Combination of T-Cell Bispecific Antibodies With PD-L1 Checkpoint Inhibition Elicits Superior Anti-Tumor Activity

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T-cell Bispecific Antibodies (TCBs) elicit anti-tumor responses by cross-linking T-cells to tumor cells and mediate polyclonal T-cell expansion that is independent of T-cell receptor specificity. TCBs thus offer great promise for patients who lack antigen-specific T-cells or have non-inflamed tumors, which are parameters known to limit the response of checkpoint inhibitors. The current study deepens the understanding of TCB mode of action and elaborates on one of the adaptive resistance mechanisms following its treatment in vivo in humanized mice and syngeneic pre-clinical tumor models. Single-agent TCB treatment reduced tumor growth compared with controls and led to a 2–10-fold increase in tumor-infiltrating T-cells, regardless of the baseline tumor immune cell infiltration. TCB treatment strongly induced the secretion of CXCL10 and increased the frequency of intra-tumor CXCR3+ T-cells pointing to the potential role of the CXCL10-CXCR3 pathway as one of the mechanisms for T-cell recruitment to tumors upon TCB treatment. Tumor-infiltrating T-cells displayed a highly activated and proliferating phenotype, resulting in the generation of a highly inflamed tumor microenvironment. A molecular signature of TCB treatment was determined (CD8, PD-1, MIP-a, CXCL10, CXCL13) to identify parameters that most robustly characterize TCB activity. Parallel to T-cell activation, TCB treatment also led to a clear upregulation of PD-1 on T-cells and PD-L1 on tumor cells and T-cells. Combining TCB treatment with anti-PD-L1 blocking antibody improved anti-tumor efficacy compared to either agent given as monotherapy, increasing the frequency of intra-tumoral T-cells. Together, the data of the current study expand our knowledge of the molecular and cellular features associated with TCB activity and provide evidence that the PD-1/PD-L1 axis is one of the adaptive resistance mechanisms associated with TCB activity. This mechanism can be managed by the combination of TCB with anti-PD-L1 blocking antibody translating into more efficacious anti-tumor activity and prolonged control of the tumor outgrowth. The elucidation of additional resistance mechanisms beyond the PD-1/PD-L1 axis will constitute an
important milestone for our understanding of factors determining tumor escape and deepening of TCB anti-tumor responses in both solid tumors and hematological disorders.

Keywords: solid tumors, immunotherapy, T-cell bispecific antibody, carcinoembryonic antigen T-cell bispecific antibody, programmed death–ligand 1, combination, humanized mice

INTRODUCTION

Targeting T-cells with antibodies that directly enhance T-cell activity, including the checkpoint inhibitory molecules (CPIs) programmed death receptor 1 (PD-1), PD-ligand 1 (PD-L1), and cytotoxic lymphocyte activated antigen 4 (CTLA4) has become an established approach in clinical practice (1–3). Antibodies to checkpoint molecules have gained broad approval in various tumor indications for the treatment of advanced cancer types such as metastatic melanoma, advanced non-small cell lung cancer, or renal cell carcinoma (4). However, despite these advances, obstacles still exist including the inability to predict treatment efficacy and patient response, the need for additional biomarkers, the development of primary and secondary resistance to cancer immunotherapies, the lack of clinical study designs that are optimized to determine efficacy and toxicity (and their relationship), and high treatment costs (5).

T-cell Bispecific Antibodies (TCBs) elicit anti-tumor responses by cross-linking of T-cells to target tumor cells (6, 7). TCB-mediated polyclonal T-cell activation is independent of the T-cell receptor specificity and does not require (at least initially) costimulatory signals. Thus, factors normally affecting the efficiency of CPIs to mount an endogenous anti-tumor immune response, including MHC downregulation, antigen presentation, the frequency of antigen-specific T-cells, T-cell receptor affinity, and T-cell avidity, are less relevant for TCB activity. Hence, TCBs are a highly attractive approach to activate T-cells regardless of their antigen specificity. Due to the increase of intra-tumor T-cell infiltration upon treatment (8–10), TCBs offer great promise in patients that lack the baseline antigen-specific T-cells (or any type of T-cells, the so called immune desert tumors), which is thought to render responses to checkpoint inhibition less likely (5).

Although more than forty different bispecific antibodies have been described to date (6, 11–14), the promising results obtained in preclinical studies do not translate directly into the clinical setting. Only two TCBs were approved for use in the clinic so far: catumaxomab and blinatumomab. Catumaxomab targets epithelial cell adhesion molecule (EpCAM) and was initially approved by the European Medicines Agency (EMA) in 2009 for the treatment of malignant ovarian ascites (15). Catumaxomab has not been marketed in the EU since 2014 and market authorization was withdrawn in 2017. Blinatumomab targets CD19 and was approved by the FDA and EMA in 2014 and 2015, respectively, for the treatment of Philadelphia chromosome negative B cell acute lymphoblastic leukemia (16). Promising clinical activity has been reported with other TCBs, particularly in hematological malignancies (6, 7, 17).

Clinical development of TCBs in solid tumors has been challenging and may be hampered by multiple constraints. These include the lack of tumor-specific antigens that are not expressed in primary epithelium (18), the local suppressive tumor microenvironment [characterized by expression of IL10, TGFβ, IDO, COX-2, adenosine, and arginase, and presence of regulatory T-cells and myeloid-derived suppressor cells (19, 20)], and the physical barriers that trap immune cells in the stroma [also called immune exclusion, (21, 22)]. These factors may limit the frequency and activation of effector cells within the tumor (5, 23). Moreover, a dysfunctional T-cell state characterized by the abundance of intra-tumoral PD-1hi T-cells hampered TCB activity ex vivo (24), providing an additional primary resistance mechanism affecting TCB activity.

TCB-induced T-cell activation is has been shown to upregulate PD-1 expression on T-cells and induce PD-L1 expression on tumor cells (IFNγ driven) (8, 9, 25–29). This may lead to adaptive immune resistance mechanisms related to the TCB mode of action, similar to what has been described for checkpoint inhibition (30, 31). The same studies provided pre-clinical evidence that blockade of the PD-1/PD-L1 axis restores TCB activity in vitro and in vivo and provided the rationale for combining TCBs with therapeutic strategies targeting T-cell dysfunction in the clinic to potentiate the activity of TCBs (13, 32). These studies led to several Phase 1 trials evaluating T-cell engaging bispecific antibodies in combination with checkpoint inhibitors, particularly anti-PD-1/ PD-L1 antibodies (6, 7, 11).

We have previously described the so-called 2:1 TCBs that carry two tumor antigen binding moieties and a single CD3 binding moiety in an IgG-based format (33, 34). This 2:1 format shows advantageous properties over classical 1:1 TCBs (9). In the current study, we deepen the understanding of TCB mode of action by characterizing molecular and cellular features of immune cells and tumors following TCB treatment in vivo in humanized mice and syngeneic tumor models, and provide additional evidence that combination with checkpoint inhibitors of the PD-1/PD-L1 axis improves TCB activity. We demonstrate that combination treatment increases the frequency of total intra-tumor T-cells, and identify the CXCL10-CXCR3 pathway as one of the potential mechanisms mediating such increase. We also show that combination treatment lowers the intra-tumor frequency of putatively exhausted T-cells. Together, the study corroborates the relevance of blocking the PD-1/PD-L1 axis to improve TCB activity and highlights the importance of exploring additional combinations that enable generation of T-cells maintaining the optimal functional status.
MATERIALS AND METHODS

Therapeutic Antibodies

The human carcinoembryonic antigen TCB (CEA-TCB; cibisatamab) monoclonal antibody was generated as described previously (8). A murine surrogate of CEA-TCB (muCEA-TCB) was generated for studies in fully immunocompetent mice on a fully silent murine IgG1 backbone. MuCEA-TCB antibody was generated using an anti-CEA binder that binds to a partially overlapping (but not competing) epitope to the human CEA binder include in CEA-TCB antibody and contains the murine-specific anti-CD3 binder (clone 2C11). The potency of muCEA-TCB is about 10-fold lower than the potency of human CEA-TCB, attributed to the lower activity of the murine anti-CD3 antibody clone and putatively to the lower cytotoxic activity of murine splenocytes in ex vivo killing assays used to profile the activity of the surrogate molecule (data not shown). The anti-PD-L1 monoclonal antibody used in the humanized NOG mouse studies is the clone YW243.55.S70 on a mlgG1 DAPG backbone. The anti-PD-L1 monoclonal antibody used in the immunocompetent mouse studies was mlgG1 anti-PD-L1 antibody (clone 6E11), which reacts to human and murine PD-L1 (35).

Cell Lines

The MKN-45 human gastric adenocarcinoma cell line was used in humanized NOG mouse studies was purchased from DSMZ (Braunschweig, Germany; Cat No.: ACC 409). The cells were cultured in DMEM containing 10% FCS and 1% Glutamine and split 1:3 to 1:5 every 2–4 days. The HPAF-II pancreatic adenocarcinoma cell line that was also used in humanized NSG mouse studies was purchased from the American Type Culture Collection (Manassas, VA 20110 USA; Cat No.: CRL-1997). The MC38-hCEA for use in the immunocompetent human CEA transgenic mouse (huCEA Tg) study were derived from a mouse colon adenocarcinoma and engineered to express human CEA, obtained from Beckmann research institute of City of Hope (Duarte, CA, USA) (36). The WSU-DLCL2 human diffuse large cell lymphoma (DLBCL) cell line used in the humanized mice studies with CD20-TCB were obtained from the European Collection of Cell Culture. MV3 is a human melanoma cell line, that was established by Ruiter DJ (Department of Pathology, Collection of Cell Culture. MV3 is a human melanoma cell line, that was established by Ruiter DJ (Department of Pathology, University Hospital Nijmegen, Netherlands) (37). The cells were cultured in DMEM, containing 10% FCS and 1% Glutamax and split 1:3 to 1:6 every 3–4 days. Skov3 (ATCC, HTB-77) is a human ovarian adenocarcinoma cell line. The cells were cultured in RPMI containing 10% FCS and 1% Glutamine and split 1:4 to 1:8 every 4 days. HT-29 (ATCC, HTB-38) is a human, female Caucasian colon adenocarcinoma cell line. The cells were cultured in McCoy’s 5A +10% FCS and 2nM GluMax and split 1:3 to 1:8 every 2–4 days. LS174T (ATCC, CL-188) is a human colon carcinoma cell line. The cells were cultured in in DMEM containing 10% FCS and 1% Glutamine and split 1:3 to 1:5 every 2–4 days.

Mouse Models

All mice were maintained under specific-pathogen-free condition with daily cycles of 12-h light/12-h darkness according to international (Federation of European Laboratory Animal Science Associations) and national [Gesellschaft für Versuchstierkunde/Society of Laboratory Animal Science (GV-Solas) and Tierschutzgesetz (TierSchG)] guidelines. The study protocol was reviewed and approved by the local government (license ZH193/2014). Animals were maintained for 1 week after arrival to get accustomed to the new environment and for observation. Daily continuous health monitoring was conducted.

Hematopoietic stem cell humanized mice were generated in house. Briefly, 4–5-week-old female NOG (NOD.Cg-Prkdcreid Il2rgtm1Sug) mice (Taconic, Cologne, Germany) or NOD scid gamma (NSG) mice (Jackson Laboratory, Sacramento, CA USA) were injected i.p. with 15 mg/kg Busulfan (Busilvex, Pierre Fabre Limited) in a total volume of 200 μl. Twenty-four hours later, mice were injected intravenously (i.v.) with 1 × 10^6 CD34+ cord blood cells (STEMCELL Technologies Inc, Grenoble, France). Fifteen weeks after cell injection, mice were bled and screened for successful humanization by flow cytometry. The generation of these mice will be reported in detail elsewhere.

Immunocompetent human CEA transgenic (huCEA Tg) C57BL/6J mice were obtained under license agreement from Beckmann research institute of City of Hope (36). Double transgenic CEA424-SV40Tag x CEACAM5 Tg mice were obtained under license agreement from LIFE-Center of “Klinikum der Universität München” (Prof. Dr. Wolfgang Zimmermann) (38, 39). Both strains were bred by Charles River Laboratories (Lyon, France).

Subcutaneous Tumor Cell Inoculation

MKN-45 cells, HPAP-II cells and WSU-DLCL2 cells were cultured in RPMI containing 10% FCS (PAA Laboratories, Pasching, Austria) and 1% Glutamax (Gibco, Zug, Switzerland) at 37°C in a water-saturated atmosphere at 5% CO2. Afterwards, 1 × 10^6 cells of MKN-45 or HPAP-II cells (1.5 × 10^6 for WSU-DLCL2 cells) were injected s.c. using a 1:1 mixture of RPMI medium and Matrigel in a total volume of 100 μl. MC38-huCEA cells were maintained in RPMI medium supplemented with 10% FCS, 50 μg/ml Genetecin (G418, Gibco). Mice were injected s.c. with 0.5 × 10^6 cells using RPMI medium and Matrigel (1:1) in a total volume of 100 μl.

Therapeutic Antibody Treatment

All mice were injected i.v. or i.p. with 200 μl of the appropriate solution. The mice in the vehicle group were injected i.v. with Histidine buffer (20 mM Histidine, 140 mM NaCl, pH 6.0) and the treatment group with the antibody diluted with Histidine buffer to a volume of 200 μl.

Tumor Volume Measurement

Tumor volume (½ [length × width^2]) was measured 3 times per week by caliper.

Necropsy

At study termination, mice were bled under anesthesia (retro-orbital) and sacrificed. Fresh blood was collected in Heparin tubes. Tumors were surgically removed from all animals and cut into three pieces. One part was snap-frozen in liquid nitrogen for RNA sequencing analysis and multiplex cytokine/chemokine
analysis, one part was fixed overnight in 4% paraformaldehyde for histological analysis, and one part stored in PBS for flow cytometric analysis.

**Whole Body SPECT/CT Imaging Technique**
CEA-TCB and untargeted TCB (DP47-TCB) antibodies were conjugated with 2-(4-isothiocyanatobenzyl)-1,4,7,10-tetraazaacyclododecane-1,4,7,10-tetraacetic acid (p-SCN-Bz-DOTA; Macrocyclic, Plano, TX, USA) and radiolabeled with $^{111}$In and $^{177}$Lu, respectively, as described previously (40, 41). The integrity of the radiolabeled antibodies was confirmed before incubation with antibodies for intracellular staining. Final cell suspension was washed and acquired using a BD LSRFortessa™ cell analyzer (BD Biosciences). Manual gating was carried out using Flowjo. Living CD45+ tumor-infiltrating immune cells were further analyzed to define specific immune cell subsets and their activation and differentiation status.

**Histological Analysis**
Briefly, tumor tissue was fixed in 4% paraformaldehyde overnight and embedded in paraffin. Sections (4 μm) were cut using a microtome (Leica) and mounted on glass slides. Samples were de-paraffinized and heat antigen retrieval was performed prior to immunostaining with antibodies specific for human CEA (Roche in house), human CD3 (Abcam, Cat No.: ab5690), and human PD-L1 (Ventana, Cat No.: 790-4905). Sections were counterstained with hematoxilin (Sigma Aldrich) and slides were scanned using Olympus VS120-L100 Virtual Slide Microscope. Quantification of percentage of positive stained tumor area for PD-L1 or CEA was performed in whole scans with Definiens software. Raw data was transferred to GraphPad software for analysis of significance. A total of five mice per treatment group was evaluated.

**Cytokine/Chemokine Analyses**
Cytokine/chemokine analyses from tumors of humanized mice were conducted using the Bio-Plex Pro™ Human Chemokine Panel, 40-Plex (Bio-Rad Laboratories AG, Cressier, Switzerland, Cat No.: 171AK99MR2). Small tumor fragments were snap frozen and whole protein was isolated in the presence of Easypack Protease Inhibitor Cocktail (Roche; ref 589297001) using the Precellys®24 Homogenizer and Bio-Plex® Cell Lysis Buffer following manufacturer instructions. Whole protein content was measured with BCA™ Protein Assay Kit (Thermo Scientific) before cytokine measurement was performed.

**RNA Sequencing Analysis**
High molecular weight RNA (>200 base pairs) was extracted and RNAseq libraries were generated and sequenced using the TruSeq® Stranded mRNA Kit (Illumina®) as per manufacturer’s instructions at Expression Analysis Inc. Reads for each sample were processed using the following steps: First, reads were aligned to the human and mouse transcriptome (based on Ensembl v60) using Bowtie2 (42) with sensitive settings. In a second step, yet unmapped reads were aligned to the Human and mouse genome (hg19), and both mappings to genome and transcriptome were combined using in-house software. Reads mapped to both transcriptomes at the
same time were discarded from further analysis. Raw counts were used to create an R DGEList object [edgeR version 3.24.3 (43)].

Normalization factors were calculated using the calcNormFactors function. Genes were normalized by Trimmed Mean of M-values (TMM), and were subjected to DE analyses using the voom and lmFit functions in the limma package [version 3.38.3; (44)]. Gene set enrichment analysis was conducted using the fgsea R package [Version 1.8.0; (45)] with minSize=55, maxSize=500 an nperm=100000. Genes were ranked by the corresponding log2 fold-change and GO gene sets (C5) from MsigDB signaling database (46) were used.

**Upregulation of PD-1/PD-L1 In Vitro**

Surface expression of PD-1 on CD4+ or CD8+ T-cells and PD-L1 on surviving tumor cells was assessed after a classical tumor cell lysis assay. Briefly, peripheral blood mononuclear cells (PBMCs) from healthy volunteers were isolated with standard techniques. MKN-45 target T-cells were plated at a density of 1.4 × 10⁶ cells/well in flat-bottom 24-well plates 1 day before the assay. CEA-TCB or untargeted TCB were then added at concentrations ranging from 6.4 pM to 100 nM and PBMCs were added to obtain a final Effector : Target (E:T) ratio of 10:1 in a final volume of 1.1 ml per well. All experiments were performed in triplicate.

Target T-cell killing was assessed after 24 and 48 h of incubation by quantification of released lactate dehydrogenase (LDH) using an LDH detection kit (Roche Applied Science, Cat No.: 11 644 793 001) according to manufacturers’ instructions. Plates were read on a Spectramax ELISA reader and EC50 values were calculated.

Surviving tumor cells were detached Cell Dissociation Buffer (Gibco) and transferred into fresh 96-round-bottom well plates with the remaining PBMCs. FACS analyses were conducted on a BD Biosciences Fortessa system using fluorescently labeled antibodies specific for CD4 (BioLegend Cat No.: 300532; BD Biosciences, Cat No.: 552838), CD8 (BioLegend, Cat No.: 301014; BD Biosciences, Cat No.: 563256), PD-1 (BioLegend, Cat No.: 300532), CD8 (BioLegend Cat No.: 344704), CD25 (Biolegend Cat No.: 302614), CD69 (BioLegend Cat No.: 310934), PD-L1 (Biolegend Cat No.: 329706) and EpCAM (Miltenyi Biotec 130-091-254). Viable and dead cells were discriminated using Zombie Aqua Fixable Viability Kit (BioLegend Cat No.: 423102).

**Statistical Considerations**

Statistics are described in the legends.

Tumor growth inhibition values were calculated according to the equation:

\[
TGI = \frac{100 - \frac{Av(T_{\text{treatment} \mid \text{day}}) - T_{\text{treatment} \mid \text{baseline}})}{Av(T_{\text{vehicle} \mid \text{day}}) - T_{\text{vehicle} \mid \text{baseline}})} \times 100
\]

**TCB-Treatment Score**

Based on the Principal Component Analysis (PCA) of the ImmunoPD data, markers of PC1 (MIP-a, CXCL10, CXCL13) and PC2 (CD8+ T-cells and PD-1+ on CD8+ T-cells) were taken into account. The TCB treatment score summarizes the relative expression levels of MIPa, CXCL10 and CXCL13 as well as the levels of intra-tumor CD8+ T-cells and PD-1 expressing CD8+ T-cells. For a robust marker development, the estimation was based on the quintiles of the corresponding distributions of the markers in the present cohort. For every sample, depending on the relative expression of the particular marker, the marker got a discrete point ranging from 0 to 3: 0 for relative expression from the first quartile, up to 3 for the values from the last quartile of the corresponding distribution of marker values in the cohort. By applying this procedure, for each sample, an inhibitory receptor score in the range of 0 ≤ inhibitory receptor ≤ 15 by summing up the points for the five corresponding markers was obtained. Finally, each inhibitory receptor score was normalized by 15. To compare the TCB-treatment score, the Wilcoxon test using JMP12 (JMP, Version 12; SAS Institute Inc., 1989–2007) have been applied.
RESULTS

Tumor Targeting of CEA-TCB

Dual-isotope SPECT/CT imaging of hematopoietic stem cell humanized NSG mice (HSC-NSG) bearing a human gastric adenocarcinoma xenograft tumor (MKN-45, displaying high CEA expression) showed tumor targeting and uptake of 111In-CEA-TCB apparent at 4 h post-antibody injection, which continued to increase up to 120 h post single antibody injection (Supplementary Figure 1A). At the same time, the signal of 111In-CEA-TCB in the blood stream and other organs gradually decreased. There was negligible tumor uptake of the 177Lu-Untargeted-TCB injected simultaneously in the same mice. Quantitative analyses of the images revealed at least five-fold greater tumor uptake of 111In-CEA-TCB than 177Lu-Untargeted-TCB at all time-points. The amount of 111In-CEA-TCB and 177Lu-Untargeted-TCB in the spleen was identical, whereas the amount of 177Lu-Untargeted-TCB in the blood pool was greater than that of 111In-CEA-TCB (Supplementary Figure 1B). Together, imaging data indicated a tumor-specific targeting and accumulation of CEA-TCB over time along with clearance from the blood stream and other organs not expressing CEA.

CEA-TCB Treatment Reduces Tumor Growth and Generates a Highly Inflamed Tumor Microenvironment

Treatment of hematopoietic stem cell humanized NOG mice (HSC-NOG) bearing human MKN-45 tumor cells with CEA-TCB 2.5 mg/kg twice weekly reduced tumor growth by 62% compared with vehicle treated controls (Figure 1A). Flow cytometry analysis of tumors harvested after seven consecutive treatments showed that CEA-TCB treatment induced >10-fold increase in intra-tumor T-cells and >3-fold increase in the intra-tumor CD8/CD4 T-cell ratio (Figures 1B, C, G). Tumor-infiltrating T-cells demonstrated an activated phenotype as detected by increased expression of 4-1BB (CD137), an activation-induced T-cell costimulatory molecule (47), and PD-1, a hallmark of T-cell activation in this context (Figure 1D). Tumor-infiltrating T-cells also displayed a cytotoxic potential, exemplified by increased frequency of granzyme B (GZMB)-expressing cells, and proliferation, as evidenced by increased frequency of the Ki67 positive cells (Figure 1D). Treatment with CEA-TCB triggered secretion of pro-inflammatory cytokines (IFNγ, TNFα, IL-2) and several chemotactic molecules (CCL9, CCL10, CCL11, CCL13) indicating the generation of a highly inflamed tumor microenvironment (Figure 1F). CEA-TCB treatment also triggered upregulation of PD-L1 expression in tumors (Figures 1G, H). There were no changes related to T-cell activation or counts in the peripheral blood upon CEA-TCB treatment, further indicating that CEA-TCB activity is restricted to areas of CEA expression, such as in tumors (Figure 1E). The activity of CEA-TCB was additionally assessed in a CEA-expressing pancreatic tumor models (HPAF-II) in humanized NSG mice resulting in 72% of tumor growth inhibition, and confirmed the previous observations related to TCB mode of action consisting of strong increase of intra-tumor T-cells displaying an activated phenotype and increase of CD8/CD4 T-cell ratio (Supplementary Figure 2). The activity of CEA-TCB was further assessed in a genetically modified CEA424-SV40 TAg transgenic model, crossed with human CEACAM5 transgenic mice that spontaneously develop gastric tumors in the pyloric region (Steinhoff N et al., in preparation). CEA-TCB treatment led to a statistically significant reduction of CEA positive tumor area accompanied by a trend towards the increase of intra-tumor T cell infiltration and improvement survival (Supplemental Figures 3A–D).

The heatmap generated by combining cell surface and secreted T-cell activation markers (generated by flow cytometry and multiplex analysis from the MKN-45 experiment; Figures 1A–G) confirmed the clear separation of the CEA-TCB-treated animals and controls. CEA-TCB-treated tumors displayed a clear upregulation of several pro-inflammatory cytokines and chemokines [MIPα (CCL3), CCL10, CCL13, CCL9, IL-16, and I-TAC (CXCL11)], an increase in intra-tumor CD3+, CD4+, CD8+ T-cells that express high levels of 4-1BB, PD-1, and upregulation of GZMB (Figure 2A). The relative Principal Component Analysis (PCA) confirmed distinct clustering of CEA-TCB-treated mice as compared to controls (Figure 2B), and further revealed the presence of two sub-clusters within the CEA-TCB-treated mice: one associated with high infiltration of activated T-cells (expressing high levels of PD-1, 4-1BB, and GZMB) and high expression of pro-inflammatory chemokines and cytokines (particularly CCL10, CCL13, and MIPα); the other associated with high infiltration of activated T-cells (expressing high levels of PD-1, 4-1BB, and GZMB) but low expression of pro-inflammatory chemokines and cytokines (CCL10, CCL13, and MIPα). The correlation of these two inflamed tumor phenotypes with tumor volume or TCB activity with regards to tumor regression did not reveal a meaningful association (data not shown).

We further defined a CEA-TCB-treatment score (methods) with the aim to identify the parameters that most robustly characterize the CEA-TCB activity. The TCB-treatment score was generated based on the PCA of the ImmunoPD data considering top markers of Principal Component 1 (PC1) (MIP-a, CXCL10, CXCL13) and Principal Component 2 (PC2) (CD8+ T-cells and PD-1+ on CD8+ T-cells). CEA-TCB-treated tumors have a significantly higher CEA-TCB- treatment score (p=0.0044) compared to vehicle treated tumors (Supplementary Figure 4).

Identification of Gene Signature Associated With CEA-TCB Treatment

To more broadly characterize the molecular parameters associated with CEA-TCB treatment we performed bulk RNA sequencing of CEA-TCB-treated (seven consecutive treatments) and untreated tumors derived from MKN-45 tumor-bearing humanized mice (experiment from Figure 1; GEO accession number GSE155887). Similar to the ImmunoPD analysis described above, RNA sequencing analysis revealed a clear distinction between CEA-TCB-treated and control animals,
A

B

C

D

E

F

G

H

FIGURE 1 | Treatment with CEA-TCB induces tumor growth inhibition and leads to increased frequency of tumor-infiltrating human T-cells and a tumor-specific T-cell activation in MKN-45-bearing hematopoietic stem cell humanized mice. Hematopoietic stem cell humanized NOG mice were inoculated subcutaneously with 1 × 10⁶ MKN-45 cells and treated with either buffer (vehicle; n=12) or with 2.5 mg/kg i.v. of CEA-TCB (n=12) twice weekly starting with a tumor volume of ~150 mm³ (Day 8). At termination (Day 32), blood and tumors were harvested for subsequent flow cytometry, histological and cytokine analysis (ImmunoPD data). (A) Tumor growth kinetics revealed a tumor growth inhibition (TGI) of 62%. Arrows indicate treatments (seven in total). (B–E) Flow cytometry analysis of tumor and blood in vehicle- and CEA-TCB-treated animals showing the frequency of tumor-infiltrating T-cells (B) and ratio of CD8+ to CD4+ T-cells in the tumor tissue (C), the expression of activation markers in tumor (D) and blood (E). (F) Cytokine/chemokine expression in tumor lysates. (G) Representative histological staining for human CEA, CD3, and PD-L1 on paraformaldehyde fixed tumor samples from vehicle (upper row) and CEA-TCB-treated animals (lower row). (H) Quantification of PD-L1 staining by IHC. (A) Data are mean ± SEM; (B–F, H) solid bars represent mean values; p-values are two-tailed unpaired t-test; *p < 0.05; **p < 0.01; ***p < 0.001; ****p<0.0001.
with several genes being upregulated in tumors treated with CEA-TCB as compared to controls (Figure 3). The list of all genes that were found to have a significantly different expression (absolute log2 fold-change > 1 and an adjusted p-value < 0.05) between CEA-TCB-treated tumors and controls is provided in Supplementary Table 1.

Among the top 15 upregulated genes in CEA-TCB-treated tumors there were many reflective of a strong T-cell activation, migration, immune cell response (CXCL13, GNLY, GZMB, CXCL10, IDO1, SLA, CD2, TRBC2, TIGIT, IL2RB, CXCL9, CX3CL1, CCL4L2, and CD3E). Among the top downregulated ones we found keratin 6A and keratin 20 (KRT6A and KRT20) suggestive of the reduction of tumor cells as the result of TCB-mediated killing (Figure 3 and Supplementary Table 1). The Gene Set Enrichment Analysis (GSEA) further enabled identification of the main biological pathways upon CEA-TCB treatment and confirmed that the main Gene Ontology families that characterize the TCB response consist of T-cell activation (Response to interferon gamma; Adaptive Immune Response; T-cell activation; Inflammatory Response; Activation of Immune response; Cytokine secretion) and migration (Leukocyte Migration; Regulation of Cell Adhesion) (Figure 3 and Supplementary Figure 5). Interestingly, we also noticed upregulation of many major histocompatibility class II molecules that are known to be expressed on antigen presenting cells (HLA-DRA, HLA-DRB5, and HLA-DRB1) along with CX3CL1 (fractalkine, a known monocyte/T-cell attractant molecule) and CSF1 (colony stimulating factor 1, macrophage) suggestive of myeloid cell recruitment and activation at tumor sites post TCB treatment. We also observed the upregulation of PD-1 and PD-L1 transcripts following CEA-TCB treatment as compared to controls (Supplementary Figure 6).

**CEA-TCB Treatment Induces Upregulation of PD-1 and PD-L1 Expression; Combination of CEA-TCB With Anti-PD-L1 Blocking Antibody Enhances Its Efficacy in Stem Cell Humanized and Fully Immunocompetent Mice**

Data shown in Figures 1D, G, H and Supplementary Figure 6 provided evidence of PD-1 and PD-L1 upregulation on T-cells and tumors upon in vivo treatment with CEA-TCB. Additional evidence of the dose-dependent PD-1 upregulation on CD4 and CD8 T-cells as well as PD-L1 upregulation on tumor cells and T-cells upon CEA-TCB treatment was obtained from in vitro experiments (Figures 4A–F; Supplementary Figure 6). Incubation of the CEA-expressing MKN-45 target cells with human PBMC in presence of increasing concentrations of CEA-TCB led to the expected tumor cell lysis (Figure 4A). Flow cytometry analysis of co-cultured cells upon treatment revealed
dose-dependent upregulation of PD-L1 on tumor cells (Figure 4B) and dose-dependent upregulation of PD-1 (Figures 4C, D and Supplementary Figures 7A, B) and PD-L1 (Figures 4E, F and Supplementary Figures 7C, D) on CD4+ and CD8+ T-cells compared with cells incubated with untargeted TCB control. IFNγ is the main mediator of PD-L1 upregulation on tumor cells (Supplementary Figure 8A) and is released by CEA-TCB activated T cells in co-culture with tumor cells (Supplementary Figures 8B, C). Treatment of tumor cells with CEA-TCB in the absence of immune cells did not lead to PD-L1 upregulation on tumor cells, further confirming the key role of activated immune cells in secreting IFNγ (Supplementary Figures 8B, C). Further experiments corroborated the key role of IFNγ demonstrating that blocking of IFNγ by means of neutralizing antibodies reduced the PD-L1 upregulation on tumor cells resulting from CEA-TCB-mediated T cell activation (Supplementary Figures 8D, E).

The upregulation of PD-1 on T-cells and PD-L1 on tumor and T-cells following in vitro and in vivo TCB treatment led us to investigate whether combining CEA-TCB with PD-L1 blocking antibody could enhance the anti-tumor efficacy of CEA-TCB. We initially investigated the activity of this combination in HSC NOG mice bearing MKN-45 tumors. Treatment with CEA-TCB and anti-PD-L1 blocking antibody improved anti-tumor activity compared with either agent alone; in addition, the onset of tumor regrowth was significantly delayed in the combination group.
compared with anti-PD-L1 single-agent treatment (Figure 4G). Although the difference of the combination treatment compared with CEA-TCB monotherapy was only close to being significant, data support a trend for improved efficacy of the combination effect. Stronger anti-tumor activity was achieved when the combination started from the beginning (e.g. from the first treatment cycle) (Figure 4G) compared with combination that started after progression to CEA-TCB monotherapy treatment (Figure 4H).

The efficacy of the combination of CEA-TCB plus a PD-L1 blocking antibody was further assessed in a fully immunocompetent model consisting of human CEA transgenic (huCEA Tg) C57BL/6
mice bearing a syngeneic colorectal tumor line (MC38) stably expressing human CEA (MC38-hCEA). Treatment with murine surrogate of CEA-TCB (muCEA-TCB) and of anti-PD-L1 blocking antibody (muPD-L1) led to more rapid, more pronounced and sustained tumor growth inhibition as compared to the respective monotherapy treatment groups (Figure 5A).

Flow cytometry analysis of treated tumors, harvested 24 h after third infusion of the molecules, revealed that, similar to studies in humanized mice, CEA-TCB monotherapy increased the frequency of intra-tumor T-cells with preferential increase of CD8 T-cells over CD4 T-cells, leading to an increased ratio of intra-tumor CD8/CD4 and CD8/Tregs (Figure 5B). The majority of CD8 T-cells displayed a cytotoxic effector and effector memory phenotype and a significant fraction of CD8 T-cells (35%) displayed triple expression of PD-1+Tim3+Lag3+ (putatively exhausted cells). In addition, CEA-TCB treatment increased the frequency of intra-tumor CD4 and CD8 T-cells expressing CXCR3, a key receptor regulating T-cell chemotaxis (Figure 5B). Interestingly, combination treatment of CEA-TCB and anti-PD-L1 blocking antibody increased the intra-tumor frequency of both CD4 and CD8 T-cells compared with monotherapies and vehicle control.

This led to a similar frequency of cytotoxic effector and effector memory cells, but a lower frequency of triple positive, putatively exhausted PD-1+Tim3+Lag3+ CD8 T-cells (15% in combination vs 35% in CEA-TCB monotherapy) and higher frequency of intra-tumor CXCR3+ CD8+ T-cells (11% in combination vs 5% in CEA-TCB monotherapy) (Figure 5B). Together, the intra-tumor T-cell phenotype upon CEA-TCB and anti-PD-L1 combination treatment is reflective of T-cells having higher propensity for recruitment and putatively lower exhaustion status, and may reflect a higher recruitment of fresh T-cells from the peripheral blood.

**Combination With Anti-PD-L1 Also Enhances the Efficacy of CD20-TCB in Stem Cell Humanized Mice**

We further demonstrated the value of combining a TCB antibody with PD-L1 blockade for hematological malignancies using CD20-TCB (glofitamab), another “2:1” format TCB targeting CD20 on B cells and CD3 on T-cells (9, 48). Glofitamab is currently under clinical development in patients with relapsed or refractory B-cell non-Hodgkin lymphoma. To

![FIGURE 5](image-url) | Blockade of PD-1/PD-L1 axis improves the efficacy of CEA-TCB in immunocompetent mice. (A) Immunocompetent human CEA transgenic (huCEA Tg) C57BL/6J mice were inoculated subcutaneously with 0.5 × 10^6 MC-38-huCEA cells and treated with i.v. buffer (vehicle), murine surrogate of CEA-TCB (muCEA-TCB; 2.5 mg/kg i.v. once weekly), murine surrogate of anti-PD-L1 (10 mg/kg i.v. initial dose followed by 5 mg/kg i.p. twice weekly), or with a combination of muCEA-TCB and a-muPD-L1 (same dose and schedule as in monotherapy groups). Treatment started with a tumor volume of 200–400 mm^3 (Day 20). Arrows indicate treatments. Tumor growth kinetics are shown as mean ± SEM for all treatment groups (n=16 mice per group). Combination group vs muCEA-TCB: p=0.023 and vs vehicle: p<0.001; one-way ANOVA with Tukey’s multiple comparison correction done after five treatments (Day 38). (B) 24 h after the third treatment (Day 29), scout mice were sacrificed, tumor tissue was obtained and analyzed for T-cell infiltration and phenotype. p-values are one-way ANOVA with Tukey’s multiple comparison correction: ns, not significant; *p < 0.05; **p < 0.01; ***p < 0.001.
better evaluate the effect of the combination, HSC NSG mice bearing an aggressive human DLBCL cell line (WSU-DLCL2) were treated with a suboptimal dose of CD20-TCB (0.15 mg/kg). This resulted in suboptimal anti-tumor activity compared to its optimal dose (9), as monotherapy or in combination with an anti-PD-L1 blocking antibody. While monotherapy treatment with CD20-TCB or anti-PD-L1 blocking antibody did not show anti-tumor efficacy, the combination treatment led to tumor growth inhibition (Supplementary Figure 9).

**DISCUSSION**

The current study was undertaken to expand our understanding of cellular and molecular features associated with TCB activity and to address one of the key adaptive resistance mechanisms related to TCB activity, namely PD-1/PD-L1 axis upregulation, similarly to what has been described for checkpoint inhibitors (30).

The efficacy and mode of action of single-agent CEA-TCB was evaluated in different preclinical CEA-expressing mouse tumor models. These comprised hematopoietic stem cell humanized NOG mice bearing human gastric and pancreatic tumors and immunocompetent human CEA transgenic C57BL/6 mice (hCEA Tg mice) bearing a murine colorectal cancer tumor line (MC38) or crossed with genetically modified CEA424-SV40 TAg transgenic mice that spontaneously develop gastric tumors in the pyloric region. The former represent a hyper-mutated and highly inflamed form of colorectal cancers (MSIhi CRC) (49) transferred to stably express human CEA (MC38-hCEA), the latter an aggressive form of murine gastric cancer with immune desert phenotype, which is poorly responsive to cancer immunotherapy treatment (Steinhoff N et al., in preparation).

In all models, single-agent CEA-TCB slowed the growth of tumors compared with controls. Treatment of mice bearing CEA-positive tumors with CEA-TCB led to a 2 to >10-fold increase in tumor-infiltrating T-cells (depending on the tumor and mouse model). The tumor-infiltrating T-cells displayed a highly activated and proliferating phenotype, with tumors displaying a highly inflamed microenvironment as evidenced by increased levels of several pro-inflammatory cytokines and chemokines. Notably, anti-tumor efficacy along with tumor inflammation and increases of activated intra-tumoral T-cells was obtained in response to CEA-TCB treatment, even in settings with low pre-existing baseline tumor immune cell infiltration. This indicates that unlike other immunotherapies, CEA-TCB has the potential to be efficacious in patients with poor pre-existing inflammation. This is particularly relevant for patients with low frequency of pre-existing intra-tumoral CD8+ cells, who respond poorly to cancer immunotherapy (50) and particularly for the vast majority of human (CEA-expressing) colorectal cancer tumors with proficient mismatch repair (MMR) or with microsatellite stable (MSS) tumors who do not benefit from immunotherapy. Colorectal tumors with microsatellite instability (MSI) are typically more antigenic and have greater infiltration of CD8+ cells than MSI-negative tumors (51, 52).

A molecular signature of TCB treatment was identified consisting of pro-inflammatory cytokines/chemokines, and higher frequency and activation of T-cells. The signature appeared to be robust, as components of the signature were confirmed using complementary techniques: RNA expression analysis and protein expression as determined by flow cytometry and multiplex analysis. In particular, CXCL9 and CXCL10 were identified by both methods as the key molecules significantly upregulated by CEA-TCB treatment compared to controls. CXCL9 and CXCL10 are potent pro-inflammatory chemokines and chemotactants for multiple immune effector cells, including NK cells, monocytes/macrophages and T-cells by binding to the CXCR3 receptor expressed on the same cells (53, 54). In line with this, CEA-TCB treatment also increased the frequency of intra-tumor CXCR3+ CD8+ and CD4+ T-cells, corroborating the relevance of the CXCL10-CXCR3 axis in mediating the attraction of T-cells leading to increase of intra-tumor T-cell infiltration upon TCB treatment (25) (and unpublished data). It will be interesting to investigate the prognostic value of the TCB-treatment score in biopsies obtained from the ongoing interventional trial of the combination of CEA-TCB (cibisatamab) and atezolizumab in previously treated metastatic colorectal adenocarcinoma patients (NCT03866239).

A clear upregulation of both PD-1 (on CD4 and CD8 T-cells) as well as PD-L1 (on tumor cells and CD4 and CD8 T-cells) was detected in response to CEA-TCB treatment, indicative of the PD-1/PD-L1 axis being one of the adaptive resistance mechanisms related to TCB activity (8, 9, 25–29). Combination of TCBs with anti-PD-L1 blocking antibody (in different tumor and mouse models and using different TCBs targeting both CEA (solid tumors) and CD20 (hematological malignancies) consistently translated into superior anti-tumor efficacy and stronger tumor growth inhibition when compared to either agent given as monotherapy. Better tumor growth inhibition was obtained when the two agents were combined simultaneously from the first treatment cycle, as compared to starting the combination when tumors progressed to CEA-TCB monotherapy. This finding is consistent with previous in vitro data with a CEA-BiTE MEDI-565/AMG 211 that showed that T-cell killing was maximized when dual blockade of PD-1 and PD-L1 was applied early (26). Interestingly, the combination of CEA-TCB and an anti-PD-L1 blocking antibody led to increased frequency of intra-tumor CD4 and CD8 T-cells displaying a cytotoxic effector and effector memory phenotype; at the same time, the combination treatment lowered the frequency of putatively exhausted T-cells (characterized by co-expression of PD-1+Tim3+Lag3+ CD8 T-cells) and increased the frequency of T-cells having migratory capacity (characterized by CXCR3+ expression on CD8+ T-cells). Taken together, the intra-tumor T-cell phenotype upon CEA-TCB and anti-PD-L1 combination treatment is reflective of T-cells having higher propensity to migrate and putatively lower exhaustion status, and may indicate a stronger recruitment of fresh T-cells from the periphery.

These pre-clinical data support the rationale for the clinical investigation of CEA-TCB and atezolizumab, which is currently...
in Phase Ib (NCT03866239). Preliminary results of clinical activity indicated promising anti-tumor efficacy in patients with CEA+ solid tumors (mostly colorectal cancer) when cibisatamab was combined with the anti-PD-L1 antibody atezolizumab (55). Comparison of pre-treatment and on-treatment patient tumor biopsies (most of which came from MSS CRC patients with a non-T-cell inflamed immunophenotype prior to treatment) indicated that cibisatamab and atezolizumab combination treatment led to the increase of intra-tumor proliferating T-cells, increase of PD-1+ T-cells, upregulation of PD-L1 expression on immune cells, and reduction of CEA expressing tumor cells (56, 57), corroborating pre-clinical findings presented in the current study.

In conclusion, the data of the current study expand our knowledge of the cellular and molecular features associated with TCB activity, and provide evidence that the PD-1/PD-L1 axis is one of the adaptive resistance mechanisms associated with TCB activity. This adaptive resistance mechanism can be managed by the combination of TCB with anti-PD-L1 (or anti-PD-1) blocking antibodies translating into more efficacious anti-tumor activity and prolonged control of the tumor outgrowth. However, the data also show that tumors continue to progress despite the anti-PD-L1 combination treatment, suggesting that additional mechanisms, beyond the PD-1/PD-L1 axis, contribute to tumor escape. The elucidation of such mechanisms, most likely contributed to by both tumor cells and different immune cell subsets, by using high dimensional single cell approaches for tumor analysis, will constitute an important milestone in our understanding of additional resistance mechanisms to immunotherapy and novel combination approaches for efficient tackling of the same.

DATA AVAILABILITY STATEMENT

The original contributions presented in the study are publicly available. This data can be found here: https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE155887.

ETHICS STATEMENT

This study involving laboratory animals was reviewed and approved by the Institutional Animal Care and Use Committee of the Preclinical Pharmacology Department, Roche Innovation Center Zurich, Schlieren, Switzerland. This study was performed in accordance with the animal research protocols approved by the local government (Kantonale Verwaltung Veterinäramt kant. Zürich, Switzerland; license ZH193/2014). All animals were handled in accordance with the guidelines of the Federation of European Laboratory Animal Science Associations, Gesellschaft für Versuchstierkunde/Society of Laboratory Animal Science (GV-Solas) and the Tierschutzgesetz (TierSchG).

AUTHOR CONTRIBUTIONS

TF, MBi, and LF were involved in the design and generation of in vitro data. JS and SC designed, supervised, and interpreted the in vivo and ex vivo studies. MP designed and supervised in vivo studies in immunocompetent mice. EB, AS, MK, ML, and NS conducted all in vivo and ex vivo experiments. VN generated all histology results. TN designed and supervised the SPECT/CT imaging study. AR supervised and analyzed the bulk RNAseq data. MBa, CK, and PU supervised the project. MBa and JS wrote the manuscript. MBa contributed to experimental design and data interpretation of all in vitro and in vivo studies. All authors reviewed the manuscript. All authors contributed to the article and approved the submitted version.

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fonc.2020.575737/full#supplementary-material

REFERENCES

1. Ribas A, Dummer R, Puzanov I, VanderWalde A, Andtbacka RHI, Michielin O, et al. Oncolytic virotherapy promotes intratumoral T cell infiltration and improves anti-PD-1 immunotherapy. Cell (2017) 170:1109–19.e10. doi: 10.1016/j.cell.2017.08.027
2. Mullard A. Top product sales forecasts for 2018. Nat Rev Drug Discovery (2018) 17:86. doi: 10.1038/s41596-018-0015
3. Emens LA, Ascierto PA, Darcy PK, Demaria S, Eggermont AMM, Redmond WL, et al. Cancer immunotherapy: Opportunities and challenges in the rapidly evolving clinical landscape. Eur J Cancer (2017) 81:116–29. doi: 10.1016/j.ejca.2017.01.035
4. Ribas A, Wolchok JD. Cancer immunotherapy using checkpoint blockade. Science (2018) 359:1350–5. doi: 10.1126/science.aar4060
5. Hegde PS, Chen DS. Top 10 challenges in cancer immunotherapy. Immunity (2020) 52:17–35. doi: 10.1016/j.immuni.2019.12.011
6. Goebeler M-E, Bargou RC. T cell-engaging therapies - BiTEs and beyond. Nat Rev Clin Oncol (2020) 17(4):481–34. doi: 10.1038/s41571-020-0347-5
7. Kaiser J. Forced into battle. Science (2020) 368:930–3. doi: 10.1126/science.368.6494.930
shelf therapy. Sci Transl Med (2019) 11:eaav5989. doi: 10.1126/scitranslmed.aav5989

49. Efremova M, Rieder D, Klepsch V, Charoentong P, Finotello F, Hackl H, et al. Targeting immune checkpoints potentiates immunoediting and changes the dynamics of tumor evolution. Nat Commun (2018) 9:32. doi: 10.1038/s41467-017-02424-0

50. Tumeh PC, Harview CL, Yearley JH, Shintaku IP, Taylor EJM, Robert L, et al. PD-1 blockade induces responses by inhibiting adaptive immune resistance. Nature (2014) 515:568–71. doi: 10.1038/nature13954

51. Le DT, Uram JN, Wang H, Bartlett BR, Kemberling H, Eyring AD, et al. PD-1 blockade in tumors with mismatch-repair deficiency. N Engl J Med (2015) 372:2509–20. doi: 10.1056/NEJMoa1500596

52. Vilar E, Gruber SB. Microsatellite instability in colorectal cancer-the stable evidence. Nat Rev Clin Oncol (2010) 7:153–62. doi: 10.1038/nrclinonc.2009.237

53. Dufour JH, Dziejman M, Liu MT, Leung JH, Lane TE, Luster AD. IFN-gamma-inducible protein 10 (IP-10; CXCL10)-deficient mice reveal a role for IP-10 in effector T cell generation and trafficking. J Immunol (2002) 168:3195–204. doi: 10.4049/jimmunol.168.7.3195

54. Tokunaga R, Zhang W, Naseem M, Puccini A, Berger MD, Soni S, et al. CXCL9, CXCL10, CXCL11/CXCR3 axis for immune activation - A target for novel cancer therapy. Cancer Treat Rev (2018) 63:40–7. doi: 10.1016/j.ctrv.2017.11.007

55. Tabernero J, Melero I, Ros W, Argiles G, Marabelli A, Rodriguez-Ruiz ME, et al. Phase Ia and Ib studies of the novel carcinoembryonic antigen (CEA) T-cell bispecific antibody (CEA CD3 TCB) antibody as a single agent and in combination with atezolizumab: Preliminary efficacy and safety in patients with metastatic colorectal cancer (mCRC). JCO (2017) 35:3002–2. doi: 10.1200/JCO.2017.35.15_suppl.3002

56. Melero I, Segal NH, Saro Suarez JM, Ros W, Martinez Garcia M, Calvo E, et al. Pharmacokinetics (PK) and pharmacodynamics (PD) of a novel carcinoembryonic antigen (CEA) T-cell bispecific antibody (CEA CD3 TCB) for the treatment of CEA-expressing solid tumors. JCO (2017) 35:2549–9. doi: 10.1200/JCO.2017.35.15_suppl.2549

57. Melero I, Segal NH, Saro J, Ros W, Martinez-Garcia M, Argiles G, et al. Pharmacokinetics (PK) and pharmacodynamics (PD) of a novel carcinoembryonic antigen (CEA) T-cell bispecific antibody (CEA-CD3 TCB) for the treatment of CEA-positive solid tumors. Ann Oncol (2017) 28: v28. doi: 10.1093/annonc/mdx363.020

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