Immunotherapy using IgE or CAR T cells for cancers expressing the tumor antigen SLC3A2


ABSTRACT

Background  Cancer immunotherapy with monoclonal antibodies and chimeric antigen receptor (CAR) T cell therapies can benefit from selection of new targets with high levels of tumor specificity and from early assessments of efficacy and safety to derisk potential therapies.

Methods  Employing mass spectrometry, bioinformatics, immuno-mass spectrometry and CRISPR/Cas9 we identified the target of the tumor-specific SF-25 antibody. We engineered IgE and CAR T cell immunotherapies derived from the SF-25 clone and evaluated potential for cancer therapy.

Results  We identified the target of the SF-25 clone as the tumor-associated antigen SLC3A2, a cell surface protein with key roles in cancer metabolism. We generated IgE monoclonal antibody, and CAR T cell immunotherapies each recognizing SLC3A2. In concordance with preclinical and, more recently, clinical findings with the first-in-class IgE antibody MOv18 (recognizing the tumor-associated Folate Receptor alpha), SF-25 IgE potentiated Fc-mediated effector functions against cancer cells in vitro and restricted human tumor xenograft growth in mice engrafted with human effector cells. The antibody did not trigger basophil activation in cancer patient blood ex vivo, suggesting failure to induce type I hypersensitivity, and supporting safe therapeutic administration. SLC3A2-specific CAR T cells demonstrated cytotoxicity against tumor cells, stimulated interferon-γ and interleukin-2 production in vitro. In vivo SLC3A2-specific CAR T cells significantly increased overall survival and reduced growth of subcutaneous PC3-LN3-luciferase xenografts. No weight loss, manifestations of cytokine release syndrome or graft-versus-host disease, were detected.

Conclusions  These findings identify efficacious and potentially safe tumor-targeting of SLC3A2 with novel immune-activating antibody and genetically modified cell therapies.

BACKGROUND

Recent years have seen the successful translation of immunotherapy strategies to the clinic with monoclonal antibodies being at the forefront of efficacious treatments. Similarly, the development of genetically modified cell therapy approaches utilizing synthetic chimeric antigen receptor (CAR) T cell technologies, has resulted in three cluster of differentiation-19 (CD19) and one B-cell maturation antigen targeting product receiving FDA approval for hematological cancers. Suitable tumor associated antigens (TAAs) for antibody and CAR T cell therapies require cell surface expression on cancer cells, and low/restricted distribution in normal tissues. These requirements critically limit the choice of TAAs suitable for monoclonal antibody and CAR T cell therapy development.

In 1988, Wilson et al. investigated the antigenic changes correlated with malignant transformation of hepatocytes: their research was focused on the detection of common antigens among tissues derived from the same germ layer that could be associated with transformed cell phenotypes. Mice were immunized with the hepatocellular carcinoma cell line FOCUS and hybridomas were developed. Subsequent screening identified 18 antibodies reactive against human colon carcinoma cell lines. The murine IgG, SF-25 antibody clone was chosen for further investigation. SF-25 bound 17 out of 17 human colon adenocarcinoma biopsies, while demonstrating no staining of normal adjacent mucosa. In vivo localization to subcutaneous tumors in nude mice was demonstrated. These data confirmed colon-cancer specificity for SF-25.
develop the SF-25 clone were undertaken, including engineering of a human/murine IgG1, chimeric antibody, development as a positron emission tomography imaging tracer, and as an antibody drug conjugate (ADC). The chimeric SF-25 antibody has also been used to improve the targeting and effector function of adoptively transferred lymphokine activated killer cells to cancer cells in a liver metastasis model of colorectal cancer. These data highlighted the potential of the SF-25 clone for cancer specific drug discovery. However, the target antigen of SF-25 remained elusive, hampering further clinical development.

Monoclonal antibodies represent a well-established platform to combat cancer. Until recently only those of the IgG class have been employed for cancer immunotherapy. Since different antibody classes function through unique Fc-receptors and induce specific immune responses at different anatomical sites, the design of new therapeutics could exploit antibody isotypes other than IgG. IgE antibodies, well known for their pathogenic roles in allergic disease, may offer multiple advantages over those conferred by IgG in treating solid malignancies. These are based on known aspects of IgE biology, commonly employed in immune protection from parasites and in allergic responses, which may translate to superior efficacy in targeting tumors. Such attributes of the IgE class include: (1) high affinity for IgE Fcepsilon receptors (FceRs) (2–5 orders of magnitude greater than that of IgGs for FcγRs), (2) expression of FceRs on a distinct spectrum of tissue-resident and tumor-resident immune effector cells, (3) long tissue residency and retention of IgEs by immune effector cells, (4) lack of an inhibitory Fc receptor and (4) active immune surveillance in Th2-biased environments in tissues such as the skin and gut. The potential efficacy of antitumor IgEs recognizing cancer antigens has been demonstrated in several in vivo and in vitro models. First in class human data for IgE utility in ovarian cancer (NCT02546921) has reported interim promising safety and biological activity.

Translating the success of CD19 targeting CAR T cells into solid tumors is hampered by numerous challenges. Namely, homing to tumor sites, persistence of adoptively transferred cells, development of exhausted phenotypes on adoptive transfer, and on-target off-tumor killing in healthy tissues. The apparent tumor specificity of the SF-25 antibody raised the potential that this antibody, and its elusive target, could be a promising axis for the development of novel strategies for IgE and cell-based immunotherapy.

Here, we undertook to identify the target of SF-25 on human cells by utilizing a bioinformatics and mass spectrometry pipeline. We then designed IgE and CAR T cell approaches to demonstrate the potential for broad cancer immunotherapy development.

### METHODS

#### Cloning and production of recombinant antibodies

To clone the SF-25 variable region into a human IgG1 backbone we performed a four-fragment Polymerase Incomplete Primer Extension (PIPE) PCR: two big fragments F2 and F4 (3000–4000bp) containing at 5' the constant regions of the light and heavy chain, respectively; and two small fragments F1 and F3 (300–400bp) composed by SF-25 Vk and VH, respectively. The PCR reactions contained 0.5μM of each primer, 25μL of PhusionTM Flash High-Fidelity PCR Master Mix, 10ng of template DNA and sterile water up to 50μL. The cycles for amplification were: 10s at 98°C, 35 cycles of 1s at 98°C, 5s at 62°C, and 10s at 72°C. PCR products were treated with DpnI, bacteria were transformed with equal amount of the digested PCR products to combine the four fragments and plated in Luria Bertani (LB) agar plates supplemented with 200μg/mL hygromycin B. Colonies were amplified overnight in LB supplemented with 200μg/mL hygromycin B and DNA was extracted with a QiAprep spin miniprep kit (Qiagen). Correct assembly of the plasmid was verified by sending the newly generated and extracted plasmid for sequencing (Source Bioscience). The sequencing output was analyzed using FinchTV.

To obtain the pVitro1-SF-25 IgE expression vector, we designed the primers for a three fragment PIPE cloning protocol, amplifying the fragment containing the epsilon heavy chain constant region (PCR1) from an in-house representative pVitro1-IgE vector and the other two fragments (PCR2 and PCR3) from our pVitro1-SF25-IgG1. The cycles for PCR1 and 2 were: 10s at 98°C, 35 cycles of 1s at 98°C, 5s at 62°C, and 10s at 72°C. PCR3 was performed with 10s at 98°C, 35 cycles of 1s at 98°C, 5s at 62°C, and 8s at 72°C. PCR products were separated on 1% agarose gels to discriminate the multiple PCR products by molecular weight. DNA was purified from the gel using the PureLinkTM Quick Gel Extraction Kit (ThermoFisher). PCR products were treated with DpnI to digest the template DNA. One-Shot TOP10 bacteria were transformed with equal amount of the digested PCR products to combine the four fragments (F1-F2-F3-F4) and generate the pVitro1-SF-25-IgG1 vector. The correct assembly of the plasmid was verified via sequencing and analyzed using FinchTV. Antibodies were produced in Sp2/0 (IgE) and Expi293F (IgG1) mammalian cells and purified using previously described methods (see online supplementary materials and methods).

#### SF-25 antigen expression screening

Cell lines were detached with Trypsin-EDTA treatment, counted and 2.5×10^6 cells were used per tube. 2mL FACS buffer [Phosphate Buffer Saline (PBS); 5% Foetal Bovine Serum (FBS); 3mM Ethylenediaminetetraacetic acid (EDTA)] was added to each tube before a 5min centrifugation at 400rcf at 4°C. Cells were resuspended in 100μL FACS buffer and incubated for 20min at 4°C with a range of concentrations from 0 to 50ng SF-25 IgG1, or 0 to 5μg...
Figure 1  Engineering and testing of SF-25 antibodies with human IgG1 or IgE Fc regions. (A) Schematic depicting the steps to obtain the variable regions of the SF-25 antibody and representative images for the visualization of the PCR products on agarose gels, followed by final sequence constructs of pVitro1 expression vectors to produce SF-25 IgE or IgG1 full length chimeric antibodies and the HPLC elution profile of SF-25 IgE and IgG1. (B) Lectin blot data reporting the glycosylation profile of SF-25 IgE pVITRO and SF-25 IgE Sp2/0 for fucose, galactose, mannose and sialic acid (detected, respectively, with AAL-biotin (Aleuria aurantia lectin), RCAI-biotin (Ricinus communis agglutinin I lectin), Con-A-biotin (concanavalin a lectin) or EBL (Sambucus nigra lectin)). (C) Flow cytometric data depicting SF-25 IgG1 (left panel) and IgE (right panel) binding to different cell lines. Values are reported as mean fluorescence intensities (MFI) detected with antihuman IgG or IgE (respectively) FITC-conjugated secondary antibodies. Representative binding profiles for cell lines targeted in the present study (top panels) (see also online supplemental figure S1). Mouse image used courtesy of MCKIBILLO. HPLC, High performance liquid chromatography.
SF-25 IgE. Cells were washed with 2 mL FACS Buffer and were then incubated with goat antihuman IgG-FITC (Fluorescein isothiocyanate) or antihuman IgE-FITC for 20 min at 4°C. Cells were washed with 2 mL FACS buffer and resuspended in 400 µL FACS buffer for cytometry analysis. Analysis of flow cytometry data was performed by FlowJo (TreeStar Inc) software.

**SF-25 antigen: bioinformatics analysis**
SF-25 antibody binding scores were generated from three experimental datasets, two binding datasets generated for this study by Flow Cytometry and one previously published radioligand binding dataset. Transcriptome RNAseq datasets E-MTAB-2706; E-MTAB-2770 and E-MTAB-3983, including several of the human cancer cell lines present in the binding datasets, were downloaded and filtered to only keep the cell lines for which binding data were available. Further filtering and sorting steps were performed to generate matching tables between one binding dataset and the transcription level files. The matched tables were then analyzed, and Spearman correlation scores were calculated for each individual gene in each binding experiment. Average Spearman scores and their variances were generated across the three different combinations made. Further details on the bioinformatic process design and coding are reported in online supplemenal materials and methods.

**Cytotoxicity/phagocytosis (antibody dependent cell-mediated cytotoxicity/ADCP) assay and Basophil activation test**
Antibody dependent cell-mediated cytotoxicity/phagocytosis (ADCC)/ADCP assays were performed according to a previously described method. The ability of SF-25 IgE to trigger primary human basophil activation was determined using an ex vivo assay in which CD63 expression on the surface of human basophils was used as an early marker of basophil activation, as described previously (see online supplemental materials and methods).

**Cytotoxicity assays with CAR T cells**
After 24 hours coculture, the viability of tumor cell monolayers was quantified. T cells were removed from the wells and MTT (3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide, Sigma) was added at 500 µg/mL in 200 µL complete DMEM medium and incubated for 1 hour at 37°C and 5% CO₂. After removal of the supernatant, formazan crystals were resuspended in 200 µL DMSO. Absorbance was measured at 560 nm using a spectrophotometric plate reader (FluoSTAR Omega) and tumor cell viability percentage was calculated as follows: (absorbance of coculture/absorbance of monolayer alone) x 100. Further details of CAR T in vitro and in vivo assays in online supplemental materials and methods.

**RESULTS**

**Engineered SF-25 antibodies with human Fc regions recognize human malignant cell lines**
We cloned the murine variable region sequences of SF-25 into human IgG and IgE antibody scaffolds (figure 1A) using previously established platforms. High performance liquid chromatography (HPLC) analyses demonstrated high antibody purity and negligible aggregation (<5%) (figure 1B). Production of high-purity intact antibody was demonstrated in different expression systems, culture media, serum content and culture vessel conditions (online supplemental figure S1A, B). Serum-free, serum depleted, or specialist serum-free (ADCF, animal-derived component-free) media all led to high purity antibody with negligible protein aggregates or non-assembled light chain (online supplemental figure S1C,D). Intact antibody can be generated at small and large scale in serum-free conditions demonstrating utility for preclinical process development and clinical testing.

IgE class antibodies are highly glycosylated (12% of molecular weight). We evaluated IgE glycosylation in two SF-25 IgEs produced using mouse (Sp2/0) and human (Expi293F) expression systems. Lectin blot analyses were performed for the most common sugars known to decorate antibodies expressed in mammalian systems: mannose, fucose, galactose and sialic acid. No significant difference in fucose and mannose was observed between murine and human expression systems. Sp2/0-derived SF-25 IgE contained 2.5 times higher galactose and sialic acid content compared with IgE engineered in human Expi293F (figure 1B). We confirmed the reactivity of each SF-25 chimeric SF-25 IgG, and IgE against a panel of human and non-human tumor cell lines. We showed largely comparable reactivity between IgG and IgE and variable binding across human tumor cell lines, likely reflective of target expression levels. Overall, 18 out of 19 human cancer cell lines from 7 different origins tested were recognized by SF-25 antibodies. Antigen density varied, but with clear population shifts demonstrated for the cell lines tested (figure 1C). No or low binding were detected to the non-human rat colon carcinoma cell line CC531.

Together, these findings confirm the generation of intact, monomeric human Fc IgG and IgE SF-25, and production of functional IgE in serum free-culture mammalian expression systems suitable for future translation. We confirmed binding of the antibodies to human tumor cells of different malignant origins in agreement with reports of the original SF-25 mouse clone.

**Identification of the target of the SF-25 antibody clone with engineered antibodies**
Three complementary approaches were employed to identify the target of SF-25. We performed immuno-precipitations with the chimeric SF-25 IgG1 on cell lysates from three different human cancer cell lines with differing SF-25 target expression levels (figure 1C): MDA-MB-231, MDA-MB-468 (breast cancer, BRCA) and
A2058 (melanoma) (figure 2A-left panel). No specific protein band emerged by comparing these three samples based on the predicted target abundance. However, bands at 27 and 42 kDa were visible across the three cell lines. They were further analyzed by trypsin digestion and mass spectrometry. Peptides belonging to 138 different proteins were identified (online supplemental table S1). A transcriptomic analysis was performed to compare binding levels on panels of human cancer cell lines against transcript levels for each gene in the same cell lines (figure 2A-right panel). Three sets of binding scores were generated reflecting the binding intensity measured for the SF-25 antibody on each cell line, counts per minute for previous radioligand binding data, and mean fluorescent intensity (MFI) for the experiments in figure 1C. The relative binding scores for the three binding panels are shown in online supplemental table S2. The binding intensities observed across different cell lines in one binding experiment were compared with transcript levels in aggregate RNA-seq data for the same cell lines. A Spearman correlation score was calculated between SF-25 binding and transcript levels for each gene and average Spearman scores and variance values were calculated from the three comparative studies. The heavy chain of CD98 (CD98hc) encoded by the solute carrier family 3 member 2 (SLC3A2) demonstrated the highest Spearman score (figure 2A-bottom panel). The gene SLC7A5, coding a binding partner of CD98hc, was the second highest hit. Both partners were identified among the candidates immunoprecipitated (figure 2A-bottom panel). SLC3A2 was also identified among 138 different proteins after immunoprecipitation, offering a weak biochemical validation. We, therefore, identified SLC3A2 as the top candidate.

In separate immuno-mass spectrometry experiments, SF-25 IgG reactivity was tested against human cutaneous and tumor antigens. This immuno-mass spectrometry method can be applied to discover proteome-wide targets of antibodies by using complex protein mixtures from human tissues as sources of candidate proteins. Tissue lysates were generated from three pooled human skin samples, two human cutaneous metastatic melanoma lesions, and the human ovarian carcinoma cell line IGROV1. In two independent experiments, applying filtering for contaminants and non-specific binding using a non-binding (hapten-specific) antibody (figure 2B), 10 and 7 peptides corresponding to SLC3A2 were identified from the three skin samples. Separately, 18 peptides from two human melanoma lesion samples and 16 peptides from the IGROV1 cell line identified recognition of SLC3A2 by the antibody (figure 2B). The average peak area of binding to melanoma cells and normal skin across two independent experiments was significantly higher in malignant cells for SLC3A2, demonstrating highest expression of the target in tumor samples (figure 2C). In concordance, the identified SLC3A2 peptides (online supplemental figure S2) were demonstrated at significantly higher levels in melanoma vs normal skin (p<0.05, data from three normal skin and two melanoma samples tested twice each in two independent experiments) (figure 2C). Finally, using single guide CRISPR Cas9 knock out of the SLC3A2 gene in A2058 cells, comparable loss of binding was seen with SF-25 IgG and a commercially available CD98hc specific monoclonal antibody (figure 2D).

These data identify the protein CD98hc, coded for by SLC3A2, as the target of the SF-25 antibody clone both bioinformatically and biochemically.

The SLC3A2 derived protein CD98hc is a tumor-associated-antigen expressed on a broad range of human tumors with limited normal tissue expression

It was previously reported that the murine SF-25 clone demonstrated normal human tissue binding against a subpopulation of cells in the distal tubules of the kidney. Using RNA-sequencing raw data from The Cancer Genome Atlas (TCGA) and the Genome Tissue Expression (GTex) we examined the expression of SLC3A2 in malignant vs normal tissues across a range of human cancers (figure 3A and online supplemental figure S3). Tumors were selected for analysis when adequate equivalent normal tissue data were available. For the majority of tumors assessed, significantly higher levels of SLC3A2 expression were observed in malignant vs normal tissues, most notably in colorectal, breast, genitourinary cancers and cutaneous melanomas (figure 3A). When considering primary vs metastatic tumors, analysis was hampered by the paucity of metastatic tumors covered by TCGA and GTex. Data were analyzed for melanoma (SKCM) and breast cancer (BRCA) where notably only melanoma had a large resource of metastatic disease available (figure 3B). These data showed that SLC3A2 overexpression was maintained in metastatic deposits of melanomas (figure 3B). True TAAs, limited only to tumor tissue, are very hard to come by in solid tumor oncology. Instead, we develop targeted immune therapy directed against antigens where expression is largely higher in malignant vs normal tissue. To explore the levels of SLC3A2 in relation to established TAAs, we compared SLC3A2 expression levels with those recorded for HER-1 and HER-2 in normal skin and breast tissues. HER-1 and HER-2 were selected as key examples of antigens targeted safely and successfully in the oncology clinic with immune therapy strategies including antibody and cell therapy approaches. SLC3A2 expression, in these normal tissues, is similar or lower than HER-1 and HER-2 expression levels (figure 3C).

We next evaluated the reactivity of the antibody clone against human malignant and non-malignant tissues by immunohistochemistry. Directly labeled SF-25 IgG, AF488 showed reactivity against melanoma but not against normal skin samples (figure 4A). Tissue microarray immunofluorescence analyses showed no/restricted reactivity in most normal tissues. Consistent
Figure 2  The SF-25 antibody clone recognizes the human CD98 heavy chain (CD98hc) coded for by SLC3A2. (A) Immunoprecipitation-mass spectrometry (IP-MS) evaluations identified candidate proteins (left panel). In parallel, transcriptomic and bioinformatics studies identified candidate proteins (right panel). Analyses were matched to identify the target of the SF-25 antibody (bottom panel). (B) Immuno-MS was performed on lysates from human skin (n=3) and human melanoma (n=2) specimens. In two independent experiments, recognition of more than one antigenic peptide from one antigen confirmed antigen reactivity. Binding results were subjected to sequential filtering criteria: first, Ig heavy and light chain contaminants were removed; then peptides that were identified in negative and positive antibody controls were excluded; peptides bound in only one of the two MS injections were removed; and finally, if the peak area of the MS detection peptide profile was zero in both tests. Proteins were ranked based on the average peak area of antibody-bound peptides identified across two independent experiments of melanoma specimens (bottom panel). (C) The average peak area and the cumulative number of SLC3A2 peptides identified via SF-25 IgG, binding to two melanoma and three normal skin specimens by immuno-MS each tested in two injections and across two independent experiments (*p<0.05). (D) The target identity of SF-25 antibody clone was confirmed on SLC3A2 CRISPR KO A2058 cells: the reactivity of the SF-25 clone and a commercially available (MEM-108) to cancer cells was impaired in transduced tumor cells (see also online supplemental figure S2).
with previous data, we confirmed kidney binding and revealed lower-level antibody bindings in testis, and at low levels in human cerebellum (figure 4B). In contrast, robust staining was observed in human malignant tissues, including melanoma, breast, ovarian, testis and soft tissue cancers (figure 4C). Our transcriptomic and immunohistochemical analyses (figures 3 and 4) are consistent with previous reports of SF-25 antibody reactivity against several cancer types versus normal tissues.

Figure 3  SLC3A2 gene expression is enhanced in different malignancies compared with equivalent normal tissues. (A) Expression of SLC3A2 in normal (N) versus tumor (T) tissues of different origins (separated by tissue type; colorectal and colon adenocarcinomas, breast, skin and genitourinary tissues). Tumor types are described in online supplemental materials and methods—SLC3A2 differential expression study. (B) SLC3A2 gene expression primary tumor (T) and tumor metastases (M) compared with normal (N) tissues in SKCM (melanoma) and BRCA (breast cancer). (C) Expression of SLC3A2 alongside two other tumor associated antigens (EGFR and HER2) in normal breast and skin samples. Mann-Whitney U t-test was performed. ns=non-significant; *p<0.05; **p<0.01; ***p<0.001; ****p<0.0001. See also online supplemental figure S3 and online supplemental materials and methods.
To investigate reports of SLC3A2 expression in human peripheral blood mononuclear cells (PBMCs) we stained PMBCs with the MEM-108 anti-CD98hc monoclonal antibody, demonstrating binding that was significantly increased by activation of PBMCs with Phytohemagglutinin (online supplemental figure S4).

The protein derivative of SLC3A2 is an established heterodimerization partner for multiple solute transporters and plays a role in the transport of amino acids and glucose. Its expression in PBMCs suggests a potential role in immune cell function.

**Figure 4** Immunofluorescence evaluations confirm SF-25 antibody reactivity with several tumor tissues versus low/restricted reactivity with normal tissues. (A) melanoma (left) and normal skin (right) frozen tissue sections were stained with SF-25 IgG1-AF488 antibody (green) or the negative isotype control NIP IgG1-AF488 to detect antigen expression. 4',6-diamidino-2-phenylindole (DAPI) (blue) was used to reveal the cell nuclei. (B, C) A normal tissue (B) and a cancer specimen tumor microarray (TMA) (C) was stained with SF-25 IgG1-AF488 antibody (green) to detect clone reactivity to tumor versus normal tissues. DAPI (blue) was used to reveal the cell nuclei.
Figure 5  SF-25 IgE can trigger Fc-mediated effector functions. (A) SF-25 IgE did not impair the formation of human ovarian IGROV1, melanoma A2058 or pancreatic PaTu8988t cancer cell colonies. An antimalanoma (CSPG4) IgE was used as negative control with IGROV1 and PaTu8988t cells and as a positive control with A2058 melanoma cells. Antifolate receptor alpha (FRα) antibody MOv18 IgE had no effect on FRα-expressing IGROV1 ovarian cancer or on FRα-negative A2058 melanoma cells. Data represent average ±SD of n=2 independent experiments. (B, C) SF-25 IgE crosslinked on immune effector cells by target antigen-expressing cancer cells triggered effector functions: (B) degranulation of rat basophilic leukemia (RBL-SX38) cells expressing human FcεRI measured by β-hexosaminidase release: SF-25 IgE was cross-linked with SF-25 antigen-expressing melanoma A2058 (n=5), breast cancer SKBR-3 (n=4), ovarian cancer IGROV1 (n=2) or colorectal cancer LS-180 cells (n=2). CSPG4 and MOv18 IgE antibodies were used as controls; Triton treatment represented total β-hexosaminidase release controls (average ±SD of independent experiments). Welch’s test was performed. **p<0.01; ***p<0.001. (C) Healthy volunteer (left) and melanoma patient (right) peripheral blood mononuclear cells (PBMCs) activated by SF-25 IgE to mediate ADCC of A2058 melanoma cells. MOv18 served as negative control (% tumor cell killing ±SEM of n=4 independent experiments). (D) the propensity of SF-25 IgE to mediate activation on human basophils ex vivo was evaluated using the basophil activation test (BAT), in unfractionated peripheral blood of cancer patients, by measuring upregulation of CD63. While IgE:FcεRI- (anti-FcεRI, anti-IgE) and non-IgE (fMLP) mediated basophil activation (fold change in the percentage of CD63-positive cells) in a cohort of n=23 ovarian cancer patients, SF-25 IgE stimulation led to NO basophil activation (NO increase of CD63-positive cells) above background set as threefold change (D, inset). Each value corresponds to a patient sample (n=23 ovarian cancer patients). Mann-Whitney U test was performed. ns=non-significant; **p<0.01; ***p<0.001; ****p<0.0001 (see also online supplemental figure S6).
carriers. SLC3A2/CD98hc and two L-type amino acid transport binding partners, LAT1 and ascAT1 (derived from SLC7A5 and SLC7A10, respectively), were analyzed independently as prognostic markers in bladder, breast, cervical, lung, renal and head and neck cancers. Survival over 10 years from diagnosis was assessed by the Kaplan-Meier method for high and low expressing tumors (online supplemental figure S5). Diversity in prognostic value of SLC3A2 and two of its many binding partners highlights the breadth of impact potentially associated with targeting this key heterodimerization TAA.

These findings, at the transcriptomic and cell surface proteomic levels, support SLC3A2 as a TAA and confirm reactivity of the human Fc engineered antibody to human tumor tissues of different origins.

**SF-25 IgE activates immune cells through the FcεRI and demonstrates tumor cell cytotoxicity in vitro**

Since IgE antibodies may offer an alternative immunotherapy approach for solid tumors, we investigated the antitumor functions of the engineered SF-25 IgE antibody in vitro.

To determine direct effects of SF-25 IgE on target cancer cells, colony formation assays with A2058, IGROV1 and PaTu8988t cells were performed. SF-25 IgE did not impair the ability of cancer cells to form colonies (figure 5A).

We next investigated whether SF-25 IgE could trigger Fc-mediated effector functions through the high affinity Fc receptor FcεRI. Mast cells and basophils are known to express FcεRI and to participate in parasite clearance through IgE.19 The rat basophilic leukemia RBL SX-38 in vitro mast cell model expressing human FcεRI, was used to examine the Fc-mediated biological activities of SF-25 IgE.20 We investigated the Fc-mediated functions of SF-25 IgE to trigger mast cell degranulation when cross-linked by multiple copies of its target antigen expressed on the surface of cancer cell lines A2058, IGROV1, SKBR-3 and LS-180. Degranulation, measured by β2-hexosaminidase release, in the presence of non-specific control IgE was minimal, while SF-25 IgE triggered significant mast cell degranulation in the presence of different target expressing cancer cells (figure 5B). ADCC was measured in coculture experiments using human effector cell and target cell fluorescence reporters (online supplemental figure S6A,B). In the presence of healthy volunteer or melanoma patient PBMCs, SF-25 IgE engendered significant A2058 melanoma cell killing above non-specific isotype IgE or no antibody controls (figure 5C).

In vitro, SF-25 IgE did not affect proliferation or the clonogenic ability of cancer cells. The antibody exerted Fc-mediated effector functions via the high affinity FcεRI receptor. When cross-linked by multiple copies of an antigen expressed on the target cell surface, SF-25 IgE mediated specific mast cell degranulation and triggered cytotoxicity of cancer cells by both healthy volunteer and cancer patient immune cells.

**SF-25 IgE does not mediate basophil degranulation in a whole cancer patient blood assay ex vivo**

An IgE antibody introduced in the human circulation could bind to FcεRI-expressing basophils in the blood. If basophil bound IgE is cross-linked by signals such as multivalent soluble circulating antigen or by antibodies recognizing circulating multivalent antigen, this could trigger basophil degranulation, potential hypersensitivity and the onset of systemic anaphylaxis. We therefore evaluated the potential of SF-25 IgE to trigger basophil degranulation in human cancer patient blood using the basophil activation test (BAT). The BAT assay is used to monitor for hypersensitivity to the first-in-class IgE immunotherapeutic as part of its clinical development.9 15 21 Whole blood samples from ovarian cancer patients were incubated with either stimulation controls: anti-FcεRI antibody to crosslink the IgE receptor, fMLP (N-Formylmethionyl-leucyl-phenylalanine), a polyclonal activator of human basophils, or polyclonal anti-human IgE to cross-link endogenous IgEs already bound to the surface of human blood basophils. Basophil populations were identified by expression of CCR3 (online supplemental figure S5A). Basophil activation was evaluated by detection of CD63 cell surface expression, a marker normally absent in resting basophils (online supplemental figure S6C,D). Marked levels of basophil activation were observed when ovarian cancer whole blood samples were incubated with the three positive stimulation controls. These data confirmed propensity for patient basophil activation by both IgE and non-IgE-mediated mechanisms (figure 5D). In contrast, the incubation of blood with SF-25 IgE did not lead to basophil activation above a threefold threshold (figure 5D inset) in any of the 23 cancer patient samples tested.

These findings suggest that SF-25 IgE could not trigger basophil activation in a functional ex vivo assay, demonstrating no hypersensitivity reaction in cancer patient whole blood. These findings provide early evidence to support safe administration of this antibody to patients.

**A SLC3A2 specific CAR derived from the SF-25 antibody clone demonstrates tumor cell cytotoxicity in vitro**

To generate an SLC3A2-specific CAR, we first sequenced the variable regions of the heavy and light chain of the SF-25 clone and performed sequence modifications to the framework 1 region of both chains. The resultant single-chain variable fragments (scFv) were fused to a modified CD28 hinge containing a myc-tag, CD28 transmembrane and costimulatory intracellular domains and CD3ζ stimulatory domain (4SFm28ζ).22 A control CAR, truncated at the intracellular domain of CD28, was also engineered (4SFm28Tr). Retroviral vector cassettes were generated with the 4αβ chimeric cytokine receptor upstream of the CARs separated by a T2A sequence (figure 6A). The 4αβ (interleukin 4, IL4/2R) enables selective expansion and enrichment of CAR-positive T cells after transduction through delivery of an IL-2 binding signal in response to
Figure 6  4SFm28ζ CAR T cells effectively target adherent cancer cell lines in vitro. (A) Plasmid maps of the second generation 4SFm28ζ CAR construct (top panel) and 4SFm28Tr control construct (bottom panel). (B) Transgene expression level, expressed as MFI, and transduction efficiency, as percentage transduction, of untransduced (black circles), 4SFm28Tr (gray squares) or 4SFm28ζ (red triangles) CAR constructs. Cells were probed after 10 days in culture for the myc-tag, incorporated in the extracellular domain of the constructs, with the anti-MYC tag antibody (BE10). MFI reported as average ± SD of n=3 independent experiments. (C) Cytotoxicity of 4SFm28ζ CAR T cells on four different tumor cell line monolayers. Plots represent the percentage of viable tumor cells after a 24-hour coculture with untransduced (black circles), 4SFm28ζ (red triangles) or 4SFm28Tr (gray squares) T cells from three different donors±SD at three different E:T ratios. (D) Interleukin-2 (bottom left panel) and interferon-γ (bottom right panel) concentrations in coculture supernatants at 24 hours. Supernatants from untransduced (black circles), 4SFm28Tr (gray squares) and 4SFm28ζ (red triangles) T cells from three different donors±SD. (E) Activation markers (CD69 and PD-1) were probed pre (open symbols) and post (closed symbols) 72-hour coculture on PC3-LN3 monolayers at a 1:2 E:T ratio. Representative dot plots are shown. CD69 expression by MFI preantigen and postantigen exposure on untransduced (black circles), 4SFm28Tr (gray squares) and 4SFm28ζ (red triangles) T cells from three different donors±SD. PD-1 percentage expression is also demonstrated. (F) T cell phenotyping pretarget and post-target cells exposure. Representative plot (top left) for untransduced T-cells. Color-coded chart (bottom left) of phenotype quadrants.
IL-4. After 10 days in IL-4 supplemented culture, robust CAR expression was seen (figure 6B).

High (melanoma—A2058), middle (prostate—PC3-LN3, breast—MDA-MB-468) and low (breast—MDA-MB-231) SF-25 expressing cell lines (figure 1C) were used for in vitro cytotoxicity analysis at effector to target (E:T) ratios of 5:1, 1:1 and 1:5. Significant CAR specific cytotoxicity was demonstrated at 24 hours at all coculture densities for the A2058 and MDA-MB-468 cell lines. For PC3-LN3 4SFm28ζ cytotoxicity was significant compared with 4SFm28Tr at the 5:1 E:T ratio and against untransduced T cells at 5:1 and 1:1. For the low density MDA-MB-231 cell line significantly superior 4SFm28ζ cytotoxicity was seen vs 4SFm28Tr at E:T 5:1 and untransduced T cells at E:T 1:1 (figure 6C). Culture supernatants probed for and IL-2 demonstrated donor variability. At an E:T of 1:1 at 24 hours IL-2 levels were significantly higher for 4SFm28ζ compared with both controls against PC3-LN3, MDA-MB-231 and A2058. Interferon gamma was raised for 4SFm28ζ compared with controls in all cocultures (figure 6D). These data demonstrate signaling CAR T restricted antigen specific in vitro efficacy of SF-25 derived CAR T cells.

Using coculture at an E:T ration of 1:2 on the prostate cancer monolayer PC3-LN3 4SFm28ζ, 4SFm28Tr and untransduced T cells were probed for phenotype and markers of activation (CD69 and PD-1) prior to and after exposure to monolayers. MFI of CD69 significantly increased on 4SFm28ζ CAR T cells after exposure to antigen compared with controls. Accordingly, PD-1 percentage increased post antigen exposure for the signaling CAR (figure 6E). On exposure to PC3-LN3, activation marker levels increased for the 4SFm28ζ signaling CAR only, commensurate with antigen specific activation (figure 6E). Viral transduction alone (for 4SFm28ζ and 4SFm28Tr) resulted in loss of naive populations and expansion of both T effector memory (TEM) and T effector memory-RA (TEMRA) populations compared with the activated but untransduced controls (figure 6F). Allowing for the natural variability in donors, the impact of engagement of antigen by CAR T cells resulted in further enrichment of TEMRA (figure 6E). These data demonstrate robust, antigen specific, activation of 4SFm28ζ CAR T cells.

SF-25 IgE and derivative CAR immunotherapies demonstrate in vivo activity

Immunodeficient NSG mice were intravenously injected with human LS-180 colorectal cancer cells which led to the development of cancer lesions in animal lungs. Freshly isolated human PBMCs were adoptively transferred in the presence or absence of SF-25 IgE. Antibody treatment alone was readministered twice more (figure 7A). After 3 weeks, tumor load in the lungs of mice was measured (figure 7B). Analyses demonstrated significantly restricted tumor growth in SF-25 IgE-treated mice, both in relation to the number of metastatic foci (p=0.0028) and with regard to the area of tumor occupancy in animal lungs (p=0.0061) compared with PBMC treated controls (figure 7C, D).

For CAR T in vivo efficacy testing, PC3-LN3-luciferase subcutaneous xenografts were established in NSG mice. Tumor engraftment was confirmed by bioluminescence imaging. Tumor growth was monitored by caliper measurements as central tumor necrosis limits bioluminescence accuracy. Tail vein injections of 4SFm28ζ and 4SFm28Tr CAR T cells were undertaken 3 days after tumor inoculation (figure 7E). A significant increase in overall survival was seen in the 4SFm28ζ group (p=0.0208) (figure 7F). Of five, two treated mice demonstrated tumor eradication after initial growth, with one showing initial response followed by tumor escape (figure 7G). The animals displayed no signs of weight loss or manifest any symptoms of cytokine release syndrome or graft-versus-host disease (online supplemental figure S7).

These findings collectively support the functional capability of novel immunotherapies targeting SLC3A2 to restrict the growth of tumors and improve survival in vivo in the absence of overt toxic effects.

DISCUSSION

We have identified the solute carrier family member SLC3A2 as the target of the SF-25 antibody clone. The bioinformatics approach described here constitutes a novel approach to identify target proteins for an antibody with unknown antigen specificity. A similar comparison method has been applied once for identifying an aptamer’s target. The SLC3A2 protein CD98hc is expressed at high levels on several solid tumors. Normal tissue distribution is restricted to kidney, testis and cerebellum. We have demonstrated the utility of SLC3A2/CD98hc targeting with two highly current immunotherapy approaches. First, human IgE class engineering of the SF-25 mouse clone and second, integration of a derivative scFv into a CAR T cell approach.

The protein product of SLC3A2 (initially known as 4F2 cell-surface heavy chain (4F2hc)) was identified as the heavy chain subunit of CD98 complexes (CD98hc). CD98hc is a component of cell surface heterodimeric complexes, stabilized by disulfide bonds, with several putative light chain subunits, including the L-type amino acid transporters LAT1 (SLC7A5), LAT2 (SLC7A8), y+LAT1 (SLC7A7), y+LAT2 (SLC7A6), ascAT1 (SLC7A10) and the cystine/glutamate antiporter xCT (SLC7A11). Each of these complexes allows specific solutes to cross the plasma membrane, the light subunit conferring the solute specificity to the complex. In concordance, our bioinformatics analysis showed that SLC3A2 and SLC7A5 expression highly correlated with binding of the SF-25 antibody. This suggests that SF-25 may be specifically, or preferentially, interacting with the
Figure 7  SF-25 IgE and SF-25-based CAR T cells restrict tumor growth in vivo. (A) Schematic representation of the experimental timeline for investigating SF-25 IgE effects on the development of LS-180 cancer lesions in the lungs of NSG mice. (B) Diagram of necropsy procedure for lung removal. The mid-line incision to expose the trachea is depicted with a purple dotted line. The black arrow shows how the Indian ink is infused into the lungs through the tracheal cannula. Lungs were removed en bloc from the thoracic cavity (red dotted line). (C) Representative images of lungs extracted from control-treated and SF-25 IgE-treated animals after staining of tumor-bearing tissues with India ink and subsequent analysis under an inverted light microscope. (D) Scatter plots of the number of tumor lesions (on the right) and of the percentage of tumor occupancy (left) in mouse lungs (each dot represents values from one mouse, average measurement ±SEM of each experimental group (PBMCs: n=8; SF-25 IgE +PBMCs: n=6) and data from two independent experiments with two human PBMC donors; non-parametric t-test; tumor occupancy, *p-value 0.0061, number of metastases **p-value 0.0028. (E) Schematic representation of the experimental timeline employed to investigate SF-25 based CAR effects on tumor growth in a PC3-LN3-fLucTdTTomato subcutaneous model in NSG mice. Total bioluminescence efflux (photons per second) was measured to verify tumor engraftment and randomize the mice at day 3. (F) Survival curves for the three different groups, mice have been culled when tumors were about to reach 1500 mm³ or had lost 15% of their weight at day 0 (statistical analysis performed with Mantel-Cox log-rank test; *p value 0.0189). (G) Individual tumor growth curves as measured with calipers in PBS treated (left), truncated CAR T cells (middle) and SF-25 second generation CAR T cells (right panel). The two long-term remission mice are highlighted with round and square symbols on the curve (see also online supplemental figure S7). CAR, chimeric antigen receptor; PBMCs, peripheral blood mononuclear cells. Mouse image used courtesy of MCKIBILLO.
CD98hc-LAT1 heterodimer. Additionally, CD98hc acts as a chaperone protein enabling translocation of heterodimerization partners from the endoplasmic reticulum to the cell surface. The resulting amino acid trafficking is central to cell functionality, providing essential amino acids for protein, vitamin and nucleotide synthesis.28–30

CD98hc has been demonstrated to stabilize the glucose transporter GLUT1,31 and to interact with galectin-3, ICAM-1, CD9, and integrins β1a, β3 and B4, suggesting a broader role in tissue architecture.30–34 It is directly involved in pathways leading to ER-stress responses,35 oxidative stress responses,35–37 B and T cell activation,35–37 cell fusion,35 mechanotransduction,36 angiogenesis,37 cell survival and migration,37 and cell proliferation.37–39 Highly proliferative tissues, in health and disease, overexpress CD98hc. It has been directly linked to tumorigenesis by mediating β1 integrin signaling50 and indirectly by affecting the mTOR activity51 through its associated light chain amino acid transporters.51

The potential to capitalize on CD98hc expression and functional importance to highly metabolic tumors has been established.51–55 A role in resistance to chemotherapy,51–54 radiotherapy,55 and to T cell-mediated killing through inhibition of ferroptosis,56 together highlight potential for integrating targeting with established therapeutic modalities to overcome resistance. Recent immunotherapy and theranostic approaches have been developed.57–59

In this study, we report the engineering, production, purification and functional evaluations of two immunotherapy approaches based on SF-25. SF-25 IgE demonstrates specific antitumor activity in vitro and in vivo. Our results are consistent with previous findings with our first in class IgE antibody MOv18, specific for the TAA folate receptor alpha (FRα). MOv18 is in phase 1 clinical testing with early data demonstrating safety and signs of biological activity in patients with ovarian cancer.60–62 We showed that SF-25 IgE could restrict the growth of tumor lesions in the lung of mice when administered with human immune effector cells. Administration of human immune effector cells was necessary in murine models to study the ability of SF-25 IgE to engender Fc Receptor interactions as the human IgE Fc does not cross-react with mouse Fc receptors. Employing a severely immunocompromised animal model could underestimate the antitumor activity of the IgE tested since the absence of a self-replenishing supply of effector cells, a lack of mature mast cells and a short human PBMC lifespan in immunocompromised mice all limit the time frame for an antibody to exert effector functions in an in vivo rodent model.62

Future studies could address this limitation through cytokine or growth factors supplementation to enhance the survival of specific human immune cell populations.63,64

By interrogating a basophil activation (BAT) test, we demonstrated that SF-25 IgE is unlikely to elicit a type I hypersensitivity reaction in cancer patient blood. These findings support early preclinical safety evaluations for this antibody. The BAT is employed alongside clinical observations, and biological parameters, to monitor and potentially predict safe administration of IgE immunotherapies.65 For SF-25 IgE, further ex vivo BAT assays using different SLC3A2+ cancer patient cohorts will confirm lack of basophil activation and provide confidence in future safe administration to humans.

Our second immunotherapy approach, with a 4SFm28ζ CAR T cells, has broad potential for clinical translation. Loss of cytotoxicity below a 1:1 E:T ratio highlights the ‘tunability’ of CAR for this target. Expression of CD98hc on activated T cells has been reported.66 This raises the possibility that ‘fratricide’ of CD98hc targeting CAR T cells could occur during manufacturing. Nonetheless, we were able to consistently generate sufficient CAR-positive T cells and there was no suggestion of increased enrichment for the 4SFm28ζ transduced cells, which one would expect if there was significant stimulation through the chimeric receptor during expansion. One Uni-CAR approach targeting CD98hc, based on the monoclonal antibody MEM-108, recently demonstrated no fratricide during coculture with tumor monolayers, despite transient expression on activation. The authors concluded that the antigen density on target cells was far higher than the CAR T cells with the threshold for cytotoxicity higher than the level of T cell expression of CD98hc.67 The upregulation of CD98hc in both activated immune cells and tumor cells suggest that these cells compete for the same metabolites as a source of energy. Targeting SLC3A2 on cancer cells could enhance the ability of immune cells to function in the tumor microenvironment.68 Loss of target expression is a key mechanism of acquired resistance to CAR T cell therapies in the clinic.69 In the case of SLC3A2, we observed that its knock-out in A2058 cancer cells stopped their in vitro proliferation, which is consistent with previously reported data both in vitro and in vivo.70 As a consequence, antigen loss is unlikely to be a resistance mechanism to SLC3A2 targeting as it would result in impaired cell proliferation in cancer cells. Future studies combining this CD98hc specific CAR with additional engineering strategies could further enhance the potential of 4SFm28ζ as a translatable cancer therapeutic.

CONCLUSION

Immune metabolism is a nascent and fertile area of therapeutic discovery. The preferential transport and utilization of key solutes by malignant cells in a tumor microenvironment is both key to tumor progression and resistance to immune-mediated killing. The role of SLC3A2/CD98hc in transporting and stabilizing multiple solute transporters and cell–cell adhesion

molecules makes it a highly attractive candidate for immune therapy development. Here, we have shown its utility for novel antibody class and CAR T cell targeting and provide early evaluations of safety. Our findings form the basis for developing new treatment options for aggressive cancers and contribute to the short but expanding panel of promising target antigens for novel immunotherapies.

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Competing interests SNK and JS are founders and shareholders of EpiSolon Ltd., and HJB is now employed through a fund provided by EpiSolon Ltd.

Patient consent for publication Not required.


SUPPLEMENTAL MATERIALS & METHODS

Human blood samples

Human samples were collected with informed written consent. Melanoma patient blood samples were collected as part of studies conducted at King’s College London, Guy’s and St Thomas’ NHS Foundation Trust (08/H0804/139 approved by London Bridge NRES committee and 16/LO/0366 approved by London-Central NRES Committee); Ovarian cancer patient blood samples were collected as part of a study conducted at King’s College London, Guy’s and St Thomas’ NHS Foundation Trust (09/H0804/45). Peripheral venous blood was collected in BD Vacutainer™ Hemogard Closure Plastic K2-EDTA Tubes (BD, Wokingham, UK). Peripheral blood was also obtained through the UK National Health System (NHS) Blood and Transplant system from anonymous donor leukocyte cones.

PBMC isolation from blood samples

Equal volumes of blood and 2% FCS/2 mM EDTA were gently mixed to a final volume of 30 mL and gently pipetted on top of 15 mL of Ficoll-Paque™ PLUS density gradient in a 50 mL conical tube. The tube was then centrifuged at 1200 x g with slow acceleration and no brake at room temperature (RT) for 20 minutes. The plasma interface was collected and washed with PBS at 600 x g at 4°C for 10 minutes. The erythrocytes present in the sample were lysed with RBC lysis buffer for 5 minutes at RT, followed by washing with PBS + 2% FCS/2 mM EDTA.

Cell lines

IGROV1 ovarian cancer cells were a gift from Prof. Silvana Canevari (Istituto Nazionale dei Tumori, Milano, Italy). PaTu-8988-T and PANC-1 human pancreatic cancer cell line were kindly provided by Dr Debasish Sarker at the department of Research Oncology, School of Cancer and Pharmaceutical
Sciences, King’s College London. The rat basophilic leukemia RBL SX-38 cells transfected to stably express the human FcεRI αβγ was kindly provided by Professor J.P. Kinet (Harvard University, Boston, USA). The PC3-LN3 (PL) cell line was kindly provided by Professor Sue Eccles (Institute of Cancer Research, Sutton, UK). All other cancer cell lines were sourced from ATCC. Expi 293-F human embryonic kidney cells were from ThermoFisher. Cell culture media for cell lines were supplemented with 10% fetal calf serum (FCS) (v/v), 100 U/mL penicillin and 100 U/mL streptomycin. PBMCs and T cells were cultured in RPMI1640 plus 5% human serum. Adherent cell lines were passaged once they reached 80-90% confluence by detachment in 0.5% Trypsin/0.53 M EDTA at 37°C for 5 minutes, washed and plated in fresh media.

**SF-25 antigen immunoprecipitation**

Pellets containing 20 to 50x10⁶ target expressing MDA-MB-231, A2058 or MDA-MB-468 cells were resuspended in 1.75mL lysis buffer (PBS, 0.1% Tween20, 1X Halt™ protease inhibitors cocktail – Thermo Scientific) in a 15mL tube, incubated at 4°C on a roller and vortexed for 20 minutes. 300µL ProteinA Dynabeads® (Invitrogen) were prepared with 100µg SF-25 humanized IgG1 in 800µL Binding Buffer as per manufacturer’s protocol. Cell lysates were centrifuged for 10 minutes at 4500rcf at 4°C and supernatants were transferred to the washed SF-25 Dynabeads®. Beads with immunoprecipitated fraction were placed in 30µL of Elution Buffer and 10µL of 4X LDS sample buffer (Invitrogen) and stored at -20°C. Samples were thawed, beads were concentrated on a magnet, supernatant were transferred to a 1.5mL microcentrifuge tube and β-mercapto-Ethanol was added to a final concentration of 5% (v/v). Samples were incubated for 10 minutes at 95°C before being resolved on a 4-12% gradient NuPAGE™ gel (Invitrogen) at 200 Volts in MOPS buffer. Migration was stopped when Coomassie G250 reached the bottom of the gel. The gel was fixed in 7% Acetic Acid/40% methanol (v/v) for 30 minutes at room temperature and stained with a 1X solution of colloidal Brilliant Blue G with 20% methanol for 1 hour at room temperature. The gel was first destained for 5 minutes with 7% Acetic Acid and
25% methanol and then overnight with 2% acetic Acid and 25% methanol and stored in distilled water. Bands of interest were cut out and sent to Aulesa Biosciences for mass spectrometry analysis.

SF-25 antigen: Mass spectrometry analysis

Proteins in gel slices were reduced (DTT), alkylated (iodoacetamide) and digested overnight with trypsin. Peptides within the tryptic digests were fractionated using an Ultimate 3000 nano-LC system in line with an Orbitrap Fusion Tribrid mass spectrometer (Thermo Scientific). In brief, peptides in 1% (v/v) formic acid were injected onto an Acclaim PepMap C18 nano-trap column (Thermo Scientific). After washing with 0.5% (v/v) acetonitrile 0.1% (v/v) formic acid peptides were resolved on a 250 mm x 75 μm Acclaim PepMap C18 reverse phase analytical column (Thermo Scientific) over a 150 min using 7 gradient segments (1-6% solvent B over 1min, 6-15% B over 58min, 15-32%B over 58min, 32-40%B over 5min, 40-90%B over 1min, held at 90%B for 6min and then reduced to 1%B over 1min) with a flow rate of 300 nL.min⁻¹. Solvent A was 0.1% formic acid and Solvent B was aqueous 80% acetonitrile in 0.1% formic acid. Peptides were ionized by nano-electrospray ionization at 2.2 kV using a stainless-steel emitter with an internal diameter of 30 μm (Thermo Scientific) and a capillary temperature of 250°C. All spectra were acquired using an Orbitrap Fusion Tribrid mass spectrometer controlled by Xcalibur 2.0 software (Thermo Scientific) and operated in data-dependent acquisition mode. FTMS1 spectra were collected at a resolution of 120 000 over a scan range (m/z) of 350-1550, with an automatic gain control (AGC) target of 400 000 and a max injection time of 100ms. The Data Dependent mode was set to Cycle Time with 3s between master scans. Precursors were filtered according to charge state (to include charge states 2-7), with monoisotopic precursor selection and using an intensity range of 5E3 to 1E20. Previously interrogated precursors were excluded using a dynamic window (40s +/-10ppm). The MS2 precursors were isolated with a quadrupole mass filter set to a width of 1.6m/z. ITMS2 spectra were collected with an AGC target of 5000, max injection time of 50ms and HCD collision energy of 35%. LC-MS/MS data was processed using Proteome Discoverer.
(ThermoFisher Scientific) with database searching against a downloaded FASTA file originating from Uniprot_SwissProt_2019_02. Results were initially visualized within the software and then exported to Excel for further review.

**SLC3A2 differential expression study**

Data for normal and associated tumor tissues were retrieve from the UCSC Toil RNA-seq Recompute dataset.[1] TPM values data from TARGET were filtered out and statistical analysis of SLC3A2, HER1 and HER2 differential expression was assessed by Mann-Whitney test in Graphpad Prism. All tumor samples were paired with the associated normal tissue samples from TCGA and the GTex normal tissues specified below. READ: Rectum Adenocarcinoma (GTex Colon); COAD: Colon Adenocarcinoma (GTex Colon); LIHC: Liver Hepatocellular Carcinoma (GTex Liver); CHOL: Cholangiocarcinoma; ESCA: Esophageal Carcinoma (GTex Esophagus); BLCA: Bladder Urothelial Carcinoma (GTex Bladder); STAD: Stomach Adenocarcinoma (GTex Stomach); OV: Ovarian Serous Cystadenocarcinoma (GTex Ovary); LGG: Brain Low Grade Glioma (GTex Brain); GBM: Glioblastoma Multiforme (GTex Brain); PAAD: Pancreatic Adenocarcinoma (GTex Pancreas); PRAD: Prostate Adenocarcinoma (GTex Prostate); ACC: Adrenocortical Cancer (GTex Adrenal Gland); BRCA: Breast Invasive Carcinoma (GTex Breast); TGCT: Testicular Germ Cell Tumor (GTex Testis); UCEC: Uterine Corpus Endometrioid Carcinoma (GTex Uterus); UCS: Uterine Carinosarcoma (GTex Uterus); CESC: Cervical & Endocervical Cancer (GTex Cervix Uteri); THCA: Thyroid Carcinoma (GTex Thyroid); LUAD: Lung Adenocarcinoma (GTex Lung); LUSC: Lung Squamous Cell Carcinoma (GTex Lung); SKCM: Skin Cutaneous Melanoma (GTex Skin); LAML: Liquid Acute Myeloid Leukemia (GTex Blood); DLBC: Diffuse Large B-cell Lymphoma (GTex Blood); HNSC: Head & Neck Squamous Cell Carcinoma; KICH: Kidney Chromophobe (GTex Kidney); KIRP: Kidney Papillary Cell Carcinoma (GTex Kidney); KIRC: Kidney Clear Cell Carcinoma (GTex Kidney).

**Retroviral constructs for CAR expression in PBMCs**

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The SF-25 scFv was subcloned into a myc-tag containing 28ζ construct downstream of the 4αβ chimeric cytokine receptor (IL4/2R).[2] The truncated CAR version 4SFm28Tr was generated by PCR by introducing a stop codon and a cloning site after the Lysine in position 3 in the cytoplasmic CD28 fragment. RD114 viral particles were produced by transiently transfecting HEK-293T cells. 1.5x10^6 cells were plated in a 10cm diameter culture dish in IMDM medium without antibiotics and allowed to grow overnight. Transfection reagents were prepared by gently pipetting 30µL GeneJuice® (Novagen) mammalian cell transfection reagent into 470µL plain IMDM medium (no serum) and incubated for 5 minutes at room temperature. Plasmids were gently added and incubated for 15 minutes before evenly dispensing the transfection reagents dropwise over the 10cm dish. Triple transfections were performed with 3µg RD114, 4.5µg pEQ-Pam3 and 4.5µg of the CAR containing plasmid. Supernatants containing the RD114 viral particles were harvested at 48 and 72 hours post transfection, pooled aliquoted, snap frozen in dry-ice cold ethanol bath and stored at -80°C. Stable packaging cell lines were established in HEK293 VECS GaIV cell by transducing 1x10^5 overnight plated cells with 2mL RD114 viral supernatant in 6 well plate. Transduction efficiency in HEK293 VECS GaIV cells were assessed by flow cytometry using the 9E10 anti-myc antibody and a polyclonal Goat anti-Mouse IgG PE secondary antibody (Agilent). Cells were transduced with RD114 virus until achieving >95% transduction. For GaIV viral production, HEK293 VECS GaIV transduced cells were grown in complete DMEM medium without antibiotics in 175cm^2 culture flask and supernatant was harvested when reaching 90% confluency. GaIV virus containing supernatants were aliquoted, snap frozen and stored at -80°C.

**Human PBMCs transduction and CAR T cell expansion**

Isolated PBMCs were counted and placed in 6 well non-tissue culture plates at a concentration of 3x10^6 cells per mL at a maximum of 3mL per well. Polyhydroxyalcanoate (PHA) was used to activate the PBMCs at a final concentration of 5µg/mL. Plates were incubated for 24 hours at 37°C and 5% CO2. Interleukin-2 is added at 24 hours to a final concentration of 100U/mL. RetroNectin® (Takara) coating
solution was prepared using polypropylene pipettes by mixing 200µg RetroNectin® with 12mL cold PBS. Non-tissue culture treated 6-well plates were coated by overnight incubation at 4°C. Coating solution was replaced with 2mL GaLV virus containing supernatant and incubated at 4°C for 4 hours before being replaced by another 2mL of pre-chilled viral supernatant. At 48 hours post-isolation and 24 hours after Interleukin-2 treatment, 1x10^6 activated PBMCs were added to each well in 500µL RPMI 5% human serum. Human interleukin-2 and interleukin-4 (R&D systems) were added at a final concentration of 100U/mL and 30ng/mL respectively. Fresh RPMI with 5% human serum containing interleukin-2 (untransduced cells) or interleukin-4 (4αβ transduced cells) was added every 2 days during the expansion phase. Enrichment of CAR positive T cells was assessed by Flow Cytometry on a Fortessa cytometer at days 10 using the 9E10 anti-myc antibody. Analysis of flow cytometry data was performed by FlowJo (TreeStar Inc) software.

**Tumor cell and CAR T cell coculture assays**

Tumor cells were seeded at 5.10^4 tumor cells per well in 48-well plate in 200µL culture medium without antibiotics and allowed to form monolayers over 24 hours at 37°C in 5% CO₂. CAR T cells were washed in PBS and 25.10^4, 5.10^4 or 1.10^4 T cells were added in 100µL culture medium to the monolayers respectively resulting in a 5:1, 1:1 and 1:5 Effector T cell:Target cell ratio. After a 24-hour coculture, 200µL of supernatant was harvested and stored at -20°C for further cytokine measurement. Monolayers viability was assessed immediately as detailed in the main text Methods.

**Cytokine detection after coculture**

Supernatants from cocultures were thawed and analyzed using human Interferon-γ (R&D systems) and human Interleukin-2 (Invitrogen) ELISA kits as described by manufacturer. Supernatants were diluted from 15 to 45-fold to fit within the standard curves. Cytokine levels were plotted using Graphpad Prism.
Phenotype and activation analysis

At day 14-post viral transduction, T cells pre-coculture were counted, washed in 2mL FACS buffer and 25,10^4 cells were stained either with a cocktail of anti-human CD69-APC (FN50 Biolegend) / anti-human PD-1-PE (EH12.2H7 Biolegend) / antihuman CD3-FITC ; anti-human CD45ra-APC (JS-83 Invitrogen) / anti-human CD197(CCR7)-PE (3D12 Invitrogen) / anti-human CD3-FITC (UCHT1 Biolegend) or corresponding isotypes (MOPC-21 -FITC -APC Biolegend / P3.6.2.8.1-PE Invitrogen) for 20 minutes at 4°C. Cells were then washed with 2mL FACS buffer and a Live/Dead staining was performed with 7-AAD reagent as per manufacturer instructions (Cambridge Bioscience). Samples were run on a BD LSFRFortessa flow cytometer and live CD3+ cells phenotype and activation pattern were analyzed using FlowJo (TreeStar Inc) Software. Anti-Mouse Igκ compensation beads (BD™) were also used. 72 hours cocultures were set the same day in 24-well plates with 1.8x10^5 PC3-LN3 overnight-grown monolayers at a 1:2 Effector:Target ratio. After coculture, cells were flushed from 4 wells/construct, pooled and distributed into 3 FACS tubes (2/5th for phenotype labelling, 2/5th for activation labelling and 1/5th for isotype labelling). Same staining, gating strategy and analysis were used in post-coculture conditions as per pre-coculture staining. Analysis of flow cytometry data was performed by FlowJo (TreeStar Inc) software and data were plotted using Graphpad Prism.

CD98hc Expression on human PBMCs

Human PBMCs were isolated by density centrifugation as describe in the Methods. Basal expression level was assessed on 1.10^5 cells stained with the 1µg anti-CD98hc antibody MEM-108 (Biolegend) for 20 minutes at 4°C, washed with 2mL cold FACS buffer, stained again with goat anti-mouse IgG AlexaFluor647 (Jackson Labs), washed again prior to analysis. Isotype control staining was utilized. Cell surface staining was assessed by Flow Cytometry on a BD LSFRFortessa cytometer. PBMCs were then activated or not with PHA as described above. Over 15 days, 1x10^5 PBMCs were retrieved from the cell
culture and stained as described above. Analysis of flow cytometry data was performed by FlowJo (TreeStar Inc) software and data were plotted using Graphpad Prism.

**Immunofluorescence study of antigen expression**

SF-25 IgG1 and NIP-IgG1 were directly labelled with Alexa Fluor 488 fluorophore using the Alexa Fluor™ 488 Antibody Labeling Kit (ThermoFisher). Frozen tissues were thawed at room temperature for 10 minutes, allowed to dry and fixed with 4% paraformaldehyde (PFA) for 10 minutes. Once PFA was removed, tissues were washed 3 times with PBS. Sections were covered with Human AB serum for 1 hour at room temperature. Tissues were incubated overnight with SF-25 IgG1-AF488 or NIP-IgG1-AF488 and kept at 4°C in the dark. Sections were washed 3 times in PBS and allowed to dry at RT. ProLong™ Gold Antifade Mountant with DAPI (ThermoFisher) was added, coverslip were applied and slides solidified overnight at room temperature in the dark.

**In vivo human cancer xenograft model to study SF-25 IgE**

On day 0, NSG mice were injected simultaneously with 5x10^5 LS-180 colorectal cancer cells, 5x10^6 PBMCs from a healthy human volunteer and 10mg/kg of SF-25 IgE or 50µl PBS, to a final volume of 200µl. Subsequent injections of antibody/PBS were performed on days 2 and 3. Mice were sacrificed on day 21 by CO_2 asphyxiation. Lungs were analyzed for tumor growth in the lungs using the following protocol: the trachea was exposed by performing a mid-line incision from below the diaphragm to the throat and the chest activity opened. An intravenous cannula was inserted into the trachea, and a 10 mL syringe was attached to it. Approximately 3 mL of a solution of 15% (v/v) Indian Ink was then injected into the lungs. The ink-stained lungs were then removed from the thoracic cage, placed in MilliQ water to remove the excess ink and then transferred to Fekete solution for 48 hours. The number of metastatic nodules/lung lobe and the proportional surface occupancy of tumor nodules were calculated. Images were acquired using a Nikon SMZ1500 Stereo Microscope (Nikon UK Ltd) with 0.75x.
and 1.0x objective. The NIS Elements Basic Research software (Nikon UK Ltd) was used to determine the total surface area of the lung section and to distinguish the white lung metastasis nodules. Data were acquired from two independent experiments using PBMCs from two human donors.

**In vivo SF-25 CAR T cell in prostate tumor model**

6 to 12 weeks old NSG™ mice have been inoculated with 2.5x10⁵ PC3-LN3 cells engineered to express ffLuciferase and TdTomato (PL-LT) by subcutaneous (s.c.) injection on the right flank. At Day 3, Bioluminescent imaging (BLI) was performed using Xenogen IVIS imaging system with Living Image software (Xenogen). Mice were injected intra-peritoneally (i.p.) with 200μL of imaging solution (15μg/mL D-Luciferin in PBS), anaesthetized under 2% Isofluorane and imaged after 20 minutes. Mice were distributed in the different groups to ensure minimal variation in mean BLI signal and standard deviation between groups. The same day, 1x10⁷ CAR T cells were adoptively transferred to mice intravenously (i.v.) in 200μL PBS. Mice were weighed and their tumors measured by calipers 2 to 3 times a week for the rest of the experiment.

**Lectin blots**

Purified IgE samples (150ng) were reduced with 50mM dithiothreitol and boiled at 95°C for 5 minutes. Samples were run at 150 V on Mini-PROTEAN TGX Gels 4-15% (Bio-Rad Laboratories, Hercules, Calif) and blotted with Trans-Blot Turbo Transfer Pack PVDF (Bio-Rad Laboratories) by using the Trans-Blot Turbo Blotting System (Bio-Rad Laboratories) according to the manufacturer’s instructions. The blotted membrane was then cut just above 35kDa to have heavy (50kDa) and light (25kDa) chains in different membranes. The heavy chain membrane was blocked with Carbo-Free Blocking Solution (Vector Laboratories) for 1 hour and then probed with Ricinus communis agglutinin I lectin (RCAI-biotin) [Vector Laboratories] specific for galactose, Aleuria aurantia lectin (AAL-biotin) [Vector Laboratories] specific for fucose, concanavalin A lectin Con-A–biotin) [Vector Laboratories] specific for
mannose, or EBL (Sambucus nigra lectin) [Vector Laboratories] specific for sialic acid at 0.2 μg/mL in Carbo-Free Blocking Solution for 30 minutes. The membranes were then washed 3 times in PBS–Tween 0.05% (T-PBS) and incubated with High Sensitivity Streptavidin-HRP (1:30000; Pierce) for 30 minutes, washed as above, and developed with ECL (Amersham, GE Healthcare). The light chain membrane was blocked with T-PBS and 5% BSA for 1 hour at room temperature and then incubated overnight at 4°C with rabbit anti-human kappa light chain antibody (1:1000 in T-PBS 5% BSA; Abcam, Cambridge, United Kingdom), followed by 3 washes in T-PBS. The membrane was incubated with anti-rabbit IgG horseradish peroxidase antibody (1:2000 in T-PBS and 5% BSA; Cell Signaling Technology, Danvers, Mass) for 1 hour at room temperature, washed as above, and developed with ECL. Densitometric quantification was performed with ImageJ software (National Institutes of Health, Bethesda, Md), and values were normalized by the loading control (kappa light chain).

Transcript-to binding-level correlation analysis

This section is divided into three steps, an example of the code used is located at: https://github.com/ramipod/Antigen-Identification-by-Binding-and-Transcriptomic-Comparison

- TPM dataset generation (TPM averaging)
- Spearman score calculation for a specific binding dataset and the TPM counts (spearman AVG1)
- Spearman scores compilation (Spearman-AVGs)

Statistical methods

All statistical analyses were performed using GraphPad™ Prism Software (version 6.0). Error bars represent SD and SEM in in vitro and in ex vivo evaluations.
REFERENCES


### Table S1

#### SF-25 IgG1 immunoprecipitated proteins

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### Table S1. Related to Figure 2A - SF-25 target candidates identified by immuno-precipitation and mass spectrometry.

Target identification was conducted by a biochemical approach using immuno-precipitation and mass spectrometry analysis of pulled-down proteins from MDA-MB-468 tumor cell lysates. Two bands corresponding to 42kDa and 27kDa were sent for identification to Aulesa Biosciences. 138 proteins were identified in total using MASCOT.
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Table S2. Related to Figure 2A - Binding panels for bioinformatical target identification.
Relative binding scores generated for three independent binding screens with the SF-25 antibodies. Binding scores for Binding Panel 1 were generated from a radioligand binding assay with the SF-25 murine IgG1 antibody in (Takahashi et al., 1988). Binding scores for Binding Panel 2 were generated from flow cytometry binding assay with the SF-25 chimeric IgE antibody [Figure 1C]. Binding scores for Binding Panel 3 were generated from flow cytometry binding assay with the SF-25 chimeric IgG1 antibody [Figure 1C].
Figure S1. Related to Figure 1A&B - Evaluation of multiple cell culture vessels and media for SF-25 IgE antibody production.

(A) Different cell culture vessels were evaluated, and measurements were taken of both the SF-25 IgE concentration in the supernatant and of the cell viability over time. The graph on the left reports the yield of SF-25 IgE antibody over time across different culture vessels, while the graph on the right depicts the % of viable cells over the time in the same vessel types. (B) SF-25 IgE antibody concentration values and cell viability values were analysed for each cell culture vessel individually. Cells were seeded in each vessel with the same density of 5x10^5 cells/ml and they were added fresh medium to the maximum working volume after three days. From day 3 onwards cells were not fed and left to starve until cell culture was harvested. (C) Sp2/0 cells can be adapted to serum-reduced condition and maintain SF-25 IgE production and secretion in the supernatant. Error bars represent SEM. (D) Cell viability was investigated over time and after every cell passage in ADCF cell culture, showing how cells keep a higher vitality rate even in serum optimised conditions. Error bars represent SEM.
Figure S2. Related to Figure 2B - Amino acid sequence of SLC3A2 – identified peptides by immuno-mass spectrometry.

Peptides of the identified SLC3A2 protein are highlighted in the amino acid sequence. The sequence was obtained from UniProt (accession number=P08195).
Figure S3. Related to Figure 3 - SLC3A2 gene expression is enhanced in different malignancies compared with equivalent normal tissues.
Expression of SLC3A2 in normal (N) versus tumor (T) tissues of different origins. Tumor types are described in Supplementary Materials and Methods - SLC3A2 differential expression study. Mann-Whitney t-test was performed; ns = non-significant; * = p-value<0.05; ** = p<0.01; *** = p<0.005; **** = p<0.001.
**Figure S4**

**SLC3A2 expression in PBMCs**

![Graph showing SLC3A2 expression in PBMCs](image)

**SLC3A2 expression in PBMCs**

![Graph showing individual curves of SLC3A2 expression in activated and unactivated PBMCs](image)

---

**Figure S4. SLC3A2 expression in PBMCs in vitro.**

Grouped (top) and individual curves (bottom panel) of SLC3A2 expression in activated (plain) and unactivated (empty symbols) PBMCs cultured 15 days in vitro (n=3 individual donors). Activation is represented by an arrow. Mann-Whitney two-tailed analysis was performed excluding values before activation; **p=0.0041.**
Figure S5. Related to Figure 3 - Survival by expression of SLC3A2, SLC7A5 and SLC7A10 in human cancers.
Kaplan-Meier curves of survival over 10 years for patients with bladder, breast, cervical, head and neck squamous cell carcinoma, renal clear cell carcinoma and lung adenocarcinoma by SLC3A2, SLC7A5 and SLC7A10 expression. High (red) and low (black) expression at baseline.
Figure S6. Related to Figure 5(C-D)- Representative dot plots of ADCC/ADCP *in vitro* and of the Basophil Activation Test *ex vivo*

(A) Representative flow cytometry dot plots depicting the gating strategy to visualise the population of engulfed cells (left) and the population of dead target cells (right). (B) Representative flow cytometry plots depicting cell populations of effector (anti-CD89-APC) and target (FITC+) cells during ADCC/ADCP assay. The loss of tumour cells can be appreciated in the sample treated with SF-25 IgE when compared to controls. (C) Flow cytometric gating strategy to analyze the basophil population in unfractionated human blood samples. Together with analysis of the physical parameters two basophil-identifying markers were included in the flow cytometry analysis: CD63, an activation marker and CCR3 a basophil cell subset-specific surface molecule. (D) Representative flow cytometry plots showing the gating strategy to detect basophil activation (conditions tested included PB=patient baseline, anti-FcεRI, fMLP, anti-IgE).
Figure S7. Related to Figure 7G - SF-25 based CAR T cell in vivo toxicity.
Individual weight curves for mice in PBS treated (top), truncated CAR T cells (middle) and SF-25 second generation CAR T cells (bottom panel). The two long-term remission mice are highlighted with round and square symbols on the curve.