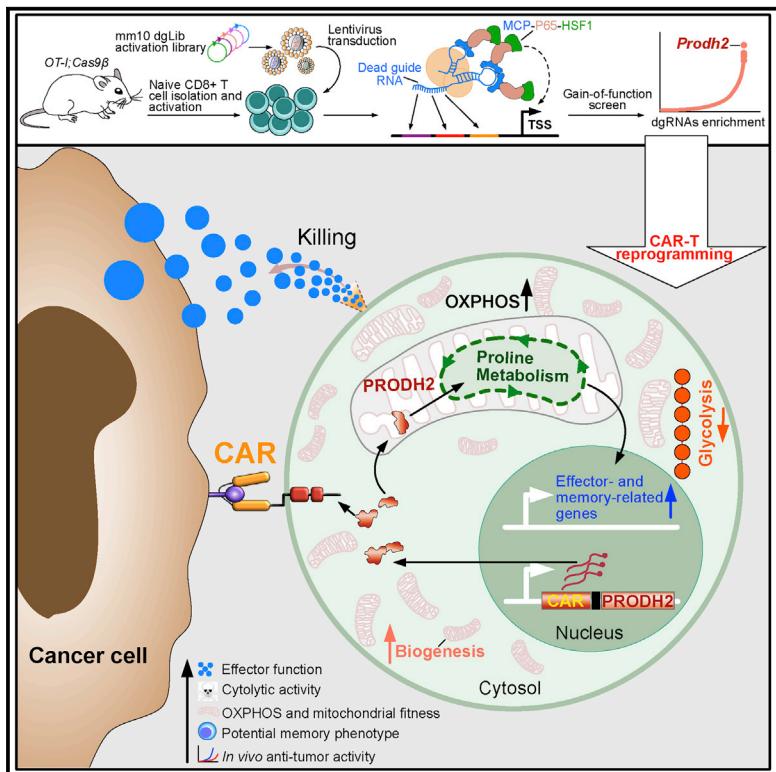


A genome-scale gain-of-function CRISPR screen in CD8 T cells identifies proline metabolism as a means to enhance CAR-T therapy

Graphical abstract



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In brief

Ye et al. develop and apply a dead-guide RNA-based genome-wide gain-of-function CRISPR activation screen, which identifies *PRODH2* as an enzyme that reprograms proline metabolism in CAR-T cells to enhance antitumor efficacy.

Highlights

- Dead-guide RNA-based CRISPR activation screen identifies targets for CAR-T engineering
- *PRODH2* gain of function reprograms proline metabolism in CD8⁺ T cells
- *PRODH2* reshapes transcriptome, metabolome, and immune functions in CAR-T cells
- *PRODH2* reprogramming enhances CAR-T cell therapy in both *in vitro* and *in vivo* settings

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A genome-scale gain-of-function CRISPR screen in CD8 T cells identifies proline metabolism as a means to enhance CAR-T therapy

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SUMMARY

Chimeric antigen receptor (CAR)-T cell-based immunotherapy for cancer and immunological diseases has made great strides, but it still faces multiple hurdles. Finding the right molecular targets to engineer T cells toward a desired function has broad implications for the armamentarium of T cell-centered therapies. Here, we developed a dead-guide RNA (dgrRNA)-based CRISPR activation screen in primary CD8⁺ T cells and identified gain-of-function (GOF) targets for CAR-T engineering. Targeted knockin or overexpression of a lead target, *PRODH2*, enhanced CAR-T-based killing and *in vivo* efficacy in multiple cancer models. Transcriptomics and metabolomics in CAR-T cells revealed that augmenting *PRODH2* expression reshaped broad and distinct gene expression and metabolic programs. Mitochondrial, metabolic, and immunological analyses showed that *PRODH2* engineering enhances the metabolic and immune functions of CAR-T cells against cancer. Together, these findings provide a system for identification of GOF immune boosters and demonstrate *PRODH2* as a target to enhance CAR-T efficacy.

INTRODUCTION

T cells are the cornerstone of adaptive immunity and therefore key players in maintaining human health. Over the past two de-

cades, an armamentarium of powerful therapies have been developed centering on T cells or T cell-mediated immunological pathways, several of which have been approved for clinical use or are in active clinical trials (Tang et al., 2018). These include

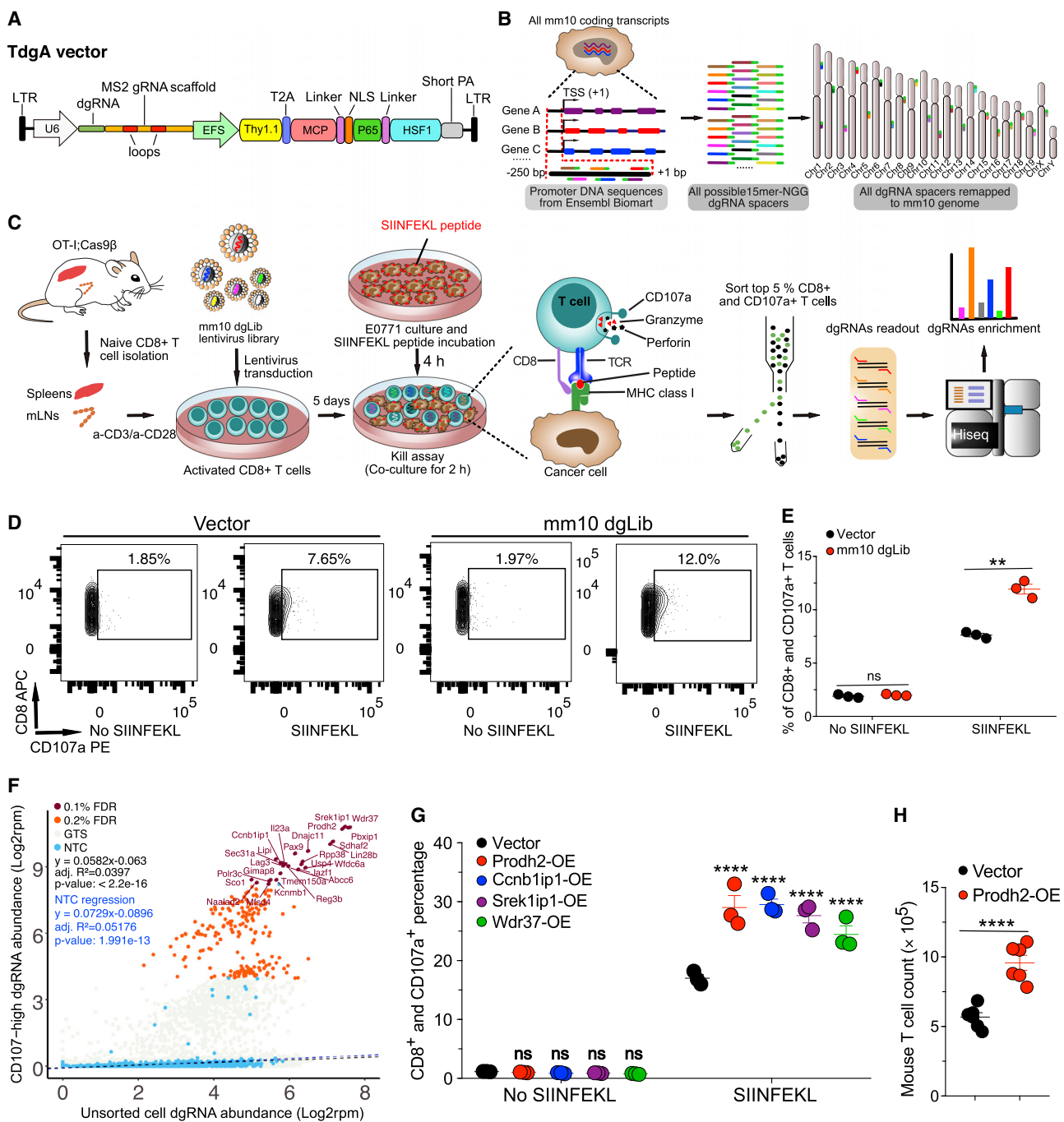


Figure 1. Genome-scale dgRNA activation screen identified genes that boost the effector function of CD8⁺ T cells

(A) Schematic representation of a T cell dead-guide RNA (dgRNA) activation lentiviral vector (TdgA), which contains a human U6 promoter, a dgRNA scaffold, and Thy1.1-MPH (MCP-p65-HSF1) expression cassette driven by an EFS promoter.

(B) Schematics of experiment: mouse genome-scale dgRNA library (mm10dgLib) design, including 15-nt proximal promoter spacer identification, on-target and off-target mapping, scoring, filtering, and prioritization of final spacers (details in STAR Methods).

(C) Schematics of genome-scale dgRNA-library-based mouse primary CD8⁺ T cell kill assay activation screen (dgTKS) to identify genes that boost effector functions of CD8⁺ T cells. The main procedure includes naive CD8⁺ T cell isolation, mm10dgLib transduction, a kill assay (CD8⁺ T cell degranulation, as measured by CD107a level in CD8 T cells in a T cell:cancer cell coculture), CD8⁺;CD107a⁺ T cell sorting, genomic DNA preparation, dgRNA library readout, and dgRNA enrichment.

(D) Representative flow cytometry results of the kill assay in the dgTKS experiment. FACS gating plot showing the percentage of CD107a⁺ cells among all CD8⁺ cells in vector and mm10dgLib transduced CD8⁺ T cells coculture with E0771 cancer cells pulsed with SIINFEKL peptide (n = 3 biological replicates). Representative data from two independent experiments.

(E) Quantification of CD107a in the mm10dgLib screen (n = 3 biological replicates).

(legend continued on next page)

Cell Metabolism

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immune checkpoint blockade that targets immunological synapses between cancer cells and T cells (Herbst et al., 2018; Pardoll, 2012; Ribas, 2012), agonist antibodies or small molecules that augment T cell functions (Moran et al., 2013) or enhance production of T cell-secreted or T cell-modulating cytokines and chemokines (Silva et al., 2019), neoantigen cancer vaccines based on T cell recognition of peptide-major histocompatibility complexes (MHCs) (Hollingsworth and Jansen, 2019), direct adoptive transfer of tumor infiltrating lymphocytes (TILs) (Rosenberg and Restifo, 2015), and chimeric antigen receptor (CAR)-T cells (June et al., 2018; Yu et al., 2019). Therefore, engineering T cells toward a desired function has a broad range of applications with therapeutic implications.

CAR-T cell-based immunotherapy is a revolutionary approach to treat cancer and is potentially applicable to various other diseases (June et al., 2018; Majzner and Mackall, 2019; Rosenberg and Restifo, 2015). Five CAR-T products have been approved by the US FDA to date for the treatment of B cell malignancies or multiple myeloma. However, current CAR-T therapies still face several major challenges leading to primary or secondary resistance and relapse of disease in hematologic cancers. These challenges are due to a variety of reasons, including loss of antigen, failure of target recognition, cancer immune escape, or insufficient persistence (June et al., 2018; Sharma et al., 2017). Therefore, it is important to identify genetic targets that can boost effector function in CD8⁺ T cells and to directly harness such factors to engineer more effective T cells for cell-based therapy, including CAR-Ts.

To date, targets for T cell engineering predominantly rely on the immunology literature, leveraging endogenous genes, such as *TRAC*, *TET2*, and *NOTCH/DELTA*, that provide significant enhancement of T cell function (reviewed in Brown and Mackall, 2019; Roybal and Lim, 2017; Sadelain et al., 2017). Loss-of-function (LOF) screens enable high-throughput identification of essential genes of T cell function using RNA interference (RNAi) (Chen et al., 2014; Zhou et al., 2014) or CRISPR-Cas9-mediated genetic knockout (Henriksson et al., 2019; Shifrut et al., 2018). In contrast, gain-of-function (GOF) screens can directly identify functional boosters that can be harnessed for T cell programming. This makes it possible to identify specific genes, which, regardless of their original physiological function, when overexpressed, knocked in, or exogenously supplied can augment the function of immune cells.

GOF screening directly in primary T cells has remained challenging thus far. This is in part due to the difficulty of introducing three separate components (RNA-guided nucleases and trans-activator and guide RNAs) simultaneously into primary T cells to achieve CRISPR activation (CRISPRa). This is important because unlike previously reported T cell CRISPR knockout

screens (Henriksson et al., 2019; LaFleur et al., 2019; Shifrut et al., 2018; Ting et al., 2018; Dong et al., 2019; Ye et al., 2019), CRISPRa screens, being a GOF approach, can identify a new class of targets that can be harnessed as functional boosters for T cell reprogramming. Here, we designed a genome-scale catalytically dead-guide RNA (dgRNA) library as a versatile CRISPRa screening tool for high-throughput identification of GOF targets in any catalytically active Cas9-expressing cells, which is particularly useful for applications in primary immune cells that are less amenable to viral transduction and genetic manipulation. Utilizing this system, we identified and subsequently validated genes that can directly augment the effector function of CD8⁺ T cells. With target-centered interrogations, we showed that GOF engineering of a top hit, *proline dehydrogenase 2* (*PRODH2*/*Prodh2*), into primary or CAR-T cells can reshape their metabolic pathways and distinct gene expression programs, significantly improve their functions, and enhance their antitumor efficacy *in vivo*.

RESULTS

Genome-scale identification of boosters of effector function for primary CD8⁺ T cells

Due to the challenges of introducing multiple components including Cas9, coactivators, and guide RNAs into primary T cells, we utilized a dgRNA system that is compatible with active Cas9 (Dahlman et al., 2015; Liao et al., 2017), whereby the immune cells can be readily isolated from Cas9 transgenic mice (Platt et al., 2014). We first designed and constructed a lentiviral T cell dgRNA activation (TdgA) vector (Figure 1A). We then designed a mouse genome-scale dgRNA library (mm10dgLib) using the promoter sequences of all annotated protein-coding transcripts from the mm10 genome assembly (STAR Methods; Figures 1B and S1A). After spacer identification, on-target and off-target mapping and filtering, proximal score ranking, and spacer per gene choices for library balancing, the final mm10dgLib consists of 84,601 dgRNAs that target 22,391 coding transcripts and 1,000 nontargeting controls (NTCs) (STAR Methods; Figures 1B and S1A; Data S1), which was synthesized as a pool and cloned into the TdgA vector. We sequenced the mm10dgLib plasmid library and verified that 82,197/83,601 (98.3%) of gene-targeting spacers (GTSs) and 988/1,000 (98.8%) of NTCs were successfully cloned, and both GTSs and NTCs showed a log-normal distribution (Figures S1B and S1C). We then packaged the plasmid library into lentiviral delivery system and performed functional titration by flow cytometry to confirm adequate viral titer (Figures S1D and S1E), enabling genome-scale activation screens for primary cells expressing catalytically active Cas9.

(F) Bulk screen scatterplot of dgTKS screen, showing relative dgRNA abundances in the entire mm10dgLib library, with CD107a⁺-high FACS sorted CD8⁺ T cells, as compared with unsorted T cells. Blue dots are NTCs; brown dots are scoring GTSs that passed FDR 0.1% cutoff, with gene name labeled; orange dots are scoring GTSs that passed FDR 0.2% cutoff but did not pass FDR 0.1%; and gray dots are remaining GTSs. Black dashed line is a regression line of all data points. Blue dashed line is a regression line of 1,000 NTCs representing a neutral baseline. Regression parameters and p values were shown. GTSs deviating from the baseline showed enrichment in the CD107a⁺-high FACS as compared with the behavior of NTCs. The points were shown at the individual gRNA level. Representative top scoring genes targeted by specific sgRNAs were shown.

(G) Quantitative analysis of flow cytometry for kill assay for individual genes overexpressed by lentiviral vectors.

(H) Mouse T cell number quantification at day 4 after IL-2 withdrawal (n = 6 in total of one independent experiments).

p < 0.01, *p < 0.0001 by multiple t tests (with adjusted p value) (E and G) or unpaired t tests (H). See also Figure S1.

Degranulation is one of the major mechanisms through which cytotoxic CD8⁺ T lymphocytes (CTLs) mediate the killing of target cells (Trapani and Smyth, 2002). CD107a (also known as LAMP-1) is a marker that can be presented on the cell surface after degranulation (Peters et al., 1991). To identify genes that when activated can enhance the degranulation ability of CD8⁺ T cells after encountering their cognate antigen presented on the cell surface, we devised and performed a genome-scale dgRNA library-based CD8⁺ T cell kill assay activation screen (dgTKS) (Figure 1C). To do this, we developed a coculture system (i.e., a kill assay), in which OT-I;Cas9β CD8⁺ T cells sensitively respond to E0771 breast cancer cells presenting SIINFEKL peptide, the cognate antigen of the CD8⁺ T cells from OT-I transgenic mice (Figure 1C). In this system, we measured CD107a⁺ expression among CD8⁺ T cells cocultured with E0771 cancer cells with or without SIINFEKL peptide pulsing and found that mm10dgLib lentivirus-transduced CD8⁺ T cells had significantly higher CD107a⁺ levels compared with empty vector-transduced cells ($p < 0.001$) (Figures 1D and 1E). Deep-sequencing data showed that the cumulative coverage in each independent experiment was between 93.3% and 98.2% (Figure S1F; Data S1). Using fluorescence-activated cell sorting (FACS), we sorted the mm10dgLib-transduced CD8⁺ T cells expressing a high level (top 5%) of CD107a, in three independent biological replicates, for genomic DNA preparation and dgRNA library readout (STAR Methods; Figure 1C). We then used Illumina sequencing to read the dgRNA cassette of both CD107a⁺-high sorted and unsorted CD8⁺ T cells, and we quantified the dgRNA abundance in the entire mm10dgLib (Data S1). As a neutral baseline, we found that the NTCs were relatively evenly distributed in the unsorted cell population but were rarely detected in the CD107a⁺-high CD8⁺ T cells (Figure 1F). In contrast, there were two distinct populations of dgRNAs that deviated from the distribution and regression lines of NTCs (Figure 1F). With an FDR of 0.1%, we identified significantly enriched dgRNAs in sorted CD107a⁺-high cells targeting 26 genes, including *Prodh2*, *Srek1ip1*, *Wdr37*, *Ccnb1ip1*, *Pbxip1*, and *Sdhaf2* (Figure 1F). To determine how a theoretically neutral cell population would behave, we utilized the 1,000 NTCs in the pool and found that they largely follow a log-linear relation. The regression is statistically significant, showing the baseline dynamics without genetic effect as theoretically random factors alone for NTCs as a population (Figure 1F). A regression using all data points generated a similar curve (Figure 1F). An observed strong shift in a substantial population of dgRNAs from the theoretical neutral regression line suggested that there are a population of dgRNAs as hits under potential selection (Figure 1F). To measure whether and how much each gene-targeting dgRNA deviates from the theoretically neutral line (gene perturbation effect, representative of potential selection), we used an outlier test to calculate the degree of shift and statistical significance and identified hits such as *Prodh2*, *Srek1ip1*, *Wdr37*, *Pbxip1*, *Sdhaf2*, *Lin28b*, *Pax9*, *Dnajc11*, *Il23a*, *Ccnb1ip1*, and *Wdr37* (Figure S1G). Another approach based on direct comparison of the mean difference of dgRNA abundance between sorted and unsorted populations uncovered similar hits including *Prodh2*, *Ccnb1ip1*, *Srek1ip1*, and *Wdr37* (Figure S1H). This screen revealed an unbiased picture of GOF effects of endogenous genes on CD8⁺ T cell degranulation and provided a ranked list of potential targets for T cell engineering.

Hits identified from the GOF screen enhance CD8⁺ T cell effector function

Based on the enrichment in the GOF screen, we hypothesized that increased expression of high-rank candidate genes from the dgTKS screen might enhance the degranulation ability of CD8⁺ T effector cells. Using qRT-PCR, we measured the natural expression levels for several high-rank candidate genes, including *Defb19*, *Schip1*, *Sox5*, *Il1rap*, *Srek1ip1*, *Wdr37*, *Sdhaf2*, *Lin28b*, *Prodh2*, *Ccnb1ip1*, and *Bdnf*. Results showed that most of these genes have low baseline levels of mRNA expression compared with the housekeeping gene *Gapdh* in primary CD8⁺ T cells (Figure S1I). The baseline expression of each of these genes is physiologically low in T cells, leaving sufficient room for gene overexpression in GOF T cell engineering. After confirming overexpression by qRT-PCR (Figures S1J and S1K), we then performed validation kill assays 4 days after lentiviral transduction of cDNA for overexpressing four top hits (*Prodh2*, *Ccnb1ip1*, *Srek1ip1*, or *Wdr37*) and found that activation of each significantly enhanced degranulation by CD8⁺ T cells upon encountering SIINFEKL-pulsed cancer cells (Figure 1G). These data validated that augmented gene expression of *Prodh2*, *Ccnb1ip1*, *Srek1ip1*, or *Wdr37* can enhance CD8⁺ T cell effector function, providing GOF candidates for engineering improved T cell capabilities.

Interestingly, among all targets, we noticed that *Prodh2*-OE T cells proliferated rapidly in culture as compared with non-transduced or vector-transduced control T cells. Under the IL-2 withdrawal condition, quantification of cell numbers showed that the bulk *Prodh2*-OE CD8⁺ T cells proliferate 1.8 times faster than the vector-transduced counterparts (Figure 1H). In addition, *Prodh2*-OE CD8⁺ T cells have lower levels of apoptosis (Figure S1L).

GOF *PRODH2* engineering by genomic knockin or lentiviral overexpression boosts CAR-T killing of cancer cells

To harness PRODH2's function to program T cells for cell therapy, we set out to co-engineer *PRODH2* and CAR together in human T cells. We generated an anti-CD22 CAR (CD22-CAR) AAV construct specifically targeted to the *TRAC* locus, with simultaneous knockin of the *PRODH2* transgene (AAV-CD22-CAR-T2A-PRODH2), to allow expression of CD22-CAR and PRODH2 in the same T cells (Figure 2A). In parallel, for functional studies—because untargeted primary cells or CAR-T cells without any other transgene are imperfect controls—we generated a matched control with a prematurely stopped *PRODH2* CDS (AAV-CD22-CAR-T2A-PRODH2(Stop)) (STAR Methods; Figure 2A). With electroporation of Cas9 ribonucleoprotein (RNP) targeting *TRAC* exon 1 followed by transduction of the AAV constructs, we introduced these transgenes into the endogenous *TRAC* locus of primary CD8 T cells to generate CD22-CAR;PRODH2 stable knockin CAR-T cells as well as the CD22-CAR;PRODH2(Stop) control knockin CAR-T cells (STAR Methods; Figure 2A). Flow cytometry analysis of both AAV knockin constructs in the CD8⁺ T cells showed highly efficient *TRAC* editing by Cas9 RNP, as evident by a population of cells with CAR⁺ and CD3[−] expression (as CD3 and TCR form a functional complex) (Figure 2B). We used FACS to purify TCR[−]CAR⁺ T cells (Figure 2B).

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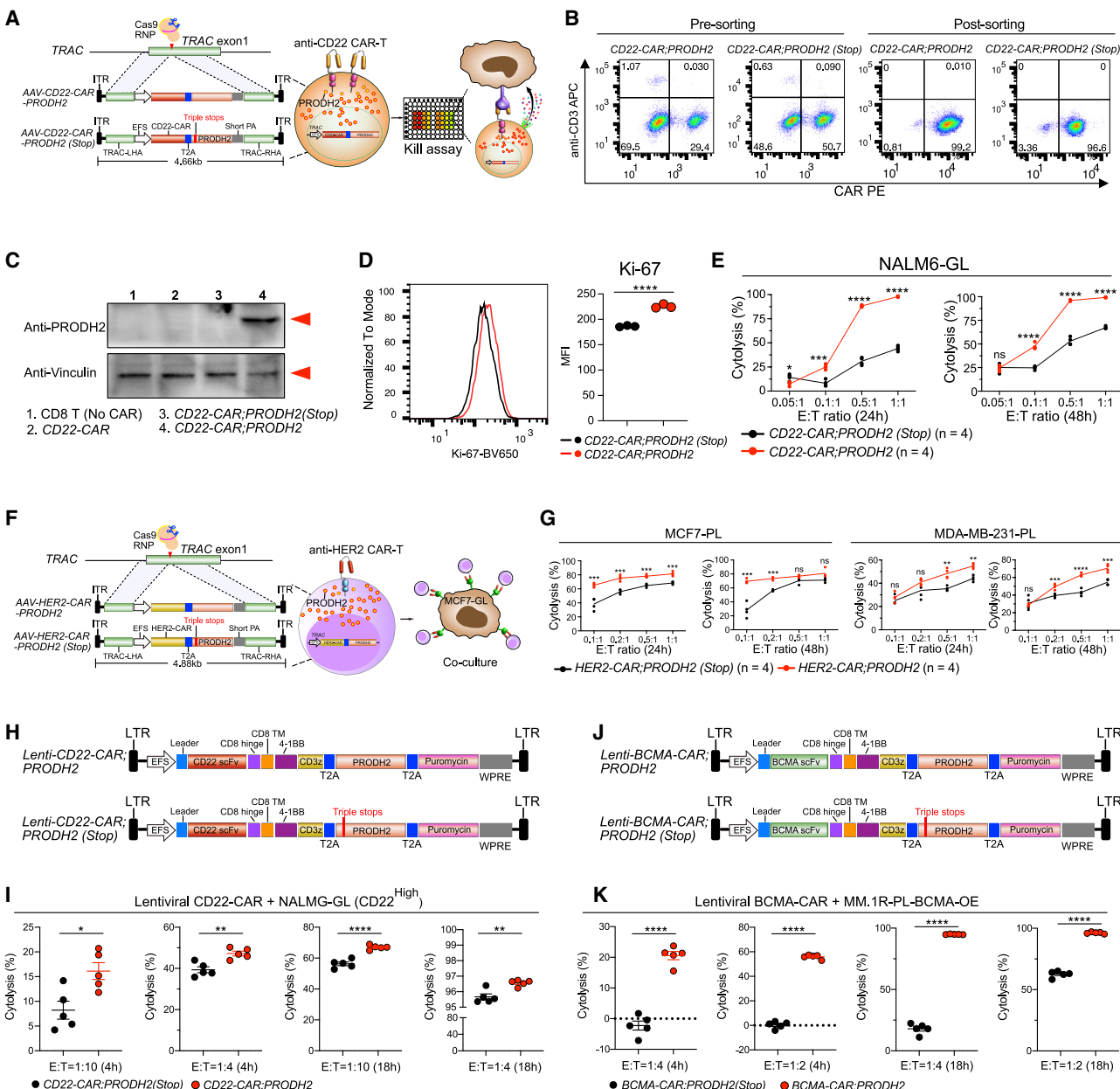


Figure 2. PRODH2 engineering by genomic knockin or lentiviral overexpression boosts cytotoxic activity of CAR-Ts against cognate cancer cells

(A) A schematic of human CD22-CAR;PRODH2 (PRODH2 KI CAR-T) and CD22-CAR;PRODH2(Stop) (PRODH2(Stop) KI CAR-T, control CAR-T) cell generation. In the CD22-CAR;PRODH2(Stop) construct, three artificial premature stop codons were inserted between the 318 and 319 bp position of the PRODH2 CDS to generate a truncated mutant version. Knockin constructs consisting of TRAC locus homology-directed repair (HDR) 5' and 3' arms, an EFS promoter, a CD22-CAR expression cassette, a T2A sequence, a PRODH2 or PRODH2(Stop) CDS, and a short polyA. AAV6-packaged KI constructs were introduced into T cells by viral transduction after TRAC first-exon targeting Cas9:crRNA RNP electroporation.

(B) Representative flow cytometry plots of PRODH2 knockin and control CAR-T cell proliferation by Ki-67 staining.

(C) Kill assay of purified PRODH2 knockin and control CAR-T cells with NALM6-GL (NALM6 with GFP and luciferase reporters) cancer cells, with a titration series of effector:target (E:T) ratios, and at two time points (24 and 48 h). The time point of CAR-T cells used for coculture was day 67 after CAR knockin. Individual replicate data points were shown ($n = 4$ biological replicates). Representative data from two independent experiments.

(D) Flow analysis of PRODH2 knockin and control CAR-T cell proliferation by Ki-67 staining.

(E) Kill assay of purified PRODH2 knockin and control CAR-T cells with NALM6-GL (NALM6 with GFP and luciferase reporters) cancer cells, with a titration series of effector:target (E:T) ratios, and at two time points (24 and 48 h). The time point of CAR-T cells used for coculture was day 67 after CAR knockin. Individual replicate data points were shown ($n = 4$ biological replicates). Representative data from two independent experiments.

(F) A schematic of human HER2-CAR;PRODH2 and HER2-CAR;PRODH2(Stop) CD8 T cell generation. HER2-CAR;PRODH2 and HER2-CAR;PRODH2(Stop) constructs were established by replacing CD22-CAR with HER2-CAR construct in CD22-CAR;PRODH2 and CD22-CAR;PRODH2(Stop) constructs.

We then performed immunoblotting to measure the baseline expression of PRODH2 protein in PRODH2-knockin CAR-T cells, along with three control T cells (CD8 T cell without CAR, CD22 CAR-T cell without additional transgene, and PRODH2(Stop)-knockin CD22 CAR-T cell). The results demonstrated that PRODH2 is highly expressed in *CD22-CAR;PRODH2* T cells but is undetectable in CD8 T cells (no CAR), *CD22-CAR*, and *CD22-CAR;PRODH2(Stop)* T cells (Figure 2C), which showed minimal baseline expression and excluded the possibility of PRODH2(Stop) transgene leakage. We analyzed CAR-T cell proliferation by flow cytometry of Ki-67 and found that PRODH2-KI (knockin) CD22 CAR-T cells had higher levels of Ki-67 (Figure 2D). We then performed a coculture assay to test the ability of CAR-T cells to kill cognate leukemic cells, NALM6-GL (GFP and luciferase positive). As compared with the *CD22-CAR;PRODH2(Stop)* control, *CD22-CAR;PRODH2* CAR-T cells have a significantly stronger ability to kill the NALM6-GL in an antigen-specific manner (Figure 2E). Similarly, we engineered a HER2-specific CAR with PRODH2 overexpression along with matched control (*HER2-CAR;PRODH2* and *HER2-CAR;PRODH2(Stop)*) by knocking the AAV constructs into the TRAC locus (Figure 2F). Coculture assays also showed that PRODH2-KI HER2 CAR-T cells had a stronger cytolytic activity against MCF7-PL (puromycin and luciferase positive) and MDA-MB-231-PL breast cancer cells (Figure 2G).

In addition, in order to test if PRODH2 can be engineered in the traditional lentiviral CAR-T system, we also generated lentiviral *CD22-CAR* and *BCMA-CAR* with cisticronic overexpression of PRODH2 or PRODH2(Stop) constructs (Figures 2H and 2J). The coculture data again showed PRODH2 overexpressed *CD22-CAR* and *BCMA-CAR* T cells significantly enhanced cancer cell killing (Figures 2I and 2K), where the cytotoxicity enhancement effect of PRODH2 in BCMA-CAR is particularly strong (Figure 2K). These data together suggest that *PRODH2* GOF engineering via either genomic knockin or lentiviral overexpression enhanced *in vitro* killing ability of antigen-specific CAR-T cells, in cancer-specific CARs in three cellular models (leukemia, multiple myeloma, and breast cancer).

PRODH2 GOF engineering enhances CAR-T *in vivo* efficacy against cancer in animal models

We then asked if PRODH2 GOF can enhance CAR-T cells' therapeutic efficacy in *in vivo* settings. We first tested a B cell leukemia model with CD22 as the cancer antigen (Figure 3A). The bioluminescence imaging data showed that control *CD22-CAR* T cells had antitumor activity, but the leukemia relapsed quickly ($p < 0.0001$) (Figure 3B), whereas *PRODH2* knockin *CD22-CAR* T cells (*CD22-CAR;PRODH2*) showed significantly

stronger antitumor activity compared with control (*CD22-CAR;PRODH2(Stop)*) T cells and can more effectively control leukemia progression, in terms of both tumor burden ($p < 0.0001$) and survival ($p < 0.05$) (Figures 3B, 3C, and S1M). We also tested PRODH2-OE CAR-T in a solid tumor model (anti-HER2-CAR), where breast tumors were induced by an intramammary fat-pad injection of an established human breast cancer line MCF7-PL-HER2OE (Figure S1N). Tumor growth kinetics showed that control HER2-CAR T cells had a small antitumor effect, whereas PRODH2-OE CAR-T cells (*HER2-CAR;PRODH2*) had a significant enhancement of efficacy compared with the control CAR-Ts ($p < 0.0001$), although all groups had tumor growth potentially due to the challenges in the solid tumor microenvironment (Figure S1O).

Because the effect of PRODH2 on BCMA CAR-T is most pronounced *in vitro* (Figure 2), we focused the subsequent *in vivo* experiments on PRODH2-engineered BCMA CAR-Ts. Using two independent CAR-T platforms (AAV-KI and lenti-OE), we established with *PRODH2* CAR-T and *PRODH2(Stop)* control CAR-T and tested them in parallel against a systemic multiple myeloma model (Figure 3D). We performed three different experiments, one with an AAV PRODH2-KI BCMA-CAR against an MM.1R multiple myeloma model, the second also with AAV PRODH2-KI BCMA-CAR against a BCMA-OE MM.1R multiple myeloma model, and the third using a lentiviral PRODH2-OE BCMA-CAR (Figure 3D). The MM.1R cells when injected into NSG mice develop into multiple myeloma as confirmed by histology (Figure S1P). Overall survival analysis showed that control AAV-KI BCMA CAR-T cells slightly extended animal overall survival; however, all animals still succumbed to disease (Figure 3E). AAV-KI PRODH2-OE CAR-T cells (*BCMA-CAR;PRODH2*) had a significantly stronger *in vivo* therapeutic effect and significantly enhanced the overall survival (median survival time, PBS, 51 days; control CAR, 63 days; PRODH2-OE CAR, not reached) ($p < 0.05$), with more than half of the animals surviving longer term (Figure 3E). We repeated the experiment in a second model with BCMA-OE MM.1R cells. The treatment effect overall is better as the animal survival is longer in the setting of antigen overexpression (Figure 3F). Again, we observed that although control CAR-Ts slightly extend the overall survival of diseased animals, AAV-KI PRODH2-OE CAR-T cells had a significantly stronger efficacy and significantly enhanced the overall survival ($p < 0.01$), with $\geq 50\%$ of the animals surviving longer term (Figure 3F).

Because turnaround time is important for CAR-T production, we also tested the *in vivo* efficacy of PRODH2-engineered CAR-Ts in a third model, using the traditional lentiviral CAR-T system that requires shorter production time. The *in vivo* tumor

(G) Kill assay of *HER2-CAR;PRODH2* and *HER2-CAR;PRODH2(Stop)* knockin T cells with MCF7-PL and MDA-MB-231-PL (MCF7 and MDA-MB-231 cells expressed with puromycin and luciferase reporters) breast cancer cells, with a titration series of effector:target (E:T) ratios, and at two time points (24 and 48 h). All CAR-T cells were used for coculture at day 17 after CAR knockin. Individual replicate datapoints were shown ($n = 4$ biological replicates).

(H) A schematic of lentiviral *CD22-CAR;PRODH2* (PRODH2-OE CAR-T) and *CD22-CAR;PRODH2(Stop)* (control CAR-T) cell generation.

(I) Kill assay of *Lenti-CD22-CAR;PRODH2* (*Stop*) and *Lenti-CD22-CAR;PRODH2* T cells with NALM6-GL (*CD22^{high}*) cancer cells at day 3 after lentiviral transduction ($n = 5$ biological replicates).

(J) A schematic of lentiviral *BCMA-CAR;PRODH2* (PRODH2-OE BCMA CAR-T) and *BCMA-CAR;PRODH2(Stop)* (Control BCMA CAR-T) cell generation.

(K) Kill assay of *Lenti-BCMA-CAR;PRODH2* (*Stop*) and *Lenti-BCMA-CAR;PRODH2* T cells with MM.1R-PL-BCMA-OE cancer cells at day 3 after lentiviral transduction ($n = 5$ biological replicates).

* $p < 0.05$, ** $p < 0.01$, *** $p < 0.001$, **** $p < 0.0001$; ns, not significant by multiple t tests (with adjusted p value) (E and G) or unpaired t tests (D, I, and K). See also Figure S1.

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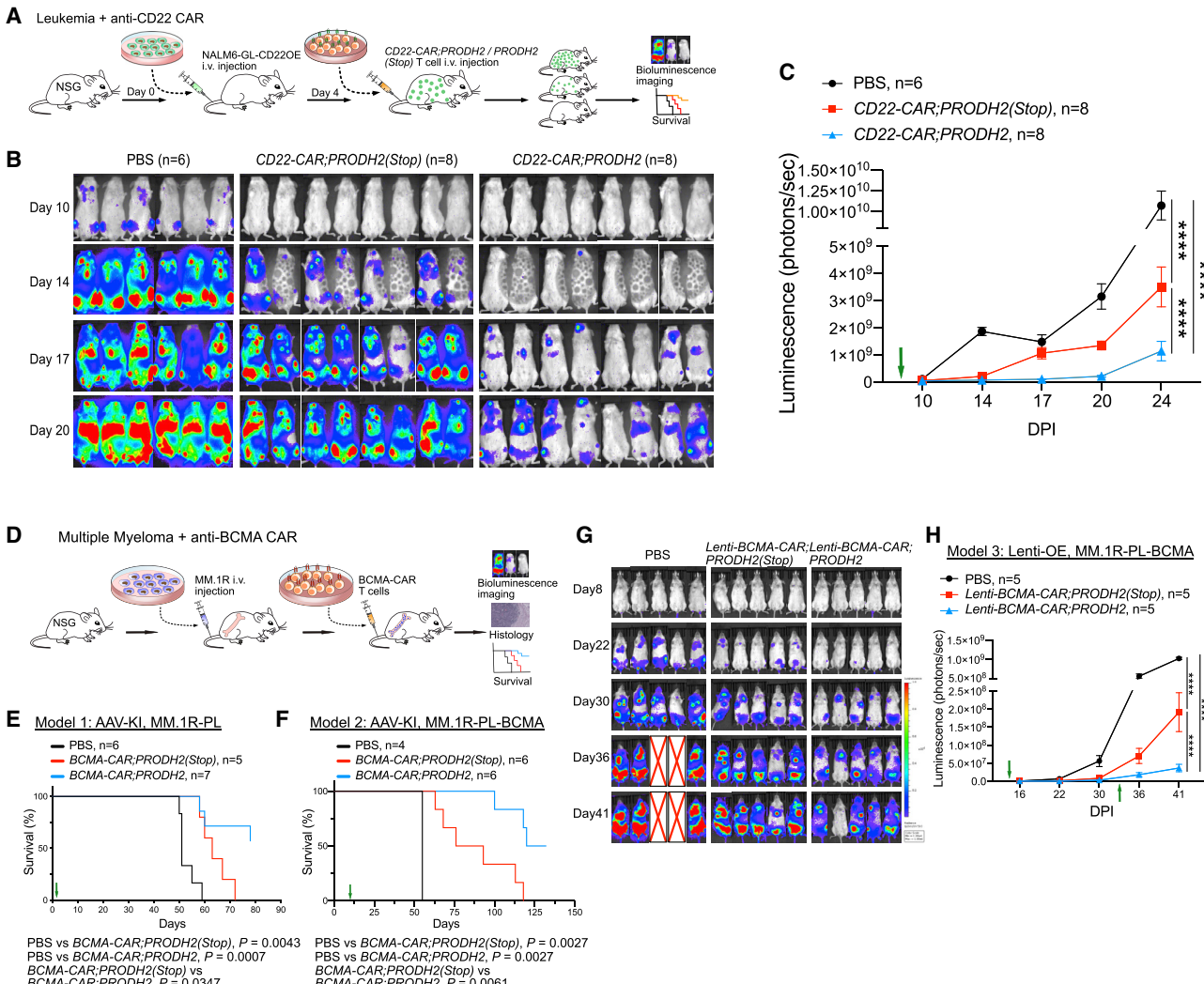


Figure 3. PRODH2 engineering by genomic knockin or lentiviral overexpression enhanced CAR-T in vivo efficacy in mouse models

(A) A schematic of the experimental design of leukemia model PRODH2 knockin CAR-T efficacy testing, showing leukemia induction, CD22-CAR intravenous injection, survival, and bioluminescence imaging. CAR-T cells were prestimulated with NALM6-GL-CD22OE cancer cells at an E:T ratio = 1:1 at day 25 before injection.

(B) IVIS imaging showing bioluminescence of NSG mice that were injected with NALM6-GL-CD22OE cancer cells and with CD22-CAR therapy. Note: the dark shadow on the mouse at day 14 was induced by the imaging machine but has no influence on bioluminescence quantification.

(C) Quantification of cancer burden by total luminescence. Green arrow indicated that CAR-T injection was performed at day 4 (n = 6–8 mice/group).

(D) A schematic of the experimental design of PRODH2 CAR-T efficacy testing in multiple myeloma models, showing induction, BCMA-CAR intravenous injection, and survival.

(E) Survival curve of MM.1R-induced myeloma-bearing NSG mice after AAV-KI BCMA-CAR;PRODH2 or BCMA-CAR;PRODH2 (Stop) T cell adoptive transfer therapy (CAR-T adoptive transfer indicated with a green arrow) (n = 5–7 mice/group).

(F) Survival curve of BCMA-OE MM.1R-induced myeloma-bearing NSG mice after AAV-KI BCMA-CAR;PRODH2 or BCMA-CAR;PRODH2 (Stop) T cell adoptive transfer therapy (CAR-T adoptive transfer indicated with a green arrow) (n = 4–6 mice/group).

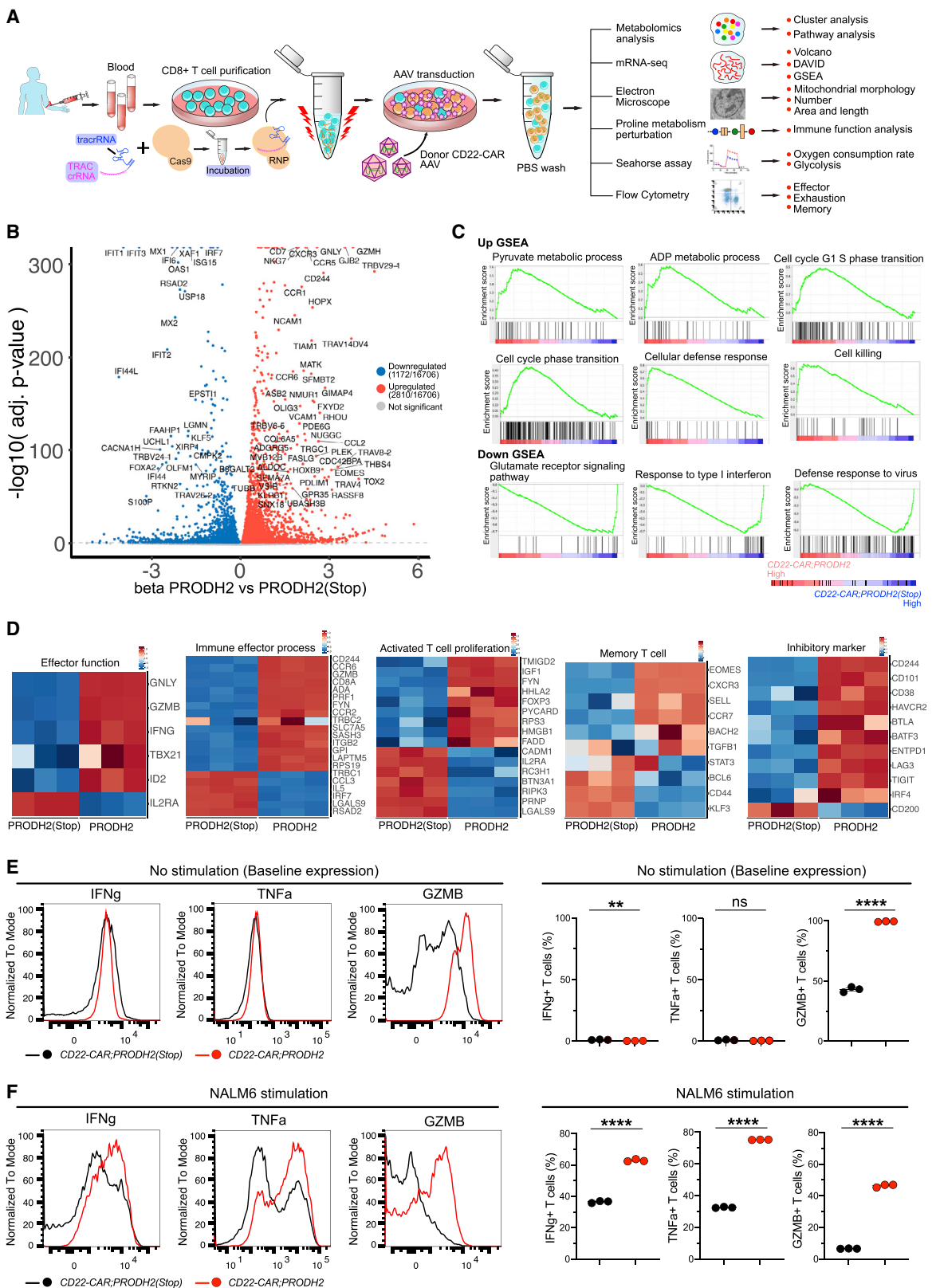
(G) IVIS imaging showing bioluminescence of multiple myeloma-bearing NSG mice after lentiviral-based BCMA-CAR;PRODH2 or BCMA-CAR;PRODH2 (Stop) T cell adoptive transfer therapy. “X” represents dead or euthanized animals (end point) (n = 5 mice/group).

(H) Quantification of cancer burden by total luminescence for (G). CAR-T injection indicated with green arrows.

*p < 0.05, **p < 0.01, ***p < 0.001, ****p < 0.0001 by two-way ANOVA (with multiple comparisons test) (C and H) or log rank (Mantel-Cox) tests (E and F). See also Figure S1.

growth data again showed that although both CAR groups had an antitumor effect (Figures 3G and 3H), PRODH2 overexpressed lentiviral BCMA-CAR T cell adoptive transfer had significantly enhanced efficacy as compared with PRODH2(Stop) control (p < 0.0001) (Figures 3G and 3H). Together, these *in vivo*

data demonstrated that PRODH2 GOF engineering, either by genomic knockin or lentiviral overexpression, significantly enhanced CAR-T cells’ efficacy against cancer in several mouse models, although the effect is more pronounced in the BCMA CAR-T as compared with other models.



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PRODH2 knockin restructures CAR-T cell gene expression and immune effector functions

We then asked why PRODH2 GOF-engineered CAR-T cells showed stronger killing and *in vivo* efficacy. To provide molecular bases on how PRODH2 GOF engineering programs the human CAR-T cells, we performed a series of mechanistic interrogations by multiomics profiling (transcriptome profiling, CyTOF, untargeted metabolomics, targeted metabolomics, and integrated analyses) as well as in-depth analyses (biochemical immunological and cellular assays) (STAR Methods; Figure 4A). To unbiasedly understand the PRODH2-mediated global cellular changes, we first performed mRNA-seq to profile the entire transcriptome of *CD22-CAR;PRODH2* as well as *CD22-CAR;PRODH2(Stop)* knockin CAR-T cells (Data S2). Differentially expressed (DE) analysis again uncovered broad transcriptomic changes as a result of PRODH2 knockin, with 2,810 upregulated and 1,172 downregulated genes (FDR adjusted p value, $q < 0.001$) (Figures 4B, 4C, S2A, and S2B; Data S2). The knockin of PRODH2 cocristronically with the CAR-T construct again showed strong changes in the transcriptomic programs of cell cycle, metabolism, apoptosis, and immune response (Figures 4B, 4C, S2A, and S2B; Data S2). With regard to immune genes and pathways, PRODH2 knockin CAR-T upregulated genes and processes including immune cell activation, leukocyte-mediated cytotoxicity, and largely proinflammatory signatures—such as T cell activation, signal transduction, and cytokine production (Figures 4B, 4C, S2A, and S2B; Data S2)—whereas the terms are not mutually exclusive because these genes often play important roles in multiple T cell pathways. These data together suggest that PRODH2 altered the gene expression programs of CD22-CAR-T cells centered on cell cycle, T cell activation, and metabolic processes.

A number of highly upregulated genes are directly related to T cell function, such as effector function and immune effector process, activated T cell proliferation, T cell memory, and T cell exhaustion (Figures 4D and S2C). Representative highly upregulated genes in these lines include those well documented in T cell activation (e.g., *CCR3*, *CCR5*, *CCR9*, *CXCR3*, *CXCR4*, *EOMES*, *ADA*, *SIRPG*, and *RHOU*), T cell signaling (e.g., *PLCG2*, *ZAP70*, *FYN*, *LCK*, *JAG2*, *PRKDC*, *PRKG2*, *PIK3CG*, and *PIK3R6*), and/or effector function (e.g., *IFNG*, *TGFBR2*, *GZMB*, *GZMH*, and *GZMK*) (Figures 4B, 4D, and S2C). Knocking out a representative gene implicated in T cell activation (Saoudi et al., 2014), *RHOU*, abolished the effect of PRODH2-mediated enhancement of cytotoxicity (Figure S2D), supporting the validity of the RNA-seq data and differential expression analysis.

To cross-validate the immunological phenotypes of PRODH2 KI CAR-T cells, we first performed an unbiased immune profiling using mass cytometry (CyTOF), with a 19-marker panel that included major T cell lineage and functional state markers. We profiled a total number of 216,501 cells across 6 samples, with 3 biological replicates from purified *CD22-CAR;PRODH2* and *CD22-CAR;PRODH2(Stop)* knockin CAR-T cells, in a baseline state without cancer stimulation (Figure S3A). We randomly sampled 5,000 cells from each sample and quantified the expression of each marker at surface protein level, and we revealed the changes in each marker between control and PRODH2 knockin CAR-T cells (Figure S3B). We found that PRODH2 knockin CAR-T cells have significantly higher levels of IFNg and TNF- α , major effector cytokines for CD8 T cells (Figures S3B and S3C). PRODH2 knockin CAR-T cells also have a significantly higher level of CD134/OX40, a costimulatory marker, as well as TIM-3, an immune checkpoint but also T cell activation marker (Figures S3B and S3C). PRODH2 knockin CAR-T cells have a significantly lower level of cleaved caspase-3, an indicator of apoptosis (Figures S3B and S3C).

Because effector function stood out as one of the strongest signals, we then performed flow cytometry to validate the findings from mRNA-seq and CyTOF. We measured the effector cytokines, such as IFNg, TNF- α , and GZMB, by intracellular staining and flow cytometry analysis of CAR-T cells before and after coculture with cognate cancer cells. Results showed that PRODH2-GOF significantly increased GZMB in CAR-T cells at baseline (Figure 4E) and substantially increased the production of all three cytokines (IFNg, TNF- α , and GZMB) under cancer stimulation (Figure 4F). Cytokine secretion of IL2, IL15, and IL7 measured by ELISA was not different between PRODH2-GOF and control CAR-T cells (Figure S3D). These data together suggest that PRODH2 GOF enhanced T cell effector function.

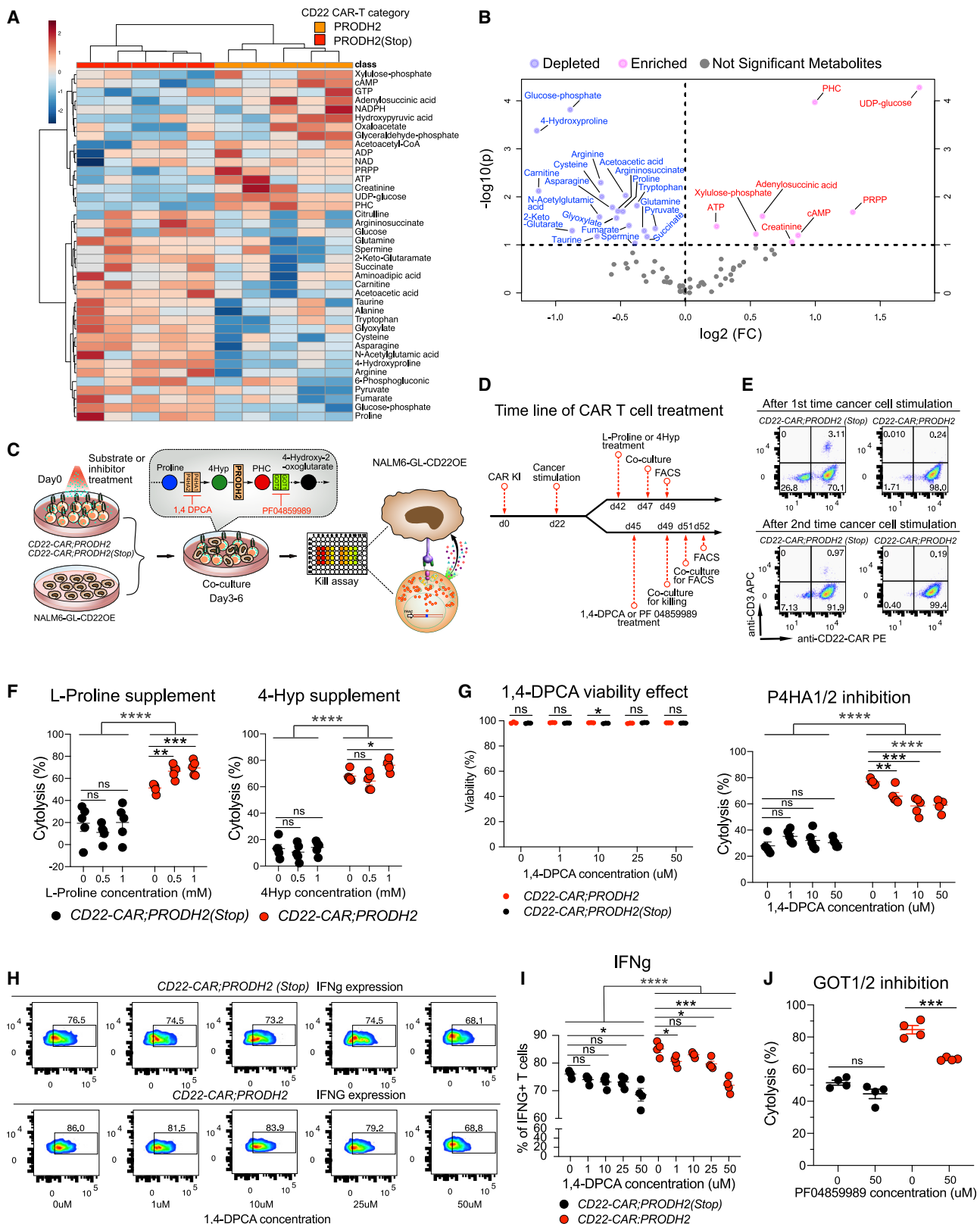
PRODH2 GOF alters metabolism of CD8⁺ T cell and CAR-T cells

PRODH2/Prodh2 encodes an enzyme that catalyzes the conversion of 4-hydroxyproline (IUPAC name: (2S,4R)-4-hydroxypyrrolidine-2-carboxylic acid; aliases: 4Hyp, hydroxyproline, L-hydroxyproline, *trans*-4-hydroxy-L-proline) into 1-pyrroline-3-hydroxy-5-carboxylate (PHC), a key step of the proline metabolic pathway without known redundancy (Phang et al., 2010). This enzyme is expressed at low levels across most organs or cell types in the human body, including primary CD4⁺ and CD8⁺ T cells (Stelzer et al., 2016; Thul et al., 2017; Uhlen et al., 2015). The low baseline expression of PRODH2 makes it a promising candidate to be harnessed for GOF T cell

Figure 4. Whole-transcriptome profiling of PRODH2 knockin CD22 CAR-T cells and FACS validation of enhanced effector function

- (A) A schematic of the experimental design of PRODH2 CAR-T mechanistic investigation: AAV- or lenti-based CAR-T generation, metabolomics, mRNA-seq, metabolic analysis, immunological analysis, transmission electron microscopy (TEM), and other experiments.
- (B) Volcano plots of mRNA-seq differential expression between PRODH2 versus PRODH2(Stop) CD22-CAR-T cells from human donor ($n = 3$ biological replicates) (FDR adjusted $q < 0.001$). CD22-CAR-T cells were collected for RNA extraction and bulk mRNA-seq at day 25 after CAR knockin.
- (C) Gene set enrichment analysis (GSEA) individual pathways for mRNA-seq of PRODH2 knockin CD22-CAR-T cells. GSEA plots of individual pathways from the representative up- and downregulated gene sets between *CD22-CAR;PRODH2* and *CD22-CAR;PRODH2(Stop)* T cell groups (cutoff criteria: $p < 0.001$).
- (D) Heatmaps of differentially expressed genes in representative pathways such as effector function, immune effector process, activated T cell proliferation, memory T cell, and inhibitory marker.
- (E and F) Intracellular staining of effector function markers, IFNg, TNF- α , and GZMB, in both PRODH2 overexpressed CD22 CAR and control CD22 CAR-T cells, without stimulation (E) or after 12 h of NALM6-GL stimulation (F).

** $p < 0.01$, *** $p < 0.0001$; ns, not significant by unpaired t tests (E and F). See also Figure S2.



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Cell Metabolism

Article



manipulation. We therefore next investigated whether augmenting *PRODH2* expression can change the characteristics of human T cells (Figure S4A). We overexpressed human *PRODH2* in human primary CD8⁺ T cells using lentivirus (hPRODH2-OE) and confirmed potent augmentation of gene expression over endogenous level (Figure S4B). Again, we found that the human CD8⁺ T cells with augmented *PRODH2* also proliferate significantly faster than vector control (Figure S4C), similar to the phenotype with mouse T cells reported above. We then performed a targeted metabolomics experiment focused on proline and arginine metabolism (STAR Methods; Figure S4D; Data S3). In hPRODH2-OE CD8⁺ T cells, we observed that the 4-hydroxyproline level was significantly decreased and the PHC level was significantly increased (Figure S4D), suggestive of an increased level of biochemical activity consistent with the augmented gene expression by GOF (on-target metabolic function in T cells). Moreover, hPRODH2-OE CD8⁺ T cells have significantly lower levels of metabolites including oxoproline, ornithine, pyruvate, aspartate, leucine, malate, and oxaloacetate and higher levels of metabolites including choline, glutamate, cysteine, and lactate.

We then performed metabolomics directly in PRODH2 GOF CAR-T cells. Metabolomics data identified a total of 75 metabolites in both *CD22-CAR;PRODH2* and *CD22-CAR;PRODH2(Stop)* knockin CAR-T cells (Figure S5A; Data S3). Among those, 8 metabolites are more abundant in *CD22-CAR;PRODH2* knockin CAR-T cells, whereas 19 metabolites are less abundant (Figures 5A and 5B). As established in the biochemical literature and Kyoto Encyclopedia of Genes and Genomes (KEGG) database, these metabolites have clearly defined metabolic pathways with enzymes that catalyze their production and catabolism (Kanehisa and Goto, 2000). By intersecting the genes encoding those enzymes that produce or break down these metabolites, we observed multiple DE genes whose upregulation or downregulation is consistent with the predicted metabolic activity, i.e., the directions of metabolite alterations (Figures S5A and S5B; Data S3). We again observed on-target activity of PRODH2

GOF, where the upregulation of *PRODH2* is in concert with decreased 4-hydroxyproline and increased PHC levels, along with alteration of other genes and metabolites (Figures 5A, 5B, S5A, and S5B). For example, upregulation of *GAMT* that encodes a guanidinoacetate N-methyltransferase is consistent with the reduction of its substrate arginine and downstream product creatinine; upregulation of glucose-6-phosphatase 3 encoded by the *G6PC3* gene is consistent with reduced glucose-6-phosphatase level, and reduced glutamine level is consistent with downregulation of *GLUL* that produces it from glutamate and upregulation of *CAD* that converts it into carbamoyl-phosphatase (Figures S5A and S5B). Furthermore, multiomics analysis with MetaboAnalyst that integrates mRNA-seq DE analysis and metabolomics data identified enriched metabolic pathways in an unbiased manner, with the note that the significant changes of these metabolic pathways can be in both directions. The second-most enriched metabolic pathway with the DE gene set in *CD22-CAR;PRODH2* knockin CAR-T cells is arginine and proline metabolism, with significantly enriched intersecting DE genes involved in these pathways as well as differentially represented (DR) metabolites (Figures S5C and S5D; Data S3), again highlighting the on-target activity of augmenting PRODH2.

Biochemical-immunology investigation demonstrates the proline metabolic pathway activity of PRODH2-enhanced T cell function

To follow on the observations of unbiased profiling by multiomics, we then investigated the details on how PRODH2 overexpression enhanced CAR-T function. We first tested manipulation of part of the proline metabolic pathway by supplying extra L-proline and 4-hydroxyproline (4Hyp) substrates for P4HA1 and P4HA2 and PRODH2, respectively (Cooper et al., 2008; Luanay et al., 2019; Summitt et al., 2015). We pretreated the CAR-T cells with substrates or inhibitor for 3–6 days depending on the specific experiment, then subjected them to coculture or flow-based immunological assays (Figures 5C and 5D). High-efficiency CAR knockin was confirmed prior to coculture assays

Figure 5. Metabolomic profiling and biochemical-immunological validation of PRODH2 GOF CAR-Ts

- (A) Heatmap of the relative abundance of top 40 QTOF/QQQ detected metabolites of PRODH2 versus PRODH2(Stop) CD22-CAR-T cells ($n = 5$ biological replicates). Representative data from two independent experiments.
- (B) Volcano plot of differentially represented (DR) metabolites between PRODH2 versus PRODH2(Stop) CD22-CAR-T cells. Blue dots indicate decreased metabolites, and pink dots indicate increased metabolites.
- (C) Schematic of biochemical-immunological validation of PRODH2 GOF CAR-Ts, including chemical compound treatment, coculture, and flow cytometry. CAR-T cells were supplied with extra L-proline (substrate for P4HA1 and P4HA2), 4-hydroxyproline (4Hyp, substrates for PRODH2), or 1,4-DPCA (inhibitor of P4HA1 and P4HA2) and PF 04859989 (inhibitor of GOT1 and GOT2) before coculture and flow cytometry analyses.
- (D) Timeline of CAR-T cells treated with L-proline, 4Hyp, 1,4-DPCA, and PF 04859989; coculture; and FACS.
- (E) Representative CAR knockin percentages after 1st and 2nd cancer stimulations. Representative data from two independent experiments.
- (F) Substrate supplement experiment. Cytolytic activity measurement by coculture of *CD22-CAR;PRODH2* and *CD22-CAR;PRODH2(Stop)* T cells with NALM6-GL-CD22OE cancer cells for 6 h after T cells pretreated with different concentration of L-proline and 4Hyp.
- (G) P4HA enzymatic inhibition experiment. Left: DPCA toxicity analysis. Cell viability measurement of *CD22-CAR;PRODH2* and *CD22-CAR;PRODH2(Stop)* T cells after treatment with different concentrations of 1,4-DPCA. Right: cytolytic activity measurement by coculture of *CD22-CAR;PRODH2* and *CD22-CAR;PRODH2(Stop)* T cells with NALM6-GL-CD22OE cancer cells for 12 h after T cells were pretreated with different concentrations of 1,4-DPCA.
- (H) Representative flow plots of IFNg production of *CD22-CAR;PRODH2* and *CD22-CAR;PRODH2(Stop)* T cells after 1,4-DPCA treatment and coculture. T cells were prestimulated with NALM6-GL-CD22OE cancer cells for 8 days, then treated with the 1,4-DPCA inhibitor for 3 days. The co-culture E:T = 0.5:1 ($n = 4$ biological replicates).
- (I) Quantification of (H).
- (J) GOT enzymatic inhibition experiment. Cytolytic activity measurement by coculture of *CD22-CAR;PRODH2* and *CD22-CAR;PRODH2(Stop)* T cells with NALM6-GL-CD22OE cancer cells for 24 h after T cells were pretreated with PF04859989.

* $p < 0.05$, ** $p < 0.01$, *** $p < 0.001$; ns, not significant by multiple t tests (with adjusted p value) (G), two-way ANOVA, and unpaired t tests (F, G, I, and J).

(Figure 5E). As a result, we found that although the baseline (no chemical treatment, PBS only) cytosis of *CD22-CAR;PRODH2* is substantially higher than that of the control, L-proline significantly increased the cancer killing ability only in *CD22-CAR;PRODH2* cells but not in control CAR-T that normally do not express PRODH2 and thereby lack the enzyme (Figure 5F). At 1 mM concentration, 4Hyp also significantly increased the cancer killing ability only in *CD22-CAR;PRODH2* cells (Figure 5F). These data together showed that supplementing a direct substrate of PRODH2 significantly improved cancer killing ability only in the presence of PRODH2 overexpressed CAR-T cells but not in control CAR-T that normally lack the enzyme, which provided direct evidence that the introduced biochemical activity by PRODH2 GOF is responsible for the enhanced cancer cell cytosis.

Furthermore, to restrict 4Hyp production, we leveraged 1,4-DPCA inhibitor to block P4HA1 and P4HA2 functions (Xiong et al., 2014, 2018). Using a similar biochemical-immunology assay, we found that blocking P4HA1/2 by its inhibitor 1,4-DPCA reduced cancer killing ability only in *CD22-CAR;PRODH2* T cells but had no influence on control CAR-T (Figure 5G). The live-dead staining data showed 1,4-DPCA treatment itself has no influence on T cell viability (Figure 5G). Consistent with the flow, RNA-seq, and CyTOF data above, we again found that the baseline IFNg production level was higher in PRODH2-OE CAR-T cells as compared with controls across all conditions (Figures 5H and 5I). In concordance with the metabolic function and cytosis phenotype, at E:T ratio of 0.5:1 where cancer cells can still saturate T cells within the assay period, IFNg level is strongly suppressed by 1,4-DPCA in PRODH2-OE CAR-T cells to a level around control CAR-T cells without treatment, whereas this effect is moderate in control CAR-T cells (Figures 5H and 5I). Because PHC, the metabolic product of PRODH2, is transported by GOT1/2 to subsequently enter the TCA cycle, we tested the usage of GOT1, and GOT2 inhibitor PF04859989 was used to block PHC downstream metabolism. The GOT1/2 inhibition-cytosis experiment showed that PF04859989 reduced cancer killing ability only in the presence of PRODH2 GOF (i.e., only in *CD22-CAR;PRODH2* T cells but not in PRODH2(Stop) control group) (Figure 5J). Blocking the one-step-upstream enzyme that produces PRODH2 direct substrate and downstream enzyme of PHC in PRODH2-OE CAR-T had an exactly opposite effect in cancer killing. These results from the proline metabolic pathway perturbation experiments suggested that the mechanism underlying enhanced CAR-T function is mediated by an on-target enhancement of PRODH2 enzymatic function.

Mitochondrial cellular and function analyses revealed PRODH2 GOF-driven energetics in CAR-T cells

We then investigated how PRODH2 engineering drives the CAR-T cellular metabolic state. Because PRODH2's enzymatic activity is primarily in the mitochondria (Mt), we performed a series of cellular and biochemical Mt analyses on PRODH2 CAR-T cells under a longer-term culture. We first used electron microscopy (EM) to examine the morphology of the CAR-T cells. Both purified *CD22-CAR;PRODH2* and *CD22-CAR;PRODH2* (Stop) knockin control CAR-T cells have similar morphology in cell shape, nucleus, and various organelles. Interestingly, as compared with control, the PRODH2 knockin CAR-T cells have

elevated mitochondria representation (Figures 6A–6H), which is a signature of memory-like metabolic state (Buck et al., 2016; van der Windt et al., 2012). This is measured in terms of higher mitochondria count per cell (Figures 6A and 6E) as well as larger total mitochondria area per cell (Figure 6G) in *CD22-CAR;PRODH2* CAR-T cells, but no difference in mitochondria length between *CD22-CAR;PRODH2* and *CD22-CAR;PRODH2* (Stop) CAR-T cells was observed (Figure 6F). Furthermore, compared with control CAR-T cells, the PRODH2 knockin CAR-T cells remodeled cristae morphology leading to cristae expansion and widening (Figure 6C), which was also previously implicated in T cell effector phenotypes (Buck et al., 2016). Another interesting phenomenon was that PRODH2 knockin CAR-T cells have more cytolytic granules (Figures 6D and 6H), a signature of cytolytic T cell function. To further evaluate the mitochondrial features by independent approaches, we measured mitochondrial mass and depolarization by MitoTracker staining (Bengsch et al., 2016; Schaping et al., 2016) and found that PRODH2 knockin and lentiviral overexpressed BCMA-CAR-T cells had higher mitochondrial mass (Figure 6I) but no difference of mitochondrial depolarization compared with control CAR-T cells (Figure S6A). We also measured mitochondrial DNA copy number and complex I and IV expression levels by qRT-PCR. The results showed that there was no significant difference between PRODH2 knockin and control CAR-T cells in Mt DNA or complex I/complex IV level (Figure S6B). The increased mitochondria load was also cross-validated in an independent CAR-T setting, with a different vehicle (lentiviral vector) and a different CAR transgene (BCMA-CAR) (Figure S6C).

We then performed quantitative Seahorse analysis to measure oxygen consumption rate (OCR), spare respiratory capacity (SRC), and extracellular acidification rate (ECAR) for these CAR-T cells. We found that compared with control CAR-T cells, the PRODH2 knockin CAR-T cells have higher OCR and SRC (Figures 6J and 6K). We confirmed this observation with an independent condition in which cell density was doubled (Figure S6D). The ECAR data showed the PRODH2 knockin CAR-T cells have low ECAR (Figure 6L), an indicator of glycolysis, which further suggested that PRODH2 GOF CAR-T cells were shifted toward the oxidative phosphorylation (OXPHOS) pathway. SRC and OXPHOS are memory-like signatures of T cells (Buck et al., 2016; van der Windt et al., 2012). It has been shown that CAR-T cells with enhanced mitochondria biogenesis and oxidative metabolism have better persistence during cancer immunotherapy (Kawalekar et al., 2016), which are metabolic and mitochondria features similarly observed in PRODH2 knockin CAR-T cells. The mitochondria EM analysis revealed the cell biology features of PRODH2 GOF CAR-T cells linked to increased memory-like features, as well as effector function.

PRODH2 GOF alters the behaviors of CAR-T cells after antigen-specific cancer stimulation in long-term coculture

Retaining long-term activity under chronic antigen stimulation is important for effective T cell-based cancer immunotherapies. A recent study showed that reprogramming T cell metabolism and mitochondria fitness can enhance both effector function and long-term memory phenotype simultaneously, for example by targeting the Regnase 1 enzyme (Wei et al., 2019). Along

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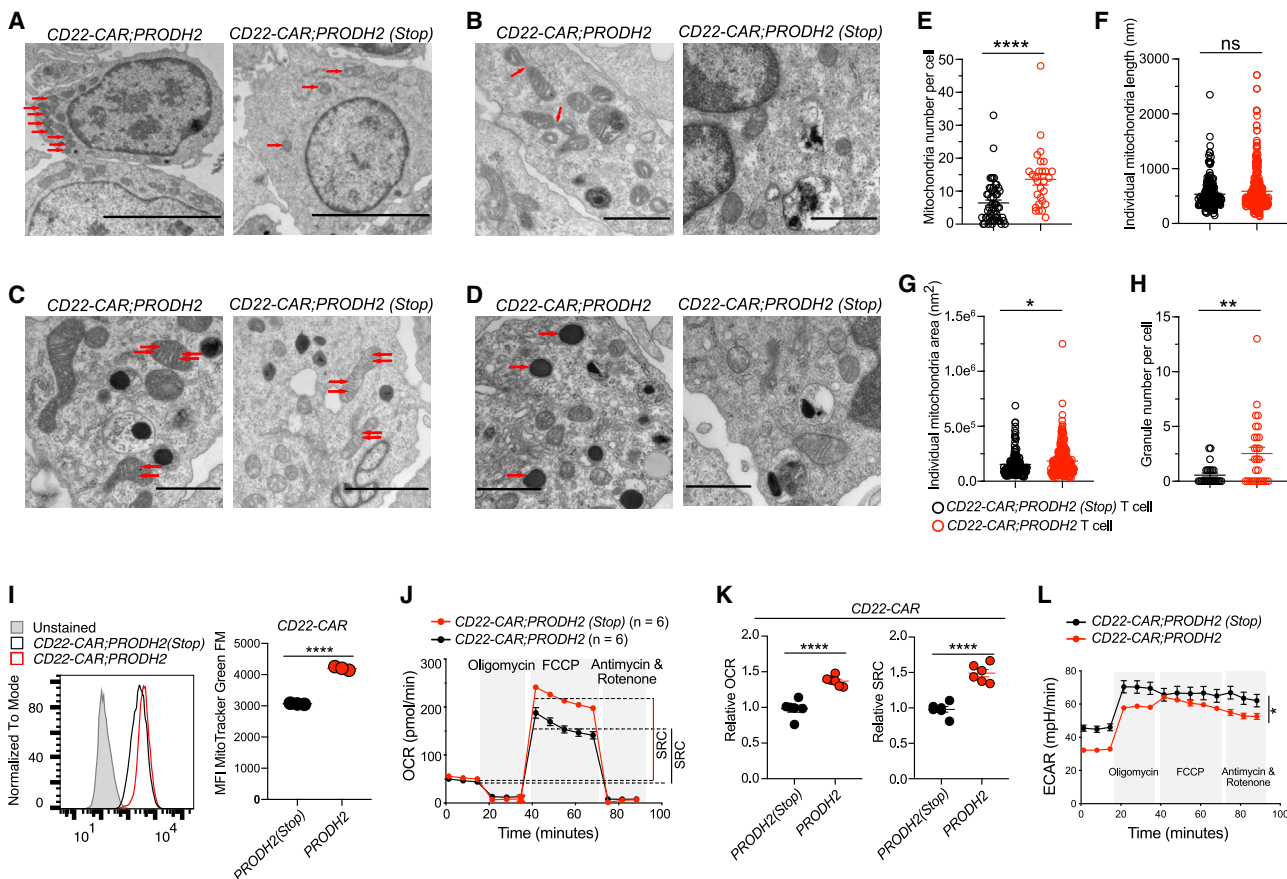


Figure 6. Characterization of long-term mitochondria energetics of PRODH2 knockin CAR-T cells

(A–D) TEM analysis of *PRODH2* knockin and control CAR-T cells. TEM images were examined for mitochondrial numbers (A) (red arrows indicate mitochondria), mitochondrial fission (B) (red arrows indicate mitochondria fission), mitochondrial cristae remodeling (C) (red arrows indicate mitochondrial cristae), and granule numbers (D) (red arrows indicate granules). Scale bars, 5 μm (A), 1 μm (B, C [left], and D), and 2 μm (C, right). Data from one experiment with independent replicates.

(E) Quantification of mitochondria number per cell.

(F) Quantification of individual mitochondria length.

(G) Quantification of individual mitochondria area.

(H) Quantification of granule number per cell.

(I) Mitochondrial mass as measured by MitoTracker Green FM staining of indicated CAR-T cells at day 36 after stimulation with NALM6-GL-CD22OE cancer cells. (J–L) Seahorse experiment of *PRODH2* knockin and control CAR-T cells, with a density of 2e5 CAR-T cells/well ($n = 6$). Data are representative of three independent experiments.

(J) Oxygen consumption rate (OCR) was measured at baseline and in response to oligomycin (Oligo), fluoro-carbonyl cyanide phenylhydrazone (FCCP), and rotenone plus antimycin A.

(K) Relative maximum OCR and relative spare respiratory capacity (SRC) were quantified.

(L) Extracellular acidification rate (ECAR) was measured at baseline and after drug treatment as the OCR measurement ($n = 6$). Representative data from three independent experiments.

* $p < 0.05$, ** $p < 0.01$, *** $p < 0.0001$; ns, not significant by unpaired t tests (E–I and K) and two-way ANOVA (L). See also Figure S6.

with the mitochondrial function observations, we set out to test *PRODH2* overexpression in CAR-T cells over a long course of coculture with cancer cells, mimicking the chronic antigen stimulation in disease settings (Figure 7A). *CD22-CAR;PRODH2* and *CD22-CAR;PRODH2(Stop)* control T cells manifested drastically different phenotypes at day 27 after CAR knockin but before encountering cancer antigen (without cancer stimulation), where *PRODH2* overexpressed CD22-CAR showed predominant effector phenotype higher than the control (Figure 7B), consistent with the multiple evidence of enhanced effector phenotypes

shown above. We then stimulated both groups with antigen-specific cancer cells every 12 days to mimic the chronic cancer antigen stimulation in therapeutic settings (Eyquem et al., 2017; Figure 7A). We used the well-established canonical CD45RA;CD62L dual marker flow analyses at day 69 after CAR-T knockin. We found that although *PRODH2* CAR-T cells have higher CD45RA-high;CD62L-low population (representing effector cells), which is consistent with the findings above, *PRODH2* CAR-T cells interestingly quickly shift away from the CD45RA-high;CD62L-low phenotype during the

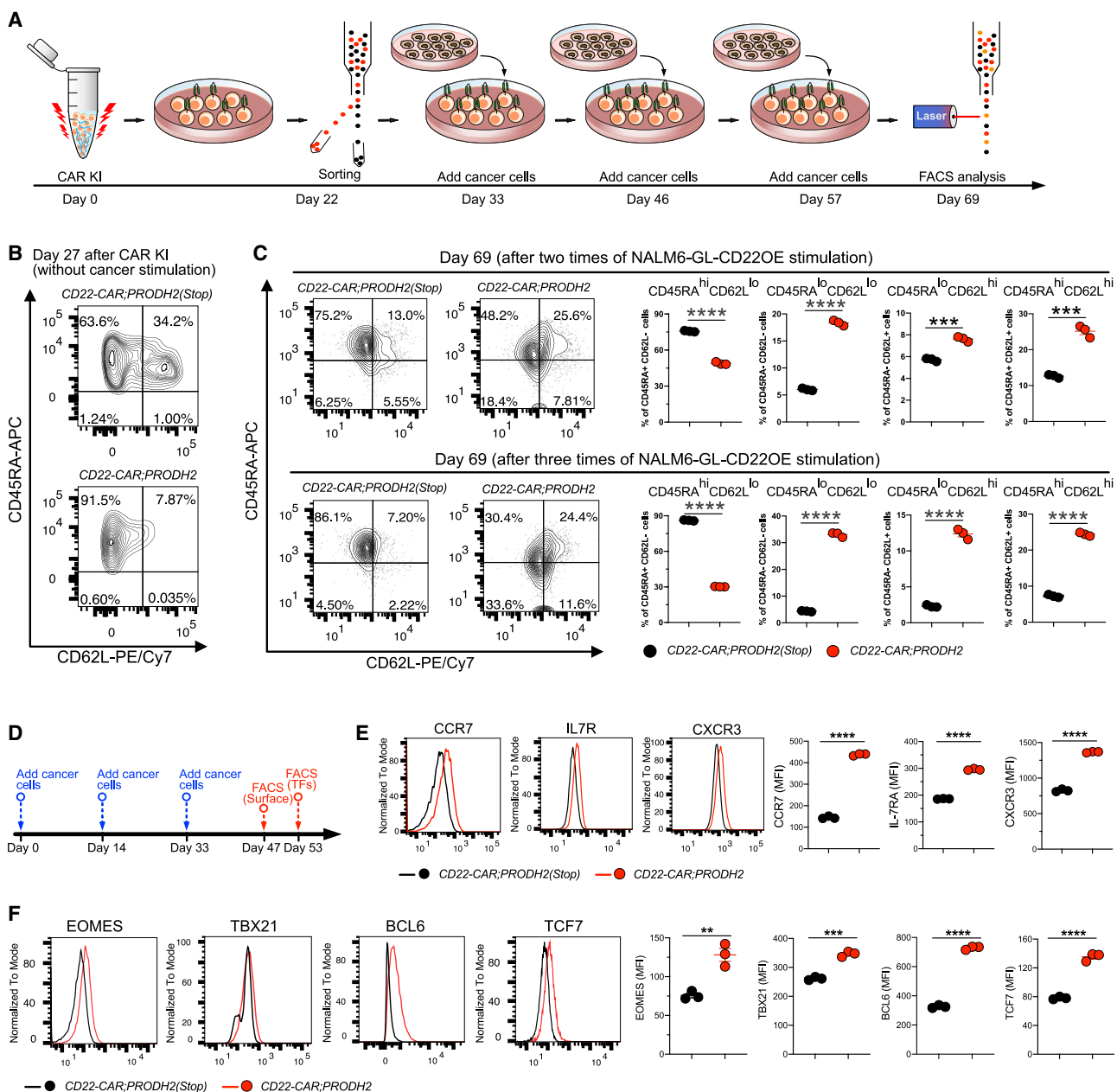


Figure 7. PRODH2 promotes CAR-T cell memory formation after cancer cell engagement in long-term coculture

(A) A schematic of the experimental design showing timeline of CAR knockin, FACS sorting, cancer stimulation, and flow analysis. *CD22-CAR;PRODH2* and *CD22-CAR;PRODH2(Stop)* T cells were purified by FACS sorting at day 22 after CAR knockin, then CAR-T cells were stimulated with NALM6-GL-CD22OE cancer cells various times, each about every 12 days with an E:T ratio = 1:1. T cells stimulated at different times were harvested for analysis at day 69.

(B) Baseline expression of CD45RA and CD62L in *CD22-CAR;PRODH2* and *CD22-CAR;PRODH2(Stop)* T cells analyzed at day 27.

(C) CD45RA and CD62L expression in *CD22-CAR;PRODH2* and *CD22-CAR;PRODH2(Stop)* T cells analyzed at day 69, stimulated either 2 or 3 times with NALM6-GL-CD22OE cancer cells (hi, high expression; lo, low expression).

(D) Timeline of cancer stimulation and flow cytometry.

(E and F) Analysis of T cell memory in PRODH2 knockin CAR-T cells after three times of cancer cell stimulation in long-term coculture.

(E) Flow analysis and quantification of human T cell memory surface markers, CCR7, IL7R, and CXCR3.

(F) Flow analysis and quantification of human T cell memory transcription regulators, EOMES, TBX21, BCL6, and TCF7.

p < 0.01, *p < 0.0001, ****p < 0.0001 by unpaired t tests (C, E, and F).

prolonged culture and cancer stimulation. After two stimulations, they exhibited significantly higher fractions in the CD45RA-low;CD62L-low, CD45RA-low;CD62L-high, and CD45RA-

high;CD62L-high populations (Figure 7C). Because these three populations resemble the less-differentiated (memory or naive) phenotypes, this observation is consistent with the EM and

Cell Metabolism

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Seahorse results. This phenotype was more pronounced after a third stimulation (Figure 7C). In this long-term coculture, the PRODH2 GOF T cells are associated with a slight increase of PD1, TIM3, and LAG3 (Figure S6E). We also measured several independent surface markers and gene expression regulators in long-term culture (Figures 7D–7F). The results showed that CCR7, IL7R, and CXCR3, which are indicators of memory-like phenotypes in T cells, were significantly increased after cancer stimulation in long-term culture (Figure 7E). BCL6 and TCF7, which are well-known master regulators of T cell memory differentiation (Chang et al., 2014; Crotty et al., 2010), were also substantially increased in PRODH2 GOF CAR-T cells with cancer stimulation in long-term coculture (Figure 7F).

Finally, we investigated the potential risk of malignant transformation of PRODH2 GOF. We first performed an IL-2 withdrawal assay and found that the majority CAR-T cells were dead at day 7 without IL-2 in the medium, suggesting that they were not immortalized or transformed (Figure S6F). Moreover, we analyzed the expressions of NOTCH1, GATA3, and RUNX1, malignant transformation markers of T cell acute lymphoblastic leukemias (T-ALL) (Van Vlierberghe and Ferrando, 2012). The flow cytometry data showed that these markers are negative or expressed at a low level across all groups, where PRODH2 GOF CAR-T cells are at levels similar to both PRODH2(Stop) control CAR-T and untransduced T cells (Figure S6G). These data together suggested that PRODH2 GOF is not associated with obvious risk of malignant transformation of the T cells themselves.

DISCUSSION

Catalytically, dgRNAs can be utilized to modulate gene expression with catalytically active Cas9 (Dahlman et al., 2015; Kiani et al., 2015). High-throughput CRISPR screens enabled unbiased discovery of therapeutic targets (Shalem et al., 2015); however, the application of dgRNA technologies has not yet been harnessed at massively parallel scale. Recently, genetic screens were performed to identify T cell regulators *in vitro* (Henriksson et al., 2019; LaFleur et al., 2019; Shifrut et al., 2018; Ting et al., 2018). T cell knockout screens have also been recently conducted *in vivo* (Dong et al., 2019; Ye et al., 2019). Although several studies demonstrated genome-scale GOF screen systems for protein-coding genes (Gilbert et al., 2014; Konermann et al., 2015) and long noncoding RNAs (lncRNAs) (Joung et al., 2017) in human melanoma cells, it is noted that in CRISPRa-based systems genes with higher fold activation are often lowly expressed. Thus, the basal expression levels of genes affect screening results with GOF screens (Gilbert et al., 2014; Konermann et al., 2015). In Jurkat cells, CRISPRa has been applied to identify stimulation-responsive enhancers (Mumbach et al., 2017; Simeonov et al., 2017), to map T cell signaling pathways (Chi et al., 2016), and to activate master regulator gene expression (Forstnerič et al., 2019). However, Jurkat is an immortalized lymphocyte cell line and not primary T cell, which is not clinically relevant. These previous studies discussed the future application of CRISPRa in primary T cells (Forstnerič et al., 2019). It is challenging to introduce the entire CRISPRa screening machinery (dCas9, transactivator, and a large-scale guide RNA library) simultaneously into primary CD8⁺ T cells.

Here, we leveraged the dgRNA system and the catalytically active Cas9 transgenic mice to achieve the screening capability. We thereby developed an orthogonal system for CD8⁺ T cell GOF screens and identified endogenous factors for boosting T cell function, by designing a catalytically dgRNA library and leveraging Cas9 transgenic mice where Cas9⁺ immune cells are readily available to be isolated and cultured at a large scale. With the development of orthogonal screening technologies, the dgRNA library system may be used in conjunction with knockout systems to perform complex intersectional screens in individual Cas9⁺ cells.

Previous studies have demonstrated that overexpression of STAT5 can improve antitumor effects (Grange et al., 2012, 2013). However, these studies are not unbiased screens in T cells. Using a systems approach, we unveiled several ranked GOF candidates enriched in T cells with superior effector function. We validated 4 genes (*Prodh2*, *Ccnb1ip1*, *Srek1ip1*, and *Wdr37*) where augmenting their expression consistently enhanced CD8⁺ T cell degranulation. These data validated the ability of the dgTKS system for discovery of new GOF targets in primary T cells. This recently developed screening system can potentially be broadly applied to discovery of GOF genes in various other immune and primary cell types.

T cell survival, activation, development, proliferation, differentiation, and antitumor effector function are virtually all driven by metabolism (Geltink et al., 2018). Naive T cells are supported by ATP that is predominantly generated from OXPHOS before activation (MacIver et al., 2013). The T cell proteome and phosphoproteome were mapped in dynamic T cell activation signaling networks during T cell activation (Tan et al., 2017). Activated T cells shifted their metabolic state from oxidative metabolism to glycolysis for cell proliferation and effector functions (Chang et al., 2013; Geltink et al., 2018; MacIver et al., 2013). Glycolysis is a relatively inefficient pathway for generating ATP, which is important for supporting rapid T cell growth and proliferation (O'Neill et al., 2016). The total biomass is dramatically increased after T cell activation for proliferation, which requires substantial nucleotide, lipid, and amino acid synthesis (Fox et al., 2005). Programming T cell metabolism using bioengineering approaches therefore may provide effective strategies for enhancement of T cell function.

Our dgTKS system and genome-scale GOF screen identified *Prodh2/PRODH2* as a top hit in cancer cell killing ability. *PRODH2* encodes an enzyme in proline metabolism by catalyzing the first step in the catabolism of *trans*-4-hydroxy-L-proline. Proline and arginine metabolism is important in T cell antitumor activity (Geiger et al., 2016), although the exact role of proline metabolism in T cell function is less well characterized. Because PRODH2 is not expressed in primary T cells, its physiological function is likely minimal, and thus, LOF perturbation will not interfere with its natural role, making it an ideal GOF target. *PRODH2/Prodh2* is a highly tissue-specific enzyme mainly expressed in livers and kidneys and only lowly expressed in most other cell types (Jiang et al., 2012; Knight et al., 2009). To the best of our knowledge, to date, there is no reported study on *PRODH2* functions in T cells. *Prodh2* played an important role in proline homeostasis, especially under stress conditions (Funck et al., 2010). *PRODH2* is important for primary hyperoxaluria (PH), as it is a disorder of cellular glyoxylate metabolism

(Harambat et al., 2011; Hoppe et al., 2009). Public databases including Immgen (Heng et al., 2008) and DICE (Schmiedel et al., 2018) showed that *Prodh2* is usually not expressed in immune cells, although it is expressed in the mitochondria of kidney, liver, and gallbladder cells. Therefore, in a normal physiological setting, PRODH2 is not expressed or detected in T cells, which explains its absence in prior LOF T cell screens (Chen et al., 2014; Dong et al., 2019; Henriksson et al., 2019; LaFleur et al., 2019; Shifrut et al., 2018; Ye et al., 2019; Zhou et al., 2014). The metabolic effects of PRODH2 makes it an interesting target for T cell metabolic programming in a setting of ectopic engineering of new elements as immune boosters.

The landmark success of CD19 CAR with US FDA approval opens a new era of cell therapy (e.g., Schuster et al., 2019). There remain many hurdles for CAR-T therapy due to various challenges (Lim and June, 2017). Cancer cells can metabolically restrict T cell antitumor functions by nutrient depletion (Chang et al., 2015). Thus, approaches to enhance T cell metabolism may improve T cell antitumor immunity. The second and third generation CARs designed with extra costimulatory domains (CD28 and/or 4-1BB) in addition to CD3ζ had a better antitumor efficacy compared with the first generation (June et al., 2018). The CD28 signaling domain has been found to have an extra function of enhancing aerobic glycolysis, and 4-1BB enhanced mitochondria biogenesis and fatty acid oxidation in the CAR-T cells (Kawalekar et al., 2016). Recently, intracellular L-arginine (involved in arginine and proline metabolism) concentration has been found to impact the metabolic fitness and survival capacity of T cells and antitumor activity (Geiger et al., 2016). Programming T cell metabolism is an emerging approach for immunotherapy (Chang and Pearce, 2016). More recently, it was shown that overexpression of canonical AP-1 factor c-Jun in CAR-T cells induced exhaustion resistance (Lynn et al., 2019). Engineering CARs with codon-optimized CD8 transmembrane domain can induce lower levels of cytokines but retain potent cytolytic activity (Ying et al., 2019). A split, universal, and programmable (SUPRA) CAR system can fine-tune T cell activation strength and sense and logically respond to multiple antigens (Cho et al., 2018). Additional approaches include restructuring of signaling domains (Sadelain et al., 2017), overexpression of boosting factors (Lynn et al., 2019), coadministration of immuno-modulating factors or viral vectors (Ma et al., 2019), or changing costimulatory domains or lowering CAR binding affinity (Ghorashian et al., 2019; Savoldo et al., 2011).

Limitations of study

The overall killing of HER2-CAR-T cells was not as striking as CD22-CAR or BCMA-CAR T cells, which was expected as solid tumor therapy using CAR-T cells is still a major challenge. In addition, the memory characterization of CAR-T cells was performed in long-term coculture *in vitro* because the *in vivo* models also have their own limitations as the study of CAR-T cells *in vivo* requires the use of immunocompromised animals. Although PRODH2's effect on effector function and cancer killing is strong and the mechanism is clear from this study, PRODH2's effect on memory establishment is worthy of further investigation in the future.

In conclusion, whole-transcriptome analysis, multiomics analyses, CAR-T cellular and immunological assays, metabolic

analyses, and mechanistic dissection together showed that GOF-engineered *PRODH2* via OE or KI can reprogram T cell metabolism; promote T cell proliferation, activation, and effector function; improve effective responses to longer-term cancer stimulation; and enhance antitumor efficacy *in vivo* (Figure S7). More generally, the dgRNA library-based CRISPRa screen technology also provides a versatile and high-throughput activation screening system for identification of new classes of GOF targets directly in T cells or other primary immune cells. Given that the dgRNA library is compatible with catalytically active Cas9, orthogonal activation and knockout screens are possible. These systems might be a broadly useful resource for the field.

STAR METHODS

Detailed methods are provided in the online version of this paper and include the following:

- KEY RESOURCES TABLE
- RESOURCE AVAILABILITY
 - Lead contact
 - Materials availability
 - Data and code availability
- EXPERIMENTAL MODEL AND SUBJECT DETAILS
 - Mouse models
 - Cell culture
 - Naive CD8⁺ T cell isolation and culture
 - Pre-clinical efficacy testing using *in vivo* models
- METHOD DETAILS
 - mm10dgLib design
 - Construction of T cell dgRNA activation (TdgA) vector
 - mm10dgLib cloning
 - Lentivirus production
 - mm10dgLib lentivirus titration
 - mm10dgLib lentivirus transduction
 - AAV-HDR based CAR and lentiviral-based CAR construction
 - AAV production
 - Flow cytometry for surface markers
 - Intracellular flow cytometry
 - Kill assay and dgLib screen
 - Kill assay screen readout and deep sequencing
 - dgRNA screen data analysis
 - Enrichment analysis of dgRNAs
 - Immunoblots
 - Cytokine secretion assays
 - RT-qPCR
 - Human primary CD8⁺ T cell electroporation and CAR-T knock in
 - Top candidate validation in kill assay
 - IL-2 withdrawal assay
 - Human CD22-CAR;PRODH2 and HER2-CAR;PRODH2 T cell kill assay
 - Lenti-CD22-CAR;PRODH2 and Lenti-BCMA-CAR;PRODH2 T cell transduction and co-culture
 - Bulk mRNA sequencing (mRNA-seq) library preparation
 - Bulk mRNA-seq data processing

Cell Metabolism

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- Gene set level pathway analysis of differentially expressed genes
- Metabolite extraction and data collection
- Metabolomics data processing
- Intersection of mRNA-seq, metabolomics and KEGG pathways
- Multi-Omics analysis
- Transmission electron microscopy (TEM) of CAR-T cells
- Seahorse assay
- Mass cytometry (CyTOF)
- CyTOF data analysis
- Mitochondria mass and depolarization measurement
- CAR T cell chemical treatment, co-culture and flow assays
- Long-term CAR-T culture with chronic antigen stimulation and memory marker analysis

● QUANTIFICATION AND STATISTICAL ANALYSIS

SUPPLEMENTAL INFORMATION

Supplemental information can be found online at <https://doi.org/10.1016/j.cmet.2022.02.009>.

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AUTHOR CONTRIBUTIONS

L.Y. and S.C. conceived and designed the experiments. L.Y. led and performed experiments with L.P. and Q.Y. J.J.P. led the NGS data analyses. R.D.C. designed dgRNA library. L.Y., L.P., and Q.Y. analyzed the data except NGS. M.B.D., S.Z.L., J.G., E.T., Y.Z., G.W., X.D., Y.D., H.R.K., H.C., Y.E., and P.C. assisted with experiments. A.B. provided technical and conceptual

advice on CAR-T. R.R.M. provided technical and conceptual advice on CyTOF. L.Y., J.J.P., L.P., and S.C. wrote the manuscript with input from all authors. S.C. secured funding and supervised the work.

DECLARATION OF INTERESTS

S.C. is a cofounder of EvolveImmune Tx and Cellinfinity Bio. A patent has been filed by Yale University related to this study (S.C., L.Y., J.J.P., L.P., R.D.C., and M.B.D. as inventors). The other authors declare no competing interests.

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STAR★METHODS

KEY RESOURCES TABLE

REAGENT or RESOURCE	SOURCE	IDENTIFIER
Antibodies		
Ultra-LEAF Purified anti-mouse CD3ε Antibody (clone: 145-2C11)	Biolegend	Cat#100340; RRID:AB_11149115
Ultra-LEAF Purified anti-mouse CD28 Antibody (clone: 37.51)	Biolegend	Cat#102116; RRID:AB_11147170
APC anti-human CD3 Antibody (Clone: UCHT1)	Biolegend	Cat#300412; RRID:AB_314066
FITC anti-human IgG Fc	Biolegend	Cat#409310; RRID:AB_2561855
PE anti-human IgG Fc (HP6017)	Biolegend	Cat#409304; RRID:AB_10895907
PE anti-DYKDDDDK Tag Antibody (L5)	Biolegend	Cat#637310; RRID:AB_2563148
APC/Cy7 anti-human CD8a antibody (HIT8a)	Biolegend	Cat#300926; RRID:AB_10613636
APC anti-human CD3 Antibody (Clone: HIT3a)	Biolegend	Cat#300312; RRID:AB_314048
APC anti-mouse CD8a Antibody (53-6.7)	Biolegend	Cat#100712; RRID:AB_312751
PE/Cy7 anti-mouse CD3ε Antibody (145-2C11)	Biolegend	Cat#100320; RRID:AB_312685
PE anti-rat CD90/mouse CD90.1 (Thy-1.1) Antibody (OX-7)	Biolegend	Cat#202524; RRID:AB_1595524
PE anti-mouse CD107a (LAMP-1) Antibody (1D4B)	Biolegend	Cat#121612; RRID:AB_2134487
PE conjugate Cleaved Caspase-3 (Asp175) Rabbit mAb (5A1E)	CST	Cat#9978S; RRID:AB_10831820
Brilliant Violet 510 anti-human CD8 Antibody (SK1)	Biolegend	Cat#344732; RRID:AB_2564624
APC/Cy7 anti-human IFN-γ Antibody (4S.B3)	Biolegend	Cat#502530; RRID:AB_10663412
PE anti-human TNF-α Antibody (MAb11)	Biolegend	Cat#502909; RRID:AB_315261
PerCP/Cyanine5.5 anti-human/mouse Granzyme B Recombinant Antibody (QA16A02)	Biolegend	Cat#372212; RRID:AB_2728379
APC anti-humanCD45RA Antibody (HI100)	Biolegend	Cat#304112; RRID:AB_314416
PE/Cy7 anti-human CD62L Antibody (DREG-56)	Biolegend	Cat#304822; RRID:AB_830801
PerCP/Cyanine5.5 anti-human CD197 (CCR7) Antibody (Clone: G043H7)	Biolegend	Cat#353219; RRID:AB_10915275
APC anti-human CD127 (IL-7R α) Antibody (Clone: A019D5)	Biolegend	Cat#351315; RRID:AB_10900814
FITC anti-human CD183 (CXCR3) Antibody (Clone: G025H7)	Biolegend	Cat#353703; RRID:AB_10962910
EOMES Monoclonal Antibody (WD1928), PECyanine7	eBioscience	Cat#25-4877-42; RRID:AB_2573456
APC anti-T-bet Antibody (Clone: 4B10)	Biolegend	Cat#644813; RRID:AB_10896913
PE anti-TCF1 (TCF7) Antibody (Clone: 7F11A10)	Biolegend	Cat#655207; RRID:AB_2728491
FITC anti-human/mouse Bcl-6 Antibody (Clone: 7D1)	Biolegend	Cat#358513; RRID:AB_2860942
APC anti-human Notch 1 Antibody (Clone: MHN1-519)	Biolegend	Cat#352107; RRID:AB_10897100

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REAGENT or RESOURCE	SOURCE	IDENTIFIER
PerCP/Cyanine5.5 anti-GATA3 Antibody (Clone: 16E10A23)	Biolegend	Cat#653811; RRID:AB_2563218
RUNX1 Monoclonal Antibody (RXDMC), PE	eBioscience	Cat#12-9816-80; RRID:AB_11151519
Anti-human CTLA4 (CD152)-161Dy	FLUIDIGM	Cat#3161004B; RRID:AB_2687649
Anti-human TIM3-153Eu	Yale CyTOF CORE	N/A
Anti-human CD278/ICOS-168Er	FLUIDIGM	Cat#3168024B; RRID:AB_2858237
Anti-human TNFRSF9/CD137-173Yb	FLUIDIGM	Cat#3173015B
Anti-human IL2-166Er	FLUIDIGM	Cat#3158007B; RRID:AB_2864735
Anti-human TNF- α -152Sm	FLUIDIGM	Cat#3152001B
Anti-human IFN- γ -165Ho	FLUIDIGM	Cat#3165002B
Anti-human CXCR3-163Dy	FLUIDIGM	Cat#3163004B; RRID:AB_2810969
Anti-human CD62L-174Yb	Yale CYTOF Core	Cat#V00751
Anti-human Perforin-176Yb	Yale CYTOF Core	N/A
Anti-human CD45RA-169Tm	FLUIDIGM	Cat#3143006B; RRID:AB_2651156
Anti-human CD8a-146Nd	FLUIDIGM	Cat#3146001B
Anti-human CD45RO-164Dy	FLUIDIGM	Cat#3164007B; RRID:AB_2811092
Anti-human CD3-170Er	FLUIDIGM	Cat#3170001B; RRID:AB_2811085
Anti-human CD45-89Y	FLUIDIGM	Cat#3089003B; RRID:AB_2661851
Anti-Cleaved Caspase3-172Yb	Yale CYTOF Core	N/A
Anti-human CD25-149Sm	FLUIDIGM	Cat#3149010B; RRID:AB_2756416
Anti-human CD134(OX40)-150Nd	Yale CYTOF Core	N/A
Anti-human LAG3-175Lu	Yale CYTOF Core	N/A
Recombinant Human Siglec-2/CD22 Fc Chimera Protein, CF	R&D	Cat#1968-SL-050
Recombinant Human BCMA/TNFRSF17 Fc Chimera Protein, CF	R&D	Cat#193-BC-050
Dynabeads Human T-Activator CD3/CD28	Invitrogen	Cat#11131D
Anti-human PRODH2	Atlas Antibodies	Cat#HPA051287; RRID:AB_2681420
Anti-human Vinculin	Abcam	Cat#129002; RRID:AB_11144129
Bacterial and virus strains		
One Shot Stbl3 Chemical Competent <i>E. coli</i>	ThermoFisher	Cat#C737303
Endura ElectroCompetent Cells	Lucigen	Cat#60242-2
Chemicals, peptides, and recombinant proteins		
PEI MAX -Transfection Grade Linear Polyethylenimine Hydrochloride (MW 40,000)	Polyscience	Cat#24765-1
DPBS, no calcium, no magnesium	GIBCO	Cat#14190136
RPMI 1640 Medium	GIBCO	Cat#11875-093
Fetal Bovine Serum	Sigma Aldrich	Cat#F4135-500ML
DMEM, high glucose, pyruvate	GIBCO	Cat#11995065
Opti-MEM I Reduced Serum Medium	ThermoFisher	Cat#31985070
Penicillin-Streptomycin (10,000 U/mL)	GIBCO	Cat#15140122
Lonza BioWhittaker L-Glutamine (200mM)	Lonza	Cat#BW17605E
2-Mercaptoethanol	Sigma Aldrich	Cat#M6250-10ML
X-VIVO 15 Serum-free Hematopoietic Cell Medium	Lonza	Cat#BE02-060F
Human AB Serum; Male Donors; type AB	MP Biomedical	Cat#092930949
ACK Lysing Buffer	Lonza	Cat#10-548E
Naive CD8a+ T Cell Isolation Kit, mouse	Miltenyi	Cat#130-096-543
LS Columns	Miltenyi	Cat#130-042-401

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Cell Metabolism

Article



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REAGENT or RESOURCE	SOURCE	IDENTIFIER
Recombinant Mouse IL-2 (carrier-free)	Biolegend	Cat#575404
Recombinant Mouse IL-7 (carrier-free)	Biolegend	Cat#577802
Recombinant Mouse IL-12 (p70) (carrier-free)	Biolegend	Cat#577008
Recombinant Mouse IL-15 (carrier-free)	Biolegend	Cat#566302
Recombinant Human IL-2 (carrier-free)	Biolegend	Cat#589104
Monensin Solution (1,000x)	Biolegend	Cat#420701
Brefeldin A Solution (1,000X)	Biolegend	Cat#420601
QuickExtract DNA Extraction Solution	Epicenter	Cat#QE09050
Proteinase K	QIAGEN	Cat#19131
Pierce Universal Nuclease for Cell Lysis	ThermoFisher	Cat#88702
RNase A	QIAGEN	Cat#19101
PEG 8000, Molecular Biology Grade	Promega	Cat#V3011
Sodium chloride	Sigma	Cat#S3014
Gibson Assembly Master Mix	NEB	Cat#E2611
Phusion Flash High-Fidelity PCR Master Mix	ThermoFisher	Cat#F548L
DreamTaq Green PCR Master Mix (2X)	ThermoFisher	Cat#K1082
E-Gel Low Range Quantitative DNA Ladder	ThermoFisher	Cat#12373031
Fixation/Permeabilization Solution Kit	BD	Cat#554714
SuperScript IV Reverse Transcriptase	Invitrogen	Cat#18090050
Random Hexamers (50 μM)	Invitrogen	Cat#N8080127
NEBNext Ultra RNA Library Prep Kit for Illumina	NEB	Cat#E7530S
NEBNext Multiplex Oligos for IlluminaR (Index Primers Set 1)	NEB	Cat#E7335S
TaqMan Fast Advanced Master Mix	Invitrogen	Cat#4444557
QIAquick Gel Extraction Kit	QIAGEN	Cat#28706
QIAamp Fast DNA Tissue Kit	QIAGEN	Cat#51404
EndoFree Plasmid Maxi Kit	QIAGEN	Cat#12362
RNeasy Mini Kit	QIAGEN	Cat#74104
Relative Human Mitochondrial DNA Copy Number Quantification qPCR Assay Kit	ScienCell	Catalog#8938
Human IL-2 ELISA Kit	Millipore Sigma	Cat#RAB0286
IL-7 Human ELISA Kit	ThermoFisher	Catalog#EHIL7
IL-15 Human ELISA Kit	ThermoFisher	Catalog#BMS2106
Agilent Seahorse XF Cell Mito Stress Test Kit	Agilent	Cat#103015-100
Agilent Seahorse XF Calibrant solution	Agilent	Cat#100840-000
Agilent Seahorse XF RPMI medium	Agilent	Cat#103576-100
Agilent Seahorse XF sensor cartridges and cell culture microplates	Agilent	Cat#101085-004
Poly-D-lysine	Sigma	Cat#27964-99-4
Seahorse XF 100 mM Pyruvate Solution	Agilent	Cat#103578-100
Seahorse XF 1.0 M Glucose Solution	Agilent	Cat#103577-100
Seahorse XF 200 mM Glutamine Solution	Agilent	Cat#103579-100
Bovine Serum Albumin	Sigma Aldrich	Cat#A9418-100G
EDTA	Sigma Aldrich	Cat#E8008-100ML
XenoLight D-Luciferin – K+ Salt Bioluminescent Substrate	Perkin Elmer	Cat#122799
Neon Transfection System 100 μL Kit	Invitrogen	Cat#MPK10025
HiFi Cas9 protein	IDT	Cat#1081061

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REAGENT or RESOURCE	SOURCE	IDENTIFIER
Cas9 tracrRNA	IDT	Cat#1072532
crRNA	IDT	Custom, sequence specific
OVA 257-264	Invivogen	Cat#vac-sin
Bpil (BbsI) (10 U/μL)	ThermoFisher	Cat#ER1012
Esp3I (BsmBI) (10 U/μL)	ThermoFisher	Cat#ER0451
4-Hydroxyproline	Sigma	Cat#H54409
Acetyl CoA	Sigma	Cat#A2056
ADP	Sigma	Cat#A2754
Amino acid standards, physiological	Sigma	Cat#A9906
AMP	Sigma	Cat#1930
Arginine	Sigma	Cat#A5006
ATP	Sigma	Cat#A26209
cAMP	Sigma	Cat#A9501
Choline	Sigma	Cat#C7017
CoA	Sigma	Cat#C4282
CTP	Sigma	Cat#C1506
Fumarate	Sigma	Cat#D95654
Glucose	Sigma	Cat#G8270
Glutathione	Sigma	Cat#G4251
Glyceraldehyde3P	Sigma	Cat#G5251
Glycerate3P	Sigma	Cat#P8877
Glyoxylate	Sigma	Cat#G4502
GTP	Sigma	Cat#G8877
IMP	Sigma	Cat#I4625
Malate	Sigma	Cat#46940
Myo-Inositol	Sigma	Cat#I5125
Ornithine	Sigma	Cat#57197
Oxaloacetate	Sigma	Cat#O4216
Palmitoyl CoA	Sigma	Cat#P9716
Proline	Sigma	Cat#P3350000
PRPP	Sigma	Cat#P8296
Putrescine	Sigma	Cat#51799
Pyruvate	Sigma	Cat#P2256
Ribose5P	Sigma	Cat#83875
Spermidine	Sigma	Cat#S2626
Spermine	Sigma	Cat#S3256
Sphingosine	Sigma	Cat#S7049
Succinate	Sigma	Cat#W327700
Tetrahydrofolic acid	Sigma	Cat#G5251
TTP	Sigma	Cat#T0251
UTP	Sigma	Cat#U6375
Xylose-5-Phosphate	Sigma	Cat#78963
α-Hydroxyglutaric acid	Sigma	Cat#90790
Deposited data		
human CAR-T cell RNA-seq	This paper	SRA: PRJNA806391
human CD8 T cell metabolomics	This paper	Metabolomics Workbench: ST002085
human CAR-T cell metabolomics	This paper	Metabolomics Workbench: ST002084
CyTOF data	This paper	Mendeley Data: http://doi.org/10.17632/pnbjtdkfg.1

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Cell Metabolism

Article



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REAGENT or RESOURCE	SOURCE	IDENTIFIER
Experimental models: Cell lines		
HEK293FT	ThermoFisher	Catalog Number: R70007
HEK293T	ATCC	Catalog Number: CRL-3216
NALM6	ATCC	Catalog Number: 3273
MCF-7	ATCC	Catalog Number: HTB-22
E0771	CH3	Catalog Number: 940001
MB-MDA-231	ATCC	Catalog Number: HTB-26
MM.1R	ATCC	Catalog Number: CRL-2975
Human Peripheral Blood CD8+ T Cells	STEMCELL	Catalog Number: 70027
Human Peripheral Blood Mononuclear Cells	STEMCELL	Catalog Number: 70025.1
Experimental models: Organisms/strains		
OT-I	Jackson Laboratory	Stock No: 003831
NSG	Jackson Laboratory	Stock No: 005557
Cas9s	Platt et al., 2014	Jackson Lab
OT-I;Cas9s	Dong et al., 2019	Sidi Chen lab
Oligonucleotides		
mm10dg library	This paper	Data S1
Taqman probes	This paper	Table S2
Amplicon primers for surveyor or Nextera	This paper	Table S2
gBlocks of U6-dgRNA-MS2	IDT	Custom, sequence specific
EFS-Thy1.1-MCP-p65-HSF1	IDT	Custom, sequence specific
Prodh2 probe	ThermoFisher	Mm00457662-m1
Actb probe	ThermoFisher	Mm00607939_s1
Gapdh probe	ThermoFisher	Mm99999915_g1
Wdr37 probe	ThermoFisher	Mm0052518-m1
Srek1ip1 probe	ThermoFisher	Mm00482686-m1
Ccnb1ip1 probe	ThermoFisher	Rn01414585-m1
PRODH2 probe	ThermoFisher	Hs00560411-m1
ACTB probe	ThermoFisher	Hs99999903-m1
GAPDH probe	ThermoFisher	Hs02786624-g1
Recombinant DNA		
psPAX2	Addgene	Cat#12260
pMD2.G	Addgene	Cat#12259
pRepCap6 (AAV6 capsid)	Addgene	Cat#110770
pAdDeltaF6 (PDF6)	Addgene	Cat#112867
TdgA Vector	This study	Sidi Chen lab
Mm10dglLib library	This study	Sidi Chen lab
Software and algorithms		
FlowJo software 9.9.6	FlowJo, LLC	https://www.flowjo.com
Cutadapt	Martin, 2011	https://cutadapt.readthedocs.io/en/stable/
DAVID	Huang et al., 2009	https://david.ncifcrf.gov
Scran R package	Lun et al., 2016	https://www.r-project.org
Rtsne R package	Maaten, 2014; Maaten and Hinton, 2008	https://www.r-project.org
Edge R package	Robinson et al., 2010	https://www.r-project.org
Kallisto	Bray et al., 2016	https://pachterlab.github.io/kallisto/
Sleuth	Pimentel et al., 2017	https://pachterlab.github.io/sleuth/
GSEA	Subramanian et al., 2005	http://software.broadinstitute.org/gsea/index.jsp
PSCAN	Zambelli et al., 2009	http://159.149.160.88/pscan/

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REAGENT or RESOURCE	SOURCE	IDENTIFIER
Bowtie 1.1.2	Langmead et al., 2009	http://bowtiebio.sourceforge.net
Agilent Mass Hunter Qualitative Analysis Software	Agilent	(Version B.07.0.0, build 7.0.7024.0).
Agilent Mass Profiler Professional	Agilent	(Version 14.5-Build 2772)
Metabolite database	HMDB	http://www.hmdb.ca/
Metabolite database	METLIN	http://metlin.scripps.edu
MetaboAnalyst 4.0	Chong et al., 2018; Xia et al., 2015	https://www.metaboanalyst.ca
Pathvisio v3.3.0	Pathvisio	https://www.pathvisio.org
KEGG	KEGG	http://www.genome.jp/kegg

RESOURCE AVAILABILITY

Lead contact

Further information and requests for resources and reagents should be directed to and will be fulfilled by the Lead Contact, Sidi Chen (sidi.chen@yale.edu).

Materials availability

Mouse lines (*OT-I*;Cas9 β mice) and cell lines (NALM6-GL, MM.1R-PL, MCF7-PL, NALM6-GL-CD22OE, MM.1R-PL-BCMAOE, and MCF7-PL-HER2OE) generated in this study are available from the lead contact upon request.

Data and code availability

- The genomic sequencing raw data, the metabolomics raw data, and CyTOF raw data have been deposited at SRA, Metabolomics Workbench, and Mendeley data, respectively, and are publicly available as of the date of publication. Accession numbers are listed in the [key resources table](#).
- The original codes of data analysis are available from the lead contact upon reasonable request.
- Any additional information required to reanalyze the data reported in this paper is available from the lead contact upon request. The unprocessed data are deposited into public archives as above.

EXPERIMENTAL MODEL AND SUBJECT DETAILS

Mouse models

All animal work was performed under the guidelines of Yale University Institutional Animal Care and Use Committee (IACUC) with approved protocols (Chen-2015-20068; Chen-2018-20068; Chen-2021-20068). The general health of the mice are in good condition (BAR: bright, alert and responsive) before the cancer-related experiments started. Mice were housed in a free access to water and food, temperature (approximately 22°C) and humidity controlled colony room, maintained on a 12h light/dark cycle (07:00 to 19:00 light on). Mice health checks were performed daily. Mice, both female and male, aged 8-12 weeks were used for experiments. Rosa26-Cas9-2A-EGFP constitutive expressed mice (Cas9 β mice) and OT-I TCR transgenic mice (Hogquist et al., 1994) were used in this study. *OT-I* and Cas9 β mice were bred to generate *OT-I*;Cas9 β mice. *NOD-scid IL2Rgamma^{null}* (NSG) mice were purchased from JAX and bred in-house for *in vivo* tumor model and T cell based therapeutic efficacy testing experiments.

Cell culture

HEK293FT, HEK293T, E0771, NALM6, MM.1R, MCF7, and MDA-MB-231 cell lines were obtained from commercial sources (ThermoFisher, American Type Culture Collection (ATCC) and CH3) and were tested negative for mycoplasma contamination. HEK293FT, E0771 and MCF7 cells were cultured in DMEM (Gibco) media supplemented with 10 % FBS (Sigma) and 200 U / mL penicillin-streptomycin (Gibco), hereafter referred to as D10. NALM6 and MM.1R cells were cultured in RPMI-1640 (Gibco) media supplemented with 10% FBS and 200 U / mL penicillin-streptomycin. NALM6-GL-CD22OE, MM.1R-PL-BCMAOE, and MCF7-PL-HER2OE cell lines were established by transducing WT cancer cells with GFP-Luciferase (GL) or Puromycin-Luciferase (PL) lentivirus first, then purified by enriching for GFP+ or Puromycin resistant cells by FACS or puromycin selection, respectively. For NALM6-GL, MM.1R-PL, and MCF7-PL cell lines, cells were transduced with CD22-Blasticidin, BCMA-Blasticidin or HER2-Blasticidin lentivirus for overexpression of specific cancer antigen transgenes where appropriate, which were established by Blasticidin selection.

Naive CD8 $^{+}$ T cell isolation and culture

Mesenteric lymph nodes (mLNs) and spleens were dissected from *OT-I*;Cas9 β mice, then placed into ice-cold PBS supplemented with 2 % FBS. Lymphocytes were released by grinding organs through a 100 μ m filter, then re-suspended with 2 % FBS. Red blood

cells (RBCs) were lysed with 1 mL ACK lysis buffer (Lonza) for 2 spleens at 1-2 min at room temperature, then neutralized with 2 % FBS PBS at 20 volumes per volume of lysis buffer. RBC-lysed lymphocyte solution was filtered through 40 μ m filters to remove cell debris. Naïve CD8a⁺ T cell purification was performed using Naïve CD8a⁺ T cell Isolation Kits (Miltenyi Biotec) according to the manufacturer's protocols. Naïve CD8a⁺ T cells were cultured at 1 x 10e6 cells / mL density in 2 μ g / mL anti-CD3e (BioLegend) treated plates or dishes, in RPMI-1640 (Gibco) media supplemented with 10 % FBS, 2 mM L-Glutamine, 200 U / mL penicillin-streptomycin (Gibco), and 49 μ M β -mercaptoethanol (Sigma), hereafter referred to as cRPMI media. For *in vitro* experiments, cRPMI media was supplemented with 2 ng / mL IL-2, 1 μ g / mL anti-CD28, and 12 ng / mL IL-12p70 cytokines or antibodies. All cytokines and antibodies were purchased from BioLegend.

Pre-clinical efficacy testing using *in vivo* models

NOD-scid IL2Rgamma^{null} (NSG) mice were purchased from JAX and bred in-house. Eight-to-twelve-week-old both sex mice were used for cancer modeling and efficacy testing.

For the leukemia model, 5e5 NALM6-GL-CD22OE cancer cells were inoculated via intravenous injection. After 4 days of cancer inoculation, 4e6 *CD22-CAR;PRODH2* or *CD22-CAR;PRODH2 (Stop)* T cells were tail vein injected as treatments.

For the myeloma models, model 1, 1.5e6 per mouse of MM.1R-PL cancer cells were injected intravenously. Then 1.5e6 per mouse of *BCMA-CAR;PRODH2* and *BCMA-CAR;PRODH2 (Stop)* T cells were injected intravenously one day after cancer injection as treatments; Model 2, 2e6 per mouse of MM.1R-PL-BCMAOE cancer cells were injected intravenously. Then 9e5 per mouse of *BCMA-CAR;PRODH2* and *BCMA-CAR;PRODH2 (Stop)* T cells were injected intravenously ten day after cancer injection as treatments. For the lenti-BCMA-CAR model, 3e6 MM.1R-PL-BCMAOE cancer cells were injected intravenously. Then 1e6 per mouse of *lenti-BCMA-CAR;PRODH2* or *lenti-BCMA-CAR;PRODH2 (Stop)* T cells were injected intravenously at day 9 after cancer injection as treatments, the second dose of CAR-T cells (1.3e6 per mouse) was injected at day 33. Treatment dose and time-point are indicated in the figure legends.

For the breast cancer model, 1e6 MCF7-PL-HER2 cancer cells were fat-pad inoculated into female NSG mice, 1.5e6 *HER2-CAR-PRODH2* or *HER2-CAR* T cells were injected intravenously one day after cancer inoculation as treatments. Leukemia progression was measured by bioluminescent imaging using the IVIS system. Solid tumor progression was evaluated by tumor volume measurement by caliper, calculated as the formula below:

$$\text{vol} = \pi/6 * \text{length} * \text{width} * \text{height}$$

Mice were euthanized by carbon dioxide asphyxiation. Animal were dissected then the collected tissues were fixed in 4% PFA for 2 days. Samples were embedded in paraffin then sectioned at 4 μ m and stained with hematoxylin and eosin (H&E). Slides were scanned using an Aperio digital slide scanner (Leica).

All mice were killed once they reached an endpoint according to the IACUC-approved protocols.

METHOD DETAILS

mm10dgLib design

The 250 bp sequences upstream of all mouse protein-coding genes were extracted by Ensembl Biomart. All possible dgRNA spacers (15mer-NGG) were identified, both on the sense and antisense strands. 18mers were then mapped to the mm10 genome by Bowtie with settings bowtie -n 2 -e 120 -best. After discarding any alignments with mismatches in the 17th or 18th position ("GG" in NGG PAM), the number of alignments was tabulated for each spacer. Each alignment was then scored based on the number of mismatches, disregarding any mismatches in the 16th position (the "N" in NGG), using the following formula: (# of 0 mismatch alignments * 1000) + (# of 1 mismatch alignments * 50) + (# of 2 mismatch alignments * 1). Larger mismatch scores indicate decreased genome-wide mapping specificity. Subsequently, all non-uniquely mapping spacers, as well as spacers with mismatch score \geq 2000, were discarded. The top 4 dgRNA spacers were then chosen based on shortest distance to the -200 position from the TSS, since transcription factors generally bind to promoters at around this distance. In total, 83,601 dgRNAs targeting 22,391 coding transcripts and 1,000 non-targeting controls (NTCs) were designed.

Construction of T cell dgRNA activation (TdgA) vector

The gBlocks of U6-dgRNA-MS2 and EFS-Thy1.1-MCP-p65-HSF1 gene fragments were synthesized from Integrated DNA Technologies (IDT), then sub-cloned into lentiviral vectors via Gibson Assembly (NEB). The resulting activation vector, lenti-U6-dgRNA (BsmbI)-MS2-EFS-Thy1.1-MPH, is referred to as TdgA vector.

mm10dgLib cloning

Designed mm10dgLib was synthesized by IDT followed by sub-cloning into TdgA vector via Gibson assembly and electroporation. Approximately > 230x coverage (~2e7 clones were obtained) was achieved after electroporation. The mm10dgLib plasmids were extracted via Maxi preparation (Qiagen), followed by Illumina sequencing confirmation to ensure library representation. A total of 82,197 / 83,601 of GTS and 988 / 1,000 of NTC were cloned.

Lentivirus production

Lentivirus was produced using low-passage HEK239FT cells. One day before transfection, HEK239FT or HEK293T cells were seeded in 15 cm-dish at 50-60 % confluence. 2 h before transfection, D10 media was replaced with 13 mL pre-warmed Opti-MEM medium (Invitrogen). For each plate, 450 µL of Opti-MEM was mixed with 20 µg mm10dgLib plasmid or vector control plasmid, 15 µg psPAX2 (Addgene), 10 µg pMD2.G (Addgene) and 130 µL polyethylenimine (PEI) (1 mg/mL) or 100 µL lipofectamine 2000 (Thermo Fisher). After brief vortex, the mixture was incubated for 15 min at room temperature and then dropwise added to cells. After 6 h of transfection, Opti-MEM media was replaced with 20 mL pre-warmed D10 media. Viral supernatant was collected at 48 h and 72 h post-transfection, then filtered using 0.45 µm filters (Fisher / VWR) to remove cell debris, and then concentrated using AmiconUltra 100 kD ultracentrifugation units (Millipore). All virus was aliquoted and stored in -80°C.

mm10dgLib lentivirus titration

Naive CD8⁺ T cells were infected with lentivirus in different volume ratios between cRPMI and lentivirus after isolation. At day 3 after T cell isolation, T cells were collected and stained with anti-CD8-APC and anti-Thy1.1-PE, the Thy1.1-positive cells represented cells successfully infected by virus and expressing the vector. The ratio of CD8⁺ and Thy1.1⁺ cells was determined by flow cytometry. Viral titer was calculated by comparing mm10dgLib-infected cells to uninfected cells, albeit following the same staining procedures. Each group has 3 independent biological replicates.

mm10dgLib lentivirus transduction

Given the lentiviral titer information, for each replicate, 1.5e7 OT-I;Cas9β naive CD8⁺ T cells were cultured in a 10-cm plate and transduced with 400 µL mm10dgLib or empty vector lentivirus at a volume ratio of 1:25 (virus : media), which can successfully infect around 75 % T cells (Figure S1).

AAV-HDR based CAR and lentiviral-based CAR construction

TRAC homologous recombination arms (HA) were cloned from the both sides of crRNA binding site located in the first exon of *TRAC* locus. CD22-CAR sequence contains a CD22 m971 scFv, CD8 hinge, CD8 transmembrane domain, 4-1BB intracellular domain, and CD3ζ intracellular domain. HER2-CAR sequence contains a 4D5 scFv, CD8 hinge, CD8 transmembrane domain, CD28 and 4-1BB intracellular domains, and CD3ζ intracellular domain. BCMA-CAR sequence contains a BCMA scFv, CD8 hinge, CD8 transmembrane domain, 4-1BB intracellular domain, and CD3ζ intracellular domain. These sequences were synthesized as gBlocks from IDT. *PRODH2* cDNA was purchased from GenScript, then subcloned into an AAV vector. The final vector was named as LHA-EFS-CD22-CAR-PRODH2-RHA and LHA-EFS-HER2-CAR-PRODH2-RHA. Stop codons were engineered at 5' coding regions of *PRODH2* to generate control CAR-T AAV vectors. For the lenti-CAR constructs, EFS-CD22-CAR-PRODH2 / PRODH2 (Stop) and EFS-BCMA-CAR-PRODH2 / PRODH2 (Stop) DNA sequences were PCR amplified from AAV-CAR plasmids then cloned into a lentiviral backbone.

AAV production

Low-passage HEK293FT cells were used for AAV production. Briefly, 2 h before transfection, D10 media was replaced by pre-warmed DMEM (FBS-free). For each 15 cm-dish, HEK293FT cells were transiently transfected with 5.2 µg transfer, 8.9 µg AAV6 serotype and 10.4 µg pDF6 plasmids using 130 µL PEI (1 mg/mL). After 6-12 h of transfection, DMEM was replaced with 20 mL pre-warmed D10 media. Cells were dislodged and transferred to 50 mL Falcon tubes after 48-72 h post-transfection. For AAV purification, 1 / 10 volume of pure chloroform was added and solution was incubated at 37 °C in the shaker with speed of 200 rpm for 1 h. NaCl was added to a final concentration of 1 M, then pelleted at 20,000 x g at 4 °C for 15 min. The aqueous layer was gently transferred to another clean tube and the chloroform layer was discarded. 10 % (w / v) of PEG8000 (Promega) was added and shaken within the tubes until dissolved. The mixture was incubated on the ice for 1 h followed by centrifuge at 20,000 x g at 4 °C for 15 min. The pellet was resuspended with 5-15 mL PBS including MgCl₂ and benzonase (Sigma). After 30-60 min digestion, one volume of chloroform was added, shaken vigorously and spun down at 15,000 x g at 4 °C for 10 min. The aqueous layer was collected carefully and AAV was concentrated using AmiconUltra 100 kD ultracentrifugation units (Millipore). Virus was aliquoted and stored in -80°C. To measure virus titer, RT-qPCR was performed using Taqman assays (ThermoFisher) targeted to EFS promoter engineered in the AAV vector.

Flow cytometry for surface markers

T cells were collected and washed once using MACS buffer (0.5 % BSA and 2 mM EDTA in PBS) before staining. T cells were stained on ice for 30 min after adding antibodies (1:200 dilution). For the CD22-CAR;PRODH2 and CD22-CAR;PRODH2(Stop) KI T cells were incubated with 0.2 µg CD22-Fc (R&D system) per 10⁶ cells in 100 µL PBS for 30 min on ice, then washed with 1 mL cold PBS and stained with FITC anti-human IgG-Fc or PE anti-human IgG-Fc (Biolegend) for 30 min on ice. Pure CD22-CAR;PRODH2 and CD22-CAR;PRODH2(Stop) T cells were established by sorting CD3 negative and CD22-CAR positive populations. For BCMA-CAR;PRODH2 and BCMA-CAR;PRODH2(Stop) KI human CD8 T cells were stained with PE anti-Flag (Biolegend) antibody for 30 min on ice, then washed with 1mL MACS buffer. All samples were run on a BD FACSAria cytometer, and analysis was performed using FlowJo software 9.9.4 (Threestar, Ashland, OR). Cells were gated by FSC/SSC plot. To distinguish between positive and negative boundaries of the stained cells, non-staining or untreated control samples were analyzed and utilized as background.

Cell Metabolism

Article



Intracellular flow cytometry

Intracellular flow cytometry was performed to detect the expression level of cytokines and immune markers. Purified *CD22-CAR; PRODH2* and *CD22-CAR; PRODH2(Stop)* T cells were co-cultured with NALM6-GL cancer cells at 1:1 ratio for 12h. X-VIVO 15 media (Lonza) supplied with 5 % human AB serum, IL-2, and 5 µg/mL Brefeldin A was used as co-culture media. After 12h, at a time point when CAR-T cells killed majority of the cancer cells, CAR-T cells were collected and PBS washed, membrane protein staining was performed prior to intracellular staining. Cells were fixed and permeabilized after membrane protein staining, and then specific antibodies were added.

A list of flow antibodies can be found in [key resources table](#).

Kill assay and dgLib screen

mm10dgLib lentivirus-infected OT-I;Cas9β CD8⁺ T cells were cultured on the 10-cm dishes pretreated with anti-CD3ε in the cRPIMI-1640 supplemented with 2 ng / mL IL-2, 1 µg / mL anti-CD28, and 2 ng / mL IL-12p70 for 4 days, the media were changed with fresh media every day. About 12 hours before the kill assay, infected OT-I;Cas9β CD8⁺ T cells were reseeded onto new dishes without treatment with anti-CD3ε antibody, and cultured with cRPIMI only supplemented with 2 ng / mL IL-2 and 2 ng / mL IL-12p70 to rest cells. At the same time, 2e6 E0771 cells were seeded in 6-well plates in D10 media. On day 5, E0771 cells were incubated with 1 ng / mL SIINFEKL peptide for 4 hours. Before start kill assay, CD8⁺ T cells were suspended with fresh cRPIMI media supplied with 2 nM monensin and anti-CD107a-PE antibody (BioLegend) (1:400 dilution), the final cell concentration was 2e6 cells / mL. After SIINFEKL peptide incubation, 3e6 CD8⁺ T cells per replicate (E0771: T cell = 1:1) were added into E0771 cells for 2 h co-culture. At the end of co-culture, T cells were gently washed down with PBS and stained with anti-CD8α-APC, anti-CD3ε-PE/Cy7 for 30 min on ice, cells were analyzed and sorted using BD FACSAria. CD107a-high cells were sorted by FACS for library readout similar to an *in vitro* T cell cytotoxicity CRISPR screen approach in a previous study ([Dong et al., 2019](#)).

Kill assay screen readout and deep sequencing

Sorted cells and cell pellets (2 x 10⁶ cells / replicate, collected before kill assay) were lysed using QuickExtract solution (Epicentre) at 65 °C for 30 min, then 85 °C for 5 min. dgRNA readout was performed using two-step PCR amplification. PCR was performed using Phusion Flash High Fidelity Master Mix (Thermo Fisher). The first round PCR (PCR #1) used primers to amplify dgRNA cassette:

Forward: 5'-aatggactatcatatgttaccgtaaacttgaaagtatttcg-3',

Reverse: 5'-cttagttgtatgtctgttattatgtctactattctttccc-3'.

The second round PCR (PCR #2) amplification used uniquely barcoded primers, from SF1 to SF12 and SNR1 to SNR12, to allow multiplexing of samples in a single Hiseq run. For PCR #1, each sample underwent 5 reactions in a 50 µL PCR volume to capture all enriched dgRNAs as deeply as possible. The cycle condition was: 98 °C for 1 min, 25 cycles of (98 °C for 1s, 62 °C for 5 s, 72 °C for 15 s), and 72°C for 2 min. PCR products of each sample were pooled together for barcoding PCR. For PCR #2, 2 µL of pooled PCR #1 products were used as template in a 50 µL PCR reaction. The PCR cycle conditions were: 98 °C for 1 min, 30 cycles of (98°C for 1s, 62°C for 5 s, 72°C for 15 s), and 72°C for 2 min. All PCR products were quantified with a gel-based method using the Low-Range Quantitative Ladder (Life Technologies), then gel purified using the QIAQuick gel extraction kit (Qiagen). Purified PCR products were equally pooled in one tube for deep sequencing. Diluted libraries with 5-20 % PhiX were sequenced with HiSeq 2500 or HiSeq 4000 systems (Illumina).

A list of barcoded primers can be found in the [key resources table](#).

dgRNA screen data analysis

Raw single-end fastq read files were filtered and demultiplexed using Cutadapt ([Martin, 2011](#)). To remove extra sequences downstream (i.e. 3' end) of the sgRNA spacer sequences, the following command was used: cutadapt -a GTTTAGAGCTAGGC -e 0.2 –discard-untrimmed. As the forward PCR primers used to readout sgRNA representation, they were designed to have a variety of barcodes to facilitate multiplexed sequencing, demultiplexing these filtered reads was performed with the following settings: cutadapt -g fbc.fasta –no-trim, where fbc.fasta contained the 12 possible barcode sequences within the forward primers. Finally, to remove extra sequences upstream (i.e. 5' end) of the sgRNA spacers, the following command was used: cutadapt -g AAAGGAC GAAACACCG -e 0.2 -m 14 -M 18. Minimum / maximum read lengths were set at 14 / 18 bp respectively to exclude non-specific reads and enriched for the 15 bp dgRNA spacer sequences. The 15 bp dgRNA spacer sequences from each demultiplexed sample were then mapped the designed dgRNA spacers in the mm10dgLib ([Data S1](#)). A bowtie index of the dgRNA library was generated using the bowtie-build command in Bowtie 1.1.2 ([Langmead et al., 2009](#)). The filtered fastq read files were mapped to the index using the following settings: bowtie -v 1 -m 1 mm10dgLib -q <fastq_file>. Using the resulting mapping output, the number of reads that had mapped to each sgRNA within the library was quantified.

Enrichment analysis of dgRNAs

Enrichment analysis was performed using custom R scripts. The library representation of each sample was quantified by the number of read counts. Raw counts were normalized and log-transformed. Mean values were averaged from biological replicates of the same group (plasmid, unsorted cell, or high-kill). FDR values were calculated from the null distribution of the 1,000 NTCs in each variable. Linear regression for the 1,000 NTCs between samples was performed using the lm() function. The points in waterfall and scatter plots were shown at the individual gRNA level.

Immunoblots

Human CD8⁺ T cells with CD22-CAR, CD22-CAR;PRODH2 or CD22-CAR;PRODH2 (Stop) KI were collected and washed with PBS to remove media. Cells were lysed with RIPA lysis buffer supplied with protease and phosphatase inhibitor cocktail (Thermo Fisher) and incubated on ice for 30 min, followed by centrifugation at $\geq 15,000 \times g$ for 15 min at 4°C. The supernatant was collected for protein quantification. The total protein concentration was quantified by performing a Bradford protein assay (Bio-Rad), a total of 10 µg protein per sample was loaded into SDS-PAGE gel (Bio-Rad), proteins in the gel were transferred into Amersham Protran 0.45 µm NC Nitrocellulose Blotting membrane (GE Healthcare) after electrophoresis. Membranes were blocked with 2 % BSA in TBST buffer for 1 h at room temperature, followed by the primary antibody incubation at 4°C overnight. Anti-PRODH2 antibody was purchased from Atlas Antibodies (HPA051287) (1:1000 dilution) and anti-Vinculin was purchased from Abcam (ab129002) (1:2000 dilution) which was used as internal control. Antibody binding was detected using horseradish peroxidase-conjugated secondary antibody and ECL substrate (Bio-Rad).

Cytokine secretion assays

CAR-T cells and cancer cells were co-cultured at a E:T = 1:1 ratio for 12h in 200 µL X-VIVO 15 T cell media. The supernatant was collected and analyzed for IL-2, IL-15, and IL-7 secretion using ELISA kits (ThermoFisher), the protocols were provided by the manufacturer.

RT-qPCR

For cDNA overexpression, 48 h and 72 h after lentivirus transduction, OT-I CD8⁺ T cells were collected for RNA preparation. Human CD8⁺ T cell PRODH2 overexpression and CD22-CAR;PRODH2 knock-in (KI) qPCR verification were performed as described in the results. All RNA preparations were performed using RNasy Plus Mini Kit (Qiagen). Total mRNA was reverse transcribed into cDNA by using SuperScript IV Reverse Transcriptase (Thermo Fisher). Gene expression was quantified using Taqman Fast Universal PCR Master Mix (Thermo Fisher) and Taqman probes (Invitrogen). RNA expression level was normalized to *Gapdh* / GAPDH or *Actb* / *ACTB* (mouse / human). Relative mRNA expression was determined via the $\Delta\Delta C_t$ method. For the relative human mitochondrial DNA copy number quantification, CD22-CAR;PRODH2 and CD22-CAR;PRODH2(Stop) T cells were collected for DNA isolation (Qia- gen). The relative human mitochondrial DNA copy number was quantified by using a qPCR assay kit (ScienCell). The mtDNA primer set provided with the kit was used to recognize and amplify one of the most conserved regions on human mtDNA and will not amplify off-target sequence on nuclear genomic DNA. The reference primer set was used for amplifying a 100 bp-long region on human chromosome 17 and serve as reference for data normalization. Twenty microliter qPCR reactions were set up and a quantification method was used based on the protocol provided by the manufacturer.

A list of probes can be found in [key resources table](#).

Human primary CD8⁺ T cell electroporation and CAR-T knock in

Human primary CD8⁺ T cells were isolated from healthy donor PBMCs (StemCell). CD8⁺ T cells were cultured in X-VIVO 15 media (Lonza) supplied with 5 % human AB serum and IL-2. Before electroporation, TRAC crRNA (5'-TCTCTCAGCTGGTACACGGC-3') and tracrRNA were mixed in 1:1 ratio (final concentration 50 µM), heated at 95 °C for 5 min in the thermal cycler, then cooled to room temperature. 3 µL Cas9 protein (61 µM) was mixed with 2 µL Buffer R for each reaction (Neon Transfection System Kit, Thermo Fisher), then mixed with 5 µL annealed crRNA:tracrRNA duplex, incubated the mixture at room temperature for 10-20 min. Human CD8⁺ T cells were collected and washed with PBS to completely remove the media. 3 x 10⁶ T cells per reaction were resuspended in 100 µL Buffer R which included 10 µL RNP complex. 100 µL of cell:RNP mixture was loaded into the Neon Pipette without any bubbles. The electroporation parameter was set at 1600 V, 10 ms for 3 pluses. Cells were immediately transferred to a 24-well plate with pre-warmed media after electroporation. A total of ~1.5e9 viral genome copy of AAV6 HDR donor was added into each electroporated T cell reaction within 1 h after electroporation.

Top candidate validation in kill assay

Candidate hits mouse *Prodh2*, *Ccnb1ip1*, *Srek1ip1*, and *Wdr37* were selected for further validation. All cDNAs were purchased from the Dharmacon, then cloned into a lentiviral overexpression vector (lenti-EF1a-Flag-WPRE vector). Before lentivirus transduction, OT-I CD8⁺ T cells were isolated and activated by anti-CD3ε and anti-CD28 antibodies for 2-3 days, then T cells were transduced with concentrated lentivirus. At day 6, infected OT-I CD8⁺ T cells were reseeded onto new 6-well plates which were untreated with anti-CD3ε antibody and cRPML only supplemented with 2 ng / mL IL-2 and 2 ng / mL IL-12p70 to rest cells. At the same time, 5e5 / well of E0771 cells were seeded in 24-well plates for kill assay. The next day, E0771 cells were incubated with 1 ng/mL or 10 ng / mL SIINFEKL peptide for 4 hours. The validation kill assay was performed in the same manner as the screen kill assay.

IL-2 withdrawal assay

Mouse OT-I;Cas9β CD8 T cells were activated with anti-CD3ε and anti-CD28 antibodies, T cells were transduced with Prodh2-OE or Vector lentivirus after T cells were completely activated. At day 3 after lentivirus transduction, T cells were collected and washed with PBS, then equal cell numbers were plated in media without IL-2. Cells were stained with PE anti-caspase 3 (Cell signaling technology), FITC anti-CD3, and APC anti-CD8 antibodies at day 1 and 4 after IL-2 withdrawal. To test PRODH2 GOF CAR-T cell potential

cancer-like transformation, the same number of *CD22-CAR;PRODH2* and *CD22-CAR;PRODH2(Stop)* T cells were plated in X-VIVO15 media without IL-2 cytokine, then T cell survival was measured after 7 days by cell counting.

Human *CD22-CAR;PRODH2* and *HER2-CAR;PRODH2* T cell kill assay

To detect CAR-PRODH2 T cell killing, NALM6-GL, NALM6-GL-CD22OE, MCF7-PL, MCF7-PL-HER2OE or MDA-MB-231-PL cancer cell lines were established as described above, cancer cells were seeded in a 96-well plate first, then different Effector : Target (T cell : cancer cell) ratio (E : T ratio) co-cultures were set up. Cytolysis was measured after co-culture by adding 150 µg / mL D-Luciferin (PerkinElmer) using a multi-channel pipette. Luciferase intensity was measured by a Plate Reader (PerkinElmer).

Lenti-*CD22-CAR;PRODH2* and Lenti-*BCMA-CAR;PRODH2* T cell transduction and co-culture

Lentivirus was produced by HEK293T cells, lentiviral supernatant was collected and precipitated by Lenti-X Concentrator (Takara). Lentiviral pellets were resuspended with X-VIVO15 media (LONZA), then aliquoted and stored in -80°C. Human CD8 T cells were transduced with lentivirus onto 1-2e6 T cells plated in a 24-well plate which was pre-coated with Retronectin (Takara) in PBS overnight in 4°C. The spin-infection was performed at 32°C at 900 x g for 90min. The CAR-positive T cells were measured at day3 after transduction, then co-culture assays were performed to determine CAR-T cell killing.

Bulk mRNA sequencing (mRNA-seq) library preparation

The mRNA library preparations were performed using a NEBNext Ultra RNA Library Prep Kit for Illumina and samples were multiplexed using barcoded primers provided by NEBNext Multiplex Oligos for Illumina (Index Primers Set 1). For the human CD8⁺ T cell RNA-seq, *CD22-CAR;PRODH2* and *CD22-CAR;PRODH2(Stop)* cassettes were site-specifically KI into *TRAC* locus, the flow cytometry, qPCR, and western blot were performed to confirm that *CD22-CAR-PRODH2* or *CD22-CAR-PRODH2(Stop)* were successfully KI and expressed before doing the mRNA-seq library preparations. Libraries were sequenced with HiSeq 4000 or Novaseq systems (Illumina).

Bulk mRNA-seq data processing

Raw FASTQ files from mRNA-seq were analyzed for transcript quantification using Kallisto quant algorithm (Bray et al., 2016) with the setting -b 100. Transcriptome references were obtained from Ensembl. Differential gene expression analysis for the effect of *PRODH2* overexpression or genomic knock in was then performed using Sleuth (Pimentel et al., 2017) with gene-level aggregation. Visualization of differentially expressed (DE) genes including volcano plots, bar plots, and Venn diagrams were performed using standard R packages including ggplot2 and VennDiagram.

Gene set level pathway analysis of differentially expressed genes

Gene set enrichment analysis (GSEA) (Subramanian et al., 2005) was performed using the Java application from the Broad Institute. The full gene set from the differential gene expression analysis was ranked by “beta” value and then used as an input for GSEA pre-ranked analysis with database reference C5 Gene Ontology - Biological Process (GO-BP). The Database for Annotation, Visualization and Integrated Discovery (DAVID) (Huang et al., 2009) was used for further annotation enrichment analysis. Upregulated and downregulated genes from the differential gene expression analysis were defined with adjusted p value cutoff of 0.001 for human CAR-T experiments and 0.05 for mouse experiments. Pathway enrichments for GO-BP annotations were used for visualization.

Metabolite extraction and data collection

For extraction of intracellular metabolites, cell culture media was first aspirated, then harvested and washed twice with PBS. 2e6 alive cells for each sample were used for metabolite extraction. After normalizing cell counts, 800 µL of 80 % (vol / vol) HPLC-grade methanol (Sigma) (precooled to -80 °C on dry ice) was added to fresh cells in a 1.5-mL microcentrifuge tube, then tubes were put on dry ice for 30 minutes (Yuan et al., 2012). The tubes were then incubated on ice for 20 minutes and centrifuged at 15,000 x g for 15 min at 4°C to pellet the cell debris. The metabolite-containing supernatant was transferred to a new 1.5-mL microcentrifuge tube on dry ice. Metabolite extraction was repeated with 400 µL of 80 % (vol / vol) HPLC-grade methanol. The cell lysate / methanol mixtures were dried by Speedvac at room temperature. The metabolites were dissolved again with 80 % (vol / vol) methanol, then centrifuged at 18,000 x g for 10 min to remove any particulates, and the metabolite mixtures were stored at -80 °C until LC-MS analysis. For the metabolite analysis, the untargeted metabolic profiling was firstly analyzed with an Agilent 6550 Q-TOF LC/MS System first, then targeted metabolites were analyzed with an Agilent 6490 Triple quadrupole (QQQ) LC/MS System. Multiple reaction monitoring (MRM) was employed for the quantitation of purified standard (Sigma). A HILIC liquid chromatograph were optimized with a bioZen 2.6 µm Glycan LC Column, 150 x 2.1 mm (Phenomenex) and a Glycan guard column, 4 x 2 mm (Phenomenex). The eluents included buffer A, 0.01 % formic acid in HPLC-grade water, and buffer B, 0.01 % formic acid in acetonitrile. The gradient was set as follows: 0-2 min 94 % B, 2-8 min 94-90 % B, 8-16 min 90-76 % B, 16-36 min 76-50 % B, 36-42 min hold at 50 % B and then back to initial conditions for 2 min for column equilibration. The flow rate was set as 0.3 mL / min. Multiple reaction monitoring (MRM) was employed for the quantitation of purified standard (Sigma).

A list of standard metabolites used in this study can be found in [key resources table](#).

Metabolomics data processing

Two metabolomics strategies were adopted, i.e. untargeted metabolomics (aiming to unbiasedly detect all detectable metabolites) and targeted approaches (aiming to detect specifically defined metabolites, such as related metabolites in the proline metabolism and T cell metabolism). For untargeted metabolomics analysis, the optimized workflow consists of automated peak detection and integration, peak alignment, background noise subtraction, and multivariate data analysis. These steps were carried out for comprehensive metabolite phenotyping of the two groups using Agilent Mass Hunter Qualitative Analysis Software (Version B.07.0.0, build 7.0.7024.0) and Agilent Mass Profiler Professional (Version 14.5-Build 2772). The metabolites were first putatively identified based on accurate mass match (accurate mass \pm 30 ppm error) and fragmentation pattern match. Putative structural annotation was carried out by searching the metabolite databases HMDB (<http://www.hmdb.ca/>) and METLIN (<http://metlin.scripps.edu>) using the mass-to-charge ratio of the metabolic features. For the targeted metabolomics, available metabolites from the significantly changed metabolites of untargeted metabolomics analysis, as well as related proline metabolism and immune system metabolism, were purchased from Sigma. Multiple reaction monitoring (MRM) was employed for the qualitative and quantitative analysis of purified standard (Sigma). The features of spectra were extracted using Agilent Mass Hunter Qualitative Analysis Software (Version B.07.0.0, build 7.0.7024.0). Each peak was manually checked and the abundances of all metabolites were exported. The retention time of the standards was cross-referenced with those detected in the untargeted method for consistency, which also confirmed the accuracy of the untargeted methods. The integration of untargeted and targeted metabolites includes the combination of non-overlapping metabolites and the selection of targeted metabolites of overlapping metabolites. Multivariate data analyses were conducted using Statistical analysis module of MetaboAnalyst 4.0 (Chong et al., 2018; Xia et al., 2015). Briefly, Log transformation and auto scaling (mean-centered and divided by the standard deviation of each variable) were used for data processing. Heatmaps and volcano plots were generated plots with MetaboAnalyst 4.0. Distance Measure was set as Euclidean and the Clustering Algorithm was set as Ward. The functions of the metabolic flowchart were constructed with the software Pathvisio v3.3.0 based on the KEGG database (<http://www.genome.jp/kegg/>). The integrated analysis of the changed metabolites and genes were done with Joint Pathway analysis module of MetaboAnalyst 4.0.

Intersection of mRNA-seq, metabolomics and KEGG pathways

Differential expressed genes from mRNA-seq analysis were cross-referenced with expected differential expressed genes from metabolomics analysis in order to obtain consensus upregulated and downregulated gene sets due to *PRODH2* overexpression or genomic knock in. For a given metabolite, genes contributing to its production or consumption were defined using the KEGG (Kyoto Encyclopedia of Genes and Genomes) Pathway Database (Kanehisa and Goto, 2000). Genes could be both "producing" and "consuming" when involved in reversible reactions. Expected upregulated genes were defined as those producing metabolites with positive logFC or those consuming metabolites with negative logFC. Expected downregulated genes were defined as those producing metabolites with negative logFC or those consuming metabolites with positive logFC. For consensus "upregulated" genes, first the union of downregulated mRNA-seq genes from human donor 1 and donor 2 from the CAR-T experiments were removed from the union of upregulated mRNA-seq genes (setdiff). Then the resulting restricted gene set was intersected with expected upregulated metabolomics genes. For consensus "downregulated" genes, the union of upregulated mRNA-seq genes was removed from the downregulated mRNA-seq genes (setdiff). Then the resulting restricted gene set was intersected with expected downregulated metabolomics genes.

Multi-Omics analysis

Consensus differential expressed genes from intersection analysis and differentially represented metabolites with fold changes from metabolomics analysis were used as inputs for joint pathway analysis using the MetaboAnalyst Portal (Xia et al., 2009). Default parameters were used, with Hypergeometric Test for enrichment analysis, Degree Centrality for topology analysis, and Gene-metabolite pathways for pathway databases. Pathways were considered statistically significant if the p values were less than 0.05. For visualization, upregulated gene set and downregulated gene set were separated, each compared against DR metabolites (both increased and decreased, as the gene activity can influence on either direction), using the MetaboAnalyst Portal.

Transmission electron microscopy (TEM) of CAR-T cells

Purified CAR-T cells were collected and washed with PBS, then fixed in 2.5% gluteraldehyde in 0.1M sodium cacodylate buffer pH7.4 for 1 hour. Buffer rinsed cells were spun down in 2% agar and the chilled blocks were trimmed and post fixed in 1% osmium tetroxide for 1 hour then the sample was rinsed in buffer and stained in aqueous 2% uranyl acetate for 1 hour. This was followed by rinsing in distilled water, dehydrating in an ethanol series and infiltrating with Embed 812 (Electron Microscopy Sciences) resin and baked over night at 60 °C in silicone molds. Hardened blocked were sectioned using a Leica UltraCut UC7, 60nm sections were collected on formvar and carbon coated nickel grids and contrast stained using 2% uranyl acetate and lead citrate. The sections were viewed FEI Tencaj Biotwin TEM at 80kV. Images for quantitation were collected randomly and viewed using a Morada CCD and iTEM (Olympus) software. Quantification of TEM data were performed in randomized fields with anonymized images.

Cell Metabolism

Article



Seahorse assay

Oxygen consumption rates (OCR) and extracellular acidification rate (ECAR) were measured in XF media (Agilent) supplied with 25mM glucose, 2mM L-glutamine, 1mM sodium pyruvate, and human IL2 after *CD22-CAR;PRODH2* and *CD22-CAR;PRODH2(Stop)* T cells culture for two months after electroporation, one month after cancer stimulation. 1 μ M oligomycin, 1.5 μ M fluoro-carbonyl cyanide phenylhydrazone (FCCP) and 50 nM rotenone / 0.5 μ M antimycin A were used for testing extra mitochondrial capacity, spare respiratory capacity (SRC), under stress, and measuring extracellular acidification rate (ECAR). Different CD22-CAR T cell densities (2e5 and 4e5 / well) were seeded into XF cell culture microplate (Agilent) that were pre-coated with poly-D-lysine (Sigma) which can facilitate cell adhesion to the plate surface for SRC detection. The microplate was incubated for 30-60 min at 37 °C in a non-CO₂ incubator before running in a Seahorse XF96 Analyzer (Agilent). Standard Seahorse program setup was used: Calibration; Equilibration; Base line reading (Loop 3 times), Mix 3 min, Measurement 3 min, End loop; Injection Port A (Loop 3 times), Mix 3 min, Measurement 3 min, End loop; Injection Port B (Loop 5 times), Mix 3 min, Measurement 3 min, End loop; Injection Port C (Loop 3 times), Mix 3 min, Measurement 3 min, End loop; End Program.

Mass cytometry (CyTOF)

Purified *CD22-CAR;PRODH2* and *CD22-CAR;PRODH2(Stop)* T cells (without cancer stimulation) were collected and washed with PBS, resuspended cell to 1e7 / mL in PBS and Cell-ID Cisplatin (Fluidigm) was added to a final concentration of 5 μ M. Cells were incubated at room temperature for 5 min, then washed with Maxpar Cell Staining Buffer (Fluidigm). 1.5e6 CAR T cells per replicate were used for staining, each group has three replicates. Cells were stained with surface marker antibody cocktail first, then fixed and permeabilized. Second round staining was performed using cytoplasmic / secreted antibody cocktail. Finally, cells were incubated in intercalation solution (Fluidigm) in a final concentration of 125 nM, then incubated overnight at 4°C. Before running on a CyTOF machine, cell concentrations were adjusted to 5-7e5/ mL with water. All data were collected on a CyTOF Helios instrument (Fluidigm). All surface and cytoplasmic / secreted antibodies were purchased from Fluidigm or Yale CyTOF core.

A list of CyTOF antibodies used in this study can be found in [key resources table](#).

CyTOF data analysis

CyTOF quality prefiltering was performed in FlowJo with CD3 and CD8 gates. Channel values were exported and analyzed with custom scripts in R. Dimensionality reduction by Uniform Manifold Approximation and Projection (UMAP) using the umap package and figures were drawn using ggplot2.

Mitochondria mass and depolarization measurement

CD22-CAR;PRODH2 and *CD22-CAR;PRODH2 (Stop)* T cells were cultured in X-VIVO15 media supplied with human IL-2. T cells were stimulated with NALM6-GL-CD22OE cancer cells at a 1:1 ratio. At day 36 after cancer stimulation, T cells were stained with MitoTracker GFP FM and MitoTracker Deep Red FM dyes (ThermoFisher) to measure mitochondrial mass and depolarization.

CAR T cell chemical treatment, co-culture and flow assays

CD22-CAR;PRODH2 and *CD22-CAR;PRODH2 (Stop)* T cells were pre-treated with P4HA substrate L-Proline (Sigma) or PRODH2 substrate 4-Hydroxyproline (or 4Hyp) (Sigma) for 5 days. T cells were then washed with PBS and used to perform co-culture assay with NALM6-GL-CD22OE cancer cells at a 0.5:1 E:T ratio. T cells also pre-treated with a P4HA1 and P4HA2 inhibitor, 1,4-DPCA (Santa Cruz), or a GOT1 and GOT2 inhibitor, PF04859989 (Axon Medchem) for 3-6 days depending on specific assays, then subjected to the co-culture as described. Flow cytometry was carried out after co-culture to detect T cell immune markers as specified. T cell viability measurement was performed by the live-dead staining after T cells were treated with different concentration 1,4-DPCA as specified for 3 days.

A list of flow antibodies can be found in [key resources table](#).

Long-term CAR-T culture with chronic antigen stimulation and memory marker analysis

To measure *CD22-CAR;PRODH2* and *CD22-CAR;PRODH2 (Stop)* T cell memory phenotypes under chronic antigen stimulation in long-term culture, purified CAR T cells were stimulated with NALM6-GL-CD22OE cancer cells every 12 days at an E:T ratio = 1:1, for up to 3 times. CAR T cells were harvested at day 69 after various times of antigen stimulation, and stained with anti-CD45RA and anti-CD62L antibodies.

A list of flow antibodies can be found in [key resources table](#).

QUANTIFICATION AND STATISTICAL ANALYSIS

Sample sizes for animal experiments were determined with estimated power to detect significance similar to prior work, cited literature, or similar approaches in the field. Sample sizes for certain experiments, such as in vitro assays, were not predetermined by power calculations, and were predetermined according to the lab's prior work, cited literature, or similar approaches in the field. Most experiments were done with at least two biological replicates. Experimental replications were indicated in detail in the relevant sections above and in each figure panel's legend. In animal experiments, mice were randomized by sex, cage and littermates. For the

in vivo validation experiments, tumor-bearing mice were normalized by tumor burden, either caliper-measured tumor size or bioluminescence signal, before being allocated into difference groups. If tumors were unmeasurable (early timepoint) at time of grouping, all mice were randomized by sex, cage, and littermates without considering tumor burden and allocated into control and treatment groups. *In vitro* cell culture experiments were not randomized. Investigators were not blinded for the *in vitro* cell culture experiments. Investigators were blinded for the TEM experiments, including sample preparation, image capture, tumor burden measurements, and data quantification. In NGS data analysis, investigators were blinded for initial processing of the original data using key-coded metadata. Various standard statistical analyses were performed. All statistical methods are described in the figure legends and/or supplementary Excel tables. The p values and statistical significance were estimated for all analyses. The unpaired *t* tests, two sided, was used to compare two groups. Multiple *t* tests and two-way ANOVA were used to compare multiple groups. Survival curves were compared using the log-rank (Mantel-Cox) tests. Most data showed normal or near-normal distribution and are continuous, in those cases parametric tests were performed. In the cases where data were not normal, transformations such as log were performed to approach normality. In the cases where data were not normal after transformation, or discrete, non-parametric tests were performed. Different levels of statistical significance were accessed based on specific p values and type I error cutoffs (0.05, 0.01, 0.001, 0.0001). Non-NGS standard analysis was performed using GraphPad Prism and RStudio. No data was excluded in this study.

Supplemental information

**A genome-scale gain-of-function CRISPR screen
in CD8 T cells identifies proline metabolism
as a means to enhance CAR-T therapy**

Lupeng Ye, Jonathan J. Park, Lei Peng, Quanjun Yang, Ryan D. Chow, Matthew B. Dong, Stanley Z. Lam, Jianjian Guo, Erting Tang, Yueqi Zhang, Guangchuan Wang, Xiaoyun Dai, Yaying Du, Hyunu R. Kim, Hanbing Cao, Youssef Errami, Paul Clark, Alexey Bersenev, Ruth R. Montgomery, and Sidi Chen

Supplemental information

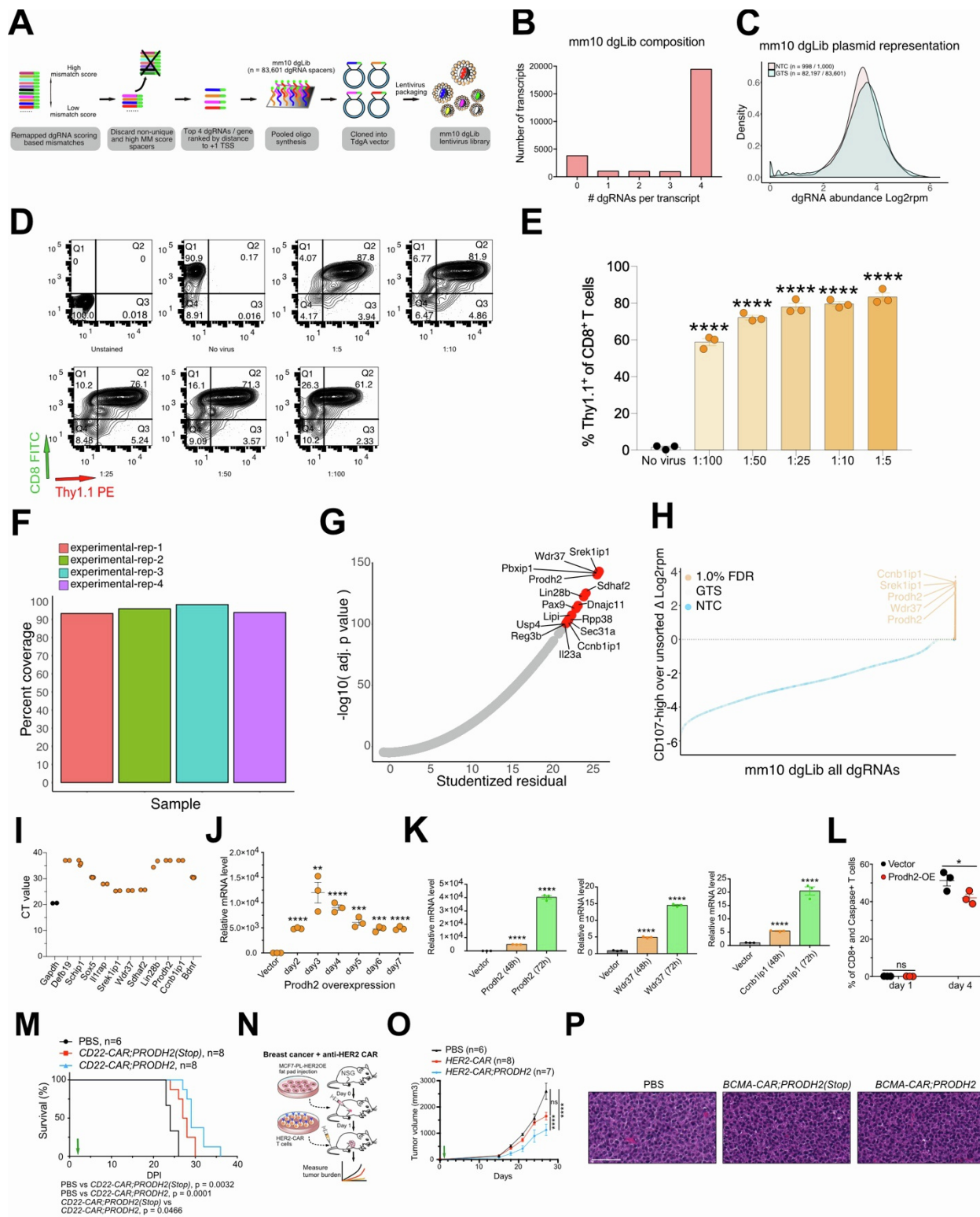


Figure S1. Additional experiments for dgRNA-based T cell CRISPRa GOF screening
(Related to: Figures 1, 2, 3)

- (A)** Schematics of detailed mm10dgLib design process, library cloning, and lentivirus production following Figure 1B.
- (B)** mm10dgLib designed composition. Most coding transcripts (19,440 / 26,205) have 4 dgRNAs. Small portions of coding transcripts only have ≤ 3 dgRNAs that pass the same criteria (3,814, 1,010, 992 and 949 transcripts have 0, 1, 2, 3 dgRNAs, respectively).
- (C)** mm10dgLib plasmid library representation analysis by Illumina sequencing of dgRNA spacers from plasmid. 82,197 / 83,601 (98.3%) of gene-targeting spacers (GTSs) and 988/1,000 (98.8%) of NTCs were successfully cloned. Both GTSs and NTCs in the mm10dgLib showed a log-normal distribution.
- (D)** Representative flow cytometry results of the mm10dgLib lentivirus titration by detecting Thy1.1 expression at day 3 after lentivirus transduction ($n = 3$ biological replicates). The results showed that the mm10dgLib lentivirus was successfully produced with high titer.
- (E)** Quantitative analysis of flow cytometry results.
- (F)** A barplot of mm10dgLib library coverage in CD8 T cell pool after lentivirus transduction in four independent experiments, at experimental replicate level.
- (G)** A volcano plot of $-\log_{10} p\text{value}$ vs. residual for mm10dgLib library based on outlier test against log-linear regression between the CD107a⁺-high FACS sorted CD8⁺ T cells and the unsorted T cells. The points were shown at the individual gRNA level. Representative top scoring genes targeted by specific sgRNAs were shown.
- (H)** A dotted waterfall plot of mm10dgLib library differentials between the CD107a⁺-high FACS sorted CD8⁺ T cells and the unsorted T cells (delta). Blue dots are NTCs; yellow dots are scoring GTSs that passed FDR 1% cutoff; grey dots are remaining GTSs. The points were shown at the individual gRNA level. Representative top scoring genes targeted by specific sgRNAs were shown.
- (I)** RT-qPCR for baseline mRNA level of high-ranked hit genes from dgTKS primary screen ($n = 2 - 3$ biological replicates, variable between genes, see figure for datapoints). Higher CT values indicate the candidates' mRNA levels as lower than house-keeping gene *Gapdh* in primary murine CD8⁺ T cells.
- (J)** Time-course qPCR detection of *Prodh2* mRNA level in lentiviral-Prodh2-OE mouse CD8 T cells ($n = 3$ biological replicates).

(K) RT-qPCR for overexpression of *Prodh2*, *Wdr37* and *Ccnb1ip1* after lentiviral transduction of mouse CD8 T cells (n = 3 biological replicates).

(L) Quantification of mouse T cell apoptosis in an IL-2 withdrawal assay. Cleaved caspase 3 was detected by flow cytometry at day 1 and day 4 after IL-2 withdrawal (n = 3 biological replicates).

(M) Overall survival of PRODH2 CAR22 in a leukemia model. Green arrow indicated CAR-T injection.

(N) A schematic of breast cancer model PRODH2 knock-in CAR-T efficacy testing, showing tumor induction, HER2-CAR T cell intravenously injection, and survival.

(O) Tumor growth curve of breast cancer bearing NSG mice after HER2-CAR;PRODH2 or HER2-CAR T therapy. Green arrow indicated CAR-T injection.

(P) H&E staining of myeloma tumor tissues from end-point mice after CAR-T therapies. Scale bar = 50 μ m.

*P<0.05, **P<0.01, ***P<0.001, ****P<0.0001, ns = not significant by unpaired t tests (E, J, K), multiple t tests (L), two-way ANOVA (with multiple comparisons test) (O) or log-rank (Mantel-Cox) tests (M).

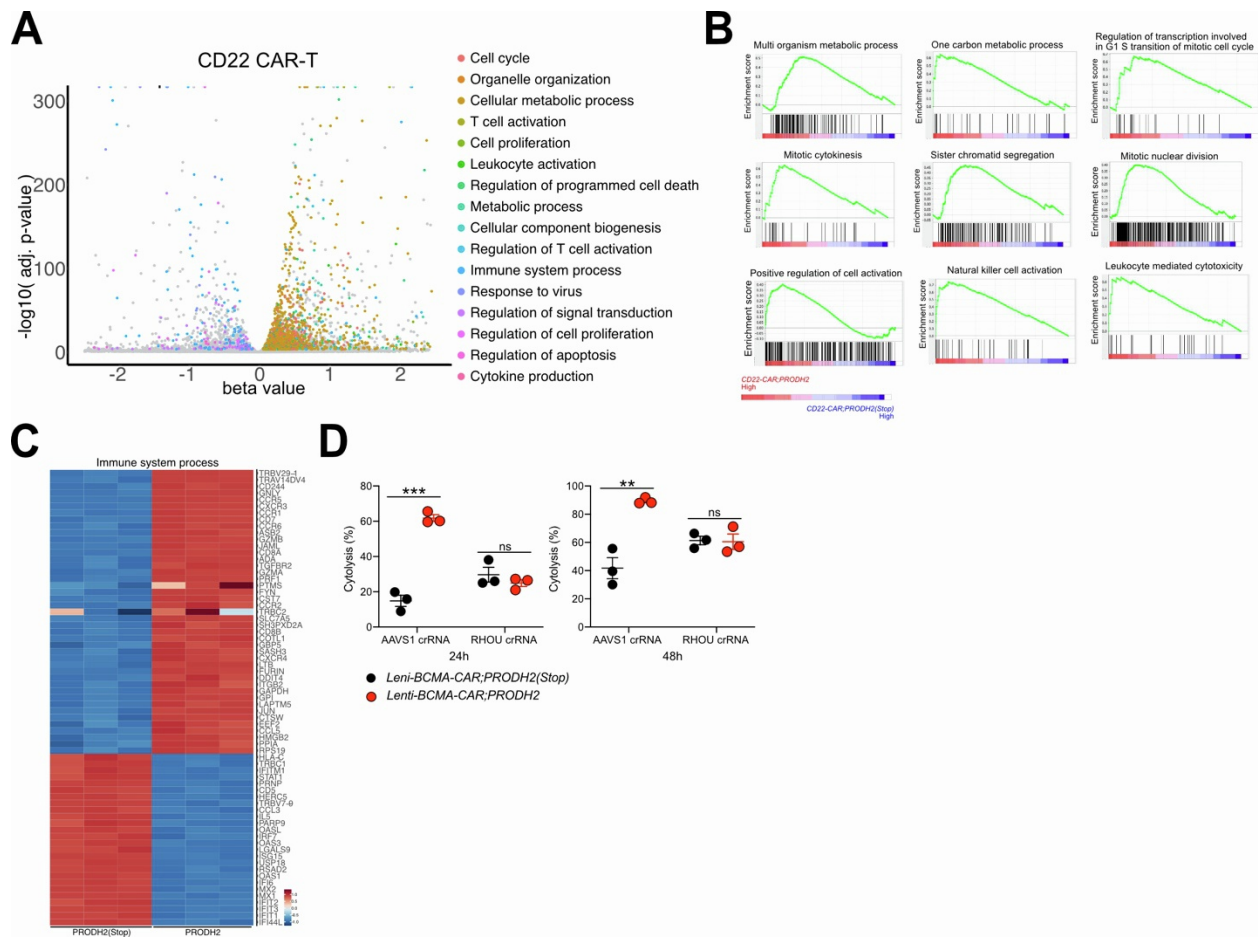


Figure S2. Additional transcriptome profiling and downstream analysis of PRODH2 knock-in CD22-CAR-T cells (Related to: Figure 4)

(A) Color-coded volcano plots of representative top enriched pathways between *CD22-CAR;PRODH2(Stop)* and *CD22-CAR;PRODH2* cells. Upregulated and downregulated pathways were annotated from DAVID analysis of the biological processes of upregulated and downregulated genes, respectively (n = 3 biological replicates).

(B) GSEA individual pathways for mRNA-seq of PRODH2 knock-in CD22-CAR-T cells. GSEA plots of individual pathways from the representative up- and down-regulated gene sets between *CD22-CAR;PRODH2* and *CD22-CAR;PRODH2(Stop)* T cell groups. (Cut-off criteria: p-value < 0.001)

(C) Heatmaps of differentially expressed genes in immune system process.

(D) Genetic testing of knockout effect of representative upregulated genes on PRODH2 overexpressed CAR-T cells. Cytolysis of PRODH2 or PRODH2(Stop) knock-in CD22-CAR-T

cells, with or without *RHOU* knockout, at two time points (left panel, 24h; right panel, 48h) (n = 3).

P<0.01, *P<0.001, ns = not significant by multiple *t* tests (D).

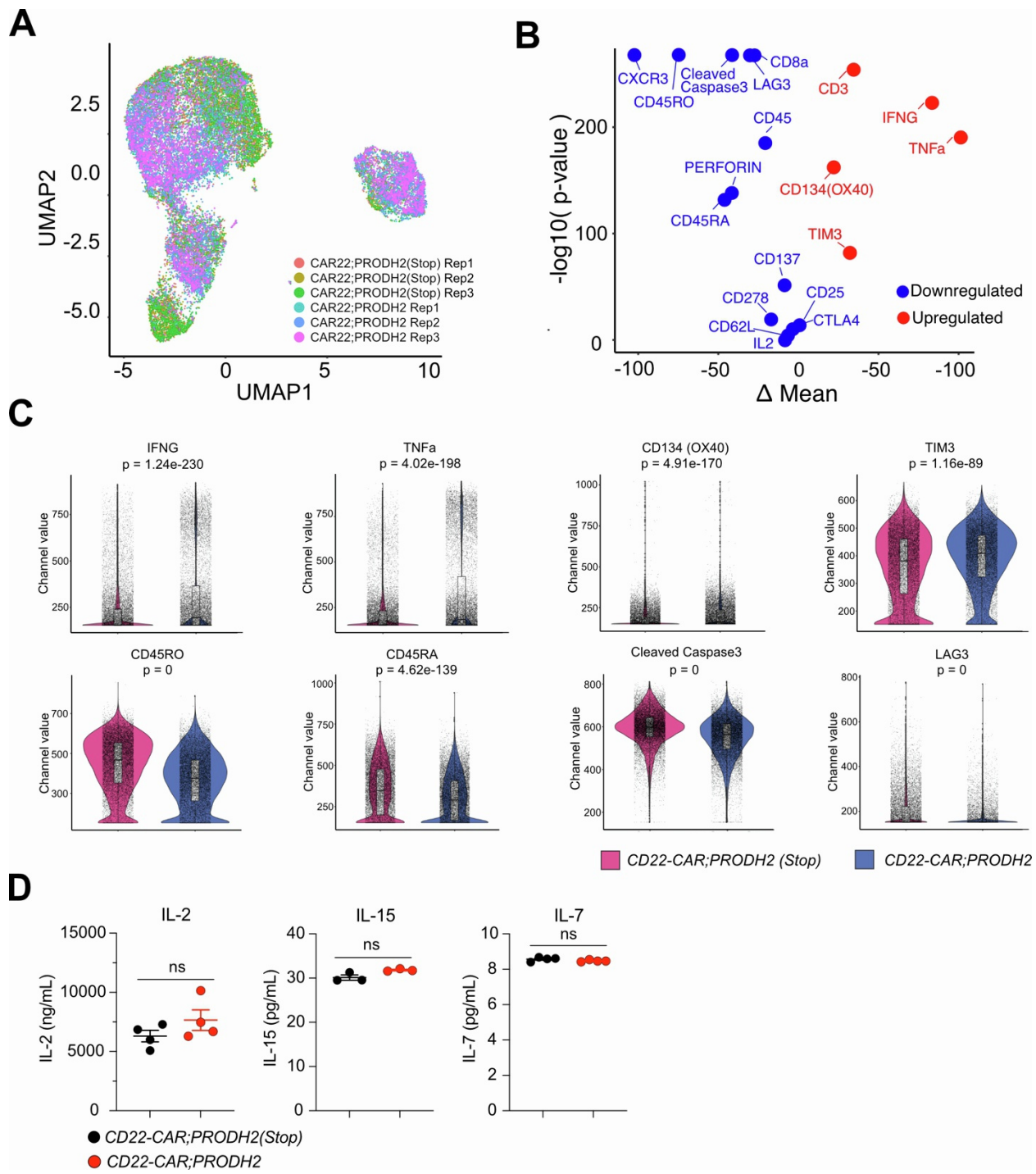


Figure S3. Baseline CyTOF analysis of CD22-CAR T cells with PRODH2 overexpression (Related to: Figure 4)

(A) UMAP plots labeled by individual biological samples. All T cells are at baseline without cancer stimulation in this assay.

(B) Volcano plot of CyTOF data of *CD22-CAR;PRODH2* vs. *CD22-CAR;PRODH2(Stop)* T cells.

(C) Violin plots of CyTOF data of *CD22-CAR;PRODH2* vs. *CD22-CAR;PRODH2(Stop)* T cells.

Each panel represents one individual marker, including IFNG, TNFa, CD134 (OX40), TIM3, CD45RO, CD45RA, Cleaved Caspase 3, and LAG3. Violin plots were used for quantification of gene expression level in single cell level. Wilcoxon Rank Sum test adjusted by Benjamini & Hochberg method was used to assess the significance.

(D) ELISA analysis of IL-2, IL-15, and IL-7 for *PRODH2* knock-in and control CAR-T cells. (n = 4 biological replicates)

ns = not significant by unpaired *t* tests (D).

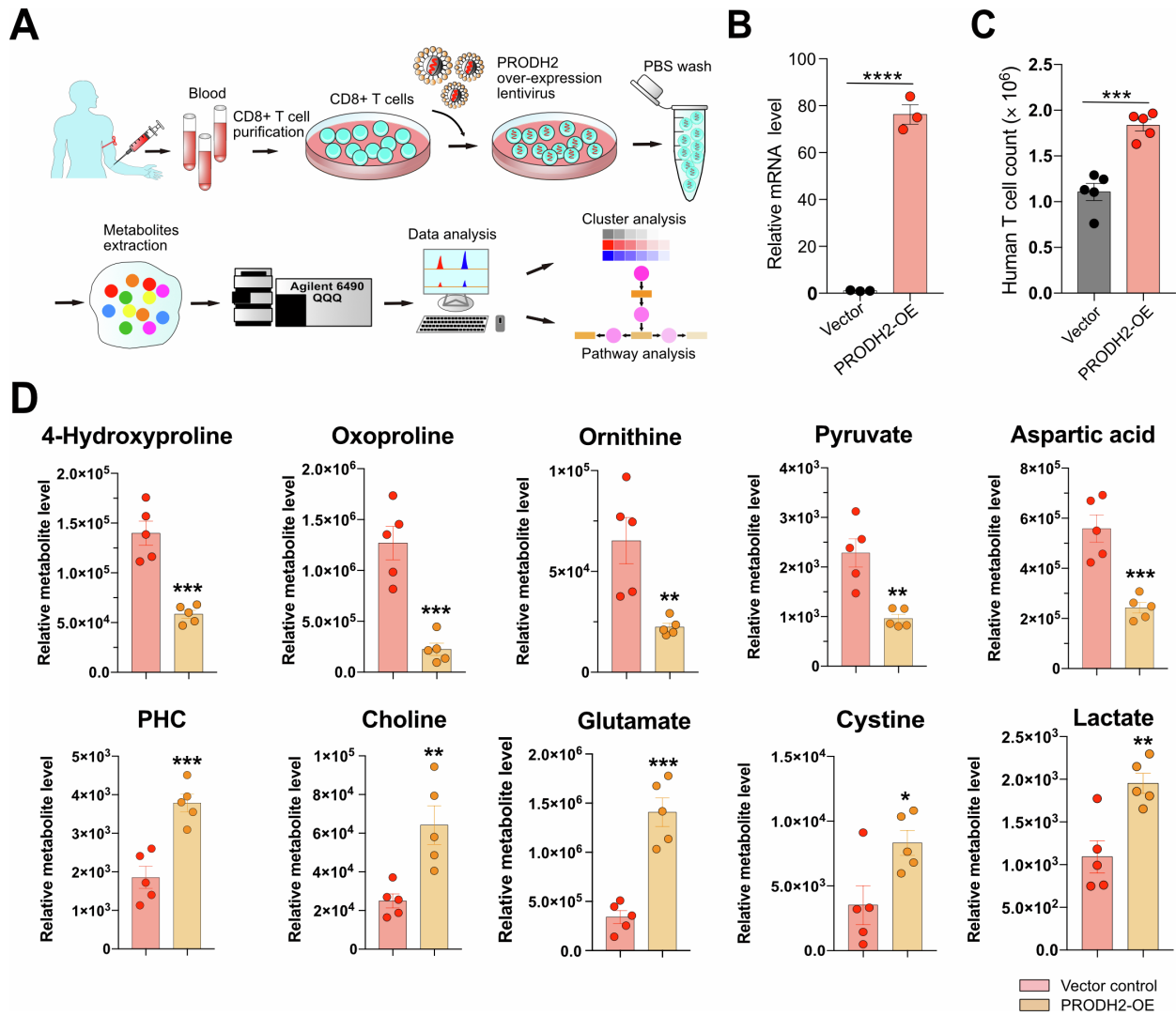


Figure S4. Metabolomic analysis of PRODH2-OE in human primary CD8 T cells (Related to: Figure 5)

- (A) A schematic of human CD8 T cell lentivirus transduction, metabolites extraction, LC/MS analysis, and data analysis.
- (B) Transcriptional level of human *PRODH2* in primary CD8 T cells (no CAR) after PRODH2-OE lentivirus transduction (n = 3 biological replicates).
- (C) Human T cell number quantification after PRODH2-OE lentivirus transduction (n = 5 biological replicates). Representative data from two independent experiments.
- (D) Quantification of significantly changed metabolites after PRODH2-OE (n = 5 biological replicates).

P<0.01, *P<0.001 by unpaired t tests (B-D).

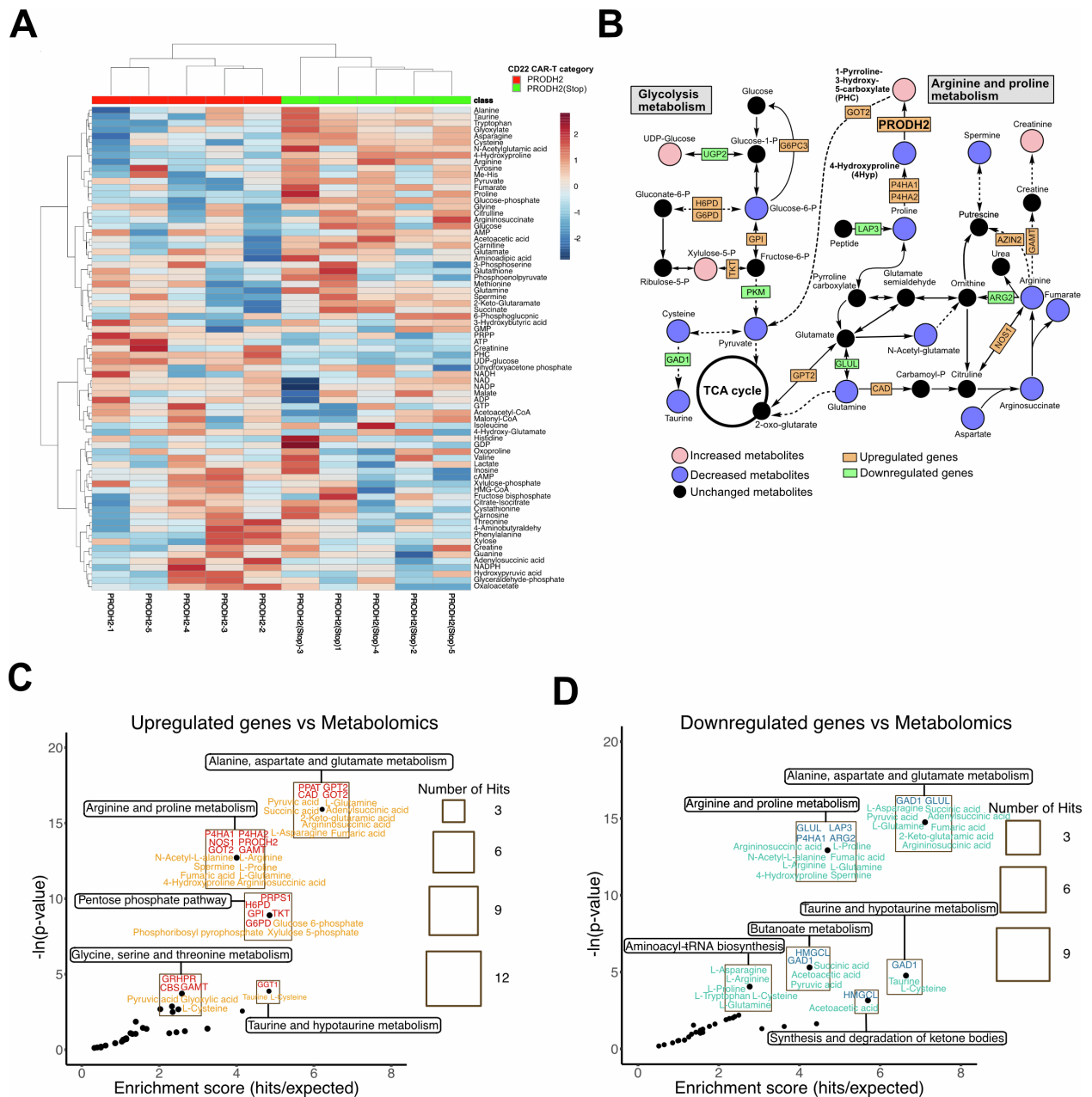


Figure S5. KEGG-guided multi-omics analysis of human CD22-CAR;PRODH2 T cells (Related to: Figure 5)

(A) A summary illustration highlighting representative enzymes encoded by DE genes, as well as DR metabolites on a metabolic map adapted from KEGG metabolic pathways. As in legend, blue circles indicate decreased metabolites; pink circles indicate increased metabolites; green squares indicate enzymes of downregulated genes; orange squares indicate enzymes of upregulated genes.

Solid arrows indicate direct biochemical reactions; dashed arrows indicate indirect or multi-step biochemical reactions; bi-direction arrows indicate reversible reactions. All arrows were drawn according to the known reactions in KEGG. Only reactions related to DE genes or DR metabolites were shown. A simplified version is shown in a main figure panel.

(B) A summary illustration highlighting representative enzymes encoded by DE genes, as well as DR metabolites on a simplified metabolic map adapted from KEGG metabolic pathways. As in legend, blue circles indicate decreased metabolites; pink circles indicate increased metabolites; green squares indicate enzymes of downregulated genes; orange squares indicate enzymes of upregulated genes. Solid arrows indicate direct biochemical reactions; dashed arrows indicate indirect or multi-step biochemical reactions; bi-directional arrows indicate reversible reactions. All arrows were drawn according to the known reactions in KEGG. Only reactions related to DE genes or DR metabolites were shown.

(C) Scatter plots of multi-omics analysis of RNA-seq upregulated genes vs. DR metabolomics metabolites (both increased and decreased) between PRODH2 vs. PRODH2(Stop) CD22-CAR-T cells. Each dot represents a metabolic pathway, with those enriched in multi-omics analysis text-labeled. The upregulated genes and the DR metabolites associated with each significant pathway were labeled inside a box scaled to the number of metabolic hits.

(D) Scatter plots of multi-omics analysis of RNA-seq downregulated genes vs. DR metabolomics metabolites (both increased and decreased) between PRODH2 vs. PRODH2(Stop) CD22-CAR-T cells.

Each dot represents a metabolic pathway, with those enriched in multi-omics analysis text-labeled. The downregulated genes and the DR metabolites associated with each significant pathway were labeled inside a box scaled to the number of metabolic hits.

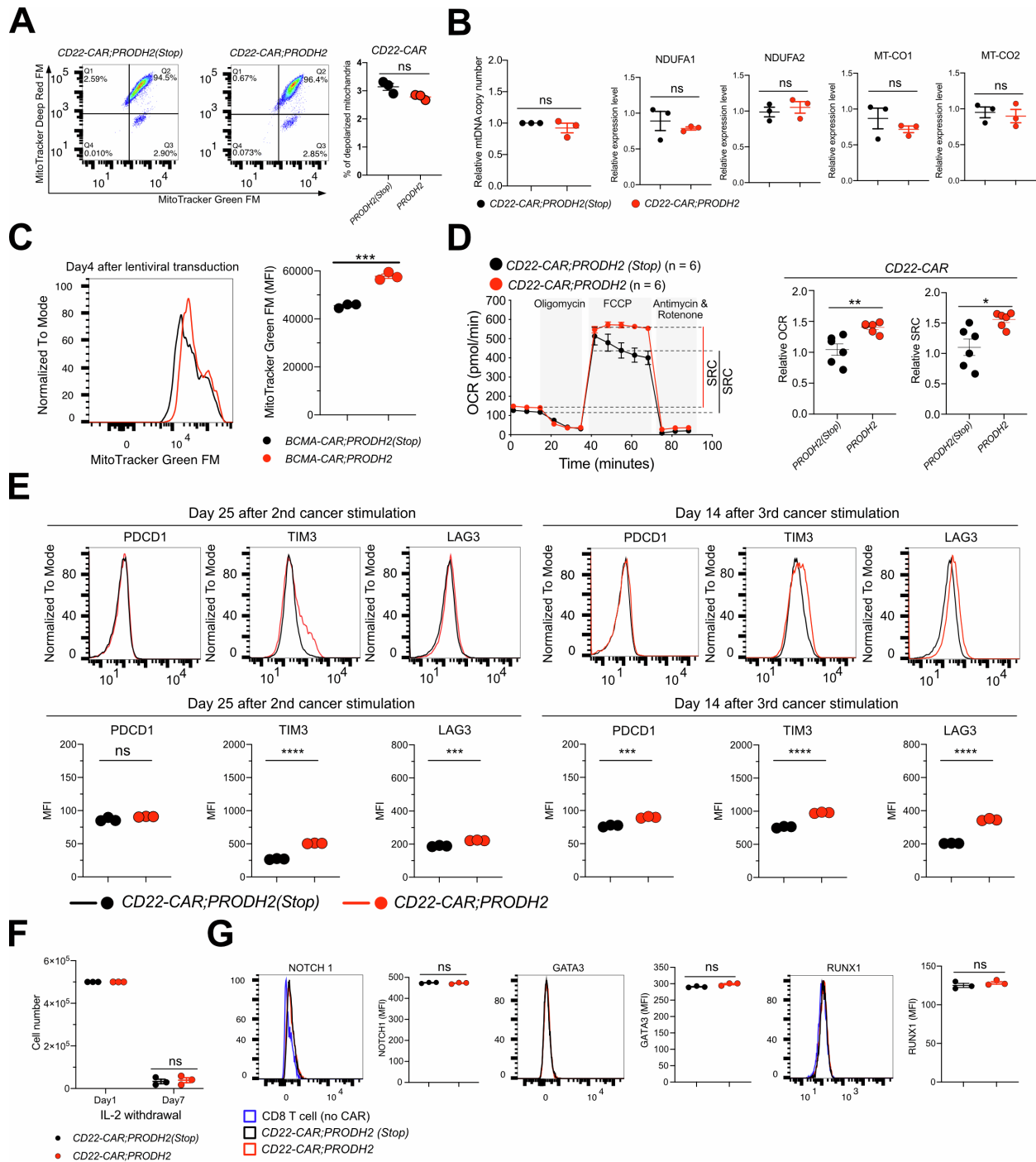


Figure S6. Additional analyses of PRODH2 engineered CAR-T, on mitochondria level, depolarization, metabolic function, Mt DNA copy number, cytokine secretion, T cell exhaustion and oncogenic potential (Related to: Figures 5, 6, 7)

- (A)** Mitochondria depolarization analysis. CAR T cells were stained with MitoTracker Green FM and MitoTracker Deep Red FM dyes to measure mitochondria depolarization (n = 3 biological replicates).
- (B)** Mt DNA analysis. Mitochondrial DNA copy and representative Mt gene expression of complex I and IV (n = 3 biological replicates).
- (C)** Mitochondria level analysis of lentiviral BCMA CAR-Ts. MitoTracker staining and quantification of *PRODH2* overexpression and Control CAR-T cells.
- (D)** Seahorse OCR analysis of Mt metabolic function. Seahorse experiment of *PRODH2* knock-in and control CAR-T cells, with a density of 4e5 CAR-T cells / well. Oxygen consumption rate (OCR) was measured at baseline and in response to oligomycin (Oligo), FCCP, and rotenone plus antimycin A. Data are representative of three independent experiments. Relative maximum OCR and relative spare respiratory capacity (SRC) were quantified (n = 6 biological replicates).
- (E)** Long-term T cell exhaustion analysis. Flow measurement of PD-1, TIM-3 and LAG-3 for CD22-CAR;PRODH2 and CD22-CAR;PRODH2 (Stop) T cells after long-term co-culture and cancer stimulation.
- (F-G)** Oncogenic potential analysis of PRODH2 in CAR-T.
- (F)** IL-2 withdrawal assay. Cell count after IL-2 withdrawal assay to evaluate the survival of *PRODH2* knock-in and control CAR-T cells.
- (G)** Flow analysis of key T-ALL oncogenic transformation markers, such as NOTCH1, GATA3, and RUNX1.

*P<0.05, **P<0.01, ***P<0.001, ****P<0.0001, ns = not significant by multiple *t* tests (with adjusted P value) (F), and unpaired *t* tests (A-E, G).

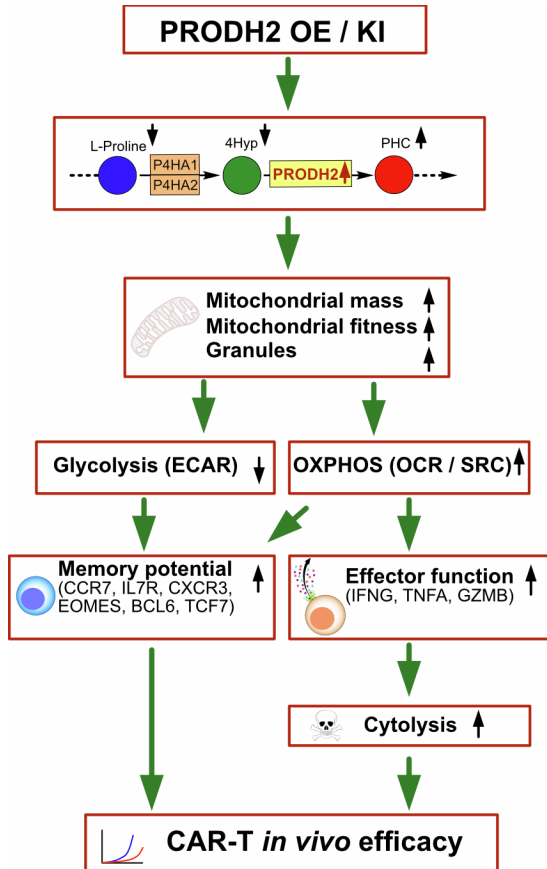


Figure S7. Brief summary of mechanism of action of PRODH2 GOF in CAR-T cell

Supplemental Datasets

Dataset S1. Design of the mm10dgLib library and readout of the dgTKS screen.

Tabs in this dataset:

Composition of the mm10dgLib library.

Readout of mm10dgLib library representation in the dgTKS screen.

Summary analysis of mm10dgLib library complexity / coverage.

dgTKS screen outlier test.

CD107a-high vs unsorted analysis of the dgTKS screen.

Ranked differential analysis of the dgTKS screen.

Dataset S2. mRNA-seq analyses of human PRODH2 knock-in CAR-T cells.

Tabs in this dataset:

Human CD22CAR-T mRNASeq PRODH2 vs. PRODH2(Stop) gene-level differential expression analysis.

Human CD22CAR-T mRNA-seq PRODH2 vs. PRODH2(Stop) upregulated genes (adj. p < 0.001)
DAVID gene-annotation analysis.

Human CD22CAR-T mRNA-seq PRODH2 vs. PRODH2(Stop) downregulated genes (adj. p < 0.001)
DAVID gene-annotation analysis.

Human CD22CAR-T mRNA-seq PRODH2 vs. PRODH2(Stop) GSEA analysis (Up-regulated)

Human CD22CAR-T mRNA-seq PRODH2 vs. PRODH2(Stop) GSEA analysis (Down-regulated)

Dataset S3. Metabolomics, multi-omics and CyTOF analyses of human PRODH2 knock-in CAR-T cells.

Tabs in this dataset:

Metabolomics analysis of human CD8 T cells with PRODH2-OE.

Metabolomics – relative abundance of detected metabolites in PRODH2 vs. PRODH2(Stop) CD22CAR-T cells.

MetaboAnalyst joint pathway analysis using consensus upregulated differentially expressed genes and differentially represented metabolites from metabolomics analysis.

MetaboAnalyst joint pathway analysis using consensus downregulated differentially expressed genes and differentially represented metabolites from metabolomics analysis.

CyTOF channel values for sampled CD22-CAR;PRODH2 and CD22-CAR;PRODH2(Stop) T cells.

UMAP coordinates for sampled CD22-CAR;PRODH2 and CD22-CAR;PRODH2(Stop) T cells.

Differential marker expression for sampled CD22-CAR;PRODH2 and CD22-CAR;PRODH2(Stop) T cells.

Supplemental Source Data and Statistics

Original data and statistics for non-high-throughput data used in figure generation.

Supplemental sequences

Additional sequences of probes and oligos were provided in an excel file.