Synthetic receptor signalling has the potential to endow adoptively transferred T cells with new functions that overcome major barriers in the treatment of solid tumours, including the need for conditioning chemotherapy. Here we designed chimeric receptors that have an orthogonal IL-2 receptor extracellular domain (ECD) fused with the intracellular domain (ICD) of receptors for common γ-chain (γc) cytokines IL-4, IL-7, IL-9 and IL-21 such that the orthogonal IL-2 cytokine elicits the corresponding γc cytokine signal. Of these, T cells that signal through the chimeric orthogonal IL-2Rβ-ECD–IL-9R-ICD (o9R) are distinguished by the concomitant activation of STAT1, STAT3 and STAT5 and assume characteristics of stem cell memory and effector T cells. Compared to o2R T cells, o9R T cells have superior anti-tumour efficacy in two recalcitrant syngeneic mouse solid tumour models of melanoma and pancreatic cancer and are effective even in the absence of conditioning lymphodepletion.

Therefore, by repurposing IL-9R signalling using a chimeric orthogonal cytokine receptor, T cells gain new functions, and this results in improved anti-tumour activity for hard-to-treat solid tumours.
IL-9R is a less-studied member of the \( \gamma \) cytokine receptor family that is naturally expressed by mast cells, memory B cells, innate lymphoid cells and haematopoietic progenitors\(^{13,15-20}\). Although T cell subsets that produce IL-9 have been described\(^{21-25}\), the effects of IL-9R signalling on T cells are not well characterized\(^{26-31}\). For example, naïve T cells are insensitive to IL-9 and T cell development is unimpaired in IL-9-deficient mice, which suggests that IL-9 is not a critical natural cytokine in T cell biology\(^{32,33,34}\). We found that activated mouse T cells did not support IL-9 signalling (Fig. 1c) owing to the absence of IL-9R expression (Extended Data Fig. 1c,d), which underscores the unorthodoxy of o9R signalling in these cells\(^{27,33,34}\). o9R signalling is a bona fide mimic of wild-type IL-9R signalling, as IL-9 treatment of mouse T cells transduced with the wild-type IL-9R resulted in a similar pattern of STAT1, STAT3 and STAT5 phosphorylation (Extended Data Fig. 1e).

As o9R cells and wild-type IL-2Rβ both use \( \gamma \) \( \delta \) signalling, we evaluated the competition between native and orthogonal signalling. Co-exposure to saturating doses of MSA-oIL-2 did not affect peak STAT5 phosphorylation by MSA-IL-2 in o9R T cells (Extended Data Fig. 2a). And although increasing doses of MSA-IL-2 partially mitigate MSA-oIL-2 induction of pSTAT1 and pSTAT3 in o9R T cells, likely due to the lower potency of the MSA-oIL-2 3A10 clone, the signalling program of o9R T cells remained active (Extended Data Fig. 2b,c).

We observed that T cells signalling through o9R (as well as o21R) enriched for a CD62L\(^{+}\) population and higher expression of Fas (CD95) and Sca-1 consistent with a TSCM phenotype, a subset known for its superior anti-tumour activity in adoptive cell therapy (ACT)\(^{6,35-37}\)(Fig. 1d–f). o9R T cells also proliferated less than o2R-expressing T cells (Fig. 1g and Extended Data Fig. 3a,b). Among orthogonal receptors, only o21R resulted in less proliferation than o9R (Extended Data Fig. 3a).

Given the unique STAT signalling profile, its lesser-known status among the \( \gamma \) cytokine receptor family and the acquisition of features of stem cell memory T (TSCM) cells, we chose to study the effect of o9R signalling in vivo using mouse models of ACT for solid tumours. We first used T cells from transgenic pmel mice, which express an endogenous T cell receptor (TCR) specific for gp100, a melanocytic antigen that is overexpressed in B16 melanoma\(^{38}\). We modified the protocol to be more stringent and omitted lymphodepleting radiotherapy, a conditioning regimen that potentiates ACT by inducing the homeostatic proliferation of adoptively transferred T cells through \( \gamma \) cytokine signalling\(^{39}\)(Fig. 2a). The STAT signalling profile and proliferation of o2R and o9R pmel T cells mirrored those of o2R and o9R T cells from C57BL/6 mice (Extended Data Fig. 4a,b).

Although both o2R and o9R pmel T cells expanded in the absence of lymphodepletion (Fig. 2b), we observed more consistent anti-tumour effects and prolonged survival in this model with o9R pmel T cells treated with MSA-oIL-2 (Fig. 2c,d and Extended Data Fig. 5a,b), achieving anti-tumour effects comparable to pmel T cells in lymphodepleted mice. And although T cells engineered to secrete native cytokines can also obviate the need for lymphodepletion\(^{40,41}\), native cytokines (unlike orthogonal cytokine signalling) induce signalling in both adoptively transferred and endogenous cells. The superior anti-tumour efficacy of o9R pmel T cells was also observed in lymphodepleted mice that were treated with MSA-oIL-2 (Extended Data Fig. 5c).

As MSA-oIL-2 clone 3A10 is highly orthogonal and does not affect endogenous cells, we hypothesized that mice would tolerate a prolonged dosing regimen of 25 days, whereas the toxic effects of MSA-IL-2 are observed after 5 days\(^{3}\). We did not observe clinical toxicity with the prolonged dosing regimen, which resulted in superior tumour control and survival with complete regressions in four of six B16 tumour-bearing mice (Fig. 2e and Extended Data Fig. 5d), exceeding the results in lymphodepleted mice treated with pmel ACT and five days of MSA-IL-2.

These results, despite the weaker proliferative effect of o9R signalling, suggested that factors beyond selective expansion underlie its enhanced anti-tumour effects. We observed greater tumour infiltration by o9R pmel T cells compared to o2R pmel T cells five days after ACT in mice that were treated with MSA-oIL-2 (Fig. 2f and Extended Data Fig. 5g). Among I8 tumour-infiltrating CD45\(^{+}\) leukocyte population...
clusters, the dominant distinguishing feature was the presence of o9R versus o2R pmel T cells (8.9% versus 1.6% of tumour infiltrating CD45+ cells; Extended Data Fig. 5e). Compared to o2R tumour-infiltrating pmel T cells, o9R pmel T cells were enriched for clusters that are associated with T cell activation, including a cluster that co-expressed CD39, PD-1 and T-bet, which o9R signalling was most active in peripheral tissues. This model provided a contrast to pmel studies, in which antigen-specific T cell activation is likely to predominate.

To examine the in vivo activity of o9R signalling, we examined the expression of CD62L in adoptively transferred pmel T cells. CD62L expression was higher in o9R than in o2R pmel T cells in the draining lymph nodes and spleens of tumour-bearing mice treated with MSA-oIL-2 (Extended Data Fig. 6c). No difference was observed intratumourally, in which antigen-specific T cell activation is likely to predominate.

To investigate o9R signalling in the context of chimeric antigen receptor (CAR)-based ACT, we used an immunotherapy-resistant model of pancreatic cancer expressing mesothelin. As the source of oIL-2 for CAR T cell studies, we chose vectored intratumoral delivery (Fig. 3a) to ensure high concentrations of oIL-2 (clone 3A10) in the tumour and to evaluate the effect of o9R signalling on T cell dysfunction in the tumour microenvironment. This model provided a contrast to pmel studies, in which o9R signalling was most active in peripheral tissues.
We transduced primary mouse T cells with retroviruses that encode a second-generation anti-mesothelin CAR together with orthogonal receptors to generate CAR-o2R and CAR-o9R T cells (Extended Data Fig. 7a,b). Stimulation of CAR-o9R and CAR-o2R T cells with MSA-olig2 reproduced the STAT phosphorylation and proliferation profiles of o9R and o2R signalling observed in Figs. 1 and 2 (Fig. 3b and Extended Data Fig. 7a). Despite their proliferative disadvantage, CAR-o9R cells showed a superior anti-tumour efficacy in vitro against the mesothelin-positive pancreatic ductal adenocarcinoma (PDA) cell line PDA7940b (Fig. 3c).

Similar to the pmel model, CAR-o9R cells were characterized by a T_{stem} phenotype after stimulation with MSA-olig2 (Fig. 3d,e and Extended Data Fig. 7c). CAR-o9R T cells also secrete higher levels of IFNγ, TNF, IL-4, IL-9, IL-10, IL-18, IL-22 and IL-23 than CAR-o2R cells in response to olig2 (Fig. 3f), mirroring the complex transcriptional signature of o9R pmel T cells. This was validated at the transcriptomic level; genes that are associated with effector functions (Ifng, Prf1, Gzmb and Gzma) and T cell activation (Pdcd1, Icos and Entpd1) were upregulated in CAR-o9R cells, along with a shift toward stemness (upregulation of Il7r, Sell, and Bcl6 and downregulation of Eomes and Ttx1; Fig. 3g).

In a recapitulating model of PDA using established subcutaneous PD7940b tumours, treatment either with an adenoviral vector encoding olig2 (Ad-olig2) or with CAR-o2R or CAR-o9R was ineffective in controlling rapidly growing tumours (Fig. 3h and Extended Data Fig. 7e). However, combination therapy with Ad-olig2 plus CAR-o2R or Ad-olig2 plus CAR-o9R resulted in complete regressions in 8 of 12 mice (67%) and 6 of 12 mice (50%), respectively (Fig. 3i). Early toxicity was observed in the Ad-olig2 plus CAR-o9R group, as 40% of mice (5 of 12) died by day 10, but notably fewer tumour relapses were seen in surviving mice in the Ad-olig2 plus CAR-o9R group (1 of 7) versus the Ad-olig2 plus CAR-o2R group (4 of 12).

The toxicity of Ad-olig2 plus CAR-o9R in lymphodepleted mice was characterized by clinical signs (tremor, delirium and seizures) of immune-effector-cell-associated neurotoxicity syndrome (ICANS). An RNA in situ hybridization (RINASH) analysis of mice with ICANS revealed that CAR T cells infiltrated into the meningeal layers of the brain, where mesothelin-expressing meningeal cells were also detected (Extended Data Fig. 8a,b). Significantly more meningeal-infiltrating CAR T cells were observed in the Ad-olig2 plus CAR-o9R group versus the Ad-olig2 plus CAR-o2R group (Fig. 8c). This was accompanied by higher expression of mesothelin, which can occur in the context of an inflammatory stimulus (Extended Data Fig. 8d). Histological examination of the brains of three long-term surviving mice that were treated with Ad-olig2 plus CAR-o9R was remarkable, including normal leptomeninges without inflammatory cells (Extended Data Fig. 8e). On the basis of these findings and the absence of similar toxicities in the pmel model, the observed ICANS appears to be specific to the CAR specificity and not inherent to o9R T cells. Still,
clinical translation of o9R must consider potential added on-target off-tumour toxicity, and may require engineering strategies (for example, on/off systems and synthetic circuits) to maximize patient safety. In the absence of conditioning chemotherapy, the superiority of the Ad-oIL-2 (3A10) plus CAR-o9R regime over Ad-oIL-2 (3A10) plus CAR-o2R was more evident, as suggested by complete regression rates of 42% (5 of 12) and 8.3% (1 of 12), respectively (Fig. 3j), as well as by prolonged survival (Extended Data Fig. 9a), as only 1 of 12 mice in the Ad-oIL-2 (3A10) plus CAR-o9R group exhibited toxicity in the absence of conditioning. 

To evaluate the translational potential for o9R signalling, we generated human orthogonal IL-2Rβ–IL-9R (ho9R) and human orthogonal IL-2Rβ–IL-9R (ho9R) (Supplementary Table 3). Human T cells were retrovirally transduced with vectors encoding either ho2R or ho9R (each containing YFP), as well as a vector encoding a TCR specific for NY-ESO-1 in the context of HLA-A0201 (NYESO1-TCR clone IG4) (ref. 4). NY-ESO-1 is a cancer-testis antigen that is overexpressed in synovial sarcoma, myoid liposarcoma, melanoma and other tumours. Transduced and sorted T cells (Extended Data Fig. 10a) were exposed to MSA-bound human orthogonal IL-2 (MSA-hoIL-2, which contains amino acid substitutions SQVLKA at positions 15, 16, 19, 20, 22 and 23 relative to the native polypeptide; Supplementary Table 4) or wild-type IL-2 (MSA-hIL-2)48,49. Consistent with observations in the mouse system, signalling through ho9R activates pSTAT1, pSTAT3 and pSTAT5 signalling (Fig. 4a). Likewise, native IL-2-STAT5 signalling through the endogenous γc is not disrupted by orthogonal receptor expression (Extended Data Fig. 10b). Finally, MSA-hoIL-2 is highly orthogonal and does not activate untransduced cells; however, wild-type MSA-hIL-2 does activate the orthogonal receptor at high concentrations (Extended Data Fig. 10b).

Similar to the mouse system, despite the weaker proliferative signal of ho9R signalling (Fig. 4b), we observed an expansion of TCM cells (CD45RA+CD27+CCR7+CD95−) and central memory T (TCM; CD45RA+CD27+CCR7+CD95−) cells by enrichment and absolute quantity among ho9R–NYESO1-TCR T cells after two and six days in culture with MSA-hoIL-2 (Fig. 4c,d and Extended Data Fig. 10c,d,e). This difference in T cell phenotype between ho9R–NYESO1-TCR and ho2R–NYESO1-TCR in vitro cytotoxicity of CAR-o9R cells (Fig. 3c) indicate that superior intratumoral potency—rather than tumour infiltration or proliferation—drives the anti-tumour efficacy of CAR-o9R T cells in the context of tumour-restricted orthogonal cytokine signalling. By contrast, systemic administration of oIL-2 in the pmel model resulted in peripheral o9R signalling (Extended Data Fig. 6c), which may explain the enhanced trafficking of o9R pmel T cells. Peripheral o9R signalling may also facilitate interactions between adoptively transferred T cells and endogenous cells (for example, antigen-presenting cells) that help to sustain anti-tumour effects.

In the absence of conditioning chemotherapy, the superiority of the Ad-oIL-2 (3A10) plus CAR-o9R regime over Ad-oIL-2 (3A10) plus CAR-o2R was more evident, as suggested by complete regression rates of 42% (5 of 12) and 8.3% (1 of 12), respectively (Fig. 3j), as well as by prolonged survival (Extended Data Fig. 9a), as only 1 of 12 mice in the Ad-oIL-2 (3A10) plus CAR-o9R group exhibited toxicity in the absence of conditioning. Efficacy was dependent on both orthogonal receptor expression in T cells and oIL-2 expression through Ad-oIL-2 (Extended Data Fig. 9b,c).

We examined the effect of Ad-oIL-2 on the quantity and quality of tumour-infiltrating CAR-o9R or CAR-o2R T cells. In contrast to the pmel model, fewer CAR-o9R than CAR-o2R T cells were observed in the tumour eight days after ACT (Fig. 3k). Comparisons between the models must be made cautiously, given the differences between CAR and TCR signalling and between tumour-restricted and systemic cytokine distribution. Nonetheless, the higher frequency of intratumoral CAR-o9R cells expressing IFNγ (Fig. 3k) and the superior direct
T cells persisted even after four antigen-specific challenges with an HLAA0201‘NY-ESO-1’ melanoma cell line, nRFP-M407, in the presence of MSA-holl-2 (Fig. 4e,f). Compared to ho2R—NYEso1 T cells, ho9R—NYEso1 T cells retained a greater frequency of CD45RA CD27+ and TCM cells, and expressed higher levels of CD62L and CXCR3. Repetitively stimulated ho9R—NYEso1 T cells expressed more IFNγ, TNF and IL-2 and exhibited greater polyfunctionality when exposed to either activating CD3/CD28 beads or the cognate melanoma cell line nRFP-M407 (Fig. 4g and Extended Data Fig. 10f). In a model of continuous antigen exposure using human T cells that co-express a mesothelin-specific CAR (MS) and either ho2R or ho9R, signalling through ho9R resulted in superior tumour cell killing after repetitive tumour challenges, with an enrichment of CD45RA CD27+ T cells with higher CRC7 expression, similar to ho9R—NYEso1 T cells (Fig. 4h and Extended Data Fig. 10g).

In conclusion, the orthogonal IL-2 cytokine receptor platform enables the redirection of signalling through modular replacement of the orthogonal IL-2Rβ ICD with the ICDs of receptors for other γc cytokines. Notably, specific in vivo stimulation of o9R signalling in TCR- and CAR-based tumour-specific T cells by the orthogonal IL-2 ligand 3A10 results in improved anti-tumour activity in two solid tumour models that are refractory to immunotherapy, and retains robust activity in a more stringent setting without conditioning lymphodepletion. This benefit is mediated by leveraging the γc cytokines to reroute the IL-2 signalling message through the IL-9Rα ICD, which results in the concomitant activation of STAT1, STAT3 and STAT5. Conveyed in T cells, the signalling message of IL-9Ra results in a unique phenotype that merges beneficial functional characteristics of stem cell memory and effector T cells, to provide improved in vivo anti-tumour activity.

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Article

Methods

Protein production
DNA encoding mouse and human wild-type and orthogonal IL-2 and wild-type IL-9 was cloned into the insect expression vector pAcGP67-A, which includes a C-terminal 8×His tag for affinity purification. DNA encoding mouse serum albumin (MSA) was purchased from Integrated DNA Technologies (IDT) and cloned into pAcGP67-A as a N-terminal fusion. Insect expression DNA constructs were transfected into Trichoplusia ni (High Five) cells (Invitrogen) using the BaculoGold baculovirus expression system (BD Biosciences) for secretion and purification from the clarified supernatant by Ni-NTA followed by size-exclusion chromatography with a Superdex-200 column and formulated in sterile phosphate-buffered saline (PBS) for injection. Endotoxin was removed using the Proteus NoEndo HC Spin column kit following the manufacturer’s recommendations (VivaProducts) and endotoxin removal was confirmed using the Pierce LAL Chromogenic Endotoxin Quantification Kit (Thermo Fisher Scientific). Proteins were concentrated and stored at −80 °C until ready for use.

Mammalian expression vectors
cDNA encoding mouse orthogonal IL-2Rβ and genebloc cDNA encoding mouse ICDs of IL-4R, IL-7R, IL-9R and IL-21R (IDT) were cloned into the retroviral vector pMSCV-MCS-ires-YPF by PCR and isothermal assembly (ITA). Human orthogonal IL-2Rβ (ho2R) and chimeric orthogonal IL-2Rβ-ECD-IL-9R-ICD (ho9R) were similarly cloned into the pMSCV vector.

Mice
Mice were housed in animal facilities approved by the Association for the Assessment and Accreditation of Laboratory Care and used under protocols approved by the Institutional Animal Care and Use Committee (IACUC) at the University of California, Los Angeles (UCLA), University of Pennsylvania and Stanford University. For experiments conducted at UCLA, C57BL/6j mice were bred and kept in the Radiation Oncology Vivarium; pmel-1 TCR/Thy1.1 transgenic mice (pmel mice) on a C57BL/6 background were obtained from The Jackson Laboratory. For experiments conducted at University of Pennsylvania, C57BL/6j and B6 CD45.1 ‘Pepboy’ mice were purchased from The Jackson Laboratory. For experiments conducted at Stanford University, C57BL/6j mice were purchased from The Jackson Laboratory.

Cell lines and cell culture
The B16-F10 mouse melanoma cell line was purchased from ATCC and cultured with RPMI 1640 with l-glutamine (Thermo Fisher Scientific) containing 10% fetal bovine serum (FBS, Omega Scientific), penicillin (100 U ml⁻¹, Omega Scientific), streptomycin (100 μg ml⁻¹, Omega Scientific) and amphotericin B (0.25 μg ml⁻¹, Omega Scientific). The mouse pancreatic cancer cell line, derived from spontaneous tumours arising in KPC (LSL-KrasG12D and LSL-Trp53R172H; Pdx-1-Cre) mice, was maintained in RPMI 1640 medium supplemented with 10% FBS and 1% penicillin–streptomycin. Human melanoma cell lines M407 and M263 were established from patient biopsies under UCLA IRB approval 11–00325 and maintained in RPMI 1640 medium supplemented with 10% FBS and 1% penicillin–streptomycin; M407 cells were stably transduced to express nuclear RFP (nRFP) for use in a live-cell imaging assay. T cells derived from C57BL/6 or pmel transgenic mice were cultured in RPMI 1640 with l-glutamine supplemented with 10% Hyclone FBS (Cytiva), antibiotics, 50 μM 2-mercaptoethanol (Gibco), 1% non-essential amino acids, 1% sodium pyruvate and HEPEPs. Primary human T cells were cultured similarly except without 2-mercaptoethanol. HEK293T cells were purchased from ATCC and maintained in DMEM supplemented with 10% FBS, 1× GlutaMax (Gibco) and penicillin–streptomycin. Cell lines were periodically authenticated and also periodically tested for mycoplasma infection using a mycoplasma detection kit (Biotool).

Retrovirus production
Production of retroviruses encoding orthogonal cytokine receptors has been previously described. In brief, HEK293T cells were seeded at 3 × 10⁶ cells per 10-cm tissue culture dish and allowed to adhere overnight. Cells were transfected with a 1:5 ratio of pMSCV retroviral vector to a pCL-Eco packaging vector using X-tremeGENE HP (Roche), Turbofect (Thermo Fisher Scientific) or TransIT Reagent (Thermo Fisher Scientific) and cultured overnight in DMEM with 5% FBS. After 24 h, the medium was replaced with fresh DMEM with 5% FBS and cultured for an additional 24 h. The medium was collected, clarified through centrifugation and flash-frozen in liquid nitrogen for storage at −80 °C. Cells were replated with fresh DMEM with 5% FBS, and cultured for an additional 24 h, and retroviral supernatant was collected and stored as described. To generate retrovirus for the transduction of pmel and human T cells, the same procedure was used with the following variations: 18 h after transfection, the medium was replaced with DMEM with 10% FBS containing 20 mM HEPEPS and 10 mM sodium butyrate and incubated for 8 h. The medium was then replaced with DMEM with 10% FBS containing 20 mM HEPEPS and no sodium butyrate and incubated overnight. The next day, the medium was collected and filtered through a 0.45μm filter. If not used immediately, virus was frozen at −80 °C for later use. Retroviruses encoding chimeric antigen receptors were produced as previously described. In brief, Plat-E packaging cells (Cell Biolabs) were transfected with pMSCV vectors using Lipofectamine 20000 (Thermo Fisher Scientific). Culture medium was replaced 24 h later and an additional 24 h the medium was collected, clarified by centrifugation and passed through a 0.45-μm filter before storage at −80 °C.

Adenovirus construction and purification
Replication-deficient EL/E3-deleted adenovirus vectors Ad5-CMV-oll-2 (Ad-oll-2) and Ad5-CMV-Null (Ad-Null) were constructed using the AdEasy XL Adenoviral Vector System (Agilent). Mouse orthogonal IL-2 clone 3A10 cDNA was synthesized (GenScript) with 5’ KpnI and 3’ HindIII restriction sites and subcloned into the multiple cloning site of pShuttle-CMV (Addgene). AdEasy-1-containing B5S13-AD-1-EScherichia coli cells were transfected with Pmel-linearized pShuttle-CMV-oll-2 for homologous recombination. The resulting recombinant Ad plasmid was sequence-verified and expanded in XL10-Gold Ultracompetent cells before PacI linearization and transfection into HEK293T cells. High-titre adenoviruses were purified by caesium chloride (CsCl) gradient centrifugation after multiple rounds of amplification. CsCl was exchanged with A195 buffer with Amicon Ultra-15 centrifugal filter units (Millipore). Viral titre (VP per ml) was determined spectrophotometrically (Nanodrop, Thermo Fisher Scientific).

Lentivirus production
Lentiviral vectors for human T cell transduction were produced in HEK293T cells. Lentiviral plasmids for anti-human anti-mesothelin M5 CAR (previously described) and human orthogonal receptors (ho2R or ho9R linked to GFP by a 2A sequence) together with packaging plasmids were transfected into HEK293T cells using Lipofectamine 3000 (Thermo Fisher Scientific). Ultracentrifugation was performed to concentrate lentiviral supernatants collected at 24 h or 48 h after transfection, and concentrated viruses were stored at −80 °C.

Activation, retroviral transduction and sorting of primary mouse T cells
Viral transduction of C57BL/6-derived mouse T cells was previously described. In brief, 12-well tissue culture plates were coated overnight with 2.5 μg ml⁻¹ solution of anti-mouse CD3ε (clone I45-2C11, Biolegend) in sterile PBS. Single-cell suspensions were prepared from spleens and lymph nodes of 6–8-week-old C57BL/6j mice by digestion through a 70-μm cell strainer followed by RBC lysis in ACK lysis buffer (Gibco). Cells were resuspended in mouse T cell medium containing 100 IU ml⁻¹ recombinant...
mouse IL-2 (mIL-2) and activated with plate-bound anti-mouse CD3ε and soluble anti-mouse CD28 (5 μg ml⁻¹; clone 37.51, BioXCell) for 24 h. Activated mouse T cells were transduced by spinfection using retroviral supernatants containing polybrene (10 μg ml⁻¹) and 100 IU ml⁻¹ mIL-2 at 2,600 rpm for 90 min at 32°C. Viral supernatant was replaced with fresh mouse T cell medium containing 100 IU ml⁻¹ mIL-2 and cultured for 24 h. Cells were collected via gentle pipetting and resuspended at 1 × 10⁶ per ml in fresh T cell medium containing 100 IU ml⁻¹ mIL-2 and expanded overnight at 37°C before further downstream cellular assays.

For retroviral transduction of pmel T cells, splenocytes from the five- to ten-week-old pmel mice were collected one to three days before transduction and activated with 50 U ml⁻¹ mIL-2 (Peptide) and 1 μg ml⁻¹ mouse gp100 peptide (Anaspec). One day before transduction, six-well tissue culture plates were coated with Retronectin (Takara) and placed in a 4°C refrigerator overnight. The following day, plates were blocked with 0.5% FBS in PBS for 30 min and washed with PBS. Viral supernatant (2 ml) was added to each well and spun at 2,000g for two hours. Activated pmel T cells (3 × 10⁷) were added to each well with 50 U ml⁻¹ mouse IL-2 and spun at 2,000g for 10 min and then cultured at 37°C for 18–24 h. Then, viable transduced cells were sorted based on expression of YFP and exclusion of 7-AAD using an Aria II cell sorter (BD Biosciences). Retroviral transduction of mouse CAR T cells was previously described³⁵. In brief, primary donor CD45.1 mouse splenocytes (from 4–6-week-old female CD45.1.B6.SJL-Ptpcr/Pepcb/BoyJ mice) were enriched for CD3⁺ cells by magnetic bead separation (STEMCELL Technologies). T cells were activated with mouse CD3/CD28 Dynabeads (Thermo Fisher Scientific) in the presence of 50 U ml⁻¹ recombinant human IL-2 (Peprotech) for 48 h before spinfection on retronectin-coated (Takara Bio) plates. Cells were collected for in vitro assays or intravenous injection two days after spinfection.

**Activation, retroviral transduction and sorting of primary human T cells**

Primary human peripheral blood mononuclear cells (PBMCs) isolated from a healthy human donor by leukapheresis were thawed and rested overnight before activation for two days with anti-human CD3/28 magnetic Dynabeads (Thermo Fisher Scientific) and human IL-2 (500 U ml⁻¹). T cells were co-transduced for 48 h on 6-well plates coated with Retronectin (Takara) and loaded with 1 ml per well of each retrovirus (encoding h2R and NYESO1-TCR clone Ig4 or Ig9R and NYESO1-TCR clone Ig4) by spinfection. Activated and transduced cells were collected and beads were removed by placing on an EasySep cell separation magnet for two minutes. Cells were stained with anti-human Vβ13.1 PE antibody (Beckman Coulter, recognizes the β-chain of the NYESO1-TCR clone Ig4), and 7-AAD live/dead dye before cell sorting based on the expression of YFP, Vβ13.1, and exclusion of 7-AAD using an Aria II cell sorter (BD Biosciences).

For transduction of human T cells with anti-mesothelin M5 CAR and human ortho-receptors, freshly isolated CD4⁺ and CD8⁺ T cells were mixed in a 1:1 ratio and activated with CD3/CD28 magnetic Dynabeads at a 3:1 bead-to-cell ratio. T cells were co-transduced 24 h later with lentiviral vectors encoding M5 CAR and h2R-GFP or ho9R-GFP. On day 5, beads were removed from the culture. T cells were maintained at 0.8 × 10⁶ per ml until reaching resting state as determined by cell size using Multisizer 4 Coulter Counter (Beckman), then cryopreserved. For flow sorting of M5 CAR–h2R-GFP and M5 CAR–ho9R-GFP double-positive T cells, the co-transduced T cells were thawed and rested overnight before staining with anti-human IgG F(ab’2) (Jackson ImmunoResearch) for M5 CAR and LIVE/DEAD Aqua for dead-cell exclusion. GFP served as surrogate marker for h2R and ho9R. Sorting was performed on a BD FACS Aria Fusion (BD Biosciences).

**Phosphoflow signalling assay**

Actively growing primary mouse or human T cells were rested in RPMI-C lacking IL-2 for 24 h before signalling assays. Cells were plated in an ultra-low-binding 96-well round bottom plate in 50 μl warm RPMI-C medium. Cells were stimulated by addition of recombinant cytokines for 20 min at 37°C, and the reaction was terminated by fixation with 1.5% paraformaldehyde (PFA) for 15 min at room temperature with agitation. Cells were washed and permeabilized with ice-cold 100% methanol for 60 min on ice or stored at −80°C overnight. Cells were washed with FACS buffer before staining with pSTAT antibodies (Supplementary Table 5) for 1 h at 4°C in the dark. Cells were washed and analysed on a CytoFlex (Beckman Coulter). Data represent the mean fluorescence intensity (MFI), and points were fit to a log(agonist) versus dose–response model using Prism 8.4 (GraphPad). For the gating strategy, see Supplementary Fig. 2.

**Western blot**

IL-2 and FBS-starved CAR T cells were stimulated for 20 min with cytokines and lysed with ice-cold RIPA buffer supplemented with protease/phosphatase inhibitor cocktail (Halt, Thermo Fisher Scientific) to extract protein. Thirty micrograms of total protein was loaded into SDS–PAGE gels (NuPage Bis-Tris, Thermo Fisher Scientific) and subsequently transferred to PVDF membranes (Immobilon-FL, Millipore). Detection of pSTAT1, pSTAT3, pSTAT5 and GAPDH was performed with respective primary antibodies followed by IRDye-labelled secondary antibodies or HRP-linked secondary antibodies. Membranes were imaged on an Odyssey CLX (LI-COR Biosciences).

**Mouse T cell proliferation assay**

Actively growing primary mouse T cells were rested in RPMI-C lacking IL-2 for 48 h before labelling with CellTrace Violet (CTV, Thermo Fisher Scientific). Labelled cells were seeded at 50,000 T cells per well in 50 μl in a 96-well round-bottom plate. Cells were cultured for two days in serial dilutions of MSA-ol-2. Cytokine was replenished on day 2. On day 4, CTV-labelled cell proliferation was evaluated by fluorescence-activated cell sorting (FACS) using the CytoFlex. Live-cell gates were based on FSC and SSC. CAR T cell proliferation was assessed by seeding 50,000 cells per well in a round-bottom 96-well plate in the presence of MSA-ol-2 or MSA-IL-2. On day 2, cells were fed with fresh medium and cytokines. Daily cell counts were acquired by staining an aliquot of cells with Calcein AM viability dye (Thermo Fisher Scientific) and analysed on the Celigo Image Cytometer (Nexcelom Bioscience).

**Cytokine assays**

For Luminex assays, transduced CAR T cells were incubated in round-bottom 96-well plates (50,000 cells per well) in triplicates for four days in the presence of cytokines after which supernatants were analysed with a Th1/Th2/Th9/Th17/Treg Cytokine 17-plex Mouse ProcartaPlex Panel (Thermo Fisher Scientific). A cytokine bead array (BD Biosciences) was used to individually measure IFNγ from supernatant of B16-F10 coculture with pmel T cells according to the manufacturer's instructions. oIL-2 expression from PDA7940b cells (10,000 cells per well) was evaluated by mouse IL-2 ELISA (Abcam) in cell culture supernatants at various time points after infection with Ad-Null or Ad-ol-2 (100 VP per cell). In vivo expression was assessed by injecting PBS, Ad-Null or Ad-ol-2 (1 × 10⁶ VP per tumour) into PDA7940b tumours and collecting 72 h later. Tumours were dissociated by three freeze-thaw cycles and homogenates were analysed for mouse IL-2 by ELISA. Terminal blood was collected by cardiac puncture and processed to serum by centrifugation. IL-2 concentrations were normalized to total protein content.

**Real-time cell killing assays**

PDA7940b tumour cells were seeded at 10,000 cells per well in a 96-well xCELLigence E-Plate (Agilent). Twenty-four hours later, transduced CAR T cells pre-incubated for 48 h in the presence of ol-2 were added at a 2:1 ratio and the target cell index was recorded every 15 min in the Real-Time Cell Analysis (RTCA) Analyzer (Agilent). T cell killing of B16-F10 cell lines
transduced with a nuclear localizing RFP was previously described. In brief, B16-F10-RFP cells pulsed with 100 ng ml⁻¹ IFNγ for 18 h were plated in a flat-bottom 96-well plate in triplicate at 5,000 cells per well for IncuCyte Live Cell Analysis (Essen Bioscience). Pmel T cells (o2R or o9R, pre-treated with MSA-IL-2 or MSA-oIL-2 for 48 h) were added at a 2:1 E:T ratio and two phase-contrast and fluorescent images were obtained of each well every two hours using the IncuCyte live imaging system and quantified by percentage confluence. The human TCR T cell repetitive killing assay was also conducted using IncuCyte Live Cell Analysis. Human melanoma cells (rNRP-M407, 5 × 10⁴) were plated per well in 6-well plates. Untransduced or co-transduced human T cells (co-transduced with either ho2R-NYESO1-TCR or ho9R-NYESO1-TCR, and pre-incubated for 48 h with MSA-ho1L-2) were added in duplicate at a 1:1 E:T ratio. Every 24 h, melanoma cells (rNRP-M407, 5 × 10⁴) were added to each well; orthogonal cytokine (MSA-ho1L-2) was replenished 24 h before every tumour rechallenge.

The human CAR T cell repetitive killing assay was conducted using the xCELLigence Analyzer. AsPC-1 human pancreatic tumour cells were seeded on a 96-well xCELLigence E-Plate at 10,000 cells per well. Twenty-four hours later, M5 CAR–ho2R or M5 CAR–ho9R cells pre-incubated for 48 h with MSA-ho1L-2 (1 μM) were added in triplicate at a 1:1 E:T ratio. Every 48 h, the CAR T cells were collected, washed, resuspended in fresh MSA-ho1L-2 and added on new wells of the E-Plate seeded with tumour cells (10,000 per well) the day before. After the last round of restimulation, the T cells were collected for phenotyping by flow cytometry.

**In vivo tumour studies**

For in vivo B16-F10 tumour growth experiments, early passage cell lines were used (fewer than 10 passages). B16-F10 cells (5 × 10⁵) were injected subcutaneously in the right flank of 6–10 week-old female C57BL/6 mice. Where indicated, mice were lymphodepleted with 500 cGy of total body irradiation one day before ACT. T cells (derived from female mice) were adoptively transferred approximately seven days after tumour inoculation, or when tumours became palpable. Specifically, 5 × 10⁵ sorted T cells were resuspended in 50 μl of PBS per mouse and administered through retroorbital injection. Where indicated, mice received treatment with cytokines: mouse serum albumin (MSA)-bound mouse IL-2 (MSA-IL-2) or MSA orthogonal IL-2 (MSA-oIL-2) (2.5 × 10⁵ units per day, intraperitoneal) for five consecutive days (or longer, where indicated) starting on the day of ACT. Tumour size (length × width) was monitored with calipers three times a week and volume based on arcsinh-scaled data gated on live, singlet CD45+ leukocytes and surface-stained. For in vivo assessments, peripheral blood was obtained by tail-vein sampling at indicated time points. At the time of necropsy, spleens were crushed and washed with PBS over a 70-μm cell strainer to collect splenocytes. Splenocytes and peripheral blood samples were treated with ACK lysis buffer before antibody staining.

B16 tumours were minced and dissociated using a mouse tumour dissociation kit (Miltenyi Biotec) and a gentleMACS Octo Dissocia-
tor (Miltenyi Biotec). Cells were then resuspended in PBS and filtered through a 70-μm cell strainer to obtain single-cell suspensions. Cells were stained with antibodies at 4 °C for 30 min in FACs buffer. Antibodies are listed in Supplementary Table 5. The 7-AAD viability dye was used to distinguish live cells from dead cells. Cells were analysed by flow cytometry using a LSRFortessa (BD Biosciences) and data were collected using BD FACS Diva (v.6.1.2). Data were analysed using FlowJo software (v.10, BD Biosciences). PDA7940b tumours were excised, weighed, minced with scalpels and dissociated using an enzyme cocktail consisting of hyaluronidase (2.5 U ml⁻¹), DNase (50 U ml⁻¹), collagen type I/II/IV (75 U ml⁻¹, 35 U ml⁻¹, 75 U ml⁻¹, respectively) in RPMI1640 supplemented with 1% penicillin–streptomycin. CD45-positive cells were isolated from single-cell tumour suspensions with CD45 (TIL) MicroBeads according to the manufacturer’s instructions (Miltenyi Biotec) and stored in liquid nitrogen. Quantification of tumour-infiltrating CAR T cells was performed using CountBright Beads (Thermo Fisher Scientific) and normalized to tumour weight. For gating strategy, see Supplementary Fig. 3.

For mass cytometry, cells were first fixed with 1.6% PFA for five minutes at room temperature. Cells were washed with 10 ml MaxPar Cell Staining Buffer (Fluidigm) and spun at 970g at 4 °C for 10 min. Next, the cells were resuspended in the surface antibody cocktail for 30 min at room temperature. Cells were washed with 5 ml of PBS and resuspended in 1 ml of ice-cold methanol for 15 min on ice. Cells were again washed with MaxPar Cell Staining Buffer and stained with the intracellular antibody cocktail for 30 min at room temperature. Finally, cells were washed with 10 ml MaxPar Cell Staining Buffer and stained with the intercalating solution (Cell-ID Intercalator-IR, 201192B) at a 1:6,000 dilution in Maxpar Fix and Perm Buffer with 1.6% PFA (Fluidigm, 201067) overnight at 4 °C. Data were acquired using the Fluidigm Helios mass cytometer. Analysis was performed using Omix based on arcing-enhanced data gated on live, singlet CD45+ leukocytes or CD8+ T cells (for gating strategy, see Supplementary Fig. 4). Cells were embedded in two-dimensional visualization using opt-SNE and clustered using FlowSOM with elbow metaclustering using Euclidean distances. Differentially abundant clusters were determined using edgeR with a P-value significance threshold of 0.05 and log-transformed fold change ≥ 1. Graphs were generated using the R package ggplot.

**Intracellular cytokine staining**

Human co-transduced T cells (either ho2R-NYESO1-TCR or ho9R-NYESO1-TCR) were collected from repetitive tumour challenge coculture 72 h after the most recent tumour challenge and 24 h after orthogonal cytokine had been replenished in the culture medium. T cells (1 × 10⁵) were cultured in a 96-well plate with anti-human CD3/CD28 Dynabeads (Thermo Fisher Scientific) or melanoma cells at a 1:1 E:T ratio (rNRP-M407 or M263) in the presence of brefeldin A and monensin. After four hours, cells were surface stained for 30 min at room temperature, fixed and permeabilized for intracellular cytokine staining for 30 min at room temperature. For intracellular cytokine staining of enriched CD45+ tumour-infiltrating leukocytes from PDA7940b tumours, cells were stimulated for six hours with Cell Activation Cocktail (with Brefeldin A) (Biologend), fixed/permeabilized in the Cyto-Fast Fix/Perm Buffer Set (Biologend) and stained with an anti-IFNγ antibody. Cells were washed and analysed by flow cytometry using a LSRII (BD Biosciences). Data were analysed using FlowJo software (v.10, BD Biosciences).
Multiplex immunohistochemistry

Formalin-fixed and paraffin-embedded tumour specimens were cut in 4-μm-thick sections onto glass slides for staining. The tyramide signal amplification (TSA)-based Opal method was used in this study for immunofluorescence staining (Opal Polaris 7-Color Automation IIIC Kit; Akoya Biosciences; NEL871001KT). The Opal fluorophores were used at a 1 in 150 dilution, as per the manufacturer’s recommendation. A fluorescent single-plex was performed for each biomarker and compared to the appropriate chromogenic single-plex to assess staining performance. Once each target was optimized with single-plex staining, the Opal 6 multiplexed assay was used to perform multiplex staining of slides. We applied primary antibodies to mouse spleen specimens as controls at optimized concentrations previously determined for single-plex staining of control tissues. Staining was performed using the BOND RX system (Leica Biosystems). The sequence of antibodies for multiplex staining was: FOXP3 (Opal 480), CD4 (Opal 520), PD-1 (Opal570), CD8 (Opal 620) and CD3 (Opal 690). Staining was performed after 20 min of heat-induced antigen retrieval using Bond Epitope Retrieval Solution 2 (Leica Biosystems). Antibodies are listed in Supplementary Table 5, and were used at a 1:200 dilution with a one-hour incubation. All fluorescently labelled slides were counterstained with DAPI and scanned on the Vectra Polaris (Akoya Biosciences) at ×20 magnification using appropriate exposure times. The data from the multispectral camera were analysed by the imaging InForm software (Akoya Biosciences) and quantification was performed using HALO image analysis software (Indica Labs).

Histopathology, clinical chemistry and RNA ISH

Mice were euthanized by means of CO₂ asphyxiation. Immediately after death, blood (n = 3 mice per group) was collected by cardiac puncture into Microvette tubes (Sarstedt) and allowed to clot at room temperature for 30 min before centrifugation at 12,000g for 10 min. Serum was stored at −80 °C before analysis. Cytokine levels in sera were measured with the Mouse 25-plex Cytokine Panel (IDEXX Bioanalytics). Serum levels of Ca, P, K and uric acid were measured with a custom clinical chemistry panel (IDEXX Bioanalytics).

Complete necropsy with macroscopic post-mortem examination was performed on all mice. Formalin-fixed tissues samples were trimmed according to the RITA guidelines (https://reni.item.fraunhofer.de/reni/trimming/) and then routinely processed for paraffin embedding, sectioning and haematoxylin and eosin (H&E) staining. The resulting slides were analysed by a board-certified veterinary pathologist blinded to experimental design.

The distribution of mesothelin expression and CAR T cell infiltration in the meninges were investigated by means of multiplex fluorescent RNA ISH (RNAscope Multiplex Fluorescent Assay, AC Diagnostics) including a custom probe designed against the mouse retrovirus used to transduce T cells with CAR and ortho-receptors. The assay was performed on formalin-fixed and paraffin-embedded brain sections. Whole-slide imaging on the resulting fluorescently labelled sections was performed using the Aperio VERSA 200 slide scanner (Leica Biosystems). CAR T cells and mesothelin-positive cells were finally counted using the object counting tool included in the Aperio ImageScope software (Leica Biosystems).

RNA-seq and analysis

For the in vitro experiments described in Fig. 2 and related supplementary material, o2R and o9R pmel T cells were stimulated with 5 μM O2L2 or 0.05 μM O2L2 for 48 h. RNA was extracted using the RNeasy mini kit (Qiagen). RNA-seq libraries were prepared using the KAPA mRNA stranded library preparation kit, according to the manufacturer’s recommendations. Libraries were pooled and sequenced on the Illumina HiSeq3000 platform (50-bp single-end reads). Reads were aligned to the mouse reference genome (mm9/GRCm38) using HISAT2 (v.2.0.4) (ref. 55). Gene expression was quantified using HTSeq-counts (v.0.6.1) (ref. 55). Differential expression analysis was performed using DESeq2 (ref. 57), and subsequent gene set enrichment analysis was performed using the fgsea (ref. 56) and misgdb (ref. 56) R packages, specifically on the TFactS annotated gene set56, and visualized using the ggplot2 R package. Differentially expressed genes were filtered to those with an adjusted P-value of less than 0.01 and a log₂-transformed fold change ≥ 1. Gene expression was visualized using the normalized gene expression (calculated using the rlog transform from DESeq2 and scaled by row) using the heat R package. Principal component analysis and sample-to-sample heatmaps were generated using the R functions pcor and dist, respectively.

For the in vitro experiments described in Fig. 3, CAR-o2R and CAR-o9R cells were stimulated for 48 h with MSA-0ll-2 or MSA-IL-2 and total RNA was extracted using the RNeasy mini kit (Qiagen). RNA expression was analysed using the nCounter Mouse Immunology Panel (Nanostring Technologies). Analysis was performed as described above.

Statistics and reproducibility

All unpaired t-tests are two-sided. Exact P-values are provided in Supplementary Table 6. For box and whisker plots (Fig. 2), box plots represent median and first and third quartiles, and whiskers extend to minima and maxima. Unless otherwise stated, n refers to biological and not technical replicates. For mouse tumour growth and survival experiments, sample size was selected on the basis of previous work using the respective ACT models (B16-pmel and PDA7940b-mesothelin CAR). Mice with tumours of equal size were randomized before treatment; tumours were measured in a blinded fashion. Data in Fig. 1b–f are representative of three independent experiments. Data in Fig. 2b,e,g are representative of two independent experiments. Data in Fig. 2c,d are representative of three independent experiments. Data in Fig. 2a are representative of three independent experiments. Data in Fig. 2f and 3a,b,c,d,e,f,g,h,i,j are representative of two independent experiments and conclusions are confirmed by three independent experiments using flow cytometry (Fig. 2f), mass cytometry (Fig. 2h) and immunofluorescence (Extended Data Fig. 5g). Data in Fig. 3a,c,e are representative of two independent experiments. Data in Fig. 3b are representative of three independent experiments. Data in Fig. 3l,j are representative of two independent experiments. Data in Fig. 4a,h are representative of two independent experiments. Data in Fig. 4b,d are representative of three independent experiments.

Reporting summary

Further information on research design is available in the Nature Research Reporting Summary linked to this paper.

Data availability

All data associated with this study are present in the manuscript or its Supplementary Information files. Gene expression data are available at https://www.ncbi.nlm.nih.gov/geo/ under accession numbers GSE199909 and GSE199956.
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Competing interests A.K. serves on the advisory board of T-Cure Therapeutics and Certis Oncology, holds stock in Certis Oncology, and receives research funding from Highlight Therapeutics. K.C.G., L.L.S., and L.K.P. are inventors on patent application PCT/US2020/050232 on the ortho-9 technology and are shareholders in Synthekine Therapeutics. K.C.G. is the founder of Synthekine Therapeutics. A.R. has received honoraria for consulting with Amgen, Bristol-Myers Squibb, Chugai, Genentech, Merck, Novartis, Roche and Sanofi, or has been a member of the scientific advisory board and holds stock in Advaxis, Arcus Biosciences, Highlight Therapeutics, Compugen, CytoDyn, Five Prime, FLX Bio, ImaginAb, IsoPlexis, Gilead Kite, Lutris Pharma, Merus, PACT Pharma, Rgenix and Tango Therapeutics. C.H.J. has received honoraria for consulting with, is a member of the board of directors for AC Immune, is a member of the scientific advisory board and holds stock in Bluesphere Bio, Cabaletta Bio, Carsima Therapeutics, Cellares, Celldex, DeCART Therapeutics, Kiasa Pharma, Trumunity Therapeutics, WIRB-Copernicus Group and Zopharm Oncology, and receives royalties for intellectual property licensed to Novartis and Trumunity.

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Extended Data Fig. 1 | Mechanisms of o9R signalling. a, pSTAT-1, -3, -5 and -6 signalling dose response curves of orthoIL-2Rβ ICD chimeric receptor expressing T cells stimulated with MSA-oIL2 for 20’. b, pSTAT-1, -3, and -5 signalling dose response of untransduced or o2R or o9R expressing T cells stimulated with MSA-oIL2 or MSA-IL2 for 20’. c,d, IL-9R expression on mock transduced or IL-9R transduced T cells from C57BL6 (c) or pmel TCR transgenic mice (d). e, Dose–response curves of STAT1, STAT3 and STAT5 phosphorylation in pmel T cells transduced with native IL-9R (pmel-IL-9R), o9R (pmel-o9R), or mock transduced (pmel) and stimulated with IL-9, MSA-IL2, or MSA-oIL2. Data are shown as mean fluorescence ± SEM, n = 3 biological replicates unless otherwise stated.
Extended Data Fig. 2 | Competition between o9R and wild-type IL-2 signalling. a–c, pSTAT5 (a), pSTAT3 (b), and pSTAT1 (c) signalling in o9R expressing C57BL/6 T cells treated with 10-fold dose titration of MSA-IL2 (starting at 100nM) in the absence (open) or presence of MSA-oIL2 [5 μM] (filled) for 20'. Data shown as mean fluorescence ± SEM, n = 3 biological replicates, YFP(+) gated.
Extended Data Fig. 3 | Proliferative effects of chimeric orthogonal receptors. a, Dose titration in vitro proliferation curves of o2R and orthogonal chimeric receptor (o4R, o7R, o9R, o21R) expressing T cells cultured for four days in MSA-IL2 (open symbol) or MSA-orthoIL2 (filled symbol). Data represent total live CD8+YFP(+) cells normalized to the maximal growth of each different chimeric receptor expressing cell cultured in MSA-IL2. Data represents mean of n = 2 biological replicates. b, In vitro proliferation of CTV-labelled T cells transduced with o2R (red) or o9R (blue) cultured for four days in MSA-oIL2 (filled curves) or MSA-IL2 (unfilled curves). YFP(+) gated; one of three representative plots shown.
Extended Data Fig. 4 | Signalling and proliferation of o2R and o9R pmel T cells. a, STAT1, STAT3 and STAT5 phosphorylation in o2R versus o9R pmel T cells after 30' stimulation with MSA-oIL2 (5μM). Shown are MFI of individual biological duplicates, gated on YFP+ cells. **, p < 0.01; ****, p < 0.0001 (ANOVA).

b, Shown are seven replicate experiments measuring in vitro proliferation (measured by fold growth) of o2R or o9R pmel T cells over 48 h in culture with MSA-oIL2, (each data point represents mean ± SD, n = 3). *, p < 0.05 (ratio paired t-test, two-sided).
Extended Data Fig. 5 | Anti-tumour efficacy, tumour infiltration, phenotype and function of o2R and o9R pmel T cells. 

a, Survival of B16-F10 tumour-bearing mice treated with (a) o2R or (b) o9R pmel T cells and MSA-IL2 or MSA-oIL2. BL/6 T cells treated with MSA-IL2 were used as an off target T cell control. Untransduced pmel T cells plus mIL-2 in tumour-bearing lymphodepleted and non-lymphodepleted mice served as controls. 

c, Effect of o9R pmel T cells in a lymphodepleted host. Tumour growth (mean ±SEM, left panel) of B16-F10 tumours in lymphodepleted C57BL/6 mice treated with o9R pmel T cells and MSA-IL2, MSA-oIL2, or no IL2. 

d, Individual B16-F10 tumour growth curves related to survival curves shown in Fig. 2e. All mice were non-lymphodepleted except one group (in purple) which received total body irradiation (5Gy). CR, complete regression. *, p < 0.05; **p < 0.01; ****, p < 0.0001 (ANOVA).

e, opt-SNE clustering of CD45+ tumour-infiltrating leukocytes seven days after adoptive transfer of o2R (left) and o9R (middle) pmel T cells (n = 4 mice/group; mice were treated without lymphodepletion and treated MSA-oIL2 2.5x10⁴ units/day for five days starting with ACT). 

f, opt-SNE clustering of the subset of tumour infiltrating o9R and o2R pmel T cells in non-lymphodepleted hosts treated with MSA-oIL2 (n = 4 mice/group, left panel), with separate plots for each treatment group illustrating only differentially abundant clusters (middle panel), and a volcano plot of differentially abundant clusters annotated with distinguishing features (right panel).

h, Tumour infiltration of CD3+CD8+ T cells and CD8+PD1+ T cells in non-lymphodepleted hosts treated with either o2R and MSA-IL2 or o9R pmel T cells and MSA-oIL2 by multiplex IHC (red = CD3, orange = CD8, yellow = PD1, green = CD4, teal = FOXP3). Images are representative of tumours from n = 3 mice/group and of one independent experiment (conclusions verified in three independent experiments by flow and mass cytometry). Quantifications (mean ± SD) shown to the right (n = 3 biological replicates/group). 

i, In vitro growth (mean ± SD, n = 3 biological replicates/group) of nRFP+B16-F10 tumour cells cocultured with o2R or o9R pmel T cells (2:1 E:T ratio) pretreated with MSA-IL2 and MSA-oIL2 by multiplex IHC (red = CD3, orange = CD8, yellow = PD1, green = CD4, teal = FOXP3). Images are representative of tumours from n = 3 mice/group and of one independent experiment (conclusions verified in three independent experiments by flow and mass cytometry). Quantifications (mean ± SD) shown to the right (n = 3 biological replicates/group).
Extended Data Fig. 6 | o9R signalling in vitro and in vivo drives a T_{SCM} phenotype in pmel T cells. a, Surface expression of CD62L, Fas (CD95) and CD44 as percentage of CD8 Thy1.1+ sorted o9R pmel T cells, o2R pmel T cells, or pmel T cells transduced with wild-type IL-9R (pmel-IL-9R) and treated with MSA-IL2 (0.05 μM), MSA-oIL2 (5 μM), or IL-9 (0.05 μM) for 48 h in vitro. Shown are mean ±SD, n = 3 biological replicates/group. ns, not significant; ****, p < 0.0001; unpaired t-tests, two-sided. b, Unsupervised transcriptomic analysis (RNA-seq) of sorted pmel-o2R (n = 3) and pmel-o9R (n = 3) T cells 48 h after exposure to MSA-oIL2 (50nM) also shown. On principal component analysis (PC1 v PC2, top right panel), samples separate by treatment group. On principal component analysis (PC1 v PC2, top right panel), samples separate by treatment group. On principal component analysis (PC1 v PC2, top right panel), samples separate by treatment group. On principal component analysis (PC1 v PC2, top right panel), samples separate by treatment group. On principal component analysis (PC1 v PC2, top right panel), samples separate by treatment group. Samples cluster by treatment group when arranged by sample-sample distances in a heat map (bottom right panel), with o9R pmel T cells treated with oIL2 most distinct among the four groups. c, Frequency of CD62L+ o2R or o9R pmel T cells (Thy1.1+YFP+) from tumour draining lymph nodes (DLN), spleen or tumours of mice treated with MSA-oIL2 (without prior lymphodepletion). Tissues were collected one or five days after adoptive transfer. Shown are mean ±SD with individual data points (n = 3-5 mice per group). **, p < 0.01; ***, p < 0.001; ****, <0.0001; unpaired t-test, two-sided.
Extended Data Fig. 7 | o2R and o9R signalling in T cells engineered with an anti-mesothelin CAR. a, Schematic of primary mouse CD3+ T cells expressing anti-mesothelin CAR and o2R (CAR-o2R) or o9R (CAR-o9R). b, Representative mesothelin CAR expression (top panel), IL-2Rβ expression (middle panel) and CAR/IL-2Rβ co-expression (bottom panel) in untransduced (UTD) or retrovirally transduced CAR-o2R and CAR-o9R T cells as determined by flow cytometry. Middle panel inset, mean fluorescence intensity (MFI) of IL-2Rβ in untransduced and transduced T cell. c,d, Expansion and phenotype of o2R and o9R CAR T cells. c, In vitro CAR T cell expansion. CAR-o2R and CAR-o9R cells were incubated in the presence of MSA-IL2 (100nM) or MSA-oIL2 (5 μM). An aliquot of cells was removed from the plate and stained with Calcein AM viability dye and counted on the Celigo Image Cytometer daily. Mean ±SEM, n = 3 replicate wells/group. d, CD44 and CD62L co-expression on CAR T cells. Full data of representative flow plots in Fig 3d. CAR-o2R and CAR-o9R cells were incubated for four days in the presence of MSA-IL2 (100nM) or MSA-oIL2 (5 μM). CD44 and CD62L surface co-expression was determined by flow cytometry on live CAR+ cells. Mean ±SEM, n = 3/group. ns, not significant. ****P < 0.0001 (ANOVA). e, Individual growth curves of PDA7940b tumours by treatment group; control groups corresponding to Fig. 3i,j.
Extended Data Fig. 8 | RNA ISH and serum markers of toxicity on day 11.

**a, b,** Representative images of brain and meningeal sections stained with fluorescent probes specific for mouse CAR (red, Cy3), mouse mesothelin (green, FITC) and counterstained with DAPI (blue). CAR-positive cells (white arrows) and mesothelin-positive meningeal cells (purple arrows) indicated in the pia-arachnoid layer of the meninges. Scale bars, 50 μm.

**c, d,** Semi-quantification of CAR T cells and mesothelin-positive cells in stained brain sections. Mean ±SEM, two sections/mouse, three mice/group. *P < 0.05 (two-sided t test with Welch’s correction).

**e,** Serum levels of calcium, phosphorus, potassium, uric acid, and f, cytokine release syndrome (CRS)-associated cytokines on day 11 of treatment. Mean ± SEM, n = 3 mice/group.

**g,** Representative photomicrographs from 3 mice (160 days post-treatment) cured of PDA7940b tumours showing histologically normal leptomeninges, without immune/inflammatory cell infiltrates, covering the temporoparietal region of the cerebral hemispheres. H&E, scale bar: 100 μm.
Extended Data Fig. 9 | See next page for caption.
Extended Data Fig. 9 | Anti-tumour efficacy of mesothelin-specific o2R and o9R CAR T cells combined with intratumoral delivery of Ad-oIL2.  

a, Survival differences between mice treated with CAR-o2R + Ad-oIL2 and CAR-o9R T cells + Ad-oIL2 with or without conditioning chemotherapy. Kaplan-Meier survival curves of treatment groups in Fig. 3i–k. **P < 0.01, ****P < 0.0001 (Log rank Mantel-Cox test).

b, Anti-tumour efficacy of Ad-oIL2 with control CAR T cells or Ad-Null and CAR-o2R T cells. b, Anti-tumour efficacy in mice treated with Ad-oIL2 and CAR T cells (without orthogonal cytokine receptor). Established subcutaneous PDA7940b tumours were treated with Ad-oIL2 on days 0 and 4 (1x10^9 VP/tumour) and with CAR T cells on day 0 (5x10^6). Mean ± SEM, n = 5 mice/group. ns, not significant (ANOVA).

c, Anti-tumour efficacy in mice treated with Ad-Null (no transgene) and CAR-o2R T cells. Established subcutaneous PDA7940b tumours were treated with Ad-Null on days 0 and 4 (1x10^9 VP/tumour) and with CAR-o2R T cells on day 0 (5x10^6). Mean ± SEM, n = 5 mice/group. ns, not significant (ANOVA).

d, Tumour volumes (mean ±SEM) of cured mice (from Fig. 3j) rechallenged with PDA7940b compared to age-matched naive mice (n = 10, 8 and 6 mice for naive, Ad-oIL2 + CAR-o2R and Ad-oIL2 + CAR-o9R groups, respectively). *P < 0.05; **P < 0.01; ns, not significant (ANOVA).

e, Quantification of CD45.1+ lymphocytes in peripheral blood of rechallenged mice (mean ±SEM, n = 10, 8 and 6 mice for naive, Ad-oIL2 + CAR-o2R and Ad-oIL2 + CAR-o9R groups, respectively). ns, not significant (ANOVA).
Extended Data Fig. 10 | Human chimeric orthogonal IL-2Rβ-ECD–IL-9R-ICD drives a stem-like phenotype and polyfunctionality in T cells even after repetitive antigen-specific tumour challenge. 

a, Flow plots of normal healthy donor T cells co-transduced with either ho2R and NYESO1-TCR (left) or ho9R and NYESO1-TCR (right), as detected by anti-human Vβ13.1 antibody that recognizes the β chain of the NYESO1-TCR clone 1G4 and YFP (internal marker of the o2R or o9R vectors). Co-transduction efficiencies are shown.

b, pSTAT signalling dose response of untransduced or ho2R (red) or ho9R (purple) or untransduced (UTD; grey) expressing activated human T cells (donor 651) stimulated with MSA-hoIL2 (top panel) or MSA-hIL2 (bottom panel) for 20'. Data are shown as mean fluorescence ± SEM, n = 2.

c, Gating strategy for immunophenotyping of YFP + population of ho2R–NYESO1-TCR and ho9R–NYESO1-TCR T cells shown in Fig. 4c.

d, Immunophenotype of ho2R–NYESO1 TCR and ho9R–NYESO1 TCR engineered human T cells after two days in culture with MSA-hoIL2 (1 μM). Shown is bar plot quantification of TSCM cells as a percentage of the YFP + population (mean ±SD, n = 3 biological replicates/group). **, p < 0.01 (unpaired t-test, two-sided).

e, T cells from Fig. 4e were collected and restimulated with either αCD3/αCD28 dynabeads, melanoma cell line M407 (HLA*0201+NY-ESO-1+) or M263 (HLA*0201+NY-ESO-1−). IFNγ, TNF and IL-2 were quantified among YFP+ CD4 T cells (CD8-) by ICS (mean ±SD, n = 3 biological replicates/group). Gating strategy for flow sorting of healthy donor T cells co-transduced with lentiviruses expressing anti-mesothelin CAR (M5) and ho2R-GFP/ho9R-GFP (left plots). Right plots indicate MFI of orthogonal IL-2Rβ in double-positive cells.
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For all statistical analyses, confirm that the following items are present in the figure legend, table legend, main text, or Methods section.

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☐ Estimates of effect sizes (e.g. Cohen’s d, Pearson’s r), indicating how they were calculated

Our web collection on statistics for biologists contains articles on many of the points above.

Software and code

Policy information about availability of computer code

Data collection

BD FACSDiva (v6.1.2);

Data analysis

HSAI2 (v2.0.4); HTSeq counts (v0.6.1); Deseq2 (v3.1.2); fgsea (v.3.12); msigdb (v.2.1.1); heatmap (v1.0.12); bowtie2 (v.2.3.5); tophat2 (v2.1.1); edgeR (v3.12); Halo (v3.1); Omiq (no version available); GraphPad Prism (v9.0.0); FlowJo (v10.7.1)

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Policy information about availability of data

All manuscripts must include a data availability statement. This statement should provide the following information, where applicable:

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All data associated with this study are present in the manuscript or supplementary materials. Gene expression data are available at www.ncbi.nlm.nih.gov/geo/.
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Life sciences study design

All studies must disclose on these points even when the disclosure is negative.

- **Sample size**
  For mouse experiments in Figure 2 and 3, sample sizes were not determined using statistical methods, but based on extensive experience in the literature (including from our group) using the pmel/B16 and the PDA/mesocAR mouse model.

- **Data exclusions**
  No data were excluded from the analysis.

- **Replication**
  For Figure 1, in vitro signaling experiments were repeated at least three times, with each data point in triplicates. In vitro proliferation experiments were repeated two times, with each data point in triplicate. For figure 2 and related supplementary material, in vitro experiments were repeated at least a total of two times, except for CyTOF, RNA-sequencing and LumineX cytokine analysis experiments. For CyTOF experiments, data was validated by flow cytometry experiments conducted in replicate experiments. For RNA-sequencing, biological triplicates were used. For lumineX cytokine analysis, biological triplicates were used.

- **Randomization**
  For mouse experiments in Figure 2 and 3, mice were selected based on similar tumor size and groups were generated randomly.

- **Blinding**
  For mouse experiments in Figure 2 and 3, tumors were measured by an individual blinded to randomization.

Reporting for specific materials, systems and methods

We require information from authors about some types of materials, experimental systems and methods used in many studies. Here, indicate whether each material, system or method listed is relevant to your study. If you are not sure if a list item applies to your research, read the appropriate section before selecting a response.

| Materials & experimental systems | Methods |
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| n/a                             | n/a     |
| Antibodies                      | Involved in the study |
| ☐ ☒ Eukaryotic cell lines       | ☐ ChiP-seq |
| ☐ ☒ Palaeontology and archaeology | ☐ Flow cytometry |
| ☐ ☒ Animals and other organisms | ☐ MRI-based neuroimaging |
| ☐ ☒ Human research participants |         |
| ☐ ☒ Clinical data               |         |
| ☐ ☒ Dual use research of concern|         |

Antibodies

Antibodies used

All antibodies used in the study, including clone and catalog number, are listed in Supplementary Table S5.

Validation

All antibodies used are commercially available and validation is noted on the manufacturer's website.

Eukaryotic cell lines

Policy information about cell lines

Cell line source(s)

- B16-F10: ATCC; PDA7940b: Dr. Gregory Beatty, University of Pennsylvania; PLAT-E: Cell Biolabs; HEK293T: ATCC; M407 and M263: Dr. Antoni Ribas, UCLA;

Authentication

Cell lines were periodically authenticated (at least once per year) using short tandem repeat analysis, with the exception of HEK293T cells which were used at low passage after freezing aliquots upon receipt from ATCC.

Mycoplasma contamination

Cell lines were periodically tested (at least once per year) for mycoplasma infection using mycoplasma detection kit (Biotool).

Commonly misidentified lines (see ICLAC register)

No commonly misidentified lines were used.
Animals and other organisms

Policy information about studies involving animals; ARRIVE guidelines recommended for reporting animal research

Laboratory animals
- (1) mouse, C57BL/6, female, 6-10 weeks of age;
- (2) mouse, pem1-1 transgenic (B6.Cg-Tm-Tm(Cg-Thy1a/Cy Tg(TcrαTcrb)8Rst/J), female, 6-10 weeks of age;
- (3) mouse, B6 CD4S.1 (B6.129-Ptprca Pepcb/Boy), female 4-6 weeks of age.

Wild animals
- The study did not involve wild animals.

Field-collected samples
- The study did not involve samples collected from the field.

Ethics oversight
- Mice were used under protocols approved by Institutional Animal Care and Use Committee at the University of California, Los Angeles (UCLA), University of Pennsylvania and Stanford University.

Note that full information on the approval of the study protocol must also be provided in the manuscript.

Flow Cytometry

Plots
- Confirm that:
  - The axis labels state the marker and fluorochrome used (e.g. CD4-FITC).
  - The axis scales are clearly visible. Include numbers along axes only for bottom left plot of group (a ‘group’ is an analysis of identical markers).
  - All plots are contour plots with outliers or pseudocolor plots.
  - A numerical value for number of cells or percentage (with statistics) is provided.

Methodology

Sample preparation
- Biological source: primary mouse cells in culture; single cell suspensions of mouse tumors after enzymatic digestion; single suspension of mouse spleen after mechanical dissociation, filtering and RBC lysis; single cell suspension of mouse lymph nodes after mechanical dissociation and filtering. Sample preparation: after exposure to experimental conditions, samples were washed with primary antibodies at 4°C for 30 minutes in PBS with 5% fetal bovine serum and 2 mM EDTA. Samples were washed and stained with 7-AAD or LiveDead Aqua viability dye to distinguish live and dead cells. For phosphohexokinase experiments, cells were washed and permeabilized with ice-cold 100% methanol for 60 minutes on ice or stored at -80°C overnight. Cells were washed prior to staining with primary antibodies.

Instrument
- Cells were analyzed by flow cytometry using a LSRFortessa (BD Biosciences) or a CytoFlex (Beckman Coulter).

Software
- Data were collected using BD FACSDiva software and analyzed using FlowJo (v10.7.1).

Cell population abundance
- For sorted o2R and o9R transduced pem1 T cells [Figure 2 and related extended data], cells were sorted based on expression of YFP and consisted of 10-30% of live cells in the sorted sample. The purity of the sort was >90% as confirmed by expression of YFP in post-sort samples.

Gating strategy
- For all analysis, dead cells and debris were gated out using FSC-A/SSC-A and viability staining. Singlets were gated based on FSC-A and FSC-H. For subsequent gating, some populations (e.g. CD45+ leukocytes, CD3+ T cells, etc.) were gated based on distinct expression of relevant markers. For populations without a clearly demarcated fluorescent signal, fluorescent-minus-one (FMO) and/or isotype staining were used to determine appropriate gating of positive and negative populations.

Tick this box to confirm that a figure exemplifying the gating strategy is provided in the Supplementary Information.