453 cells in the absence of NKAR-ErbB2 (11.2% vs 2.1% of specific killing at an E/T ratio of 5:1), which was markedly enhanced to more than...
60% specific lysis in the presence of 0.16–0.64 nM (25 to 100 ng/mL) of NKAB-ErbB2 (figure 6A). As observed with donor-derived PBMCs, cytotoxic activity of NKR-NK-92 cells decreased again gradually at NKAB-ErbB2 concentrations above 0.64 nM, likely due to competition of productive cross-linking of effector and target cells by free antibody molecules. Addition of NKAB-ErbB2 also increased cytotoxicity of parental NK-92 cells against MDA-MB453 cells in this short-term assay, although not to meaningful levels. This was likely due to the limited amount of endogenous NKG2D expressed by NK-92 (see figure 5C). Demonstrating a large degree of flexibility of the NKAB protein design, the same synergy in the killing of MDA-MB453 cells seen with NKR-NK-92 and the original NKAB-ErbB2 (rev), a molecule in which the positions of NKG2D- and ErbB2-specific binding domains relative to each other were exchanged (online supplemental figure S4).

To investigate whether the combined effect of NKAB-ErbB2 and NKR-expressing effector cells can consistently be achieved with ErbB2-positive cells of solid tumor origin, we tested specific cytotoxicity of NKR-NK-92 cells at increasing E/T ratios with MDA-MB453 and JIMT-1 breast cancer cells, and LNT-229 glioblastoma cells in the absence or presence of 0.16 nM (25 ng/mL) NKAB-ErbB2. For comparison, MDA-MB468 breast cancer cells were included, which also harbor NKG2DLs but are negative for ErbB2 (online supplemental figure S1). Even at the highest E/T ratio applied, the four tested cell lines proved largely resistant to parental NK-92 cells, which was not changed significantly by the addition of NKAB-ErbB2 (figure 6B). In contrast, NKR-NK-92 cells killed the NKG2DL-positive targets with high efficiency, which in the case of the ErbB2-expressing tumor cells was further enhanced in a synergistic manner by NKAB-ErbB2. Importantly, while also displaying markedly enhanced sensitivity to NKR-NK-92 cells, the addition of NKAB-ErbB2 did not further increase cytotoxicity of NKR-NK-92 against ErbB2-negative MDA-MB468 cells, underscoring the specificity of the NKAB-ErbB2 effect (figure 6B). In the case of ErbB2-expressing targets, NKAB-ErbB2-mediated activation of the NKGBD-CAR did not only trigger selective cytotoxicity but also induced marked upregulation of proinflammatory cytokines such as IFN-γ, which is

Figure 4 Effect of NKAB-ErbB2 on the cell killing activity of donor-derived NK cells. (A) Expression of NKG2D and CD16 by the ex vivo expanded pNK cells from healthy donors (D4–D6, B) was analyzed by flow cytometry with anti-NKG2D and anti-CD16 antibodies as indicated. (B) Cytotoxicity of the pNK cells (A) against ErbB2-expressing MDA-MB453 breast carcinoma cells in the absence of bispecific antibody (gray bars) or in the presence of increasing concentrations of recombinant NKAB-ErbB2 (blue bars) or ErbB2-specific FRP5-Fc IgG1 mini-antibody (red bars) was investigated in flow cytometry-based cytotoxicity assays after incubation at an E/T of 5:1 for 3 hours. (C) Normalized cytotoxicity data for pNK cells in the presence of NKAB-ErbB2 or FRP5-Fc (B) are plotted as fold increase in specific lysis in comparison to pNK alone. Mean values±SD are shown; n=3 individual donors. Data were analyzed by two-tailed unpaired Student’s t-test. *P<0.05, **P<0.01. ns: p>0.05. E/T, effector to target ratio; NK, natural killer; NKG2D, natural killer group 2D; ns, not significant; pNK, peripheral blood NK.
a hallmark of NK-cell activation (online supplemental figure S5).

Initially, the concept of targeting tumor cells through an NKG2D-based CAR was developed using genetically engineered T cells. To investigate whether the cytotoxicity of such CAR-T cells can also be enhanced by the NKAR-ErbB2 molecule, donor-derived T cells were transduced with the NKAR construct, and cytotoxicity of the resulting cell population against ErbB2-positive MDA-MB453 cells was tested in the absence or presence of 0.64 nM (100 ng/mL) NKAB-ErbB2. Thereby, similar to our findings with NK-92-derived NKAR-NK cells, the expression of NKAR on its own already increased cytotoxicity of NKAR-T cells against NKG2DL-positive targets,
Figure 6  Enhancement of NKAR-NK-92 cytotoxicity by NKAB-ErbB2. (A) The effect of NKAB-ErbB2 on specific cytotoxicity of NKAR-NK-92 cells against ErbB2-positive MDA-MB453 breast carcinoma cells was determined in flow cytometry-based cytotoxicity assays after coincubation at an E/T of 5:1 for 3 hours in the absence or presence of increasing concentrations of NKAB-ErbB2 (red bars). Parental NK-92 cells were included for comparison (blue bars). Mean values±SD are shown; n=3 technical replicates from a representative experiment. Data were analyzed by two-tailed unpaired Student’s t-test (shown for NKAR-NK-92 vs NK-92 in the absence of NKAB-ErbB2 and NKAR-NK-92 in the absence vs the presence of increasing concentrations of NKAB-ErbB2). *P<0.05, **P<0.01, ***P<0.001, ****P<0.0001, ns: p>0.05. (B) Cytotoxicity of NKAR-NK-92 (orange circles) and NK-92 cells (light blue triangles) in the absence, and NKAR-NK-92 (red circles) and NK-92 cells (dark blue triangles) in the presence of 0.16 nM (25 ng/mL) of NKAB-ErbB2 against MDA-MB453, MDA-MB468 and JIMT-1 breast carcinoma cells and LNT-229 glioblastoma cells was investigated in flow cytometry-based cytotoxicity assays after coincubation at different E/T ratios for 3 hours. Mean values±SD are shown; n=3 technical replicates from a representative experiment. Data were analyzed by two-tailed unpaired Student’s t-test (shown for NKAR-NK-92+NKB-ErbB2 vs NKAR-NK-92). *P<0.05, **P<0.01, ***P<0.001, ns: p>0.05. (C) The ability of NKAB-ErbB2 to compete binding of sMICA to NKAR-NK-92 cells was determined by flow cytometry with APC-conjugated anti-His-tag antibody after incubation of cells with 2.5 µg/mL of His-tagged sMICA in the absence (solid black line) or presence of 1.6 nM (0.25 µg/mL) or 16 nM (2.5 µg/mL) of NKAB-ErbB2 (dashed red lines) as indicated. Cells treated only with secondary antibody served as control (filled area). (D) Inhibition of NKAR-NK-92 cell killing activity by 2.5 µg/mL of sMICA-Fc protein and restoration by addition of 0.16 nM (25 ng/mL) of NKAB-ErbB2 were investigated in flow cytometry-based cytotoxicity assays after coincubation with MDA-MB453 target cells at an E/T ratio of 5:1 for 3 hours. Recombinant human IgG4 protein (25 ng/mL) served as isotype control. Mean values±SD are shown; n=3 technical replicates from a representative experiment. Data were analyzed by two-tailed paired Student’s t-test. *P<0.05, ns: p>0.05. E/T, effector to target ratio; ns, not significant; sMICA, soluble MICA.

while enhanced cytotoxicity against ErbB2-positive tumor cells lacking human NKG2DLs was dependent on the simultaneous presence of NKAB-ErbB2 (online supplemental figure S6).

Restoration of sMICA-inhibited NKAR functionality by NKAB-ErbB2

Proteolytic shedding of NKG2DLs like MICA has been identified as a mechanism for cancer cells to evade NKG2D-mediated immune surveillance.11 23 To test whether this could also affect NK cells expressing the NKG2D-based CAR, interaction of sMICA with NKAR-NK-92 cells was investigated by flow cytometry. Thereby, strong binding of sMICA to the surface of the CAR-NK cells was found, which was blocked in a concentration-dependent manner by NKAB-ErbB2 (figure 6C), indicating that the bispecific antibody can shield NKG2D. Occupation of the ligand binding site of the NKG2D-CAR by sMICA was also relevant for cytotoxic activity of NKAR-NK-92 cells, which was readily triggered by NKG2DLs naturally expressed by MDA-MB453 breast cancer cells, but markedly inhibited in the presence of competing sMICA (figure 6D). In contrast, the enhanced cell killing activity of the combination of NKAR-NK-92 cells and NKAB-ErbB2 was not significantly affected by an excess of sMICA, suggesting that this strategy could overcome immune evasion due to ligand shedding.

In vivo activity of NKAR-NK-92 cells and NKAB-ErbB2 antibody against ErbB2-positive glioblastoma tumors

To investigate the combined effect of NKAR-NK cells and NKAB-ErbB2 antibody in vivo in a setting where tumor cells similar to cancer stem cells lack NKG2DLs that could trigger the NKG2D-CAR directly,20 a subcutaneous tumor
model based on syngeneic GL261/ErbB2 glioblastoma tumors in immunocompetent C57BL/6 mice was established. Since GL261/ErbB2 cells are of murine origin, endogenous NKG2DLs expressed by these cells are not recognized by the NKAR molecule, which is based on human NKG2D. Consequently, in vitro cytotoxicity assays no difference in sensitivity of GL261/ErbB2 cells to NKAR-NK-92 and parental NK-92 was found (figure 7A). For NK-92 cells, this remained unchanged in the presence of NKAB-ErbB2. However, when NKAR-NK-92 cells were combined with NKAB-ErbB2, even at low E/T ratios a marked increase in cytotoxicity against GL261/ErbB2 was observed. Thereby, intact homodimeric NKAB-ErbB2 was more active against these target cells than a mutated NKAB-ErbB2 (C106S and C109S) derivative which cannot form dimers due to the lack of intermolecular disulfide bridges within the IgG hinge region (online supplemental figure S7). Likewise, NKAB-ErbB2-mediated recognition of murine melanoma cells genetically modified to express human ErbB2 induced cytokine secretion and specific lysis by NKAR-NK-92 cells (online supplemental figure S5).

For in vivo analysis, 7 days after tumor cell inoculation, mice were treated by peritumoral injection of 1×10⁷ NKAR-NK-92 or parental NK-92 cells with or without 5 μg of NKAB-ErbB2 antibody admixed to the injection medium twice per week for 3 weeks. Thereby, combination treatment with the NKAR-NK cells and NKAB-ErbB2 was highly effective. Tumor outgrowth was controlled in eight out of nine animals in this group during therapy, and complete tumor regression was seen in seven of the mice thereafter, leaving no measurable tumors 3 months after the last treatment at termination of the experiment on day 115 (figure 7B). Since the NKAR receptor cannot recognize GL261/ErbB2 cells on its own, treatment with NKAR-NK-92 cells alone had no effect on tumor development during or after therapy, with only one out of eight mice in this group being tumor-free at endpoint analysis. This was most likely due to spontaneous rejection not related to the treatment. While combination therapy with parental NK-92 cells and the NKAB-ErbB2 antibody showed some effect and resulted in delayed tumor growth, still only two of nine animals in this group presented as tumor-free on day 115 (figure 7B). The different kinetics in tumor growth among the treatment groups were also reflected in symptom-free survival. While mice in the NKAR-NK-92 only group had to be sacrificed due to disease progression earlier than in the group receiving the NK-92/NKAB-ErbB2 combination (median survival of 36.5 vs 46.0 days), this difference was not statistically significant (figure 7C). In contrast, with seven animals surviving and only two mice showing delayed tumor development, median survival in the NKAR-NK-92/NKAB-ErbB2 combination group was not reached (>115 days).

Sera from animals of this group contained IgG antibodies strongly reactive with GL261/ErbB2 as well as ErbB2-negative parental GL261 glioblastoma cells (figure 7D), suggesting that an endogenous antitumor immune response induced by the NKAR-NK-92/NKAB-ErbB2 combination treatment likely contributed to tumor rejection. To evaluate that further, the expression of PD-1 by T cells from two tumor-bearing animals of each treatment group was compared. While in comparison to splenic T cells and T cells from peripheral blood higher PD-1 expression was found for tumor-infiltrating CD4⁺ and CD8⁺ T cells in all three treatment groups, especially for CD8⁺ T cells this was much more pronounced in mice after NKAR-NK-92/NKAB-ErbB2 combination therapy (figure 7E), which is indicative of a higher proportion of previously activated and antigen-experienced T cells in their tumor tissues.

**DISCUSSION**

Activation of endogenous NKG2D and NKG2D-based CARs can be inhibited or prevented by shedding and release of soluble NKG2DLs or downregulation of NKG2DL expression, which can contribute to immune escape. To circumvent this problem but still employ activation of NKG2D as a potent mechanism for the elimination of cancer cells, here we developed a bispecific antibody which simultaneously recognizes NKG2D and the tumor-associated antigen ErbB2, resulting in targeted activation of NKG2D and NKG2D-CAR expressing effectors against ErbB2-positive cancer cells.

Recombinant NKAB-ErbB2 antibody displayed specific binding to NK2D and ErbB2, thereby crosslinking lymphocytes and tumor cells. The bispecific molecule improved cytotoxicity of donor-derived bulk PBMCs and ex vivo expanded primary NK cells endogenously expressing NKG2D and markedly enhanced cell killing activity of NK cells carrying an NKG2D-based CAR (NKAR) against otherwise NK-resistant ErbB2-positive targets. This included JIMT-1 breast cancer cells, which express NKG2DLs and elevated levels of ErbB2 (online supplemental figure S1) but are resistant to the clinically approved ErbB2-targeted therapeutics trastuzumab and lapatinib. Only picomolar concentrations of NKAB-ErbB2 were needed for maximum activity against ErbB2-expressing target cells in titration experiments with effector lymphocytes carrying natural NKG2D or transduced with the NKAR vector. Similar results were obtained when NKAB-ErbB2 was combined with T cells harboring an NKG2D-CAR (online supplemental figure S6). Importantly, while sMICA used as a model for a shed NKG2DL blocked activation of the NKAR molecule by endogenous NKG2DLs on the tumor cell surface, even a 100-fold excess of sMICA did not significantly affect the combined activity of NKAR and NKAB-ErbB2. This suggests that also in a clinical situation NKAB-ErbB2 may overcome the detrimental effects of ligand shedding on NKG2D-dependent immune surveillance.

Activation by natural NKG2DLs or NKAB-ErbB2 towards tumor cells was much more pronounced for NKAR-expressing effector cells than lymphocytes that only harbored endogenous NKG2D, which, in...
Figure 7  Combined in vivo antitumor activity of NKAR-NK-92 cells and NKAB-ErbB2 antibody against syngeneic glioblastoma in immunocompetent C57BL/6 mice. (A) Cytotoxicity of NKAR-NK-92 (orange circles) and NK-92 cells (light blue triangles) in the absence, and NKAR-NK-92 (red circles) and NK-92 cells (dark blue triangles) in the presence of 0.16nM (25ng/mL) of NKAB-ErbB2 against murine GL261/ErbB2 glioblastoma cells expressing human ErbB2 was investigated in flow cytometry-based cytotoxicity assays after coincubation at different E/T ratios for 3 hours. Mean values±SD are shown; n=3 technical replicates from a representative experiment. Data were analyzed by two-tailed unpaired Student’s t-test (shown for NKAR-NK-92+NKAB-ErbB2 vs NKAR-NK-92). **P<0.01. (B) GL261/ErbB2 cells (1x10^6) were subcutaneously injected into the right flank of C57BL/6 mice. Seven days later, the mice were treated by peritumoral injection of 1x10^7 NKAR-NK-92 cells without (n=8) or with 5µg of NKAB-ErbB2 antibody (n=9) admixed to the cells two times per week for 3 weeks. Control mice received parental NK-92 cells with NKAB-ErbB2 (n=9). Tumor growth in the individual animals was followed by caliper measurements. (C) Symptom-free survival of the mice. Data were analyzed by Kaplan-Meier plot and log-rank test. *P<0.05, **P<0.01. ns: p>0.05. (D) Induction of antibodies against glioblastoma cells in animals from the experiment shown in (B) and (C) treated with NKAR-NK-92 and NKAB-ErbB2 (n=6) was investigated by determining cell surface binding of IgG from serum to GL261/ErbB2 and ErbB2-negative parental GL261 cells as indicated (red circles) by flow cytometry. Sera from naïve mice (n=6) served as controls (gray squares). MFI: geometric mean. Mean values±SD are shown. Data were analyzed by two-tailed unpaired Student’s t-test. **P<0.01. (E) PD-1 expression by CD4+ and CD8+ T cells from TILs, splenocytes (spleen) and PBMCs of animals from the experiment (B,C) treated with NKAR-NK-92 (orange bars, n=2), NKAR-NK-92 and NKAB-ErbB2 (red bars, n=2), or NK-92 and NKAB-ErbB2 (blue bars, n=2) was determined by flow cytometry with anti-PD-1 antibody. Cells were gated using anti-CD4 and anti-CD8 antibodies as depicted in online supplemental figure S8. Mean values±SD are shown. Data were analyzed by two-tailed unpaired Student’s t-test (shown for TILs). *P<0.05. ns: p>0.05. MFI, mean fluorescence intensity; ns, not significant; PBMC, peripheral blood mononuclear cell; PD-1, programmed cell death protein 1; TIL, tumor-infiltrating lymphocyte.
addition to different expression levels of NKG2D and the NKG2D-CAR, was likely due to distinct signaling capabilities of the receptors. To be active, human NKG2D requires prior association with the adaptor molecule DNA-activating protein of 10 kDa (DAP10). In contrast, in the NKG2D-CAR employed in our study, extracellular target recognition and intracellular signaling functions are directly linked in one molecule, likely accelerating kinetics of effector cell activation. In addition, DAP10 only contains a single YXXM tyrosine-based motif similar to the ones found in the costimulatory molecules CD28 and ICOS for interaction with downstream pathways.

Instead, the NKG2D-CAR harbors CD3ζ, which contributes three potent immunoreceptor tyrosine-based activation motifs. On their own or in combination, ectopic expression of the NKG2D-CAR and application of the bispecific NKAB molecule may also bypass reduced responsiveness and downregulation of endogenous NKG2D observed on chronic stimulation by NKG2DLs and during NK cell exhaustion.

The bispecific NKAB-ErbB2 antibody mimics the general structure of a human IgG molecule, with the hinge, CH2 and CH3 domains of the IgG Fc region separating NKG2D-specific and ErbB2-specific antibody domains. Accordingly, in contrast to other bispecific molecules with only one binding site for NKG2D and a target antigen, the NKAB-ErbB2 protein was expressed as a homodimer with two binding sites each for NKG2D and ErbB2. For comparison, we also generated a mutated NKAB-ErbB2 (C106S and C109S) derivative which lacked the cysteine residues within the IgG hinge region and was therefore only produced as a monomer (online supplemental figure S7). In short-term assays, dimeric and monomeric NKAB-ErbB2 proteins similarly enhanced cytotoxicity of NKG2D-CAR NK cells against target cells coexpressing NKG2DLs and ErbB2. However, when target recognition was entirely dependent on NKAB-ErbB2 due to the lack of endogenous ligands for human NKG2D, homodimeric NKAB-ErbB2 was more effective, suggesting that the tetravalent design may be of particular advantage in a setting of escape from natural NKG2D immune surveillance (online supplemental figure S7). Furthermore, with around 180 kDa, the apparent molecular mass of intact NKAB-ErbB2 is close to that of regular IgG molecules, which similar to complete antibodies will likely translate to a longer in vivo half-life when compared with monomeric scFv-scFv or scFv-Fc molecules.

We based the NKAB-ErbB2 molecule on the structure of IgG, which does not bind to CD16 with high affinity, allowing to distinguish NKG2D- from CD16-mediated effects. This NKAB design proved highly flexible, allowing to switch the positions of NKG2D- and ErbB2-specific binding domains within the molecule without loss of activity (online supplemental figure S4), indicating that the overall NKAB structure will be applicable also for similar bispecific molecules targeting alternative tumor antigens and/or activating lymphocyte receptors. Likewise, exchanging the IgG Fc region with the respective sequence of IgG1 led to a fully functional molecule, although without a clear benefit of the possible simultaneous or alternative interaction of the resulting NKAB-ErbB2 (IgG1) antibody with NKG2D and CD16 (online supplemental figure S3). Nevertheless, engineering of the IgG Fc domain to include mutations that enhance affinity for CD16 may still increase activity of such IgG1-based NKAB molecules.

In vivo antitumor activity of NKAB-ErbB2 in combination with NKAR-NK-92 cells was investigated in a murine glioblastoma model. In this disease, activity of the NKG2D/NKG2DL system can be negatively affected by limited expression of NKG2DLs. To mimic the situation of low or absent NKG2DL expression as a mechanism of immune evasion, we used murine GL261/ErbB2 glioblastoma cells growing as subcutaneous tumors in C57BL/6 mice. While GL261 cells express moderate levels of murine NKG2DLs, these are not recognized by human NKG2D or the human NKG2D-CAR as exemplified by the results from our in vitro cytotoxicity assays with NK-92 and NKAR-NK-92 cells. Hence, as expected, repeated treatment of established GL261/ErbB2 tumors with NKAR-NK-92 cells alone was ineffective, with rapid tumor growth continuing in most animals while still under therapy. While not resulting in a statistically significant survival benefit in comparison to NKAR-NK-92 monotherapy, tumor outgrowth was delayed in mice treated with a combination of NKAB-ErbB2 and parental NK-92 cells, where the bispecific antibody can act via endogenous NKG2D expressed at moderate levels. However, when NKAR-NK-92 cells were applied together with NKAB-ErbB2, the glioblastoma tumors were readily rejected in the majority of the animals, resulting in cures with seven out of nine mice in this group staying tumor-free during the remaining course of the experiment.

In addition to direct tumor cell lysis, activated NK cells play a critical role in dendritic cell (DC) recruitment and maturation, thus indirectly inducing or enhancing tumor-specific adaptive immune responses. This likely contributed to the durable responses observed in immunocompetent mice after treatment with the NKAB-ErbB2/NKAR-NK-92 combination, as seen in previous studies with NK-92 cells which express a more classical scFv-based CAR and are currently evaluated in a phase I clinical trial in patients with glioblastoma. On NKAB-ErbB2-mediated activation, NKAR-NK-92 cells markedly upregulated the secretion of proinflammatory chemokines and cytokines pivotal in DC recruitment (online supplemental figure S5), including RANTES (CCL5), MIP-1α (CCL3) and MIP-1β (CCL4) or DC maturation and function, such as GM-CSF, IFN-γ, TNF-α and TNF-β. Indeed, sera from mice cured of their glioblastoma tumors after combination therapy with NKAB-ErbB2 and NKAR-NK-92 contained IgG antibodies reactive with GL261 and GL261/ErbB2 cells. While we did not investigate whether combined treatment with NKAB-ErbB2 and NKAR-NK-92 also resulted in the activation of tumor-reactive T cells, TILs from the two animals in this group that were not
cured expressed higher levels of the immune checkpoint molecule PD-1 than TILs from mice treated with NKAR-NK-92 alone or a combination of NKAB-ErbB2 with parental NK-92 cells. Despite the failure in these mice to reject the tumors, this is indicative of more effective TCR-mediated stimulation of tumor-infiltrating T cells on NKAB-ErbB2/NKAR-NK-92 therapy.56 Importantly, these data also suggest that combining NKAB-ErbB2 further with an immune checkpoint inhibitor may be warranted.

CONCLUSION
Taken together, our data show that the bispecific NKAB antibody efficiently redirects effector lymphocytes to cancer cells expressing the respective tumor-associated antigen, thereby overcoming the inhibitory effects of low expression or shedding of NKG2DLs and resulting in enhanced antitumor activity and therapy-induced adaptive antitumor immune responses. In particular, the combination of an NKAB antibody with NK or T cells engineered with an NKG2D-based chimeric antigen receptor represents a versatile approach to simultaneously enhance tumor-antigen-specific and natural NKG2D-mediated cytotoxicity, which may become very useful to target tumors with heterogeneous target antigen expression not amenable to classical CAR-T or CAR-NK cell therapies.

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