Non-small cell lung cancer (NSCLC) carries a high mortality, and efficacious therapy is lacking. Therapy using chimeric antigen receptor (CAR) T cells has been used efficaciously against hematologic malignancies, but the curative effect against solid tumors is not satisfactory. A lack of antigen targets is one of the main reasons for this limited efficacy. Previously, we showed that lung-specific X (LUNX; also known as BPIFA1, PLUNC, and SPLUNC1) is overexpressed in lung cancer cells. Here, we constructed a CAR-T-cell-based strategy to target LunX (CAR\textsuperscript{LunX} T cells). CAR T cells were developed so that, upon specific recognition of LunX, they secreted cytokines and killed LunX-positive NSCLC cells. In vitro, CAR\textsuperscript{LunX} T cells displayed enhanced toxicity toward NSCLC lines and production of cytokines and showed specific LunX-dependent recognition of NSCLC cells. Adoptive transfer of CAR\textsuperscript{LunX} T cells induced regression of established metastatic lung cancer xenografts and prolonged survival. CAR\textsuperscript{LunX} T cells could infiltrate into the tumor. Also, we constructed a patient-derived xenograft model of lung cancer. After therapy with CAR\textsuperscript{LunX} T cells, tumor growth was suppressed, and survival was prolonged significantly. Together, our findings offer preclinical evidence of the immunotherapeutic targeting of LunX as a strategy to treat NSCLC.

INTRODUCTION

Non-small cell lung cancer (NSCLC) is an aggressive disease and an efficacious therapy is lacking. Therapy using chimeric antigen receptor (CAR) T cells has been developed against hematologic malignancies, but their curative effect against solid tumors has not been satisfactory. Also, targets for solid tumors (especially NSCLC) are limited.\textsuperscript{2}

Several types of antigen are overexpressed on the surface of tumor cells. However, “on-target/off tumor” side effects can occur because of the high sensitivity of CAR T cells for low-level antigen expression. Thus, a good antigen target must be restricted only to tumor cells, and be expressed at very low levels in normal tissues.\textsuperscript{3}

For treatment of lung cancer using CAR T cells, the candidate target antigens that have been reported recently are mesothelin,\textsuperscript{19} disialo-ganglioside (GD2), human epidermal growth factor receptor (HER) 2 and epidermal growth factor receptor (EGFR).\textsuperscript{5,7–20}

Lung-specific X (LunX) is a member of the palate, lung, and nasal epithelium clone protein family.\textsuperscript{21} The latter shows high expression on NSCLC cells but shows no expression on human normal lung tissues or adipose tissues, or tissues from the liver, brain, pancreas, skeletal muscle, adrenal glands, kidneys, prostate gland, heart, stomach, spleen, mammary glands, or thyroid gland.\textsuperscript{22} Previously, we demonstrated that LunX antibody-based therapy could suppress the growth, metastasis, and invasion of LunX-positive tumor cells by inducing downregulation of expression of LunX and blockade of the downstream pathways of LunX.\textsuperscript{23,24} In another study by our research team using mice, we showed that treatment with LunX antibody led to no cases of sudden death, pulmonary inflammation, or lung injury, which demonstrated an absence of toxic side effects of LunX antibody.\textsuperscript{25}

In the present study, we searched for a target antigen specific for lung cancer that has high expression among NSCLC patients. We constructed human T cells to express a CAR that was specific to LunX (CAR\textsuperscript{LunX} T cells), and we incorporated a CD28-4-1BB-CD3ζ co-stimulatory signaling domain. We evaluated the anti-NSCLC function of these cells in vitro and in an in vivo orthotopic xenograft and patient-derived xenograft (PDX) mouse model of NSCLC.
Our results showed that CAR\textsuperscript{LunX} T cells could eradicate LunX-expressing NSCLC cells specifically and efficiently \textit{in vitro} and \textit{in vivo}. Also, CAR\textsuperscript{LunX} T cells could infiltrate into a solid tumor, and this eradication was dependent upon LunX. Our data suggest that treatment using CAR\textsuperscript{LunX} T cells could be promising against NSCLC.

\section*{RESULTS}

\subsection*{Construction of a LunX-Targeted CAR}

Previously, we demonstrated that LunX is overexpressed in NSCLC cells and that LunX mRNA could be a diagnostic marker for NSCLC. Furthermore, we selected one antibody (S-35-8) and showed that it could bind specifically to LunX.\textsuperscript{24} Here, we also used surface plasmon resonance to analyze kinetic affinity and confirmed that single chain variable fragment (scFv) does have strong affinity for LUNX (Figure 1A). Therefore, S-35-8 could be an "ideal" antibody to construct the CAR. It has been reported\textsuperscript{26,27} that, compared with the CD28 CD3\zeta signaling domain, the 4-1BB signaling domain not only mediates the killing and proliferative capacity of CAR T cells, it also increases their persistence \textit{in vivo}. We generated a specific CAR\textsuperscript{LunX} with a backbone of a scFv, CD137, and CD247 was increased remarkably in the PCDH-CAR\textsuperscript{LunX} lentiviral-vector backbone and transduced 293T cells compared with the empty vector. To detect the transmembrane part, we used the c-myc tag. Using immunofluorescence, C-myc was detected and localized on the cytomembrane of transduced 293T cells, which indicated that CAR\textsuperscript{LunX} had been constructed.

\subsection*{LunX Antigen Sensitizes CAR\textsuperscript{LunX} T Cells to Secrete Cytokines}

Next, we stimulated normal human donor T cells with anti-CD3 and anti-CD28 monoclonal antibody (mAb)-coated beads and then transduced T cells with a lentiviral vector encoding CD19 CAR or LunX CAR. Forty-eight hours after transduction, the efficiency of transfection was observed by inverted fluorescence microscopy (Figure 2A) and measured by flow cytometry (Figure 2B). By measuring expression of EGFP and C-myc tag, high transfection efficiency was shown. Furthermore, immunofluorescence revealed that C-myc was localized on the membrane of CAR\textsuperscript{LunX} T cells, indicating that the CAR was expressed on the surface of T cells after transduction with the CAR construct. Next, we coated 96-well plates with LunX-antigen peptides (1,000 ng/mL), left the
plates for 12 h at 4°C, and then co-cultured with $10^4$ CAR$^{\text{LunX}}$ T cells, CAR$^{\text{CD19}}$ T cells, or anti-CD3ε mAb as control (1 mg/mL) for 24 h; OKT3 was used as a positive control for T cell stimulation (Figure 2D). Expression of interferon (IFN)-γ, interleukin (IL)-2, and tumor necrosis factor (TNF)-α was measured by enzyme-linked immunosorbent assays (ELISAs) after co-culturing the peptides of LunX antigen and CAR T cells (Figure 2E). As expected, CAR$^{\text{LunX}}$ T cells secreted high quantities of IFN-γ, IL-2, and TNF-α in response to the peptides of LunX antigen. In contrast, CAR$^{\text{CD19}}$ T cells demonstrated no reactivity to the peptides of LunX antigen but showed similar cytokine secretion in response to OKT3 stimulation. Collectively, we constructed CAR$^{\text{LunX}}$ T cells that could produce cytokines specifically in response to the peptides of LunX antigen.

**Targeted Killing of Lung Cancer Cells by CAR$^{\text{LunX}}$ T Cells**

After generation of CAR$^{\text{LunX}}$ T cells, we constructed CAR$^{\text{CD19}}$ T cells as control cells. We determined if LunX-positive cells were killed more efficiently by CAR$^{\text{LunX}}$ T cells than by CAR$^{\text{CD19}}$ T cells. We measured LunX expression in the NSCLC cell lines NCI-H292, NCI-H1650, and A549 by immunofluorescence. In accordance with previous work,24 all three NSCLC cell lines showed high expression of LunX. Conversely, the lung fibroblast cell line HFL1 showed no expression of LunX, which demonstrated that the antibody S-35-8 had good specificity (Figure 3A; Figure S1A). In long-term killing assays at a ratio of effector cell:target cell of 10:1, CAR$^{\text{LunX}}$ T cells killed NCI-H292, NCI-H1650, and A549 cells more quickly than that observed using CAR$^{\text{CD19}}$ T cells. As the control, CAR$^{\text{CD19}}$ T cells did not show good killing ability (Figure 3B).

In addition, we measured toxicity with different ratios of effector cell:target cell of CAR T cells and NSCLC cell lines. After co-culture of CAR T cells and NSCLC cell lines for 20 h, CAR$^{\text{LunX}}$ T cells exhibited better specific toxicity than that seen for CAR$^{\text{CD19}}$ T cells. However, CAR$^{\text{CD19}}$ T cells killed NSCLC cell lines at very low killing efficiency, so this toxicity may have been caused by T cells themselves (Figure 3C).

**Specific Recognition of LunX-Positive Lung Cancer Cells Enhances the Immune Response of CAR$^{\text{LunX}}$ T Cells**

We showed above that LunX-antigen peptides could stimulate CAR$^{\text{LunX}}$ T cells to secrete several cytokotins. Next, we co-cultured CAR T cells and NSCLC cell lines to ascertain if LunX-positive cell lines could induce a stronger response from CAR$^{\text{LunX}}$ T cells than from CAR$^{\text{CD19}}$ T cells. We analyzed CAR T cells after a 24-h challenge with LunX-positive cells; as controls, we used CAR$^{\text{CD19}}$ T cells. As anticipated, in the presence of LunX-positive tumor cells, CAR$^{\text{LunX}}$ T showed a significant increase in expression of IFN-γ, IL-2, and TNF-α. In contrast, CAR$^{\text{CD19}}$ T cells showed low expression of IFN-γ and IL-2 and no expression of TNF-α (Figures 4A–4C). When testing proliferation against NSCLC cell lines (NCI-H292, NCI-H1650, and A549), CAR$^{\text{LunX}}$ T cells proliferated more (as determined by flow cytometry using cell trace violet staining) than CAR$^{\text{CD19}}$ T cells. The latter showed only slight proliferation in the presence of LunX-positive tumor cell lines (Figure 4D). Therefore, LunX-positive tumor cell lines could stimulate only CAR$^{\text{LunX}}$ T cells (but not CAR$^{\text{CD19}}$ T cells) to secrete IFN-γ, IL-2, and TNF-α and proliferate.
To evaluate the efficacy of CARLuN xenografts, we examined their antitumor activity in A549-xenografted B-NDG mice. Luciferin-expressing A549 cells (5 × 10⁶) were injected subcutaneously (s.c.) into the tail veins of 6- to 8-week-old female B-NDG mice (n = 30). Three days after the xenograft model had been established, mice were divided into three groups of 10 randomly (CARLuN T, CARCD19 T, and phosphate-buffered saline [PBS]) and received first-time treatments. We set three times of transfusion treatments for each group (Figure 5A). Tumor burden was measured using bioluminescence imaging. After three courses of treatment, abdominal organs showed metastatic colonization of CARCD19 T cells and PBS on day 17, but not CARLuN T cells. From day 28 to day 33, CARLuN T cells could suppress the growth of lung cancer cells and improve survival compared with that observed in the CARCD19 T cell group and PBS group (Figure 5B). Additionally, we analyzed the dynamic growth of tumors of each mouse in the three groups. Mice treated with CARLuN T cells showed reduced proliferation of tumor cells. In contrast, CARCD19 T cells or PBS in the respective mice groups could not suppress tumor growth, and the mice died (Figure 5C). Statistical analyses of the dynamic growth of tumors in each group demonstrated that CARLuN T cells suppressed tumor growth better than that in the other two groups throughout the experiment and did not cause toxicity by T cells themselves (Figure 5D). Mice treated with CARLuN T cells showed significantly (p < 0.0001) increased survival compared with that observed with control mice (Figure 5E). To investigate if CARLuN T cells could infiltrate into the tumor, we carried out immunofluorescence analyses using EGFP to indicate CAR T cells. Infiltration by CARLuN T cells was observed, whereas CARCD19 T cells showed little infiltration into the tumor after first treatments (Figure 5F; Figure S3). These results showed that treatment with CARLuN T cells not only suppressed LuNX-positive tumor growth but also increased survival significantly among mice in the CARCD19 T cells group and PBS group.

CARLuN T Cells Inhibit Growth of Lung Cancer Cells in a PDX Model in Mice

PDX models have been shown to mimic the microenvironment of primary tumors. We obtained post-resection lung-cancer tissue from patients admitted to the First Affiliated Hospital of the University of Science and Technology of China (Hefei, China). Immunohistochemical analyses showed that the tumor tissue of patients had high expression of LuNX (Figure 6A). To determine the ability of CARLuN T cells to target patient-derived tumors, we evaluated the effect of treatment on a LuNX-positive PDX model. Patient-derived lung tumor tissue was separated into 25-mm³ portions and transplanted (s.c.) into B-NDG mice (n = 21). After 3 days, an identical mean tumor size was assigned to all three groups of seven mice randomly

**Figure 3. CARLuN T Cells Are Toxic against LuNX-Positive Lung Cancer Cells**

(A) Immunofluorescent staining for LuNX in NSCLC (NCI-H292, NCI-H1650, and NCI-H358) and lung fibroblasts (HFL1). Cell nuclei were stained with 4',6-diamidino-2-phenylindole (DAPI). Scale bar, 50 μm. (B) Toxicity of LuNX-based CAR T cells against the lung cancer cell lines NCI-H292, NCI-H1650, and A549 at a ratio of effector cell:target cell of 10:1. The control indicates that effector cells were not added. One of three independent experiments is shown. (C) Toxicity of CARLuN T cells and CARCD19 T cells against the lung cancer cell lines NCI-H292, NCI-H1650, and A549 at different ratios of effector cell:target cell for 40 h. One of three independent experiments is shown.
CARLunX T cells could not only slow down the symptoms of weight loss caused by the tumor, but also had no side effects on the body-weight of mice. Under the condition that each group had an identical mean tumor size before adoptive transfer on day 3 (Figure 6D), we monitored the time to tumor growth. On day 12, data showed that treatment with CARLunX T cells reduced tumor growth significantly (p < 0.05) (Figure 6E) and that the differences in the mean tumor size between the three groups had increased (Figure 6F). Furthermore, improved survival was observed after treatment with CARLunX T cells (Figure 6G). To determine if CARLunX T cells could infiltrate into PDX tumor tissue, we carried out immunofluorescence analyses to detect GFP-positive CAR T cells. CARLunX T cells were located mainly at tumor edges, whereas CARCD19 T cells were not detected in the tumor (Figure 6F, Figure S4). These results showed that therapy using CARLunX T cells could suppress growth of lung cancer cells and improve survival in a PDX model in mice.

DISCUSSION

Here, we report that LunX-specific CARs can target LunX-positive lung cancer cells. CARLunX T cells could be stimulated by LunX and secrete a series of cytokines. Upon co-culture of CARLunX T cells with LunX-positive cells, excellent killing efficacy could be observed. After establishment of a metastatic xenograft model, CARLunX T cells were shown to inhibit growth of LunX-positive tumor cells and prolong the survival of mice. Also, tumor-infiltrating CARLunX T cells could be found within tumor niches. In addition, we measured LunX expression in patient-derived lung-tumor tissue and established a PDX model. Treatment with CARLunX T cells led to not only suppressed growth of LunX-positive tumors, but also prolonged survival in mice.

It is difficult for immune cells to infiltrate solid-tumor niches due to an absence of sufficient co-stimulated signals to activate immune cells. Therefore, targeted therapies have opened a new era of cancer treatment and improved the prognosis of patients. Therapy using CAR T cells has shown excellent outcomes against hematologic malignancies such as acute lymphocyte leukemia, non-Hodgkin’s lymphoma, and multiple myeloma. However, CAR T cells have shown limited efficacy against solid tumors in several preclinical studies. With regard to lung cancer, few candidate target antigens have been being investigated in clinical trials. In addition, the target antigens have been being investigated in clinical trials.
antigens for solid tumors are limited and clinical treatment unsatisfactory.

Of the target antigens available for lung cancer, most show some shortcomings because the expression range of target proteins is not specific (e.g., mucin1, Her2). The best-known target antigen for NSCLC is the EGFR, but the proportion of patients with EGFR-positive lung cancer is low.7,11,20,33 Previously, we demonstrated that of 150 total samples of adenocarcinoma and squamous cell carcinoma, 80.6% (121 of 150) displayed increased expression of LUNX.24 Hence, therapy using CARLunX T cells could be promising treatment for NSCLC. In the present study, LunX represented a unique antigen for NSCLC and showed a considerable range of expression among NSCLC patients.

LunX has been reported to exert anti-inflammatory effects during microbial infections and agonist stimulation with Toll-like receptors. Also, studies have indicated that LunX expression is upregulated during infection in nasopharynx epithelium.34,35 Sung et al.36 used Northern blotting to measure PLUNC expression and found no signals in the brain, spleen, liver, muscles, kidneys, or testes. Iwao et al.22 demonstrated, using northern blotting, that LUNX mRNA was undetectable in the liver, brain, pancreas, skeletal muscles, adrenal glands, kidneys, prostate gland, heart, stomach, spleen, mammary
Figure 6. CAR\textsuperscript{LunX} T Cells Suppress Growth of Lung Cancer Cells and Prolong Survival in a PDX Model in Mice

(A) Representative immunohistochemical-stained LunX-positive lung cancer tissue of patient number 1. Results are representative of three independent experiments. (B) Experimental protocol for the PDX-xenografted model used in (C)–(H). (C) Bodyweight of mice in (B) (n = 7 [CAR\textsuperscript{LunX} T cells], n = 7 [CAR\textsuperscript{CD19} T cells], or n = 7 [PBS]) at various times (horizontal axis) after PDX tissue had been xenografted. (D) Tumor size in mice in (B) (n = 7 [CAR\textsuperscript{LunX} T cells], n = 7 [CAR\textsuperscript{CD19} T cells], or n = 7 [PBS]) on days 3. (E) Tumor size in mice in (B) (n = 7 [CAR\textsuperscript{LunX} T cells], n = 7 [CAR\textsuperscript{CD19} T cells], or n = 7 [PBS]) on days 12. (F) Tumor size in mice in (B) (n = 7 [CAR\textsuperscript{LunX} T cells], n = 7 [CAR\textsuperscript{CD19} T cells], or n = 7 [PBS]) on days 15. (G) Percent survival from days 0 to 30 post-first T cell infusion in mice in (B) (n = 7 [CAR\textsuperscript{LunX} T cells], n = 7 [CAR\textsuperscript{CD19} T cells], or n = 7 [PBS]). ** p<0.01, *** p<0.001, Log Rank (legend continued on next page)
glands, adipose tissue, or thyroid tissues but was highly expressed in lung cancer, in humans.

Previously, we showed that LunX was overexpressed in most samples of NSCLC and was high in lymph-node metastases. Importantly, benign lung disease and cancer in other organs (colon, liver, and breast) were not detected.33 LunX has been reported to be a multifunctional protein that is secreted by airway epithelia and has antimicrobial activity and regulates ion channels. Meanwhile, we also observed LunX expression in the membranes of NSCLC cells by immunofluorescence staining and flow cytometry. In our previous study, we generated a therapeutic LunX antibody, S-35-8, which slowed the growth and blocked the local invasion and metastasis of multiple established NSCLC xenografts without side effects. Taken together, LunX appears to be a promising target to construct CAR T cells.

A case report by Feng et al.33 provided quantitative analyses of EGFR-targeting CAR T cells in the circulation. They found that, after the second infusion, levels of CAR T cells were detectable for ≤24 weeks, compared with 8 weeks following the initial infusion of EGFR-targeting CAR T cells. Hence, their case report revealed that more than one dose of CAR T cells is necessary for treatment of solid tumors. In our study, we employed three doses of CAR\textsubscript{LunX} T cells in the xenografted model and PDX model. We showed that LunX-positive tumor growth was suppressed and survival was increased after three infusions of CAR\textsubscript{LunX} T cells.

Our data show that LunX is a promising target antigen for CAR-T-cell-based therapy against NSCLC. We demonstrated that CAR\textsubscript{LunX} T cells could kill LunX-positive tumor cells and induce regression of established LunX xenografts and a PDX model. Thus, we propose CAR\textsubscript{LunX} T cells are attractive and promising targeted therapy against NSCLC.

MATERIALS AND METHODS
Ethical Approval of the Study Protocol
All experimental procedures involving mice were approved by the ethics committee of the University of Science and Technology of China (Hefei, China).

Cell Lines
293T cells were obtained from the Shanghai Cell Bank (Chinese Academy of Sciences, Shanghai, China) and were maintained in Dulbecco’s modified Eagle’s medium (DMEM) with 10% fetal bovine serum. A549, HFL1, NCIH292, NCI-H1299, and NCI-H358 lines were purchased from the Shanghai Cell Bank and passaged in our laboratory for fewer than 6 months after receipt or revival. DNA fingerprinting and the isoenzymes test were used to authenticate the cell lines maintained in RPMI 1640 medium with 10% fetal bovine serum.

Isolation of T Cells and Gene Transfer
Blood products were obtained from healthy donors recruited by the Hefei Central Blood Bank (Hefei, China). Peripheral blood mononuclear cells were isolated by density gradient centrifugation. T cells were isolated using a CD3\textsuperscript+ MACS kit (Miltenyi Biotec, order no.130090874) and were stimulated with Dynabeads Human T Expander CD3/CD28 (Thermo Scientific, Waltham, MA, USA) at a ratio of T cell/bead of 1:3. Two days after activation, T cells were incubated with freshly prepared lentivirus at 32°C in the presence of 10 μg/mL polybrene (Sigma-Aldrich, St. Louis, MO, USA). After centrifugation (780 × g, 2 h, room temperature), cells were cultured for 6 h at 37°C, followed by culture for an additional 7 days in complete 1640 medium with human recombinant IL-2 (50 U/mL) at 37°C.

Construction of a Lentiviral Vector and Virus Production
293T cells cultured in DMEM were co-transfected with PCDH-MSCV-LunX-scfv-CD137-CD3\textsubscript{g} or PCDH-MSCV-CD19-scfv-CD137-CD3\textsubscript{g} together with the packaging constructs, psPAX.2, and pMD2G. After 8 h, DMEM was replaced with DMEM containing 10% fetal bovine serum. Forty-eight hours after transfection, the medium containing lentivirus was harvested and passed through a 0.45-μm filter (Millipore, Billerica, MA, USA) to remove cell debris. Virus in the supernatant was concentrated by polyethylene glycol-8000.

Flow Cytometry
To assess the transfection efficiency of T cells, a single suspension of transduced T cells was incubated for 1 h at 4°C with anti-Myc tag mouse mAb 9E10 (Cell Signaling Technology, Danvers, MA, USA). Cells were washed thrice with PBS and incubated with PE-conjugated goat anti-mouse immunoglobulin (Ig)G1 secondary antibody (Invitrogen, Carlsbad, CA, USA). The transfection efficiency of T cells was also examined by GFP expression. LunX expression on the surface of lung cancer cells was assessed after labeling with fluorescein isothiocyanate-anti-LunX mAb (s-35-8) and examined by fluorescence-activated cell sorting using a LSRII analyzer (Beckton Dickinson, Franklin Lakes, NJ, USA). Data were analyzed using FlowJo (Tree Star, Ashland, OR, USA).

Cytokine Generation In Vitro and Cell Killing
LunX-antigen peptides were plated at 1,000 ng/mL. Control T cells were stimulated with OKT3 (10 μg/mL) for 12 h, but not transfected by lentivirus, and the previous culture medium was changed into the fresh one. After culturing control T cells for 4 days, we co-cultured
them with LunX protein in the fresh culture medium. Human CAR T cells (10^5) were co-cultured with lung cancer cell lines (A549, NCI-H292, NCI-H1650; 10^5) or LunX. After 24 h, supernatants were harvested, and cytokine expression was measured using ELISAs for IL-2, IFN-γ, and TNF-α (Dakewe Biotech, Shenzhen, China).

Cytotoxicity assays were run on an xCELLigence real-time cell analysis instrument according to manufacturer’s (ACEA Biosciences, San Diego, CA, USA) instructions. In brief, lung cancer cells were seeded at 10^4 cells per well. After 20 h (when cells had reached exponential growth), CAR_C14 T cells or CAR_C19 T cells were resuspended in fresh complete medium without IL-2 and added to target cells at different ratios of effector cell:target cell. Then, cell growth was measured.

Histology
Samples of lung cancer tissue from the First Affiliated Hospital of the University of Science and Technology of China were placed in 12% neutral buffered formalin overnight and then dehydrated and embedded in paraffin. Paraffin-embedded tissues were sectioned at a thickness of 7 μm, followed by incubation with primary antibodies to anti-LunX (S-35-8) and IgG1. The signal was detected using a DAB peroxidase substrate kit (SK-4100; Vector Laboratories, Burlingame, CA, USA).

Immunofluorescence Analyses
CAR T cells or frozen sections of lung tumor tissues were fixed with 4% paraformaldehyde for 15 min, followed by blockade with 5% goat serum for 1 h at room temperature. Then, sections were incubated with primary antibodies overnight at 4°C. Secondary antibodies were added, followed by staining with 4',6-diamidino-2-phenyldole. Stained sections were imaged using a microscope (LSM 880; Zeiss, Oberkochen, Germany).

Generation of an A549 Cell Line that Expresses Luciferase Stably
Luciferase was PCR amplified from a pGL3-Basic plasmid (Promega, Fitchburg, WI, USA) and inserted into a PCDH lentiviral vector to generate a PCDH-luciferase construct. Lentivirus production and infection of A549 cells were undertaken using the methods described above.

Model of Metastatic Lung Cancer in Mice and Bioluminescence Imaging
Female B-NDG (NOD-Prkdc^scid^Il-2rg^null^/Bgcen) mice (6–8 weeks old; Jiangsu Biocytek, Jiangsu, China) were used to establish a PDX model and metastatic lung cancer xenografts. The A549 line with stable expression of luciferase (5 × 10^6 cells) was injected in the tail vein of B-NDG mice. Tumors were allowed to develop for 3 days. Before treatment, the tumor burden was assessed through luminescence imaging in vivo by the standard circular regions of interest on mouse lungs to ensure that CAR_C14 T cells, CAR_C19 T cells, and PBS carried an equivalent initial tumor burden. CAR T cells were delivered at 10^5 transduced cells in 200 μL of PBS through injection into the tail veins of mice for the first treatment. After 7 days, mice received a second treatment with 5 × 10^6 CAR T cells. Then, 7 days later, mice received the final treatment of 5 × 10^6 CAR T cells.

In the PDX model, lung tumor tissues (25 mm^3) were transplanted (s.c.) into B-NOG mice. Three days later, the tumor volume was calculated ([width^2 × length]/2) to ensure that each group had an identical mean size of tumor. Then 10^6 CAR_C14 T cells, CAR_C19 T cells, or PBS were adoptively transferred to PDX mice for the first treatment. Ten days later, mice received the second treatment of 5 × 10^6 CAR T cells. Then, 10 days later, mice received the final treatment of 5 × 10^6 CAR T cells. The size of tumors was measured every 3 days. Death was recorded when moribund animals were euthanized in accordance with the requirements of the IACUC protocol.

Statistical Analyses
Data are the mean ± SEM. Plots were generated using Prism 5 (GraphPad, San Diego, CA, USA). Differences between two groups were determined using two-tailed t tests. Differences among three or more groups were determined using one-way analysis of variance followed by two-tailed t tests. Survival curves were analyzed by log rank tests. p < 0.05 was considered significant.

SUPPLEMENTAL INFORMATION
Supplemental Information can be found online at https://doi.org/10.1016/j.omto.2020.04.008.

AUTHOR CONTRIBUTIONS
Conception and design, Z.H., X.Z., R.S., Z.T., and H.W.; Development of methodology, Z.H., X.Z., and H.W.; Acquisition of data (provided animals, acquired and managed patients, provided facilities, etc.), Z.H., X.Z., R.S., and Z.T.; Analysis and interpretation of data (e.g., statistical analysis, biostatistics, computational analysis), Z.H., X.Z., Z.T., and H.W.; Writing, review, and/or revision of the manuscript, Z.H., X.Z., Z.T., and H.W.; Administrative, technical, or material support (i.e., reporting or organizing data, constructing databases), Z.H., X.Z., D.J., Y.Z., R.S., and Z.T.; and H.W.; Study supervision, Z.T. and H.W.

CONFLICTS OF INTEREST
The authors declare no competing interests.

ACKNOWLEDGMENTS
This work was supported by the key project of the Natural Science Foundation of China (reference numbers 81872318 and 81602491).

REFERENCES
1. Reck, M., Heigener, D.F., Mok, T., Soria, J.C., and Rabe, K.F. (2013). Management of non-small-cell lung cancer: recent developments. Lancet 382, 709–719.
2. Chen, J., López-Moyado, J.F., See, H., Lio, C.J., Hempleman, L.J., Sekiya, T., Yoshimura, A., Scott-Browne, J.P., and Rao, A. (2019). NR4A transcription factors limit CAR T cell function in solid tumours. Nature 567, 530–534.
3. Morello, A., Sadelain, M., and Adusumilli, P.S. (2016). Mesothelin-Targeted CARs: Driving T Cells to Solid Tumors. Cancer Discov. 6, 133–146.
4. Ye, L., Lou, Y., Lu, L., and Fan, X. (2019). Mesothelin-targeted second generation CAR-T cells inhibit growth of mesothelin-expressing tumors in vivo. Exp. Ther. Med. 17, 739–747.
