

Bispecific antibodies redirect synthetic agonistic receptor modified T cells against melanoma

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ABSTRACT

Background Melanoma is an immune sensitive disease, as demonstrated by the activity of immune check point blockade (ICB), but many patients will either not respond or relapse. More recently, tumor infiltrating lymphocyte (TIL) therapy has shown promising efficacy in melanoma treatment after ICB failure, indicating the potential of cellular therapies. However, TIL treatment comes with manufacturing limitations, product heterogeneity, as well as toxicity problems, due to the transfer of a large number of phenotypically diverse T cells. To overcome said limitations, we propose a controlled adoptive cell therapy approach, where T cells are armed with synthetic agonistic receptors (SAR) that are selectively activated by bispecific antibodies (BiAb) targeting SAR and melanoma-associated antigens.

Methods Human as well as murine SAR constructs were generated and transduced into primary T cells. The approach was validated in murine, human and patient-derived cancer models expressing the melanoma-associated target antigens tyrosinase-related protein 1 (TYRP1) and melanoma-associated chondroitin sulfate proteoglycan (MCSP) (CSPG4). SAR T cells were functionally characterized by assessing their specific stimulation and proliferation, as well as their tumor-directed cytotoxicity, in vitro and in vivo.

Results MCSP and TYRP1 expression was conserved in samples of patients with treated as well as untreated melanoma, supporting their use as melanoma-target antigens. The presence of target cells and anti-TYRP1 × anti-SAR or anti-MCSP × anti-SAR BiAb induced conditional antigen-dependent activation, proliferation of SAR T cells and targeted tumor cell lysis in all tested models. In vivo, antitumoral activity and long-term survival was mediated by the co-administration of SAR T cells and BiAb in a syngeneic tumor model and was further validated in several xenograft model.

Conclusion The SART cell-BiAb approach delivers specific and conditional T cell activation as well

WHAT IS ALREADY KNOWN ON THIS TOPIC

⇒ Melanoma has been shown to be sensitive to immunotherapy, though significant subsets of patients do not respond or relapse after initial response.

WHAT THIS STUDY ADDS

⇒ A modular and controllable adoptive T cell therapy approach to the treatment of melanoma.

HOW THIS STUDY MIGHT AFFECT RESEARCH, PRACTICE OR POLICY

⇒ This study puts forward the simultaneous and sequential targeting of melanoma-associated chondroitin sulfate proteoglycan and tyrosinase-related protein 1 as a strategy for the targeted therapy of melanoma.

as targeted tumor cell lysis in melanoma models. Modularity is a key feature for targeting melanoma and is fundamental towards personalized immunotherapies encompassing cancer heterogeneity. Because antigen expression may vary in primary melanoma tissues, we propose that a dual approach targeting two tumorassociated antigens, either simultaneously or sequentially, could avoid issues of antigen heterogeneity and deliver therapeutic benefit to patients.

BACKGROUND

Due to its high tumor mutational burden, likely driven by ultraviolet radiation, melanoma possesses a high number of neoantigens, making it one of the most immunogenic tumor types. ^{1 2} Melanoma treatment has been revolutionized by immune checkpoint blockade (ICB), reactivating T cells or preventing T cell dysfunction. ³ Despite these successes, many patients still either do not



respond or relapse after an initial response. For patients not carrying targetable mutations such as BRAFV600E or having already exhausted targeted treatments, limited options remain, resulting in an urgent need for innovative effective treatments.

While ICB-mediated prevention of T cell dysfunction has entered clinical practice in a wide array of indications beyond melanoma, the direct therapeutic use of T cells in non-hematological cancer entities has been largely ineffective.⁵ Melanoma, however, has been an exception in this regard. Tumor infiltrating lymphocytes (TIL) are a prognostic factor in melanoma and correlate with response to ICB.6 In fact, investigation using isolated, non-modified, and ex vivo expanded TIL as a treatment modality for patients with melanoma has already been explored in the pre-ICB era. There, about 50% of treated patients were sensitive to TIL therapy, a fraction of which exhibiting complete and durable responses.⁸⁹ The success of ICB therapy then suspended further development of TIL-based therapies for some time. Recently, clinical studies have explored the potential of TIL therapy after ICB failure in patients with melanoma. Consistent with pre-ICB reports, TIL therapy yielded substantial response rates of up to 32%, indicating that even in such clinically challenging situations, TIL therapy might still be of benefit to patients (NCT00937625, NCT02379195 and NCT02354690). 10 Along these lines, ICB and TIL therapy impressively demonstrate the utility of T cells in melanoma treatment regardless of treatment line. TIL therapy, however, comes with significant challenges which limit its application: (1) requirement for accessible target lesions for resection and TIL selection and expansion, (2) failure to select and expand TIL, (3) heterogeneity of TIL products with often undefined specificities and consequently and (4) heterogeneous response patterns both in extent and in duration.¹¹

T cells can be rendered tumor-specific through genetic engineering of a synthetic receptor, so called chimeric antigen receptor (CAR), that can recognize antigens on the cell surface independent of major histocompatibility complex molecules. Anti-CD19 CAR T cells have entered clinical routine after transformative results in the treatment of hematological malignancies. 12 13 CD19 as a target antigen allows for targeting of lymphoma and leukemic cells along with healthy B cells. The deleterious effects of depleting the entire B-cell compartment are clinically manageable. 14 In contrast to CD19 hematological malignancies, dispensable lineage-specific tumor-associated antigens are rarely found in solid cancer types.⁵ Additionally, an immunosuppressive milieu and target antigen heterogeneity, among other factors, have resulted in melanoma CAR T cell therapy trials faring poorly thus far. 15 Detailed analysis of TIL therapy failure highlights that loss of dominant antigens under therapeutic pressure happens quite frequently, suggesting that a successful T cell product will need to target more than one antigen.¹⁶ These results support the need for advances in melanoma cell therapy considering such a limitation.

As most of the targetable antigens are not entirely specific to melanoma but shared with other cells as well, we reasoned that any T cell therapeutic strategy would need to be controllable to allow application with a safety net. We have previously described a synthetic agonistic receptor (SAR) platform composed of the extracellular domain of epidermal growth factor receptor variant III (EGFRvIII), fused to intracellular T cell-activating domains (later referred to as E3 construct). The construct can be specifically activated by bispecific antibodies (BiAb) simultaneously targeting the SAR and the tumor antigen. The major advantages of modular adoptive cell therapy (ACT) platforms are the possibility to stop administration of the adaptor molecule in case of undesired therapy-associated toxicities and the ability to target multiple antigens by administering different T cell adaptor molecules. ^{17 18} In particular, we previously demonstrated that SAR-transduced T cell activity is conditional to the presence and binding of the BiAb, enabling a tunable activity that is advantageous in case of toxicities. 19 20 We hypothesized that this SAR platform could serve as a safe and effective way of targeting melanomaassociated antigens for melanoma treatment.²⁰

For the present study, using the CrossMab technology, ²¹ ²² we developed trivalent BiAb binding melanoma antigen and the E3 SAR. We demonstrate that SAR-transduced T cells were selectively and reversibly activated through the BiAb, solely in the presence of antigen-positive melanoma cells. We showcase substantial activity of the platform in primary melanoma cultures and in several xenograft and syngeneic mouse models, supporting further clinical development.

MATERIAL AND METHODS Patient and healthy donor material

Frozen, primary and metastatic tumor samples from 13 patients with a histologically confirmed diagnosis of melanoma were used for this study. The samples were cultured in MCDB 153 medium (Merck) complemented with 20% Leibovitz's L-15 medium (Thermo Fisher), 2% fetal calf serum (Gibco), $10\,\mu\text{g/mL}$ human insulin (Merck) and 2 M CaCl $_2$ solution and expanded until further use. Biological and clinical information were obtained from electronic medical records. Patient characteristics are summarized in online supplemental table 1. Human peripheral blood mononuclear cells (PBMCs) for the generation of human CAR and SAR T cells were isolated from healthy donors by Ficoll density gradient separation.

Mice

Female C57BL/6N and NSG (NOD.Cg-Prkdcscid Il2rgt-m1WjI/SzJ) mice were purchased from Charles River or Janvier Labs. Animals were housed in specific pathogen-free facilities in groups of 2–5 animals per cage. All experimental studies were approved and performed with mice aged 2–4 months and in accordance with guidelines and regulations implemented by the Regierung von



Oberbayern (ROB-55.2–2532.Vet_02-20-208 and ROB-55.2–2532.Vet_02-17-135). In accordance with the animal experiment application, tumor size, behavior, breathing, body weight and posture of mice were monitored three times per week. For survival analyses, the above-described criteria (in particular: curved back, apathy, weight loss >20%, piloerection, pronounced abdominal breathing and cyanosis, spasms, paralysis, tumor size >225 mm² or one of the two measure dimensions >15 mm or open wound in the tumor area) were taken as humane surrogates for survival and recorded in Kaplan-Meier plots.

Animal experiments

MV3, A375 and patient-derived (patient sample 2) xenograft models were established in NSG mice (in total n=96) following the subcutaneous (s.c.) injection of 0.2, 1 or 0.4×10⁶ tumor cells, respectively, in 100 μL phosphate buffered saline (PBS) into the right flank of NSG mice. Syngeneic tumor model was established in C57BL/6 mice (in total n=84) by intravenous injection of YUMM1.1 overexpressing luciferase (and tyrosinase-related protein 1 (TYRP1) where mentioned) (2×10^6) into the tail vein following a partial lymphodepletion of the B-cell compartment, using 250 µg murine IgG2a anti-CD20 monoclonal antibody (18B12, Roche). Animals were randomized into treatment groups according to tumor burden. Experiments were performed by a scientist blinded to treatment allocation and with adequate controls. No time points or mice were excluded from the experiments presented in the study. For s.c. models, tumor burden was measured three times per week and calculated as mm³ given by volume=(length×width²)/2. Tumor burden of intravenous models were measured using a luciferase-based IVIS Lumina X5 imaging system. For ACT studies, 10⁷T cells with transduction efficiencies of 50-90% were injected intravenously in 100 µL PBS.

Cell line generation, culture and validation

A375, MV3, PANC-1 and B16 tumor cell lines were purchased from American Type Culture Collection. The ovalbumin overexpressing murine pancreatic cancer cell line Panc02-OVA has been previously described. The murine YUMM1.1 cell line was kindly provided by Dr Bosenberg (Yale University, USA). YUMM1.1 tumor cells were stably transduced using retroviral pMP71 vector expressing TYRP1 protein (UNIPROT entry P17643) to generate YUMM1.1 TYRP1 tumor cells. Luciferase-eGFP (LUC-GFP) overexpressing cell line YUMM1.1 TYRP1-LUC-GFP and YUMM1.1 LUC-GFP were generated according to a previously described protocol. All tumor lines were grown as previously described, and used for experiments when in the exponential growth phase.

Virus production

293Vec-Galv, 293Vec-Eco and 293Vec-RD114 were a kind gift of Manuel Caruso, Québec, Canada, and have been previously described.²³ For virus production, retroviral pMP71 (kindly provided by C. Baum, Hannover) vectors

carrying the sequence of the relevant receptor were stably introduced in packaging cell lines. Single cell clones were generated and indirectly screened for virus production by determining transduction efficiency of primary T cells. This method was used to generate the producer cell lines 293Vec-RD114 for EGFRvIII-CD28–CD3 ζ (E3), EGFRvIII with CD28 transmembrane domain lacking intracellular signaling domains (E3del) and anti-HER2–CD28–CD3 ζ (HER2 CAR). 293Vec-Galv, 293Vec-Eco and 293Vec-RD114 were grown as previously described. ²⁴ All cell lines used in experiments were regularly checked for mycoplasma species with the commercial testing kit MycoAlert (Lonza). Authentication of human cell lines by STR DNA profiling analysis was conducted in house.

T cell generation, retroviral transduction and culture

Human and murine SAR construct generation was previously described. SAR-transduced T cells will be referred to as SAR T cells. The HER2 CAR was generated with a humanized single-chain variable fragment against HER2 (4D5). Murine T cells were differentiated from splenocytes from donor mice. T cell isolation and transduction protocols have been previously described. T cells were expanded or directly expanded with T cell medium supplemented with human interleukin (IL)-15 (PeproTech) every second day. Human T cells have been differentiated and transduced using previously described protocols or directly taken into culture with human T cell medium in concentrations of 10 T cells per mL medium.

Cytotoxicity assays

For impedance-based real-time killing assays using a xCELLigence system (ACEA Bioscience), previously described,²⁰ 10⁴ tumor cells were seeded per well in a 96-well plate. Cell number was monitored over the time frame of 10 hours for every 20 min. 10⁵ T cells transduced with the indicated receptors were added to the tumor cells. For lactate dehydrogenase (LDH)-based killing assays, T cells were incubated with tumor cells and BiAb at indicated effector to target ratios and concentrations. Transduced T cells were added to the adherent tumor cells and co-cultured as indicated. LDH levels were measured according to the manufacturer's protocol (Promega). Additionally, the killing of melanoma patient samples was assessed using a flow cytometry-based readout after 48 hours of co-culture with human SAR T cells in the presence of either the anti-TYRP1/anti-EGFRvIII (αTYRP1/ αE3) which is cross-reactive to human and murine TYRP1 or the anti-human melanoma-associated chondroitin sulfate proteoglycan (MCSP, also known as CSPG4)/anti-EGFRvIII BiAb (αMCSP/αE3). Tumor cells were stained with the cell proliferation dye eFluor 450 according to the manufacturer's protocol (eBioscience). Depending on the tumor cell size $2-4\times10^4$ cells per well were co-cultured with SAR T cells in an effector to target cell ratio of 2:1 in a 96-well plate. Tumor cells were detached using trypsin. Dead cells were stained using the violet fixable

viability dye (BioLegend) for 15 min at room temperature. Following this, cell surface proteins were stained for 20 min at 4°C. For the characterization and quantification of the SAR T cells antibodies against CD3 (OKT3), CD4 (OKT4), CD8a (RPA-T8), PD-1 (EH12.2H7), 4-1BB (4B4-1), CD69 (FN50) and EGFR (A-13) (all from BioLegend) were used. Tumor and T cell counts were normalized to counting beads (Invitrogen).

Proliferation assays

SAR T cell proliferation was measured using a flow cytometry-based assay that compared fold proliferation of CD3⁺ (17A2, BioLegend) T cells over a period of 48 hours normalized to the number of T cells per bead at indicated concentrations and effector to target ratio.

Biodistribution study

For the biodistribution study of the anti-TYRP1/anti-E3 BiAb (α TYRP1/ α E3 BiAb), 2×10^6 YUMM1.1 TYRP1-LUC-GFP tumor cells were intravenously injected into C57Bl/6 mice. IVIS imaging was used to verify tumor engraftment and distribution after 13 and 20 days. αTYRP1/αE3 or αMesothelin/αE3 control BiAb (5 μg/ mouse or 10 mg/kg) were injected intraperitoneally (i.p.) into tumor-bearing (for each antibody n=3) and non-tumor-bearing mice (for each antibody n=2) on day 20. Experimental readout was taken 48 hours later. In addition to the metastasis in the lung, organs with the highest TYRP1 expression relative to baseline (skin and heart) were also harvested. Organ tissue was embedded and frozen in optimal cutting temperature compound before preparation for immunofluorescence staining and imaging.

Immunofluorescence

The 5 µm tissue cryosections were stained on chipcytometry slides (Zellkraftwerk) with an antibody (polyclonal, AF555, Thermo Fisher Scientific) against the human IgG1-based $\alpha TYRP1/\alpha E3$ BiAb, a rabbit anti-GFP antibody (polyclonal, Novus Biologicals), a secondary antibody against rabbit IgG (polyclonal, PerCP, Jackson ImmunoResearch) and Hoechst 33 342 (Thermo Fisher Scientific). The fluorescence was measured using the ZellScannerONE (Zellkraftwerk).

PCR and quantitative real-time PCR

All DNA constructs were generated by overlap extension PCR²⁴ and recombinant expression cloning into the retroviral pMP71 vector²⁰ using standard molecular cloning protocols.²⁶ RNA was extracted from cells using the InviTrap Spin Universal RNA extraction Kit (Stratec). Complementary DNA was synthesized using the Super-Script II kit (Life Technologies). Real-time PCR reactions were performed using SYBR Green PCR Master Mix (Applied Biosystems) and sequence specific primers for human MCSP, human and murine TYRP1.^{28–30} The amplification was performed with CFX Connect Real-Time PCR Detection System (Bio-Rad Laboratories) running up to 50 cycles of 5 s at 95°C followed by 30 s at 60°C after

an initial step of 95°C for 2 min. Melting curves from 65°C to 95°C were performed to evaluate the specificity of the PCR. The messenger RNA (mRNA) expression levels of human TYRP1 and MCSP were normalized to the expression of phosphoglycerate kinase. The mRNA expression levels of murine TYRP1 were normalized to the expression of β -actin.

Cytokine release assays

Murine and human SAR T cell stimulation assays were set-up at indicated concentrations and effector to target ratios. Murine SAR T cells were co-cultured with B16, YUMM1.1 TYRP1 and Panc02-OVA cell lines. Human SAR T cells were co-cultured with MV3, A375 cell lines and human melanoma samples. Cytokine quantification was measured by ELISA for the following: interferon (IFN)- γ (BD), IL-2 (BD), tumor necrosis factor (TNF)- α (R&D Systems) and granzyme B (R&D Systems).

Statistical analysis

Two-tailed student's t-test was used for comparisons between two groups, while two-way analysis of variance with Bonferroni post-test (multiple time points) was used for comparisons across multiple groups. A log-rank (Mantel-Cox) test was used to compare survival curves. All statistical tests were performed with GraphPad Prism V.8 software, and p<0.05 was considered statistically significant and represented as *p<0.05, **p<0.01 and ***p<0.001. No statistical methods were used to predetermine sample size. Investigators were blinded to treatment allocation during experiments and outcome assessment.

RESULTS

MCSP and TYRP1 are differentially expressed in melanoma

To identify suitable target structures, we assessed the expression of TYRP1 and MCSP in human melanoma cell lines as well as melanoma samples from treated and untreated patients. Both genes were shown to be highly expressed in melanoma relative to PBMC and human pancreatic cancer cell line PANC-1 control samples both at RNA and protein level (figure 1A-C, online supplemental figure 1A, online supplemental table 1). Analysis of The Cancer Genome Atlas RNA sequencing (RNA-seq) expression data also revealed MCSP to be differentially regulated in cutaneous melanoma tissue relative to skin tissue from healthy donors (figure 1D). Although the median expression of TYRP1 in cutaneous melanoma tissue was similar to the expression in skin from healthy donors, there was a far greater variability in its expression in patients with melanoma with a clear differential expression in a subset of patients (figure 1D). The expression of the targets was also analyzed across different cell types within the same patient (figure 1E), taking advantage of a single cell RNA-seq data set-GSE72056 of 3993 cells from 19 patients,³¹ which revealed a distinct, only partially overlapping pattern of expression for each antigen in tumor tissue (figure 1F and G). Furthermore,

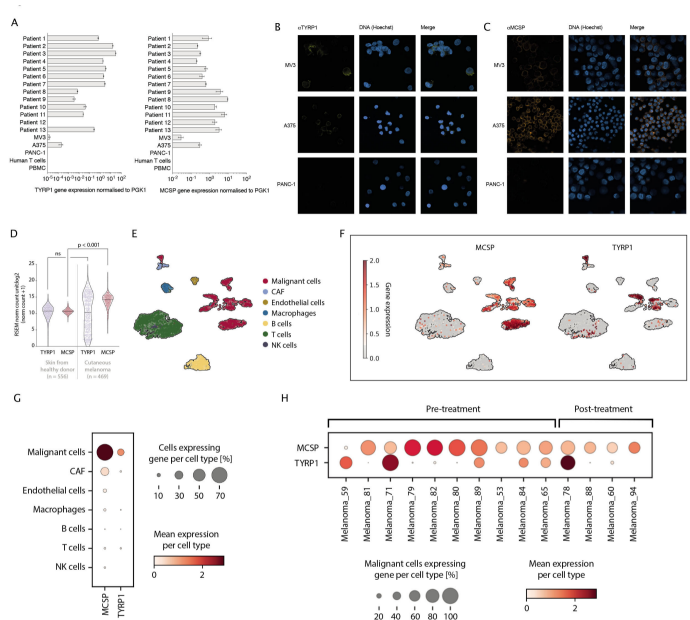


Figure 1 MCSP and TYRP1 are differentially expressed on melanomas. (A) RT-PCR MCSP and TYRP1 gene analysis of human melanoma cell lines, patient-derived melanomas and controls. (B) Microscopic analysis of TYRP1 expression on permeabilized MV3, A375 and PANC-1 cells using αTYRP1/αE3 BiAb (αTYRP1) and anti-human IgG secondary antibody. (C) Microscopic analysis of MCSP expression on MV3, A375 and PANC-1 cells using αMCSP/αE3 BiAb (αMCSP) and anti-human IgG secondary antibody. (D) TCGA analysis of RNA-seq expression of TYRP1 and MCSP in skin from healthy donors and cutaneous melanoma (cutaneous melanoma: n=469; skin from healthy donor: n=556). Scales are depicted in a log2 scale and messenger RNA normalization was estimated by the TCGA using the RSEM (RNA-seg by expectation maximization) method. (E) UMAP showing 3993 (following quality control) healthy and malignant cells from 19 previously published patients (GSE72056). Normalized gene expression values were logarithmized. Colors highlight the different cell types. Annotations of cells were provided by the authors of the respective study. (F) Expression of MCSP and TYRP1 in different cell types. Normalized gene expression values were log-transformed and visualized in a UMAP embedding. (G) Expression of MCSP and TYRP1 per cell type. Color intensity indicates mean gene expression per cell type, dot size indicates the proportion of cells expressing the respective gene per cell type. Normalized expression values were log-transformed. (H) Expression of MCSP and TYRP1 across 14 samples from melanoma patients pre-treatment or post-treatment. Color intensity indicates mean gene expression per patient, dot size indicates the proportion of malignant cells expressing the respective gene per patient. Normalized expression values were log-transformed. Statistical analysis in (D) was performed with the unpaired two-tailed Student's t-test. Experiments in subfigure (A) show mean values±SD calculated from three replicates, violin plots and the median values in (D) calculated from n independent biological replicates. Experiments in subfigures (B) and (C) show one representative of two independent experiments. CAF, cancer-associated fibroblast: MCSP, melanoma-associated chondroitin sulfate proteoglycan; NK, natural killer; PBMC, peripheral blood mononuclear cell; PGK1, phosphoglycerate kinase 1; RNA-seq, RNA sequencing; TCGA, The Cancer Genome Atlas; TYRP1, tyrosinase-related protein 1; RSEM, RNA-seg by expectation maximization.

MCSP expression on malignant cells was well-maintained in patients that received treatment (figure 1H). TYRP1 expression pattern was similar in pre-treatment and post-treatment samples but characterized by a high spread (figure 1H).

$\alpha MCSP/\alpha E3$ and $\alpha TYRP1/\alpha E3$ BiAb bind MCSP* and TYRP* melanoma cells

BiAb-mediated T cell activation is dependent on antibody aggregation on the target cell before their presentation to T cells in a polyvalent form. Our previous work on the SAR platform could show that BiAb must have a single specificity for E3 to ensure conditional SAR T cell activation in the presence of the target antigen.²⁰ This informed the BiAb design used in this study, with a trivalent and bispecific format with two specificities for the tumor antigen (TYRP1 or MCSP) and a single specificity for E3 (online supplemental figure 1B,C). The binding properties and apparent dissociation constant (K_D) of BiAb (anti-MCSP/anti-E3 BiAb (αMCSP/αE3) and anti-TYRP1/anti-E3 BiAb (αTYRP1/αE3)) to both their targets (online supplemental figure 1D,E) and EGFRvIII (online supplemental figure 1F) were analyzed by flow cytometry. The previously characterized aMesothelin/ αE3 BiAb binding the SAR and mesothelin was used as a non-melanoma targeting control construct in subsequent experiments.²⁰

$\alpha MCSP/\alpha E3$ and $\alpha TYRP1/\alpha E3$ BiAb can mediate SAR T cell activation, proliferation and differentiation

SAR constructs could be retrovirally transduced into primary murine and human T cells with high efficiencies (figure 2A). Following transduction and expansion protocols, CD4⁺ and CD8⁺ human SAR T cells were shown to be of similar frequencies and to predominantly have an effector memory phenotype (figure 2B,C). We assessed SAR T cell activation and cytokine release in both murine and human T cells. For murine T cells, we incubated SAR T cells with two TYRP1-expressing cell lines, B16 and YUMM1.1 TYRP1 and with the antigen-negative, pancreatic cancer cell line Panc02-OVA (online supplemental figure 1G-I) in the presence of $\alpha TYRP1/\alpha E3$ BiAb. Murine SAR T cells specifically released IFN- γ , IL-2, TNF- α and granzyme B and expressed the activation markers programmed cell death protein 1 (PD-1) and CD69 on co-culture with TYRP1+ melanoma cell lines, unlike in co-culture with antigen-negative Panc02-OVA tumor cells (online supplemental figure 2A,B). SAR T cells only proliferated in the presence of TYRP1-expressing tumor cells and BiAb (online supplemental figure 2C). Human SAR T cells were incubated with the MCSP⁺ and TYRP1⁺ cell lines, A375 and MV3, respectively. Only in the presence of the MCSP-targeting BiAb molecule and the target antigen, human SAR T cells released IFN-y, IL-2, TNF-α and granzyme B. In contrast, untransduced T cells (Unt) and control-E3del-transduced T cells remained inactive and did not produce cytokines or cytotoxic granules regardless of the presence of the BiAb and target

antigen (figure 2D). The frequency of CD69, PD-1 and 4-1BB-expressing SAR T cells was increased in the presence of either one of the two BiAb molecules and antigen-expressing target cells (figure 2E). CD4⁺ and CD8⁺ T cell proliferation was congruent with the activation observed, as stimulated SAR T cells proliferated more than control T cells or SAR T cells in the absence of BiAb (figure 2F).

SAR T cells can target and lyse MCSP-expressing and TYRP1expressing melanomas

Using flow cytometry-based and impedance-based assays, we evaluated whether SAR T cells could selectively lyse MCSP-expressing and TYRP1-expressing melanoma cells in the presence of a bridging BiAb. Human SAR T cells specifically eliminated antigen-positive A375 and MV3 melanoma cells when co-cultured together with either an αMCSP/αE3 or αTYRP1/αE3 BiAb whereas no lysis was detected with the antigen-negative pancreatic cancer cell line PANC-1 (figure 2G). Similarly, murine SAR T cells only lysed TYRP1⁺ B16 and YUMM1.1 TYRP1 melanoma cells in the presence of an αTYRP1/αE3 BiAb (online supplemental figure 2D,E). TYRP1-specific and MCSPspecific BiAb conditionally activated human SAR T cells in co-culture with patient-derived melanoma samples. SAR T cells showed increased expression of the activation markers CD69, PD-1 and 4-1BB, secretion of IFN-γ and proliferation, relative to E3 only or Unt T cell in presence of either of the BiAb (figure 3A-C). Also, TYRP1-specific and MCSP-specific BiAb redirected SAR T cells to target and lyse all patient-derived melanoma samples tested, whereas Unt and BiAb controls had no effect on tumor cell lysis (figure 3D).

Cleavable proteins do not impact SAR-BiAb platform efficacy and safety

With elevated levels of MCSP having been reported in the sera of patients with melanoma, ³² we sought to better understand the potential impact of soluble MCSP or TYRP1 on the SAR T cell-BiAb approach. Therefore, soluble recombinant MCSP and TYRP1 proteins were used. Proteins were added in ascending concentrations to a T cell-tumor cell co-culture to study T cell killing efficiency and kinetics. Ascending concentrations of MCSP and TYRP1, including concentrations at a physiological level, did not impair SAR T cell killing (figure 4A and online supplemental figure 3A). We also sought to test whether free soluble protein targets would induce unwanted offtumor SAR T cell activation. We found that soluble MCSP and TYRP1 did not induce SAR T cell activation in the presence of either relevant BiAb, both at physiological and supraphysiological concentrations that were tested. There, no significant changes in IFN-y levels were observed when comparing E3 and BiAb conditions to controls containing soluble recombinant MCSP or TYRP1 (figure 4B and online supplemental figure 3B). It should be noted that in this setting a higher basal SAR T cell activation was



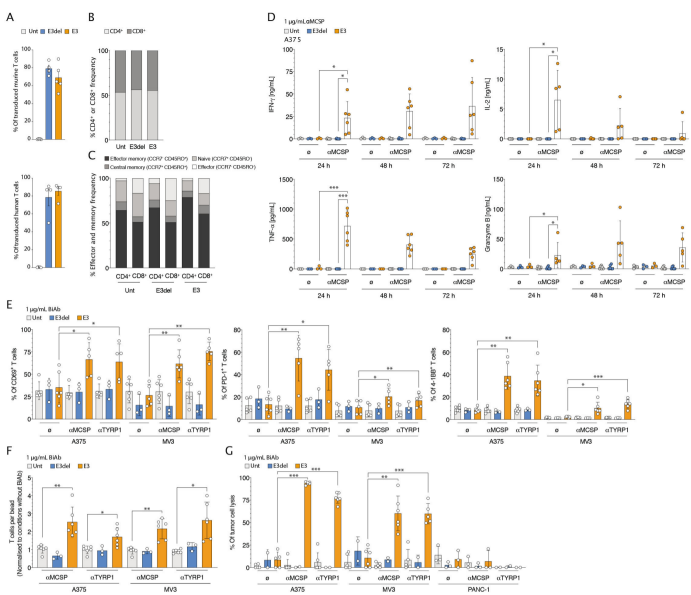


Figure 2 α MCSP/ α E3 and α TYRP1/ α E3 BiAb activate SAR T cells to mediate specific cytotoxicity against human melanoma cell lines. (A) Expression of the constructs E3 and E3del on murine T cells and human T cells from healthy donors (n=4-5). E3, E3 SAR-transduced T cells. E3del, T cells transduced with E3 SAR lacking intracellular signaling domains. (B) Frequency of CD4 and CD8 expression on human T cells. (C) Frequency of effector memory (CCR7⁻ and CD45RO⁺), central memory (CCR7⁺ and CD45RO⁺), naïve (CCR7⁺ and CD45RO⁻) and effector (CCR7⁻ and CD45RO⁻) phenotype on human T cells. (D) ELISA for granzyme B, IFN-γ, IL-2 and TNF-αon supernatant of human T cells in co-culture with human melanoma cell line A375 (E:T 2:1) and with or without αMCSP/αE3 BiAb (αMCSP, 1 μg/mL). Supernatant was taken after 24, 48 and 72 hours (n=3-6). (E) Frequency of CD69, PD-1 and 4-1BB expression on T cells after 48 hours of co-culture with A375 or MV3 (E:T 2:1) and either with or without α MCSP/ α E3 or α TYRP1/ α E3 BiAb (α MCSP or α TYRP1, 1 µg/mL) (n=3-6). (F) Following 48 hours of coculture, the CD3⁺ T cell count per bead was assessed by flow cytometry. Counts were normalized to conditions without BiAb (n=3-6). (G) The percentage lysis of melanoma cell lines A375, MV3, and antigen-negative, pancreatic cancer cell line PANC-1 by SAR T cells and either of the two BiAb was calculated using a flow cytometry-based readout after 48 hours of co-culture (n=3-6). The values shown were normalized to the Unt T cells without BiAb control condition which was taken as 0% lysis. Statistical analysis was performed using the paired two-tailed Student's t-test. Statistics shown in (D) were calculated based on the 24-hour time points. Experiments show mean values±SD calculated from n independent biological replicates. BiAb, bispecific antibodies; E:T, effector to target ratio; IFN, interferon; IL, interleukin; MCSP, melanoma-associated chondroitin sulfate proteoglycan; PD-1, programmed cell death protein-1; SAR, synthetic agonistic receptor; TNF, tumor necrosis factor; TYRP1, tyrosinase-related protein 1; Unt, untransduced T cells.

observed with the $\alpha TYRP1/\alpha E3$ compared with the $\alpha MCSP/\alpha E3$ BiAb. It appears that the SAR T cell-BiAb platform is not easily impacted by alternative soluble sources of targeted proteins and requires

immobilization of these on the tumor cell surface, as previously described for other targets in one of our previous studies.²⁰ These findings align with the fact that the BiAb was designed only to bind a

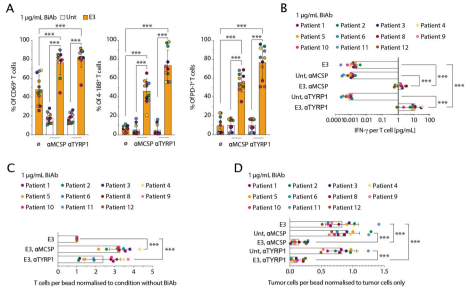


Figure 3 α MCSP/ α E3 and α TYRP1/ α E3 BiAb activate SAR T cells to mediate specific cytotoxicity against patient-derived melanoma samples. (A) Human T cells were co-cultured with patient-derived melanoma samples (effector to target ratio 2:1) and either α MCSP/ α E3 or α TYRP1/ α E3 BiAb (α MCSP or α TYRP1, 1 μg/mL) for 48 hours. The frequency of CD69, PD-1 and 4-1BB on T cells was assessed using flow cytometry. (B) Supernatant was taken and analyzed with ELISA for IFN- γ . The values were normalized to the numbers of plated T cells. (C) The CD3+ T cell count per bead was measured and normalized to conditions without BiAb. (D) The percentage lysis of the patient-derived melanoma samples by SAR T cells and either of the two BiAb was calculated based on flow cytometric readout after 48 hours of co-culture. The values shown were normalized to the tumor cells only control conditions. Statistical analysis was performed using the paired two-tailed Student's t-test. Experiments show mean values±SD. Each data point represents the mean of 2–3 biological replicates. BiAb, bispecific antibodies; IFN, interferon; MCSP, melanoma-associated chondroitin sulfate proteoglycan; PD-1, programmed cell death protein 1; SAR, synthetic agonistic receptor; TYRP1, tyrosinase-related protein 1; Unt, untransduced T cells.

membrane-proximal epitope that remains on the cell surface following cleavage.

Modular, selective and reversible activation of SAR T cells against melanoma

Melanomas are heterogeneous and stand to benefit from a modular and controllable therapeutic approach. Use of the melanoma differentiation antigens shared with other cells calls for control over these effects to antagonize potential unwanted excessive toxicities. While classic CAR T cell activity is maintained in the presence of a target antigen, SAR T cell activation is modular and controllable (figure 4C). 19 To demonstrate this in the melanoma setting, we used an in vitro stimulation assay to show how BiAb-dependent SAR activation enables greater control over T cell function (online supplemental figure 3C). As expected, following a 24-hour co-culture with MCSP⁺ TYRP1⁺ A375 tumor cells, SAR T cells could be activated in the presence of either of the two BiAb molecules (figure 4D). The same SAR T cells were then transferred to a new plate containing freshly plated A375 cells where they were co-cultured for a further 24 hours under different stimulation conditions. We found IFN-y expression was maintained when SAR T cells were redosed with either one of the two BiAb molecules (figure 4E). However, the concentration of IFN-y decreased in the absence of BiAb redosing, indicating the reversibility of SAR T cell activation. This was distinct from the lack of controllability seen with human anti-HER2 CAR T cells

when targeting HER2⁺ A375 tumor cells, which continued to sense HER2⁺ tumor cells³³ (figure 4E).

At the same time, sequential targeting of multiple antigen types would allow for more refined patient-specific tailoring of treatment and prevention of antigen-negative relapse. Through redosing with α MCSP/ α E3 BiAb (first dosing with α TYRP1/ α E3), the transferred SAR T cells remained activated, as shown by an elevated IFN- γ concentration after 48 hours of co-culture (figure 4E).

By sequentially redirecting SAR T cells towards different melanoma targets with high efficiency, the modularity of the platform was demonstrated (figure 4D, E). Overall, this approach has the potential to target a variety of melanoma-associated antigens with a level of flexibility and controllability that is superior to that of CAR T cells.

SAR T cell-BiAb combination mediates effective tumor control in vivo

To probe the in vivo function of the SAR T cell-BiAb combination, we established and used both syngeneic and xenograft melanoma models. We engrafted the YUMM1.1 TYRP1-LUC-GFP murine melanoma cell line into C57BL/6 mice. The MV3 and A375 human melanoma cell lines and a sample from a patient with primary melanoma (sample 2) were implanted into NSG mice. In the syngeneic model, following adoptive transfer, SAR T cells were shown to persist well, where SAR⁺ T cells could be tracked in the peripheral blood of mice at 7, 14 and 19 days post transfer (online supplemental figure 4A).

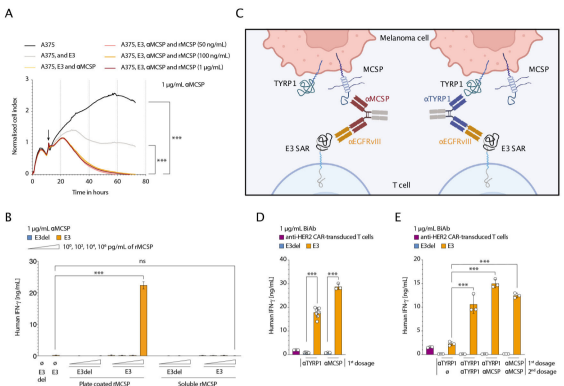


Figure 4 Modular, selective and reversible activation of SAR T cells, irrespective of soluble forms of MCSP tumor antiqen. (A) A375 melanoma cells were plated and co-cultured with human SAR T cells (E:T 2:1) and αMCSP/αE3 BiAb (αMCSP, 1 μg/ mL). Different concentrations of soluble, recombinant MCSP (rMCSP) were added. The tumor cell lysis over time was assessed using xCELLigence (n=3). The cell index was normalized to the respective time point of T cell addition as indicated by an arrow. (B) Human SAR or E3del control T cells and αMCSP/αE3 BiAb (1 μg/mL) were plated in wells either coated with different concentrations of rMCSP or where different concentrations of soluble rMCSP were added to the medium. After 48 hours the supernatant was taken and analyzed for IFN-γ using ELISA (n=3). (C) Schematic overview of SAR-transduced T cells targeting TYRP1⁺ MCSP⁺ melanoma cells via an αTYRP1/αE3 or αMCSP/αE3 BiAb. (D and E) A modularity stress test was carried out using αMCSP/αE3 or αTYRP1/αE3 BiAb (αMCSP or αTYRP1, 1 μg/mL). SAR or E3del control T cells were co-cultured with A375 tumor cells (E:T 2:1). HER2 CAR T cells were used as a control and co-cultured with HER2+ A375 tumor cells (no BiAb was added). At assay start, co-cultures received either α MCSP/ α E3 or α TYRP1/ α E3 BiAb (first dosage). At 24 hours, the T cells were washed to remove residual BiAb and transferred to freshly plated A375 tumor cells. Co-cultures were then either redosed with the same BiAb, redosed with the BiAb against the other target, or not redosed after initial dosing (second dosage) and incubated for another 24 hours. At 24 (D) or 48 hours (E), supernatants were taken and ELISA for human IFN-γ were performed (n=3). Analyses of differences between groups for (A) were performed using two-way analysis of variance with correction for multiple testing by the Bonferroni method. For statistical analysis of (B), (D) and (E), the unpaired two-tailed Student's t-test was used. Experiments show mean values±SD calculated from at least three biological replicates and are representative of three independent experiments. BiAb, bispecific antibodies; E:T, effector to target ratio; IFN, interferon; EGFRvIII, epidermal growth factor receptor variant III; MCSP, melanoma-associated chondroitin sulfate proteoglycan; SAR, synthetic agonistic receptor; TYRP1, tyrosinase-related protein 1; HER2, human epidermal growth factor receptor.

Mice that were treated with SAR T cells and repeated αTYRP1/αE3 BiAb dosing were able to clear the disease and achieved long-term survival (4 out of 10 mice with a complete response) (figure 5A,B, online supplemental figure 4B). In an endpoint experiment (19 days after T cell transfer) the lungs were harvested and analyzed using flow cytometry. SAR T cell and repeated αTYRP1/αE3 BiAb combination treatment led to a complete tumor clearance in four out of five mice based on a flow cytometry and IVIS readout (figure 5C and online supplemental figure 4C). In contrast, a single dose of the αTYRP1/ αE3 BiAb in combination with SAR T cells only showed transient tumor control and did not lead to tumor clearance nor prolonged survival of the treated mice (online

supplemental figure 4D,E), indicating the necessity for redosing for maintained SAR activity in vivo. This necessity for redosing demonstrates a reversibility in SAR T cell activity on dosing cessation.

In order to analyze the antigen-specificity of the approach in vivo, mice bearing antigen-negative YUMM1.1 LUC-GFP tumors were treated with SAR T cell-BiAb combination. Treatment of antigen-negative tumors did not impact tumor growth and survival compared with control mice treated with SAR T cells or the vehicle solution (online supplemental figure 4F,G) underpinning the necessity of target antigen expression for the functionality of the approach. To further analyze the specificity of the αTYRP1/αE3 BiAb, TYRP1-expressing organs

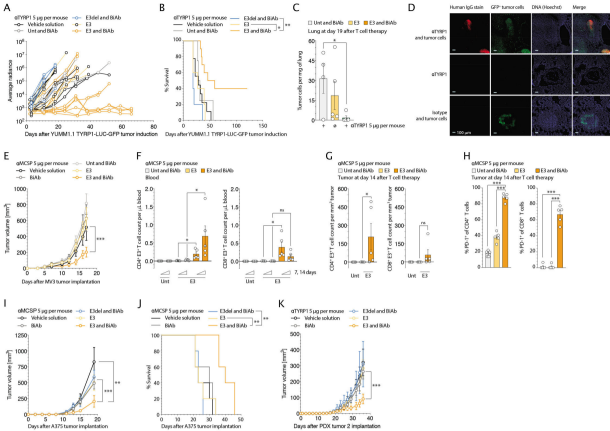


Figure 5 Treatment with the SAR T cell-BiAb combination is effective in syngeneic and xenograft melanoma models and enhances survival in vivo. (A) C57BL/6 mice were injected i.p. weekly with an anti-CD20 depleting antibody (250 µg/injection) starting 7 days before tumor cell injection. Mice were inoculated intravenously with 2×10⁶ YUMM1.1 TYRP1-LUC-GFP tumor cells. Mice were treated with a single intravenous injection of T cells 4 days after tumor cell injection. Simultaneously, antibody treatment was given by i.p. injections of the α TYRP1/ α E3 BiAb (α TYRP1, 5 µg/injection) which was redosed two times per week. In vivo luminescent signal imaging was performed one time per week using IVIS. Treatment groups were as follows: SAR T cells and αTYRP1/αE3 BiAb (n=10), E3del T cells and αTYRP1/αE3 BiAb (n=5), SAR T cells (n=5), Unt T cells and αTYRP1/ αE3 BiAb (n=4), and the vehicle solution (n=9), (B) Percentage survival readout, (C) Tumor burden per mg of lung tissue 19 days after T cell therapy using a flow cytometry-based readout. Mice were treated with either SAR T cells and α TYRP1/ α E3 BiAb (n=5), Unt T cells and αTYRP1/αE3 BiAb (n=4) or SAR T cells only (n=5) 4 days after tumor induction. (D) Immunofluorescence imaging of the αTYRP1/αE3 BiAb and tumor cell-derived GFP in lung tissue was carried out with anti-human IgG and anti-GFP stainings. Mice were injected intravenous either with 2×10⁶ YUMM1.1 TYRP1-LUC-GFP cells or with vehicle solution. After 20 days the mice were injected either with 5 μg αΤΥΡΡ1/αΕ3 BiAb, an isotype BiAb or with the vehicle solution. Following 48 hours of incubation, the lung, heart and skin were harvested, stained and imaged using ZellScannerONE. The groups were as follows: Tumor and αTYRP1/αE3 BiAb (n=3), αTYRP1/αE3 BiAb only (n=2), tumor and isotype BiAb (n=3). (E) NSG mice were inoculated s.c. with 1×10⁶ MV3 tumor cells. Mice were treated with a single intravenous injection of T cells 5 days after tumor cell injection. Simultaneously, antibody treatment was given by i.p. injections of the αMCSP/αE3 BiAb (αMCSP, 5 μg/injection) which was redosed two times per week. Treatment groups were as follows: SAR T cells and the α MCSP/ α E3 BiAb (n=5), Unt T cells and αMCSP/αE3 BiAb (n=4), SAR T cells only (n=5), BiAb only (n=5), and the vehicle solution (n=5). (F) In an endpoint experiment. SAR T cell persistence in the blood and in the tumor was analyzed using a flow cytometry-based readout 14 days after T cell transfer. The mice were treated with either SAR T cells and α MCSP/ α E3 BiAb, Unt T cells and α MCSP/ α E3 BiAb or SAR T cells only (for each group n=5). (G) SAR T cell infiltration per mm³ tumor. (H) Frequency of PD-1 expression on human CD4⁺ and CD8⁺ T cells in the tumor. (I) NSG mice were injected s.c. with 0.2×10⁶ A375 tumor cells. The mice were treated according to the experiment in (E) 11 days after tumor induction (for E3del and αMCSP/αE3 BiAb: n=4, for other groups: n=5). (J) Percentage survival readout. (K) NSG mice were injected s.c. with 0.4×10⁶ patient-derived melanoma cells (patient sample 2). The mice were treated according to the experiment in (E) 12 days after tumor induction with $\alpha TYRP1/\alpha E3$ BiAb (for E3del and αTYRP1/αE3 BiAb, and vehicle solution: n=4, for other groups: n=5). For statistical analysis of survival data, the log-rank test was applied. Analyses of differences between groups for (E), (I) and (K) were performed using two-way analysis of variance with correction for multiple testing by the Bonferroni method. For statistical analysis of (C), (F), (G) and (H) the unpaired twotailed Student's t-test was used. Experiments show mean values ±SEM calculated from n biological replicates, one experiment for (B), (C), (F), (G), (H) and (K), and one representative of two independent experiments in (A), (E), (I) and (J). BiAb, bispecific antibodies; i.p., intraperitoneally; MCSP, melanoma-associated chondroitin sulfate proteoglycan; PD-1, programmed cell death protein 1; PDX, patient-derived xenograft; SAR, synthetic agonistic receptor; s.c., subcutaneously; TYRP1, tyrosinase-related protein 1; Unt, untransduced T cells.



(heart and skin) and tumor-bearing lungs were harvested 48 hours after BiAb injection and the BiAb was stained using immunofluorescence. $\alpha TYRP1/\alpha E3$ BiAb specifically bound to YUMM1.1 TYRP1-LUC-GFP tumor cells in the lung, while no comparable binding of the BiAb could be detected in other TYRP1-expressing tissues, indicative of its selectivity for melanoma cells (figure 5D and online supplemental figure 4H).

In the human MV3 xenograft model, a strong antitumoral response was again observed in the group treated with SAR T cells and αMCSP/αE3 BiAb compared with all controls (figure 5E). In this group, durable persistence of SAR-transduced T cells was seen by flow cytometry at 7 and 14 days post transfer (figure 5F). At the experimental endpoint, tumors were harvested and analyzed by flow cytometry. Higher numbers of tumor infiltrating SAR T cells were found in mice that received the SAR T cell-BiAb treatment combination (figure 5G), with CD4⁺ T cells persisting better than CD8⁺ T cells in contrast to any of the controls. Phenotyping of transferred T cells at the experimental endpoint revealed maintenance of the effector memory phenotype they had prior to adoptive transfer (as determined by CD45RO and CCR7 expression) (online supplemental figure 4I). Furthermore, PD-1 expression was increased in SAR T cell-BiAb treated mice compared with SAR T cell only or Unt T cells and BiAb control conditions (figure 5H). Similar to the MV3 model, the A375 xenograft model, demonstrated comparable sensitivity to SAR T cells and BiAb resulting in improved tumor control and prolonged survival (figure 5I and I). Additionally, in a patient-derived xenograft model, treatment with the $\alpha TYRP1/\alpha E3$ BiAb and SAR T cells resulted in reduced tumor growth in mice receiving SAR T cell and BiAb combination (figure 5K).

DISCUSSION

We could demonstrate that MCSP and TYRP1 expression remains differentially expressed on primary melanoma samples of patients pre-treatment or post-treatment. We reasoned that their targeting using a modular and controllable T cell therapy platform, in the form of the SAR T cell-BiAb approach, could be an effective strategy and probed this hypothesis in vitro and in vivo. In order to demonstrate the translational relevance of the approach, we selected a series of relevant in vitro and in vivo models. We used immunocompetent syngeneic mouse models to better control for the immune system's impact on the treatment approach and primary human melanoma models and cell lines to test the efficacy of the approach in treated as well as untreated melanoma models.

Our results demonstrate the efficacy of the approach in several in vitro and in vivo models. SAR-engineered T cells could be redirected towards MCSP-expressing and TYRP1-expressing melanoma cells in the presence of the α MCSP/ α E3 or α TYRP1/ α E3 BiAb. SAR T cells efficiently targeted and lysed MCSP-expressing and TYRP1-expressing melanoma cell lines. Targeted specificity and

killing capacity were retained when targeting patientderived melanomas, using in vitro co-cultures, as well as several syngeneic and xenograft in vivo mouse models.

The potent cytolytic effects of the platform in the syngeneic model resulted in 4 out of 10 mice completely curing the tumor in the observed period. We also observed strong treatment effects in xenograft models which are comparable to results shown in preclinical approaches using melanoma-specific CAR T cells. ^{33–36} In these studies, HER2 and GD2 were the targeted antigens. HER2 CAR T cells have been shown to pose a risk of lethal toxicity, with cytokine release syndrome from on-target off-tumor recognition of HER2. ³⁷ While efforts are being made to create safer HER2-targeting CAR T cells, ³⁸ we demonstrate herein that the SAR T cell-BiAb approach, with its controllable and reversible facets, can be a ready-made solution for lowering and controlling toxicity. ³⁹

MCSP is differentially expressed on the surface of over 85% of melanomas.⁴⁰ It provides tumorigenic signals to melanoma cells that stimulate growth, motility, and tissue invasion. 41-44 It was therefore unsurprising to discover its expression was retained on patient with primary melanoma samples irrespective of the treatment history. Furthermore, we could show expression on human melanoma cell lines on transcript and protein level which was in line with the functional readouts. MCSP has been described as potential target for CAR T cell therapy in melanoma and glioblastoma. 39 45 TYRP1, a transmembrane glycoprotein naturally involved in melanin production, 46 has been identified as an antigen highly expressed in melanoma and stably expressed during disease progression. 47 TYRP1 expression could also be observed on primary patient samples and human melanoma cell lines. Discrepancies between transcript and protein expression could be partly caused by internalization of TYRP1. 48 49 The influence of target antigen internalization on treatment outcome must be investigated in further studies. Nonetheless, potent treatment effects were shown when targeting TYRP1⁺ cells with the SAR-BiAb approach in syngeneic and human xenograft models. In a phase I study an anti-TYRP1 monoclonal antibody was administered in patients with relapsed or refractory melanoma, where no serious adverse events were observed, indicating the potential safety of targeting.⁵⁰

With low off-tumor expression detected in some healthy tissues for both targets, it was necessary to design antibodies that would bind MCSP and TYRP1 with sufficient avidity, while minimizing the potential for off-tumor toxicity. Melanoma-specific BiAb were designed with two binding arms for the tumor target. This increased the binding avidity of the BiAb to melanoma cells showing higher target expression compared with healthy cells. Thus, the binding strength could be increased while minimizing the risk of on-target off-tumor toxicity that is often associated with increased binding affinities. Similarly, the risk of targeting endogenous T cells is limited since no expression of the target antigens was observed on human T cells. The modular facets of the SAR platform were

previously demonstrated in an acute myeloid leukemia xenograft model, ¹⁹ and again substantiated herein. The simultaneous, or in the event of antigen loss, sequential targeting of multiple tumor antigens has proven to be an effective approach in the treatment of B-cell malignancies, ⁵¹ and is further evidenced by several approaches attempting to render CAR T cells more adaptable. ⁵² We thereby reason, that a platform allowing change or simultaneous targeting of multiple antigens, with one cellular construct comes with clear advantages of feasibility and flexibility.

MCSP levels have been shown to be elevated in the sera of patients with advanced melanoma.³² We experimentally tested the potential impact of soluble MCSP on the conditional specificity of the SAR T cell-BiAb approach. At physiological levels, free-MCSP did not activate SAR T cells in the presence of BiAb. Free targeted protein in the tumor microenvironment must be considered as a potential hindrance to the efficacy of the SAR T cell-BiAb combination, as well as other adoptive T cell therapies. This could become especially problematic in targeting cleavable proteins, such as MCSP, and the targeting of membrane-associated forms of proteins could maximize efficiency in this regard.³² In fact, shed glypican-3 was shown to induce a blocking effect on CART cells targeting glypican-3-expressing hepatocellular carcinoma.⁵³ Probably because of the peculiarities of the SAR T cell-BiAb platform, we have not found such mechanism to impact efficacy or activity.

Anti-melanoma CAR T cell therapy has shown limited efficacy in the clinic thus far.⁵⁴ Results from the CARPETS phase I trial (NCT02107963) showed limited persistence of GD2 CAR T cells in patients with metastatic melanoma, with CAR transgenes only detected at low levels in patient peripheral blood after 4 months. T cell exhaustion and activation induced cell death have been shown to hinder the persistence and function of adoptively transferred T cells. 55 56 Recent work from Weber and colleagues showed that transient rest, using enforced CAR molecule downregulation via a drug-regulatable system, could restore CAR T cell functionality.⁵⁷ An advantage of the SAR T cell-BiAb system is that it is an adaptor platform inherently regulatable, via its BiAb facet. The reversibility of SAR T cell activation was demonstrated with the cessation of BiAb dosing in vitro and in vivo. This makes it very straightforward to incorporate a transient rest period simply by modifying the dosing schedule of the BiAb, thus recapitulating the previously mentioned regulatable system. Recent work by Phillipp and colleagues also showed that transient BiAb dosing reduced T cell exhaustion and improved CD3⁺ T cell engager efficacy.⁵⁸ The BiAb molecules we used have an IgG format, which extends their half-life in comparison to Fc-deficient BiAb. 59 Engineering the half-life of BiAb to offer greater flexibility towards patient needs would be a straightforward approach, that could also be beneficial. Importantly, we could previously demonstrate the use of different antibody formats and half-lives to activate and redirect SAR T cells, ¹⁹ 20 opening the door to

such optimizations. Inadequate T cell infiltration into the tumor and a suppressive milieu therein are the subject of ongoing investigation that could broadly improve the therapeutic success of adoptive T cell therapies in solid tumors,⁵ including melanoma. Equipping SAR T cells with relevant chemokine receptors while shielding them from local immune suppression could further improve treatment efficacy and warrant further investigation.²⁴ ²⁶

Harnessing the apparent advantages of the SAR T cell-BiAb platform for melanoma therapy has yielded very promising results in our preclinical models. With evident potential for improved clinical benefit, we believe these findings warrant further characterization in more advanced preclinical models and ultimately clinical studies.

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Contributors Conceptualization: M-RB, FM, CK and SK. Methodology: FM, M-RB, DB, PJM, JJ, MT, JK, LM, CK and MS. Investigation: M-RB, FM, HO and MT. Validation: HO, MS, A-KK, TL, SS, SM and RG. Writing—original draft: M-RB and FM. Writing—review and editing: M-RB, FM, BLC, JK, MG, MS, MT, MVH, SE, CK and SK. Funding acquisition: SE and SK. Resources: DB, AG, CM, SR, MPL, MVH, SE, CK and SK. Supervision: RK, CM and SK. Guarantor: M-RB and FM.

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