

1 **Title:** Sense-and-respond payload delivery using a novel antigen-inducible promoter
2 improves suboptimal CAR-T activation

3
4

5 **Authors:** Tingxi Guo^{1*}, Dacheng Ma^{1*}, Timothy K. Lu^{1,2,3}

6

7 ¹Synthetic Biology Group, Research Laboratory of Electronics, Massachusetts Institute of
8 Technology, Cambridge, MA 02139, USA.

9 ²Synthetic Biology Center, Department of Biological Engineering, Massachusetts Institute of
10 Technology, Cambridge, MA 02139, USA.

11 ³Department of Electrical Engineering and Computer Science, Massachusetts Institute of
12 Technology, Cambridge, MA 02139, USA.

13

14 *These authors contributed equally.

15 Correspondence and requests for materials should be addressed to T.K.L. (timlu@mit.edu).

16 **ABSTRACT**

17 Chimeric antigen receptor (CAR)-T cell therapies demonstrate the clinical potential of
18 lymphocytes engineered with synthetic properties. However, CAR-T cells are ineffective in
19 most solid tumors, partly due to inadequate activation of the infused lymphocytes at the site of
20 malignancy. To selectively enhance anti-tumor efficacy without exacerbating off-target
21 toxicities, CAR-T cells can be engineered to preferentially deliver immunostimulatory
22 payloads in tumors. Here, we report a novel antigen-inducible promoter and single-vector
23 sense-and-respond circuit for conditional payload expression in primary human T cells. In
24 therapeutic T cell models, the novel NR4A-based promoter induced higher transgene
25 expression than the conventional NFAT-based promoter under weakly immunogenic
26 conditions, where payload expression is most needed. Minimal activity was detected from the
27 inducible promoters in the absence of antigen and after withdrawal of stimulation. As a
28 functional proof-of-concept, we used the NR4A-based promoter to express cytokines in an
29 anti-mesothelin CAR-T model with suboptimal stimulation, and observed improved
30 proliferation compared to T cells engineered with the conventional NFAT promoter or CAR
31 alone. Our single-vector circuit achieves CAR-directed payload expression under weakly
32 immunogenic conditions and could enable the next generation of cell therapies with enhanced
33 anti-tumor efficacy.

34 Genetically programming cell functions with synthetic components holds promise for a
35 variety of clinical applications. A notable example is the adoptive transfer of T lymphocytes
36 engineered with a chimeric antigen receptor (CAR) to treat cancer^{1, 2}. However, the consistent
37 clinical benefit of these therapies has been largely limited to hematological malignancies.
38 Most carcinomas remain non-responsive to CAR-T cells because the suppressive tumor
39 microenvironment and variable antigen density prevent adequate activation of lymphocytes^{3, 4,}
40 ⁵. Amplifying the suboptimal responses of therapeutic T cells without exacerbating immune-
41 mediated toxicity is a major unmet need for the treatment of solid tumors.

42 Beyond antigen receptors, adoptively transferred anti-tumor T cells can also be
43 engineered to produce immunostimulatory payloads⁶. This strategy to augment immune
44 responses can enhance the therapeutic properties of the infused T cells and reinvigorate
45 endogenous immune cells. In preclinical models, T cells engineered to secrete common
46 gamma chain cytokines IL-2, IL-7, IL-15, and IL-21⁷; inflammatory cytokines IL-12⁸, IL-18⁹,
47 and IL-23¹⁰; or other protein-based therapeutics^{11, 12} have demonstrated superior tumor
48 control compared to non-producers. The continuous secretion of stimulatory payloads,
49 however, may counteract their beneficial effects. In one case, human T cells engineered to
50 constitutively produce IL-15 resulted in the transformation of transductants in an IL-15
51 receptor-dependent manner¹³. Constitutive production of potent cytokines such as IL-2 or IL-
52 18 also caused toxicities in pre-clinical CAR-T models^{7, 9}. These observations highlight the
53 need to tightly control recombinant payload production and, ideally, restrict it to the tumor site
54 to maximize its clinical benefit and prevent unwanted side-effects¹⁴.

55 A synthetic biology approach could leverage the antigen receptor signaling in
56 engineered lymphocytes to specify where the stimulatory payloads should be produced. A
57 sensitive, antigen-inducible promoter with low background activity could localize payload
58 delivery. The conventional approach for antigen-dependent transgene expression has been to
59 use an NFAT-based promoter¹⁵ encoding an NFAT/AP1 response element derived from the
60 human IL-2 enhancer¹⁶. This NFAT promoter was tested in the clinic to drive inducible
61 expression of IL-12 and was transduced to *ex vivo* expanded tumor-infiltrating lymphocytes¹⁷.
62 Toxicities were still observed after infusion, possibly because of non-localized production of
63 IL-12 by T cells with unknown antigen specificities. Subsequent pre-clinical developments
64 have focused on combining the antigen-inducible NFAT promoter with a recombinant
65 receptor^{9, 14, 18} to better control the input signal for conditional payload expression. Despite its
66 broad use, the standard NFAT promoter may not be the optimal choice for payload delivery.

67 Here, we identified a novel synthetic promoter based on an NR4A-binding motif that
68 induced greater responses than the conventional NFAT promoter under weakly stimulatory
69 conditions, which is when immune-enhancing molecules are most needed. We also describe
70 a single-vector circuit that incorporates this synthetic promoter to achieve an automated
71 payload response via CAR sensing of the cognate tumor antigen. The engineered T cells
72 respond to targets in an antigen-dependent manner and conditionally express a transgene of
73 choice upon antigen engagement. The inducible promoter and vector design described here
74 could enable future generations of synthetic lymphocytes, with controllable input and output to
75 safely enhance therapeutic responses.

76

77 RESULTS

78 ***A novel antigen-inducible promoter encoding an NR4A-binding motif***

79 We previously generated a synthetic promoter library termed Synthetic Promoters with
80 Enhanced Cell-State Specificity (SPECS), based on transcription factor (TF) binding motifs
81 found in public databases. SPECS vectors were constructed by encoding repeated TF
82 binding sites upstream of a minimal promoter derived from the adenoviral major late promoter
83 (MLP) and mKate as the fluorescent reporter¹⁹. In the present study, to identify novel antigen
84 receptor-inducible promoters from this library, we selected individual candidate promoters
85 encoding binding sites for known TFs directly downstream of T cell receptor (TCR) signaling
86 pathways (i.e., NFκB and MAPK targets)^{16, 20, 21, 22, 23, 24, 25, 26}, or TFs upregulated upon TCR-
87 induced activation^{27, 28, 29, 30, 31, 32, 33, 34}. TF binding site sequences ranged from 77-126 base
88 pairs (bp) (Supplementary Table 1). Individual promoter vectors were transduced into primary
89 human T cells by lentivirus and stimulated with plate-bound CD3 agonist OKT3, or left
90 untreated as a control (Supplementary Fig. 1A). Among the tested promoters, the one
91 encoding an NR4A-binding motif induced the highest percentage of reporter positive cells
92 compared to other candidates (Supplementary Fig. 1B and 1D) and was selected for
93 characterization. An AP1-based promoter was also chosen for comparison, because the AP1
94 pathway is well-established in T cell activation³⁵. As an internal positive control, CD137
95 upregulation³⁶ was measured in all assays to ensure similar activation among experiments
96 (Supplementary Fig. 1C).

97 Next, we compared the SPECS-derived NR4A and AP1 promoters with the
98 conventional NFAT promoter for OKT3-inducible responses. To facilitate quantitative
99 comparisons, we introduced a second downstream transcription module into the lentiviral
100 vector. In this module, the constitutive EFS promoter drives expression of truncated CD271
101 (tCD271) to mark transduced cells. We also tested an additional synthetic minimal promoter
102 (SMP)³⁷ in combination with each of the three response elements (Fig. 1A). The SMP and a
103 similar variant enabled robust inducible promoter activity in human cells^{18, 38}. All of the
104 promoter vectors transduced cells with comparable efficiency at ~60-80% (Supplementary
105 Fig. 2A). As a negative control vector, we cloned the EFS-tCD271 module alone, without
106 inducible promoters or mKate. All of the tested promoters responded similarly to the CD3
107 agonist among CD4 and CD8 subsets of primary human T cells after one day of stimulation
108 (Fig. 1B). In certain donors or in a TF-dependent context, SMP performed better than MLP,
109 although no consistent differences were observed (Fig. 1B and 1C). Thus, subsequent
110 experiments were performed with SMP as the minimal promoter. No significant baseline
111 activity in the absence of stimulation was observed with any of the inducible promoters (Fig.
112 1B). Using a set of vectors with only a minimal promoter sequence upstream of mKate, we did
113 not detect enhancer-like activity from the constitutive EFS promoter at the steady-state or
114 after activation, regardless of the choice of minimal promoter (Supplementary Fig. 3).

115 To investigate whether the TCR-inducible promoters could be activated by non-CD3
116 dependent mechanisms, we cultured the promoter-transduced cells in conditioned media
117 derived from strongly activated T cells to mimic an inflammatory milieu (Supplementary Fig.

118 4A). The NFAT, AP1, and NR4A promoters were activated at similarly low levels (~10%)
119 when the transduced cells were cultured in the conditioned media compared to normal media.
120 Reporter activity induced by the conditioned media was substantially lower than CD3-induced
121 responses (Supplementary Fig. 4B and 4C). Thus, we have identified a novel NR4A-based
122 promoter with anti-CD3 inducible activity, and a single lentiviral vector system that permits
123 stringent conditional gene expression alongside constitutive gene expression.

124

125 ***Inducible promoters demonstrate reversible and repeatable activation***

126 We next investigated the activity of the inducible promoters after withdrawal of antigen
127 receptor stimulation. Using the vectors shown in Figure 1A, we observed that it took up to 5
128 days after removing the source of stimulation for the mKate fluorescence to dissipate
129 (Supplementary Fig. 2B), suggesting high stability of the fluorescent protein. To measure the
130 reversibility and repeatability of inducible promoter activation, we changed the reporter to a
131 destabilized enhanced yellow fluorescent protein (dEYFP) encoding an additional PEST motif,
132 which reduces the half-life of fluorescent proteins³⁹. In this system, the reporter fluorescence
133 is more closely coupled to promoter activity. After transducing the dEYFP vectors in human T
134 cells, we stimulated the cells for one day with OKT3 as above, then transferred the cells to a
135 fresh well without the agonist for three days of rest. This process was repeated three times
136 (Fig. 2A). The EFS-tCD271 vector again served as a negative control. Across three
137 sequential stimulations, the NFAT, AP1, and NR4A promoter activities consistently returned
138 to baseline after three days of rest in CD8 T cells. In fact, the fluorescence intensity for all
139 reporters was reduced by at least 50% after only one day (Fig. 2B-D).

140 Interestingly, normalized responses were moderately higher after the second
141 stimulation (Fig. 2B, 2C, and Supplementary Fig. 5A), akin to a recall response in adaptive
142 lymphocytes. The lower responses observed after the third stimulation (Fig. 2D and
143 Supplementary Fig. 5A) were likely the result of activation-induced cell death. Reversible
144 responses were also observed in CD4 T cells with at least one round of stimulation
145 (Supplementary Fig. 5B). Repeated OKT3 stimulation biased the outgrowth of CD4⁺ cells and
146 decreased the overall viability of most samples (Supplementary Fig. 5C); thus, the promoter
147 responses in the CD4 subset after multiple rounds of activation could not be reliably
148 measured. Throughout the course of the experiment, the proportion of CD271⁺ cells did not
149 change significantly (Supplementary Fig. 5D), indicating that repeated activation of the
150 promoters was well tolerated and did not cause a growth disadvantage.

151

152 ***NR4A promoter induces higher responses than NFAT and AP1 in weakly immunogenic, 153 therapeutically relevant models***

154 Although OKT3 is a potent activator of T cells, it is not representative of therapeutically
155 relevant receptor-antigen interactions. To characterize the inducible promoter responses in
156 more appropriate models, we first selected a CAR based on the humanized single-chain
157 variable fragment (scFv) M5, targeting the widely expressed mesothelin tumor antigen⁴⁰. The
158 M5 CAR, which encodes 41BB and CD3z signaling domains (M5-BBz), is currently being
159 tested in clinical trials for treating a variety of solid tumors (NCT03054298, NCT03323944).

160 Mesothelin-targeting CAR-T strategies have yet to yield consistent objective responses⁴¹ and,
161 therefore, could benefit from the addition of inducible payloads. To investigate the inducible
162 promoter activity in a CAR setting, the M5-BBz CAR and tCD271, separated by a porcine 2A
163 (P2A) sequence, were encoded downstream of the EFS promoter for constitutive expression.
164 In these constructs, inducible promoters that drive the expression of mKate as a reporter were
165 cloned upstream of the CAR. tCD271 with the receptor alone served as a control vector (Fig.
166 3A). T cells were transduced with the vectors and stimulated with HEK293T (no mesothelin),
167 A549 (low mesothelin), or OVCAR8 (high mesothelin) target cells^{42, 43}.

168 Among the transduced CD8 and CD4 T cells, we observed no mKate fluorescence
169 when effector cells were cultured with HEK293T, again demonstrating minimal antigen-
170 independent promoter activity (Fig. 3B). Similar levels of reporter expression were observed
171 with the strong OVCAR8 stimulation. Notably, the NR4A promoter induced significantly more
172 mKate expression than the NFAT and AP1 promoters when cultured with the weakly
173 stimulatory A549 targets (Fig. 3B and Supplementary Fig. 6A). OVCAR8 was indeed more
174 immunogenic than A549 in the M5 model based on CD137 upregulation (Fig. 3G and
175 Supplementary Fig. 6A). Doubling the number of NFAT binding sites only marginally improved
176 the response, which was still lower than that of NR4A in the M5-BBz/A549 system
177 (Supplementary Fig. 7). A similar trend was observed when the promoters were tested with a
178 CD28-based M5 CAR (M5-28z, Fig. 3C): the NR4A promoter responded at higher levels than
179 the standard NFAT in response to A549 stimulation (Fig. 3D and Supplementary Fig. 6A). In
180 the M5-28z model, AP1 demonstrated higher activity than NR4A in response to OVCAR8
181 (Fig. 3D). Inducible promoter vectors for both CARs were transduced at ~50-70% efficiency
182 (Supplementary Fig. 6B).

183 Next, we constructed a similar set of vectors using the affinity-matured HLA-
184 A2/NYESO1-specific 1G4 TCR⁴⁴ as the model antigen receptor. 1G4 TCR has demonstrated
185 clinical efficacy in treating melanoma and synovial sarcoma^{45, 46}. The two P2A sequences
186 between tCD271, TCR α , and TCR β genes were codon-modified to avoid repetition in the viral
187 genome (Fig. 3E). TCR-T cells were stimulated with HEK293T or A375. Both of these cell
188 lines express HLA-A2 but only A375 expresses the cognate antigen^{44, 47}. In the TCR model,
189 NR4A also responded with consistently higher reporter positivity than NFAT or AP1 after co-
190 culture with A375, although the overall responses were lower than those seen with the CAR
191 models. The promoters induced higher responses in CD8 T cells than in CD4 cells (Fig. 3F),
192 which was expected since the 1G4 TCR is HLA class I restricted. The lower levels of CD137
193 upregulation in the TCR model were consistent with the weaker promoter activity (Fig. 3G),
194 which may have been caused by the insufficient formation of the correct TCR α / β pairing (see
195 Discussion). The TCR constructs were ~400bp larger than the CAR vectors and were
196 transduced less efficiently (Supplementary Fig. 6B). Nevertheless, across all of the receptor
197 models tested in our study, we observed significantly higher responses with the NR4A-based
198 promoter compared to the responses observed with the standard NFAT promoter under
199 poorly stimulatory conditions — precisely the context where payload expression is needed.

200

201 ***Recombinant cytokines expressed under the NR4A promoter amplify weak anti-tumor*** 202 ***responses***

203 As a proof-of-concept, we generated inducible constructs to conditionally express IL-2
204 and IL-21 in the clinically relevant M5-BBz model. The mKate reporter gene of the M5-BBz
205 CAR vectors (Fig. 3A) was replaced with recombinant IL-2 and IL-21, separated by a P2A
206 sequence (Fig. 4A). Constitutive expression of either cytokine alone has been reported to
207 enhance the proliferation of CD19 CAR-T cells⁷. Although both of these common gamma
208 chain cytokines can be produced endogenously by activated human T cells, cytokine
209 production is poor when the cells are suboptimally stimulated at low antigen density^{48, 49, 50}.
210 Therefore, we reasoned that the NR4A-based synthetic promoter system could supplement
211 these beneficial cytokines under conditions that preclude endogenous production.

212 In a proliferation assay without cytokine supplementation in the media, CAR-T cells
213 transduced with the control M5-BBz or inducible IL2/IL21 vectors demonstrated low levels of
214 proliferation in the absence of stimulation. In contrast, the majority of the cells divided after
215 co-culture with the immunogenic OVCAR8 cells. When stimulated with the weakly
216 immunogenic A549 targets, more of the NR4A-IL2/IL21 transductants proliferated compared
217 to cells engineered with other vectors. The improvement in proliferation was more
218 pronounced for the CD8 than CD4 subset (Fig. 4B and 4C), consistent with a past study
219 showing that IL-2 improves the proliferation of suboptimally stimulated CD8 but not of CD4
220 murine T cells⁵¹. Based on additional experiments in which the NR4A promoter induced
221 expression either of cytokines or mKate as a control (Fig. 4D), we determined that the
222 enhanced proliferation in response to A549 was payload-dependent, at least in CD8+ CAR-T
223 cells (Fig. 4E and 4F). Transduction efficiencies of the inducible cytokine constructs were
224 similar (Supplementary Fig. 8 and 9C).

225 Consistent with the above data, more of the CAR-T cells encoding the NR4A-IL2/IL21
226 module produced IL-2 when stimulated with A549, compared to cells transduced with other
227 inducible modules or the control vector (Supplementary Fig. 9A and 9B). Moreover, in A549
228 co-cultures, proliferation of NFAT-IL2/IL21 CAR-T cells was not increased compared to
229 control CAR-T cells (Fig. 4C and 4F), in line with the low responses observed with the NFAT
230 promoter in Figure 3B. Across these experiments, inducible expression of the cytokines did
231 not significantly enhance proliferation compared to CAR alone when cells were cultured with
232 OVCAR8 (Fig. 4C and 4F). In summary, these data demonstrate a proof-of-concept that
233 payloads delivered via the single-vector system using the NR4A promoter can augment
234 suboptimal CAR-T responses.

235

236 **DISCUSSION**

237 In this study, we identified a novel antigen-inducible transcriptional response element
238 in human T cells based on a TF binding site of the NR4A family. Notably, the NR4A-based
239 promoter outperformed the conventional NFAT-based promoter under poorly stimulatory
240 conditions. Given that at least NR4A1 expression is partly dependent on calcium and MAPK
241 signaling⁵², it is interesting that our NR4A-based promoter was more responsive than NFAT
242 and AP1-based promoters. The NR4A family of TFs could simply be more potent

243 transcriptional activators in primary human T cells. Alternatively, the choice of TF binding
244 motif or vector design used here may promote transcription for NR4A more than for NFAT or
245 AP1. Future molecular and biochemical studies are required to elucidate the mechanism
246 underlying the differential activity between respective promoters at various signaling
247 intensities. NFAT and AP1-based synthetic promoters may also be disadvantageous because
248 they sequester critical TFs away from endogenous response elements that are needed to
249 potentiate activation, especially with weak signals. Meanwhile, NR4A TFs have been
250 implicated as negative regulators of T cell activation^{53, 54} and thus the cognate synthetic
251 promoter would be unlikely to interfere with endogenous activation pathways.

252 TCR-induced activation of the NR4A pathway has been characterized previously^{31, 52,}
253 ⁵⁵. This pathway has been used to monitor TCR signaling by knocking-in a reporter at the
254 NR4A1 locus^{55, 56, 57}. In theory, payload transgenes could also be knocked-in at the NR4A1
255 site to achieve inducible expression, and improvements in site-specific integration
256 technologies for primary lymphocytes^{58, 59, 60} will facilitate the practical implementation of this
257 approach. With the knock-in method, however, transcriptional output will be dictated by
258 endogenous elements, which lacks the flexibility afforded by a synthetic promoter system that
259 can be tuned for a variety of applications. Our promoter platform similarly leverages the NR4A
260 pathway; instead of relying on endogenous response elements to drive NR4A1 transcription,
261 a short sequence encoding an NR4A-binding motif is used to drive conditional gene
262 expression, which is readily implemented using a standard lentiviral vector. Compared to
263 other systems in which the inducible module is transduced via a vector separate from the
264 antigen receptor¹⁴, our single-vector platform simplifies the manufacturing process and avoids
265 the heterogeneity in the final therapeutic product generated by multi-vector transduction.

266 The EFS promoter was used here to constitutively express the recombinant antigen
267 receptor. At ~200bp, it is one of the most compact constitutive promoters; the small size is
268 advantageous for incorporation into the large vectors designed in this study. Other compact
269 constitutive promoters, such as MND, tend to possess enhancer-like activity and can
270 influence the transcriptional activity of neighboring promoters^{61, 62, 63, 64}. Such cross-distance
271 activity would interfere with the upstream promoter for achieving stringent inducible
272 transcription. Although the EFS promoter is weaker than MND⁶⁵, we did not detect enhancer-
273 like properties, consistent with published data⁶⁴. However, stronger constitutive promoters
274 may be needed to practically implement this system with recombinant TCRs, which must
275 compete with endogenous TCR hemichains to form antigen-specific surface receptors⁶⁶. The
276 low level of response observed in our 1G4 model is likely caused by insufficient expression of
277 TCR α/β chains by the EFS promoter. Efforts to engineer potent and compact constitutive
278 promoters without enhancer-like attributes, or inducible promoters that are resistant to distal
279 enhancers, are needed.

280 In our proof-of-concept experiment, the inducible expression of common gamma chain
281 cytokines by the NR4A promoter improved proliferation of otherwise poorly responsive CAR-T
282 cells. Activation at a lower immunogenic threshold is a critical feature of the NR4A antigen-
283 inducible promoter that could widen the therapeutic index for a wide range of molecular
284 therapeutics compared to systemic or constitutive delivery. Some examples of applications

285 include site-specific production of: bispecific engagers to trigger bystander lymphocyte
286 responses; chemokines to promote infiltration of immune effectors; or cell-intrinsic regulators
287 (e.g., TFs) to conditionally re-program engineered cells in an autonomous manner. These and
288 other applications can be explored in future pre-clinical studies. In conclusion, the platform
289 described in this study could empower a variety of synthetic biology approaches to overcome
290 current challenges in adoptive cell immunotherapies.

291

292 **METHODS**

293 ***Cell culture***

294 HEK293T, A375, A549, and Jurkat.E6 cell lines were obtained from the American Type
295 Culture Collection (ATCC, Manassas, VA). OVCAR8 was a gift from Dr. Sangeeta N. Bhatia
296 (Massachusetts Institute of Technology, Cambridge, MA). HEK293T, A375, A549, and
297 OVCAR8 cell lines were cultured in DMEM (Thermo Fisher Scientific, Waltham, MA; catalog
298 #10569010). Primary human T cells and the Jurkat.E6 cell line were cultured in RPMI-1640
299 (Thermo Fisher Scientific; catalog #11875119). All media were supplemented with 10% fetal
300 bovine serum (Corning; catalog #35-010-CV) and 1% penicillin/streptomycin (Thermo Fisher
301 Scientific; catalog #15140122).

302

303 ***Generation of lentiviral vectors***

304 Truncated CD271 (tCD271)-CAR or TCR, EYFP destabilized with a PEST motif (dEYFP), and
305 IL-2/IL-21 fragments were synthesized as gBlocks by Integrated DNA Technologies
306 (Coralville, IA). The M5 scFv sequence was derived from the patent WO2015/090230.
307 Sequences of 28z and BBz signaling domains, and the affinity-matured 1G4 TCR were as
308 previously described^{44, 67}. Except NFAT, all TF binding site and mKate sequences were
309 derived from SPECS plasmids (Addgene #127842). The NFAT response element was
310 subcloned from the pSIRV-NFAT-eGFP plasmid⁶⁸ (a gift from Peter Steinberger, Addgene
311 plasmid # 118031). The EYFP sequence was derived from Addgene plasmid #51791 and the
312 PEST sequence was derived from Addgene plasmid #69072. The EFS promoter was
313 subcloned from the lentiCRISPRv2 plasmid (a gift from Feng Zhang, Addgene plasmid #
314 52961). The EFS promoter and tCD271-CAR/TCR fragments were first assembled into a
315 lentiviral backbone vector (derived from pFUGW in-house) in the reverse orientation of the
316 long-terminal repeats. Inducible promoter and reporter or payload genes were then inserted
317 upstream of the EFS promoter. Inserted sequences were confirmed by Sanger sequencing
318 (GENEWIZ, South Plainfield, NJ). Each set of NFAT, AP1, or NR4A vectors only differed at
319 the TF binding sequence.

320

321 ***Lentivirus production***

322 Lentivirus was generated by transfecting HEK293T cells of less than 10 passages and grown
323 to ~80% confluency in T25 flasks, with 1.5µg of pMD2.G (a gift from Didier Trono, Addgene
324 plasmid # 12259), 3.5µg of psPAX2 (a gift from Didier Trono, Addgene plasmid # 12260), and
325 5µg of respective transfer plasmid using the TransIT-2020 reagent (MirusBio, Japan). Media
326 was changed to fresh complete DMEM 16-24 hours post-transfection, and lentivirus was

327 collected after another 24 hours to be used immediately or stored at -80°C. Virus was titered
328 by infecting Jurkat.E6 at limiting dilutions.

329

330 ***Lentiviral transduction of primary human T cells***

331 Peripheral blood mononuclear cells (PBMCs) were isolated by density gradient centrifugation
332 from apheresis products of healthy donors (Brigham and Women's Hospital Crimson Core,
333 Boston, MA). Primary human T cells were purified from PBMCs by Pan T Cell Isolation Kit
334 (Miltenyi Biotec, Germany). Purified T cells were stimulated with anti-CD3/CD28 Dynabeads
335 (Thermo Fisher Scientific) at a T cell:bead ratio of 1:2. After 24 hours, Dynabeads were
336 removed and stimulated T cells were seeded on Retronectin (Takara Bio, Japan) coated non-
337 tissue culture treated plate with virus, and centrifuged at 1200xg, 32°C for 30 min. One time
338 infection was carried out for the smaller vectors shown in Figures 1 and 2 at a multiplicity of
339 infection (MOI) of 5-10. Larger vectors shown in Figure 3 and 4 were infected at a MOI of 10-
340 20, spread out over 2 days of daily infection. During stimulation and infection, T cells were
341 supplemented daily with 100U/ml of IL-2 and 10ng/ml of IL-15 (NCI Preclinical Repository,
342 Frederick, MD). After infection, T cells were maintained by supplementing with 100U/ml of IL-
343 2 and 10ng/ml of IL-15 every 3 days. T cells were expanded for another 5-6 days after the last
344 infection prior to use in experiments.

345

346 ***Flow cytometry***

347 The following monoclonal antibodies (BioLegend, San Diego, CA) were used in this study:
348 anti-human CD3 (clone UCHT1), anti-human CD4 (clone RPA-T4), anti-human CD8 (clone
349 RPA-T8), anti-human CD271 (clone ME20.4), anti-human CD69 (clone FN50), anti-human
350 CD137 (clone 4B4-1), and anti-human IL-2 (clone MQ1-17H12). Surface staining was carried
351 out at 4°C for 15 min with a master mix of antibodies. For intracellular staining of IL-2, cells
352 were fixed and permeabilized after surface staining using the Cytotfix/Cytoperm kit (BD
353 Biosciences). Stained cells were analyzed with a FACSCantoll,
354 FACSCelesta, or LSRFortessa flow cytometer (BD Biosciences). Data analysis was
355 performed with FlowJo. All data shown were gated on singlets and live cells, determined by
356 Aqua fixable dye (Thermo Fisher Scientific) for mKate-expressing and intracellular
357 experiments, or 7-aminoactinomycin D (BioLegend) for all other experiments.

358

359 ***T cell stimulation assays***

360 For plate-bound stimulations, non-tissue culture treated plates were coated with 2µg/ml anti-
361 CD3 clone OKT3 (NCI Preclinical Repository) with or without 2µg/ml anti-CD28 clone CD28.2
362 (BioLegend) by incubating at 4°C overnight. The same volume of PBS (Thermo Fisher
363 Scientific) was used as a control treatment. Antibody or PBS solution was removed and T
364 cells were seeded to the treated wells and cultured for 24 hours. Cells were analyzed post-
365 stimulation or transferred to fresh tissue-culture treated wells to rest. For cell-based co-culture
366 stimulations, T cells were mixed with HEK293T, A375, A549, or OVCAR8 targets at an
367 effector:target (ET) ratio of 3:1 and cultured for 24 hours to measure reporter fluorescence.
368 For intracellular IL-2 detection, CAR-T cells were cultured with targets at a 3:1 ET ratio for 18

369 hours, followed by treatment with 1500x diluted monesin (BD Biosciences) for 6 hours. To
370 measure proliferation, T cells were washed with PBS and labeled with 1 μ M of
371 carboxyfluorescein succinimidyl ester (CFSE, Thermo Fisher Scientific) by incubating them at
372 37°C for 5 min. Labeled cells were washed with complete media and co-cultured with A549 or
373 OVCAR8 target cells at an ET ratio of 10:1. CFSE dilution was assessed after 4 days of co-
374 culture.

375

376 **Statistics**

377 Comparisons between more than two groups were performed by two-way analysis of variance
378 (ANOVA) with Tukey's multiple comparisons test. Differences were considered significant at
379 an adjusted P value of less than 0.05. All statistical analyses were performed using GraphPad
380 Prism 6. Error bars denote one standard deviation.

381

382 **ACKNOWLEDGEMENTS**

383 This work was supported in part by the Department of Defense (W81XWH-17-1-0159,
384 W81XWH-16-1-0565) and National Institutes of Health (5-R33-AI121669-04). T.G. was
385 supported by a Postdoctoral Fellowship from the Natural Sciences and Engineering Research
386 Council of Canada. We thank the flow cytometry core facility at the Koch Institute for
387 Integrative Cancer Research at Massachusetts Institute of Technology. We thank Karen
388 Pepper for editing the manuscript.

389

390 **AUTHOR CONTRIBUTIONS**

391 T.G., D.M., and T.K.L. designed the experiments, analyzed the data, and wrote the
392 manuscript. T.G. and D.M. performed the experiments. T.K.L. supervised the study.

393

394 **COMPETING INTERESTS**

395 T.K.L. is a co-founder of Senti Biosciences, Synlogic, Engine Biosciences, Tango Therapeutics,
396 Corvium, BiomX, Eligo Biosciences, Bota.Bio, and Avendesora. T.K.L. also holds financial
397 interests in nest.bio, Ampliphi, IndieBio, MedicusTek, Quark Biosciences, Personal Genomics,
398 Thryve, Lexent Bio, MitoLab, Vulcan, Serotiny, and Avendesora. Other authors declare no
399 competing interests.

400 **REFERENCES**

401

- 402 1. Lim WA, June CH. The Principles of Engineering Immune Cells to Treat Cancer. *Cell*
403 **168**, 724-740 (2017).
404
- 405 2. Sadelain M, Riviere I, Riddell S. Therapeutic T cell engineering. *Nature* **545**, 423-431
406 (2017).
407
- 408 3. Anderson KG, Stromnes IM, Greenberg PD. Obstacles Posed by the Tumor
409 Microenvironment to T cell Activity: A Case for Synergistic Therapies. *Cancer cell* **31**,
410 311-325 (2017).
411
- 412 4. Rafiq S, Hackett CS, Brentjens RJ. Engineering strategies to overcome the current
413 roadblocks in CAR T cell therapy. *Nature reviews Clinical oncology* **17**, 147-167
414 (2020).
415
- 416 5. Shah NN, Fry TJ. Mechanisms of resistance to CAR T cell therapy. *Nature reviews*
417 *Clinical oncology* **16**, 372-385 (2019).
418
- 419 6. Yeku OO, Brentjens RJ. Armored CAR T-cells: utilizing cytokines and pro-inflammatory
420 ligands to enhance CAR T-cell anti-tumour efficacy. *Biochem Soc Trans* **44**, 412-418
421 (2016).
422
- 423 7. Markley JC, Sadelain M. IL-7 and IL-21 are superior to IL-2 and IL-15 in promoting
424 human T cell-mediated rejection of systemic lymphoma in immunodeficient mice. *Blood*
425 **115**, 3508-3519 (2010).
426
- 427 8. Koneru M, Purdon TJ, Spriggs D, Koneru S, Brentjens RJ. IL-12 secreting tumor-
428 targeted chimeric antigen receptor T cells eradicate ovarian tumors in vivo.
429 *Oncoimmunology* **4**, e994446 (2015).
430
- 431 9. Hu B, *et al.* Augmentation of Antitumor Immunity by Human and Mouse CAR T Cells
432 Secreting IL-18. *Cell reports* **20**, 3025-3033 (2017).
433
- 434 10. Ma X, *et al.* Interleukin-23 engineering improves CAR T cell function in solid tumors.
435 *Nature biotechnology* **38**, 448-459 (2020).
436
- 437 11. Choi BD, *et al.* CAR-T cells secreting BiTEs circumvent antigen escape without
438 detectable toxicity. *Nature biotechnology* **37**, 1049-1058 (2019).
439
- 440 12. Rafiq S, *et al.* Targeted delivery of a PD-1-blocking scFv by CAR-T cells enhances
441 anti-tumor efficacy in vivo. *Nature biotechnology* **36**, 847-856 (2018).
442
- 443 13. Hsu C, *et al.* Cytokine-independent growth and clonal expansion of a primary human
444 CD8+ T-cell clone following retroviral transduction with the IL-15 gene. *Blood* **109**,
445 5168-5177 (2007).
446

- 447 14. Chmielewski M, Abken H. TRUCKs: the fourth generation of CARs. *Expert opinion on*
448 *biological therapy* **15**, 1145-1154 (2015).
449
- 450 15. Zhang L, *et al.* Improving adoptive T cell therapy by targeting and controlling IL-12
451 expression to the tumor environment. *Molecular therapy : the journal of the American*
452 *Society of Gene Therapy* **19**, 751-759 (2011).
453
- 454 16. Fiering S, Northrop JP, Nolan GP, Mattila PS, Crabtree GR, Herzenberg LA. Single cell
455 assay of a transcription factor reveals a threshold in transcription activated by signals
456 emanating from the T-cell antigen receptor. *Genes & development* **4**, 1823-1834
457 (1990).
458
- 459 17. Zhang L, *et al.* Tumor-infiltrating lymphocytes genetically engineered with an inducible
460 gene encoding interleukin-12 for the immunotherapy of metastatic melanoma. *Clinical*
461 *cancer research : an official journal of the American Association for Cancer Research*
462 **21**, 2278-2288 (2015).
463
- 464 18. Zimmermann K, *et al.* Design and Characterization of an "All-in-One" Lentiviral Vector
465 System Combining Constitutive Anti-G(D2) CAR Expression and Inducible Cytokines.
466 *Cancers (Basel)* **12**, (2020).
467
- 468 19. Wu MR, *et al.* A high-throughput screening and computation platform for identifying
469 synthetic promoters with enhanced cell-state specificity (SPECS). *Nature*
470 *communications* **10**, 2880 (2019).
471
- 472 20. Rincón M, Flavell RA. AP-1 transcriptional activity requires both T-cell receptor-
473 mediated and co-stimulatory signals in primary T lymphocytes. *The EMBO journal* **13**,
474 4370-4381 (1994).
475
- 476 21. Kim HP, Leonard WJ. CREB/ATF-dependent T cell receptor-induced FoxP3 gene
477 expression: a role for DNA methylation. *The Journal of experimental medicine* **204**,
478 1543-1551 (2007).
479
- 480 22. Muthusamy N, Barton K, Leiden JM. Defective activation and survival of T cells lacking
481 the Ets-1 transcription factor. *Nature* **377**, 639-642 (1995).
482
- 483 23. Youn HD, Sun L, Prywes R, Liu JO. Apoptosis of T cells mediated by Ca²⁺-induced
484 release of the transcription factor MEF2. *Science* **286**, 790-793 (1999).
485
- 486 24. Li Y, Sedwick CE, Hu J, Altman A. Role for protein kinase C θ (PKC θ) in
487 TCR/CD28-mediated signaling through the canonical but not the non-canonical
488 pathway for NF- κ B activation. *The Journal of biological chemistry* **280**, 1217-1223
489 (2005).
490
- 491 25. Charvet C, Auberger P, Tartare-Deckert S, Bernard A, Deckert M. Vav1 couples T cell
492 receptor to serum response factor-dependent transcription via a MEK-dependent
493 pathway. *The Journal of biological chemistry* **277**, 15376-15384 (2002).
494

- 495 26. Malcolm T, Chen J, Chang C, Sadowski I. Induction of chromosomally integrated HIV-1
496 LTR requires RBF-2 (USF/TFII-I) and Ras/MAPK signaling. *Virus genes* **35**, 215-223
497 (2007).
498
- 499 27. Safford M, *et al.* Egr-2 and Egr-3 are negative regulators of T cell activation. *Nature*
500 *immunology* **6**, 472-480 (2005).
501
- 502 28. de la Roche M, *et al.* Hedgehog signaling controls T cell killing at the immunological
503 synapse. *Science* **342**, 1247-1250 (2013).
504
- 505 29. Glasmacher E, *et al.* A genomic regulatory element that directs assembly and function
506 of immune-specific AP-1-IRF complexes. *Science* **338**, 975-980 (2012).
507
- 508 30. Ho IC, Hodge MR, Rooney JW, Glimcher LH. The proto-oncogene c-maf is responsible
509 for tissue-specific expression of interleukin-4. *Cell* **85**, 973-983 (1996).
510
- 511 31. Woronicz JD, Calnan B, Ngo V, Winoto A. Requirement for the orphan steroid receptor
512 Nur77 in apoptosis of T-cell hybridomas. *Nature* **367**, 277-281 (1994).
513
- 514 32. Wu Z, Kim HP, Xue HH, Liu H, Zhao K, Leonard WJ. Interleukin-21 receptor gene
515 induction in human T cells is mediated by T-cell receptor-induced Sp1 activity.
516 *Molecular and cellular biology* **25**, 9741-9752 (2005).
517
- 518 33. Chikuma S, Suita N, Okazaki IM, Shibayama S, Honjo T. TRIM28 prevents
519 autoinflammatory T cell development in vivo. *Nature immunology* **13**, 596-603 (2012).
520
- 521 34. von Essen MR, Kongsbak M, Schjerling P, Olgaard K, Odum N, Geisler C. Vitamin D
522 controls T cell antigen receptor signaling and activation of human T cells. *Nature*
523 *immunology* **11**, 344-349 (2010).
524
- 525 35. Foletta VC, Segal DH, Cohen DR. Transcriptional regulation in the immune system: all
526 roads lead to AP-1. *Journal of leukocyte biology* **63**, 139-152 (1998).
527
- 528 36. Wolf M, *et al.* Activation-induced expression of CD137 permits detection, isolation, and
529 expansion of the full repertoire of CD8+ T cells responding to antigen without requiring
530 knowledge of epitope specificities. *Blood* **110**, 201-210 (2007).
531
- 532 37. Merlet E, *et al.* A calcium-sensitive promoter construct for gene therapy. *Gene therapy*
533 **20**, 248-254 (2013).
534
- 535 38. Ede C, Chen X, Lin MY, Chen YY. Quantitative Analyses of Core Promoters Enable
536 Precise Engineering of Regulated Gene Expression in Mammalian Cells. *ACS Synth*
537 *Biol* **5**, 395-404 (2016).
538
- 539 39. Li X, *et al.* Generation of destabilized green fluorescent protein as a transcription
540 reporter. *The Journal of biological chemistry* **273**, 34970-34975 (1998).
541
- 542 40. Morello A, Sadelain M, Adusumilli PS. Mesothelin-Targeted CARs: Driving T Cells to
543 Solid Tumors. *Cancer Discov* **6**, 133-146 (2016).

544

- 545 41. Haas AR, *et al.* Phase I Study of Lentiviral-Transduced Chimeric Antigen Receptor-
546 Modified T Cells Recognizing Mesothelin in Advanced Solid Cancers. *Molecular*
547 *therapy : the journal of the American Society of Gene Therapy* **27**, 1919-1929 (2019).
548
- 549 42. Ho M, *et al.* Mesothelin expression in human lung cancer. *Clinical cancer research : an*
550 *official journal of the American Association for Cancer Research* **13**, 1571-1575 (2007).
551
- 552 43. Tang Z, *et al.* A human single-domain antibody elicits potent antitumor activity by
553 targeting an epitope in mesothelin close to the cancer cell surface. *Molecular cancer*
554 *therapeutics* **12**, 416-426 (2013).
555
- 556 44. Robbins PF, *et al.* Single and dual amino acid substitutions in TCR CDRs can enhance
557 antigen-specific T cell functions. *Journal of immunology* **180**, 6116-6131 (2008).
558
- 559 45. Robbins PF, *et al.* A pilot trial using lymphocytes genetically engineered with an NY-
560 ESO-1-reactive T-cell receptor: long-term follow-up and correlates with response.
561 *Clinical cancer research : an official journal of the American Association for Cancer*
562 *Research* **21**, 1019-1027 (2015).
563
- 564 46. D'Angelo SP, *et al.* Antitumor Activity Associated with Prolonged Persistence of
565 Adoptively Transferred NY-ESO-1 (c259)T Cells in Synovial Sarcoma. *Cancer Discov*
566 **8**, 944-957 (2018).
567
- 568 47. Zuo J, *et al.* The Epstein-Barr virus-encoded BILF1 protein modulates immune
569 recognition of endogenously processed antigen by targeting major histocompatibility
570 complex class I molecules trafficking on both the exocytic and endocytic pathways.
571 *Journal of virology* **85**, 1604-1614 (2011).
572
- 573 48. Abu-Shah E, *et al.* Human CD8(+) T Cells Exhibit a Shared Antigen Threshold for
574 Different Effector Responses. *Journal of immunology* **205**, 1503-1512 (2020).
575
- 576 49. Itoh Y, Germain RN. Single cell analysis reveals regulated hierarchical T cell antigen
577 receptor signaling thresholds and intracлонаl heterogeneity for individual cytokine
578 responses of CD4+ T cells. *The Journal of experimental medicine* **186**, 757-766
579 (1997).
580
- 581 50. Majzner RG, *et al.* Tuning the Antigen Density Requirement for CAR T-cell Activity.
582 *Cancer Discov* **10**, 702-723 (2020).
583
- 584 51. Au-Yeung BB, *et al.* IL-2 Modulates the TCR Signaling Threshold for CD8 but Not CD4
585 T Cell Proliferation on a Single-Cell Level. *Journal of immunology* **198**, 2445-2456
586 (2017).
587
- 588 52. Ashouri JF, Weiss A. Endogenous Nur77 Is a Specific Indicator of Antigen Receptor
589 Signaling in Human T and B Cells. *Journal of immunology* **198**, 657-668 (2017).
590
- 591 53. Chen J, *et al.* NR4A transcription factors limit CAR T cell function in solid tumours.
592 *Nature* **567**, 530-534 (2019).

593

594

595

596

597

598

599

600

601

602

603

604

605

606

607

608

609

610

611

612

613

614

615

616

617

618

619

620

621

622

623

624

625

626

627

628

629

630

631

632

633

634

635

636

637

638

639

640

641

54. Liu X, *et al.* Genome-wide analysis identifies NR4A1 as a key mediator of T cell dysfunction. *Nature* **567**, 525-529 (2019).
55. Moran AE, *et al.* T cell receptor signal strength in Treg and iNKT cell development demonstrated by a novel fluorescent reporter mouse. *The Journal of experimental medicine* **208**, 1279-1289 (2011).
56. Guo XZ, Dash P, Calverley M, Tomchuck S, Dallas MH, Thomas PG. Rapid cloning, expression, and functional characterization of paired $\alpha\beta$ and $\gamma\delta$ T-cell receptor chains from single-cell analysis. *Molecular therapy Methods & clinical development* **3**, 15054 (2016).
57. Smith EL, *et al.* GPRC5D is a target for the immunotherapy of multiple myeloma with rationally designed CAR T cells. *Science translational medicine* **11**, (2019).
58. Eyquem J, *et al.* Targeting a CAR to the TRAC locus with CRISPR/Cas9 enhances tumour rejection. *Nature* **543**, 113-117 (2017).
59. Roth TL, *et al.* Reprogramming human T cell function and specificity with non-viral genome targeting. *Nature* **559**, 405-409 (2018).
60. Wang J, *et al.* Highly efficient homology-driven genome editing in human T cells by combining zinc-finger nuclease mRNA and AAV6 donor delivery. *Nucleic acids research* **44**, e30 (2016).
61. Koldej RM, *et al.* Comparison of insulators and promoters for expression of the Wiskott-Aldrich syndrome protein using lentiviral vectors. *Human gene therapy Clinical development* **24**, 77-85 (2013).
62. Robert-Richard E, Richard E, Malik P, Ged C, de Verneuil H, Moreau-Gaudry F. Murine retroviral but not human cellular promoters induce in vivo erythroid-specific deregulation that can be partially prevented by insulators. *Molecular therapy : the journal of the American Society of Gene Therapy* **15**, 173-182 (2007).
63. Weber EL, Cannon PM. Promoter choice for retroviral vectors: transcriptional strength versus trans-activation potential. *Human gene therapy* **18**, 849-860 (2007).
64. Zychlinski D, *et al.* Physiological promoters reduce the genotoxic risk of integrating gene vectors. *Molecular therapy : the journal of the American Society of Gene Therapy* **16**, 718-725 (2008).
65. Larson SM, *et al.* Pre-clinical development of gene modification of haematopoietic stem cells with chimeric antigen receptors for cancer immunotherapy. *Human vaccines & immunotherapeutics* **13**, 1094-1104 (2017).
66. Thomas S, Stauss HJ, Morris EC. Molecular immunology lessons from therapeutic T-cell receptor gene transfer. *Immunology* **129**, 170-177 (2010).

- 642 67. Kagoya Y, *et al.* A novel chimeric antigen receptor containing a JAK-STAT signaling
643 domain mediates superior antitumor effects. *Nature medicine* **24**, 352-359 (2018).
644
- 645 68. Jutz S, *et al.* Assessment of costimulation and coinhibition in a triple parameter T cell
646 reporter line: Simultaneous measurement of NF-kappaB, NFAT and AP-1. *Journal of*
647 *immunological methods* **430**, 10-20 (2016).
648
649

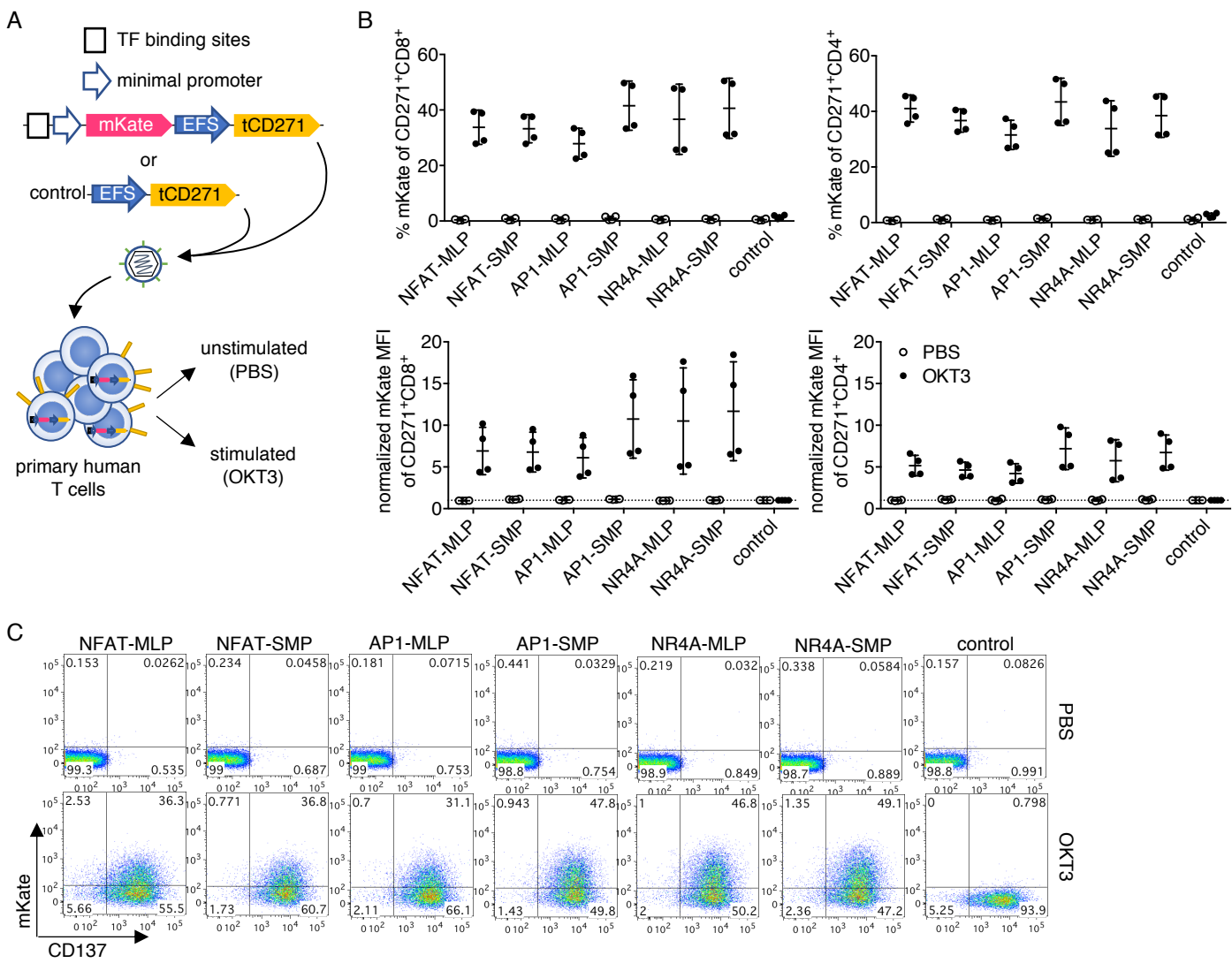


Figure 1. NFAT, AP1, and NR4A-based promoters are activated by anti-CD3 stimulation with minimal background. **a** Vector and experimental schematics. Response elements encoding NFAT, AP1, or NR4A-binding sites were cloned with either a core promoter from the adenovirus-derived major late promoter (MLP) or a synthetic minimal promoter (SMP). Lentiviral vectors were transduced into primary human T cells, and cells were treated with PBS or anti-CD3 clone OKT3. **b** Reporter fluorescence in transduced (CD271⁺) CD8⁺ or CD4⁺ cells was measured after 24 hours. **c** Representative flow plots gated on CD271⁺CD8⁺ T cells are shown. Lines and error bars denote mean \pm standard deviation. n=4 from 2 independent donors tested in 2 technical replicates.

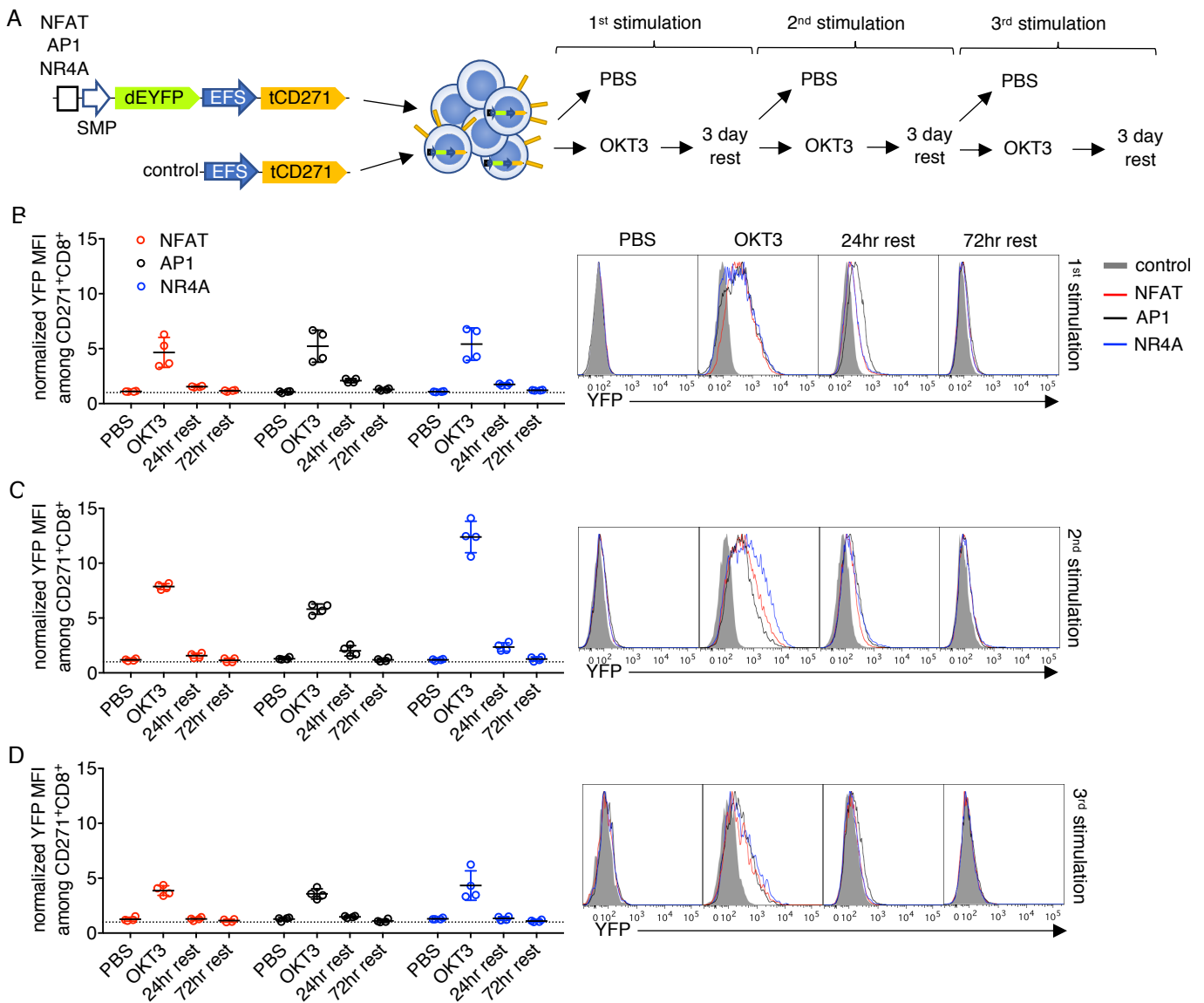


Figure 2. Inducible promoter responses are reversible and repeatable. **a** Vector and experimental schematics. Response elements encoding NFAT, AP1, or NR4A-binding sites with the synthetic minimal promoter (SMP) were used to drive destabilized yellow fluorescence protein (dEYFP). Lentiviral vectors were transduced into primary human T cells, and cells were treated with PBS or anti-CD3 clone OKT3 for 24 hours, then transferred to a fresh plate to rest for up to 3 days before repeating the process two more times. **b-d** Reporter fluorescence in CD271⁺CD8⁺ cells was measured after 24 hours of stimulation, then after 24 and 72 hours of rest, following the first (**a**), second (**b**), and third (**c**) round. Fluorescence intensity was normalized to that of control vector transductants. Representative histograms gated on CD271⁺CD8⁺ T cells are shown. Lines and error bars denote mean \pm standard deviation. $n=4$ from 2 independent donors tested in 2 technical replicates.

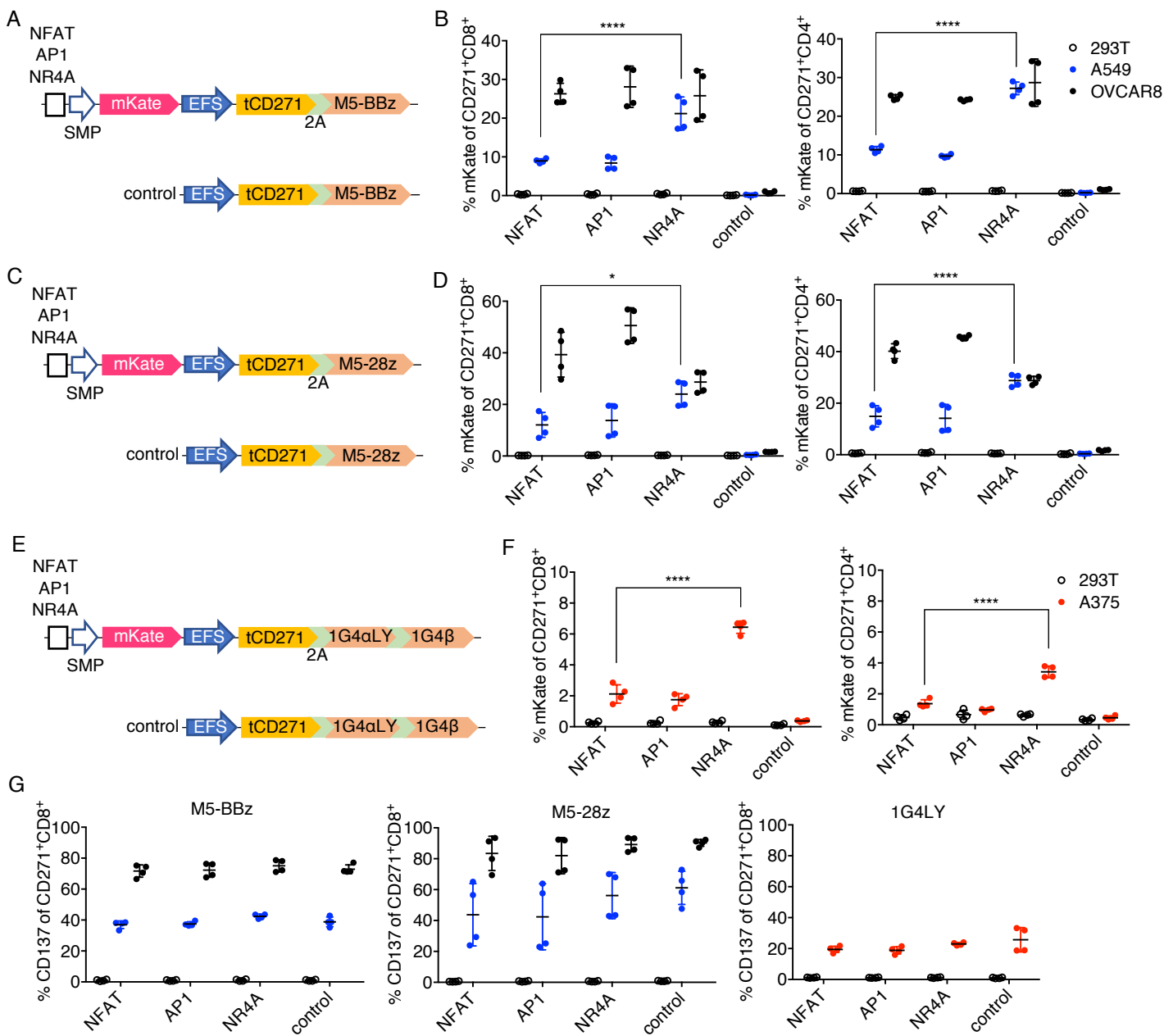


Figure 3. NR4A-based promoter induces higher responses than NFAT or AP1 in weakly stimulatory TCR/CAR-T models. **a,c,e** Schematics for CAR or TCR and inducible module encoded within a single vector. **b,d,f** Primary human T cells transduced with the vectors shown on the left of respective graphs were co-cultured with the indicated target cells, and mKate fluorescence was measured after 24 hours for CD271⁺CD8⁺ or CD4⁺ subsets. **g** CD137 expression on CD271⁺CD8⁺ T cells from the same experiments shown in panels **b**, **d**, and **f**. Lines and error bars denote mean \pm standard deviation. * $P < 0.05$, **** $P < 0.0001$ by two-way ANOVA adjusted for all possible comparisons using Tukey's test. $n = 4$ from 2 independent donors tested in 2 technical replicates.

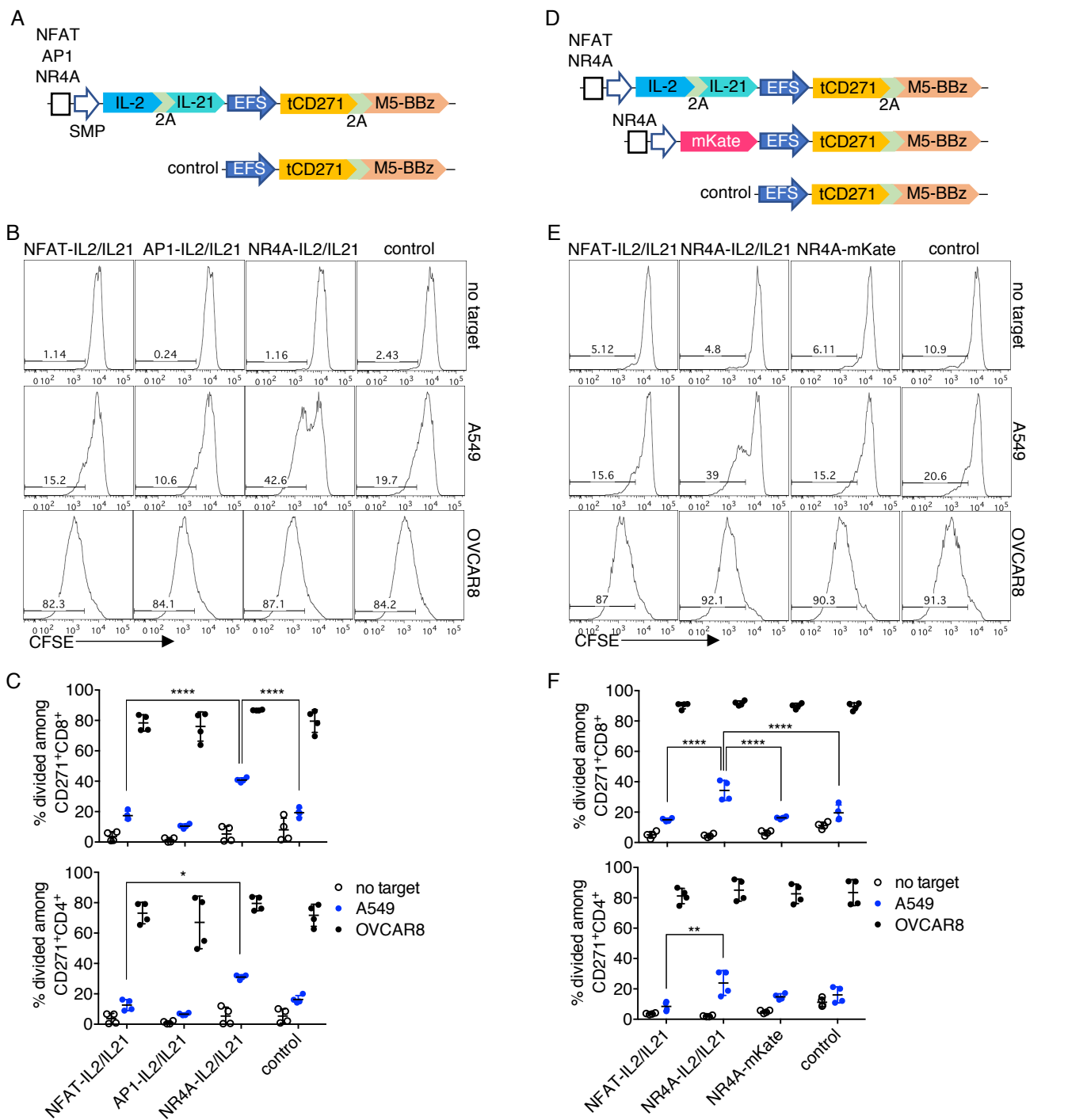
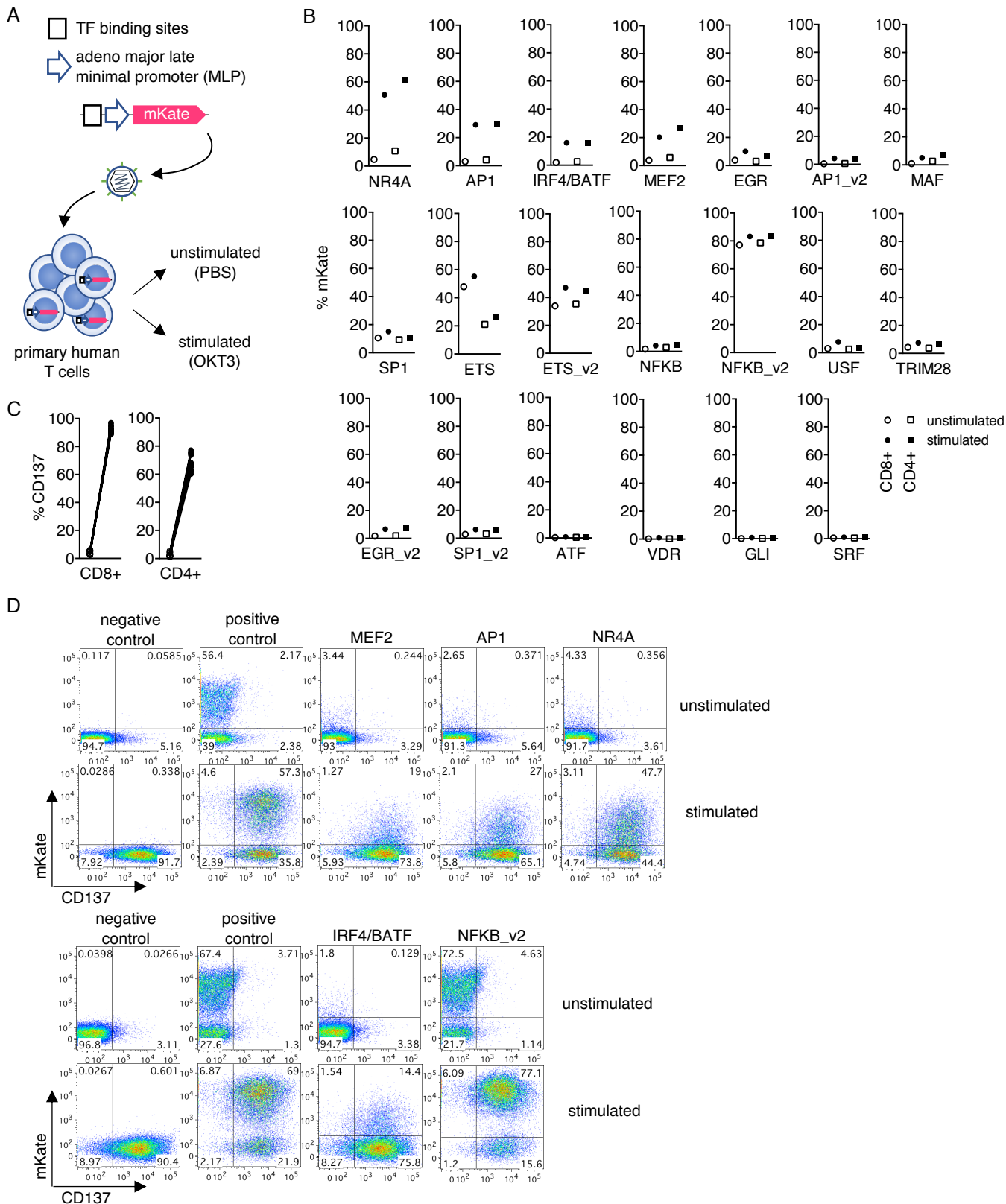
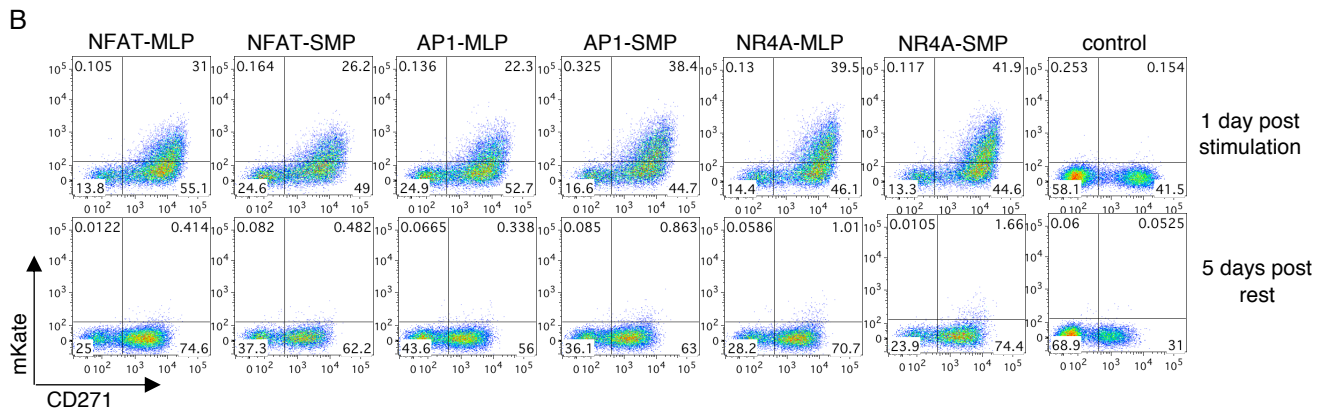
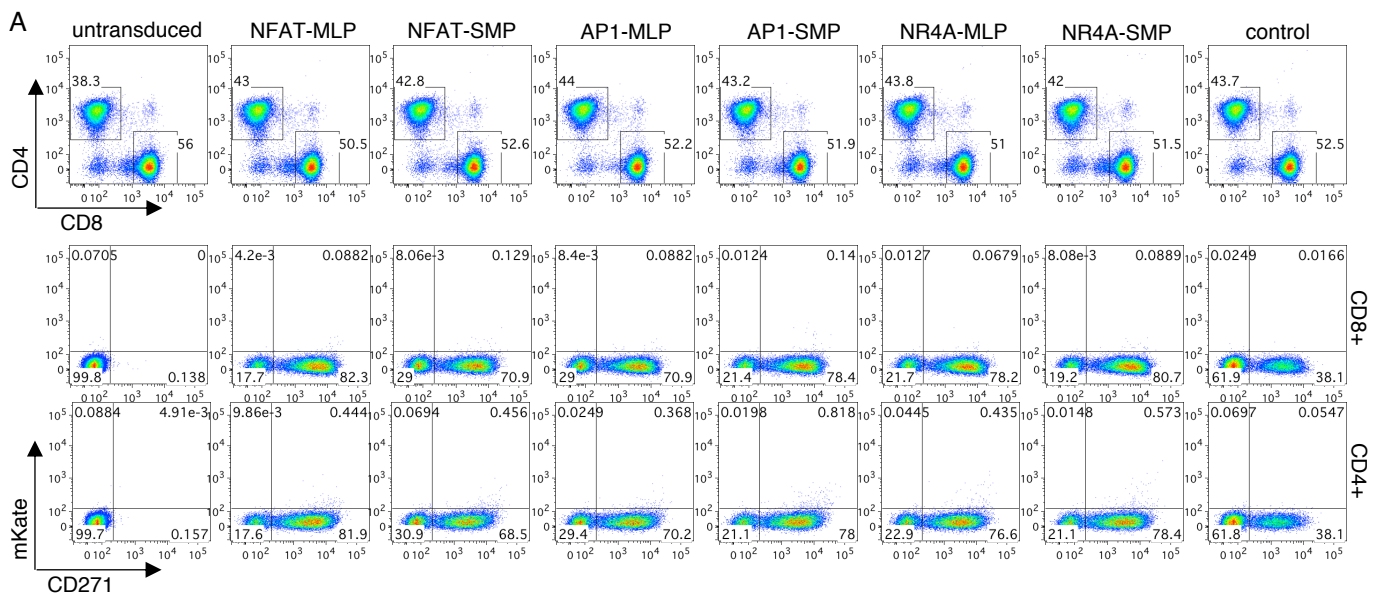


Figure 4. Cytokines delivered by the NR4A-based promoter amplify weak antigen-specific responses. **a** Schematics for vectors encoding M5-BBz CAR with or without IL-2 and IL-21 as inducible payloads. **b** Primary human T cells transduced with the vectors shown in **a** were labeled with CFSE and co-cultured with the indicated target cells. Dye dilution was measured after 4 days of co-culture. Representative histograms gated on CD271⁺CD8⁺ T cells are shown. **c** Quantification of CFSE dilution in CD271⁺CD8⁺ or CD4⁺ subsets. **d** Schematics for vectors encoding IL-2 and IL-21 as inducible payloads or mKate as a control. **e** Experiment was performed as described in **b**. Representative histograms gated on CD271⁺CD8⁺ T cells are shown. **f** Quantification of CFSE dilution in CD271⁺CD8⁺ or CD4⁺ subsets. Lines and error bars denote mean \pm standard deviation. * $P < 0.05$, ** $P < 0.01$, **** $P < 0.0001$ by two-way ANOVA adjusted for all possible comparisons using Tukey's test. $n = 4$ from 2 independent donors tested in 2 technical replicates.

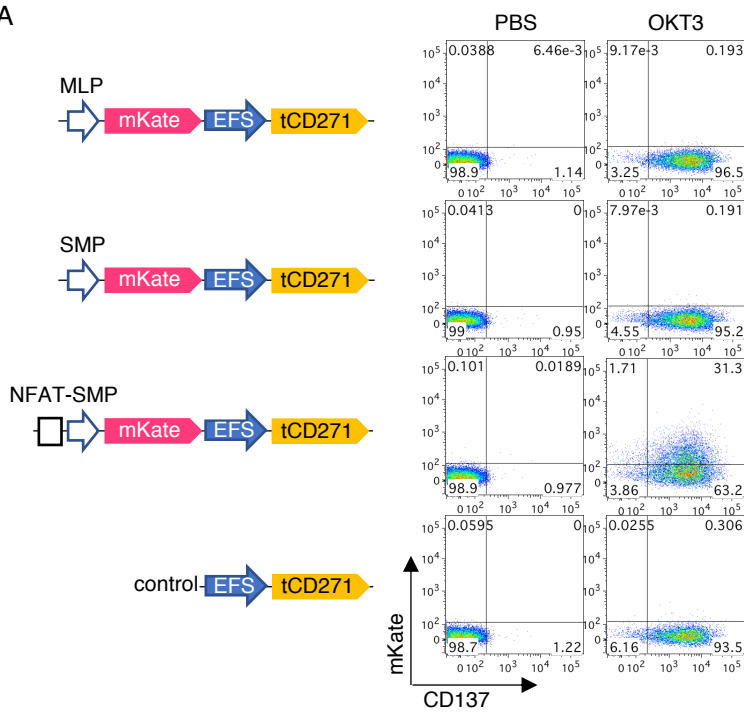


Supplementary Figure 1. Screening for novel TCR-inducible response elements. **a** Schematics for vector design from the SPECS library and experimental setup. **b** Primary human T cells transduced with a panel of vectors were treated with PBS (unstimulated, open symbols) or OKT3 (stimulated, filled symbols) for 24 hours, and mKate fluorescence in the CD8⁺ (circles) or CD4⁺ (squares) subset was quantified. **c** CD137 expression after PBS (open) or OKT3 (filled) treatment. Measurements are matched by the same vector. **d** Raw flow plots gated on CD8⁺ T cells for several notable promoters are shown. The positive control vector encodes the constitutive UBC promoter to express mKate, whereas the negative control vector lacks mKate.

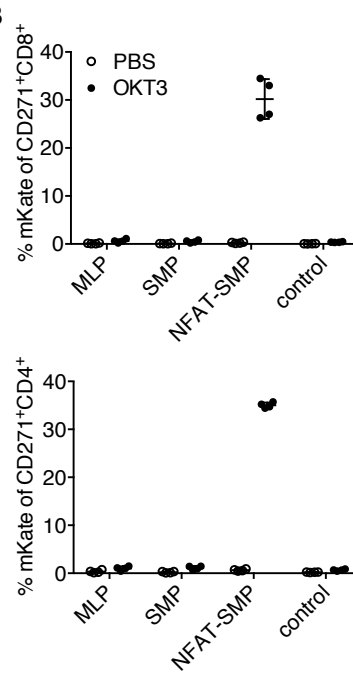


Supplementary Figure 2. Gating strategy, transduction efficiency, and mKate stability. **a** CD4⁺ or CD8⁺ T cells are gated as shown on the top row. Transduction efficiency measured by CD271 positivity is shown on the bottom rows. **b** Fluorescence of the mKate reporter after one day of OKT3 stimulation and after 5 days of rest upon transfer to fresh wells is shown for representative CD8⁺ cells.

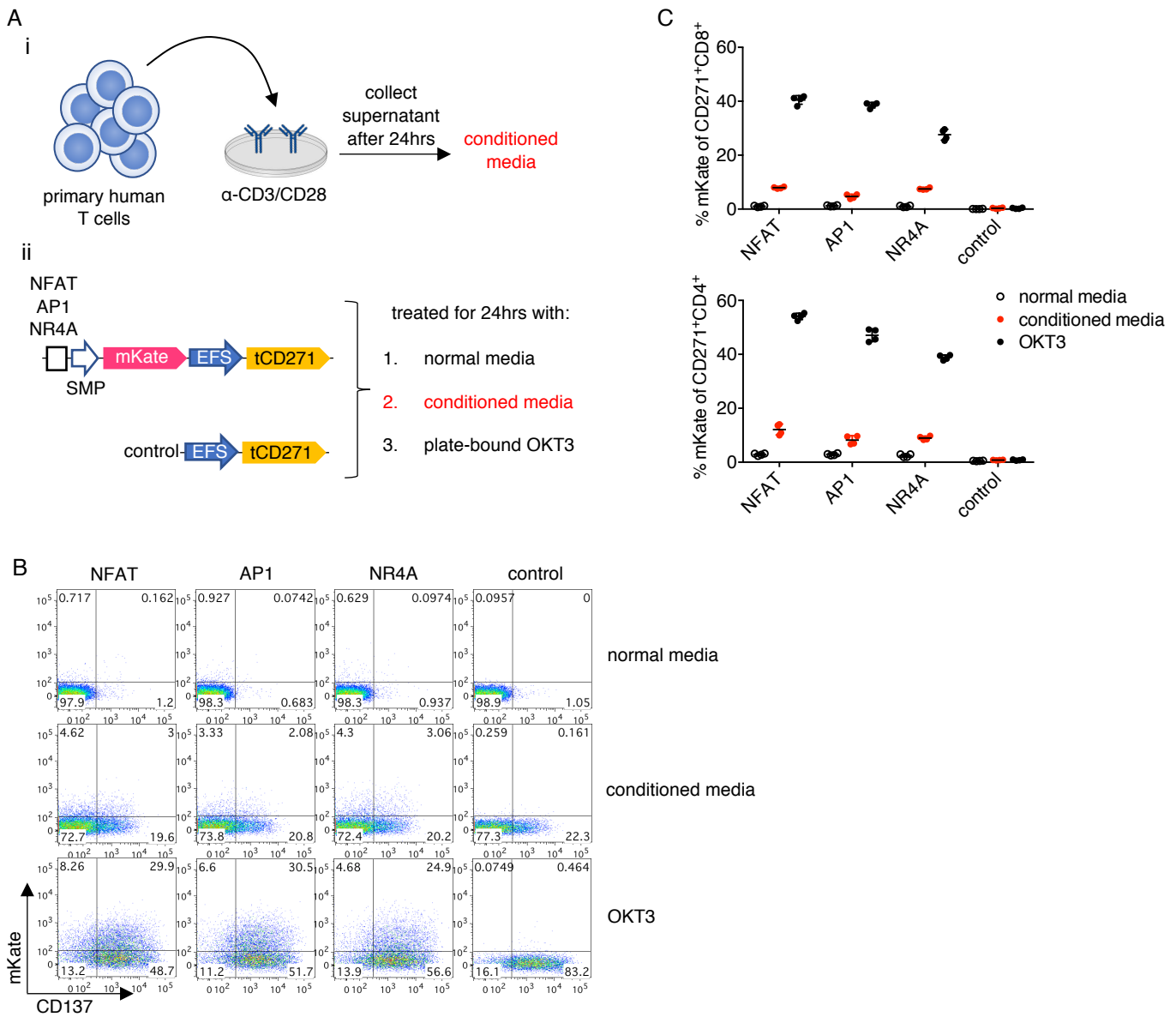
A



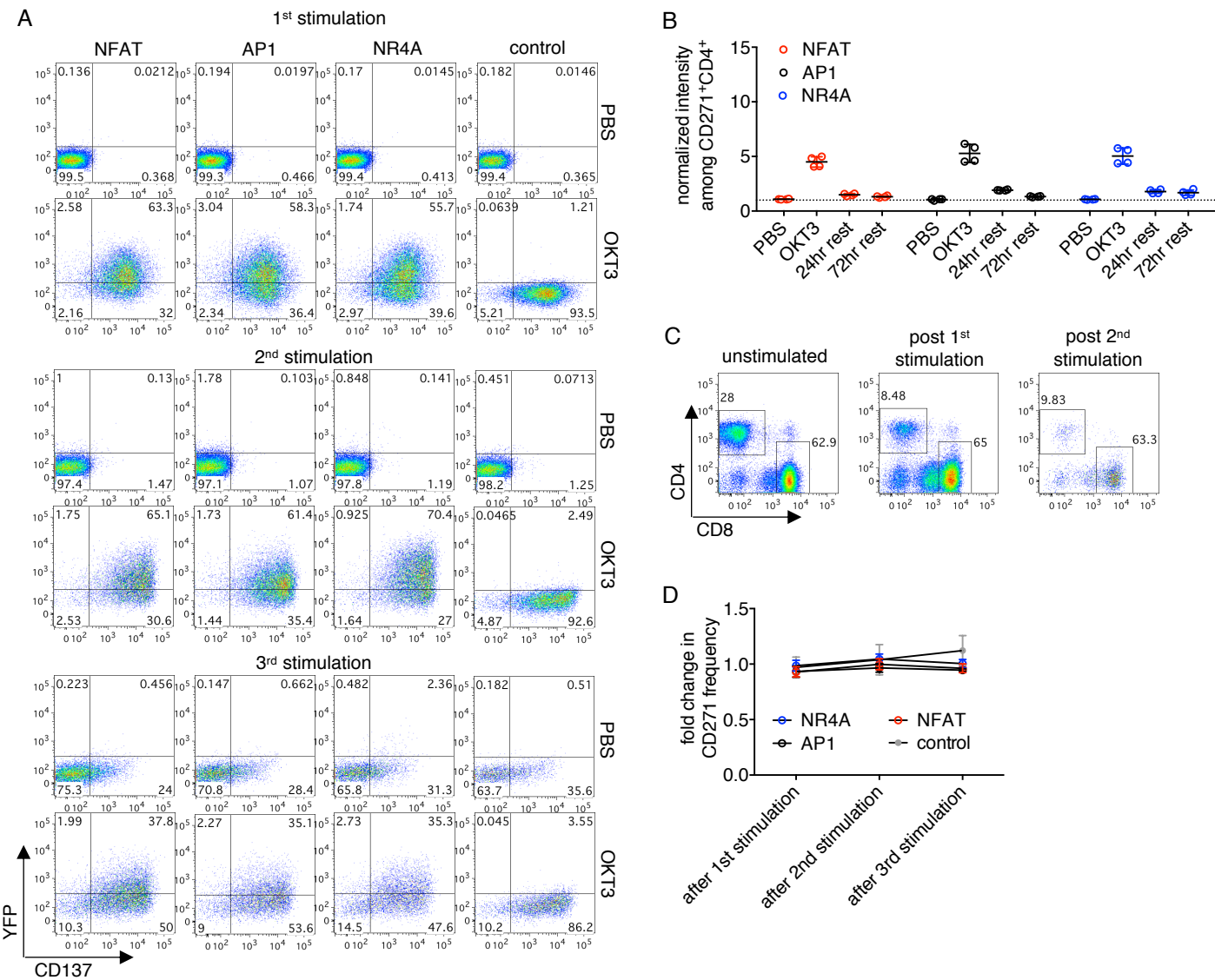
B



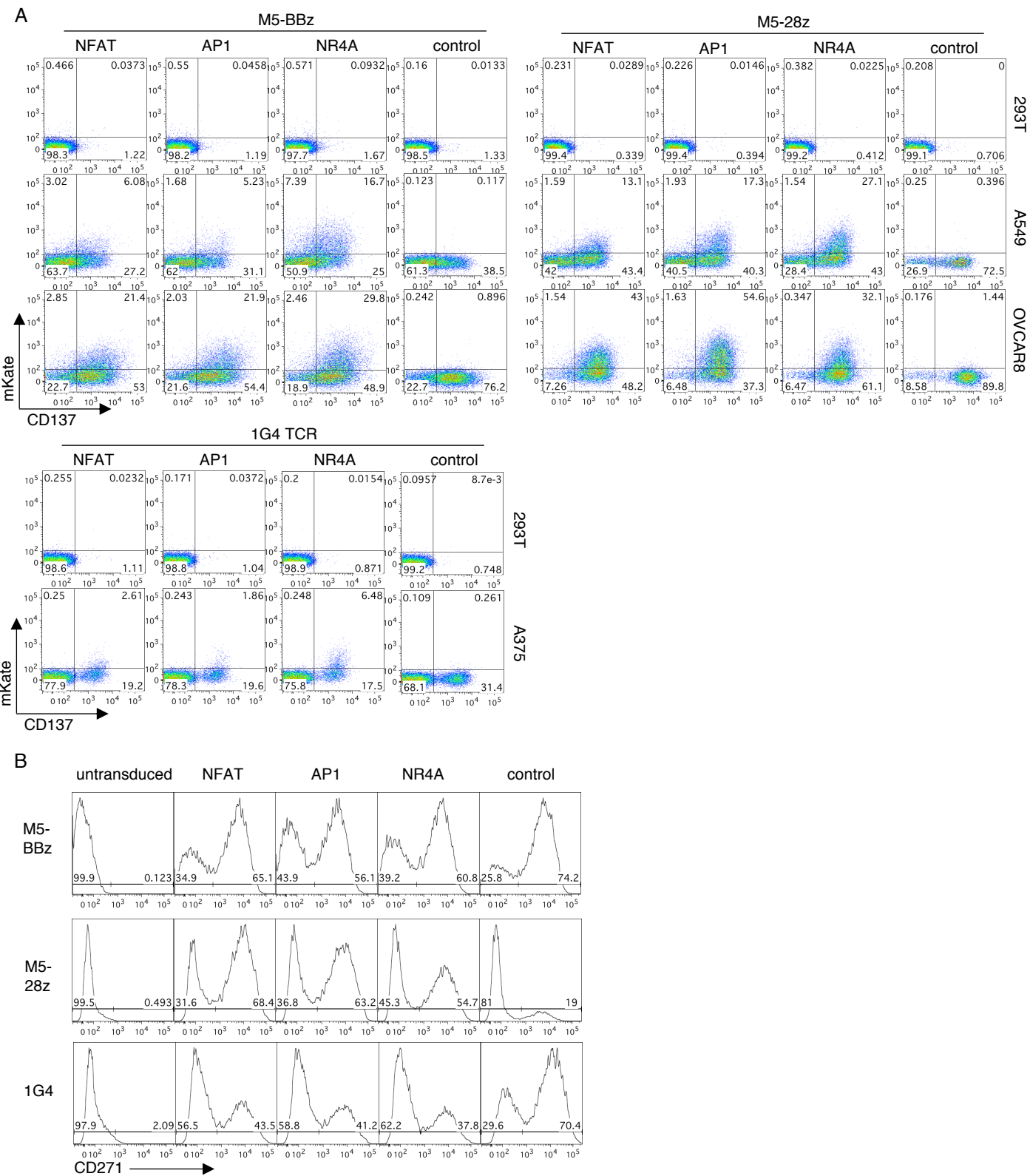
Supplementary Figure 3. The EFS constitutive promoter do not possess detectable enhancer-like activity. **a** Primary human T cells transduced with the indicated vector shown on the left were treated with PBS or OKT3. CD137 and mKate upregulation was measured after 24 hours. Representative flow plots gated on CD271+CD8+ cells are shown on the right. **b** Quantification of data shown in panel **a**. Lines and error bars denote mean \pm standard deviation. n=4 from 2 independent donors tested in 2 technical replicates.



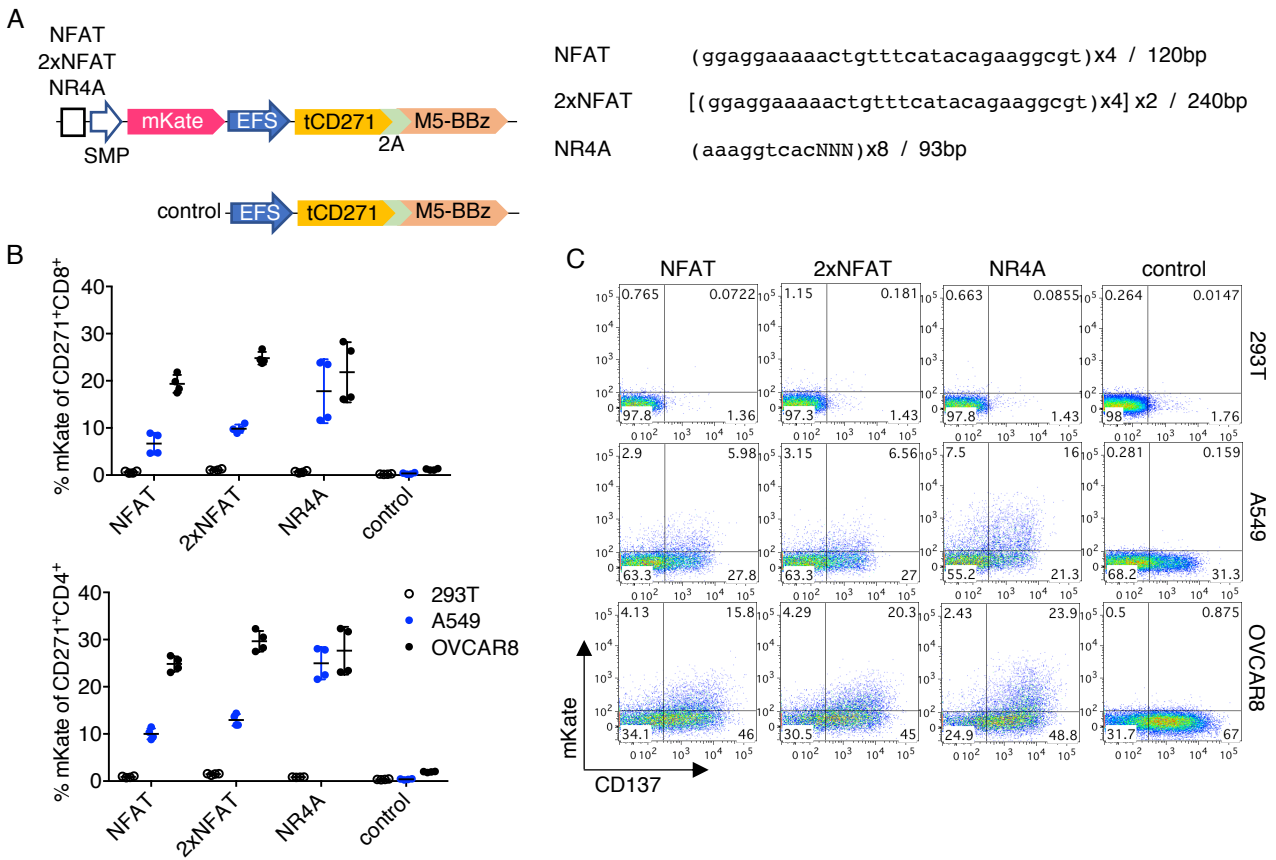
Supplementary Figure 4. TCR-inducible promoters are weakly activated by an inflammatory milieu. **a** Experimental design. **i**) Conditioned media was generated by collecting the supernatant from expanded primary human T cells restimulated with plate-bound anti-CD3/CD28 monoclonal antibodies. **ii**) Promoter or control transduced T cells were cultured with normal media, conditioned media, or plate-bound OKT3. **b** Representative flow plots gated on CD271⁺CD8⁺ cells are shown. **c** Quantification of data shown in panel **b**. Lines and error bars denote mean \pm standard deviation. $n=4$ from 2 independent donors tested in 2 technical replicates.



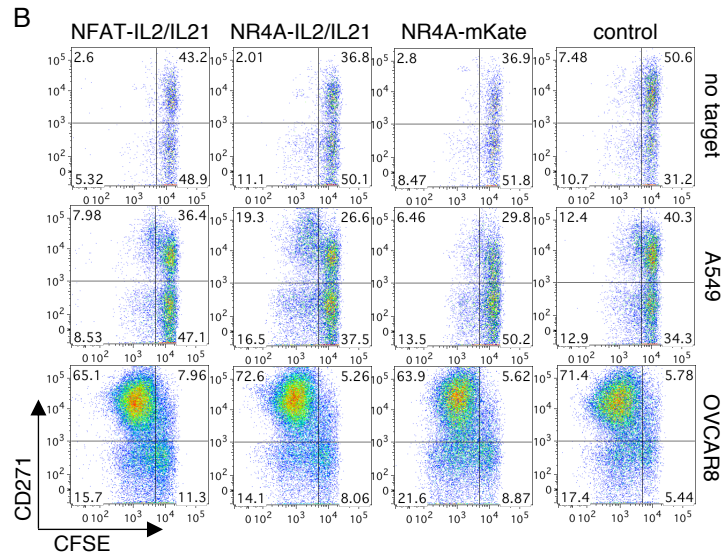
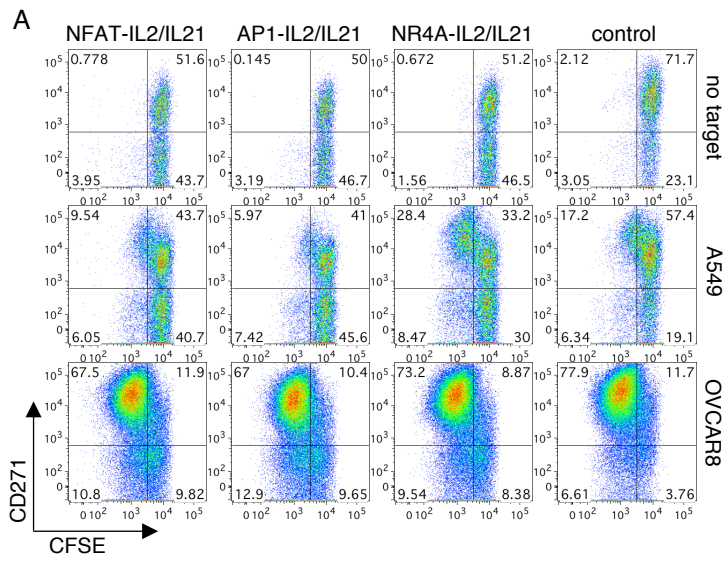
Supplementary Figure 5. Inducible promoters can be repeatedly stimulated without causing cellular toxicity. **a** Representative flow plots gated on CD271⁺CD8⁺ cells for the PBS and OKT3 conditions in Figure 2B-D are shown. **b** Reversible promoter responses among CD271⁺CD4⁺ T cells were measured, as described in the caption of Figure 2, after the first round of stimulation. **c** Representative frequency of CD4⁺ and CD8⁺ subsets in culture after repeated stimulation. Gated on live cells. Due to the activation-induced cell death and biased outgrowth of CD4⁻ cells, CD4⁺ cells could not be reliably analyzed after the first round of stimulation. **d** Change in percent of transduced (CD271⁺) cells was tracked through the three rounds of stimulation and remained constant. Frequency was normalized to that at the start of the experiment (prior to first stimulation). Lines and error bars denote mean \pm standard deviation. $n=4$ from 2 independent donors tested in 2 technical replicates.



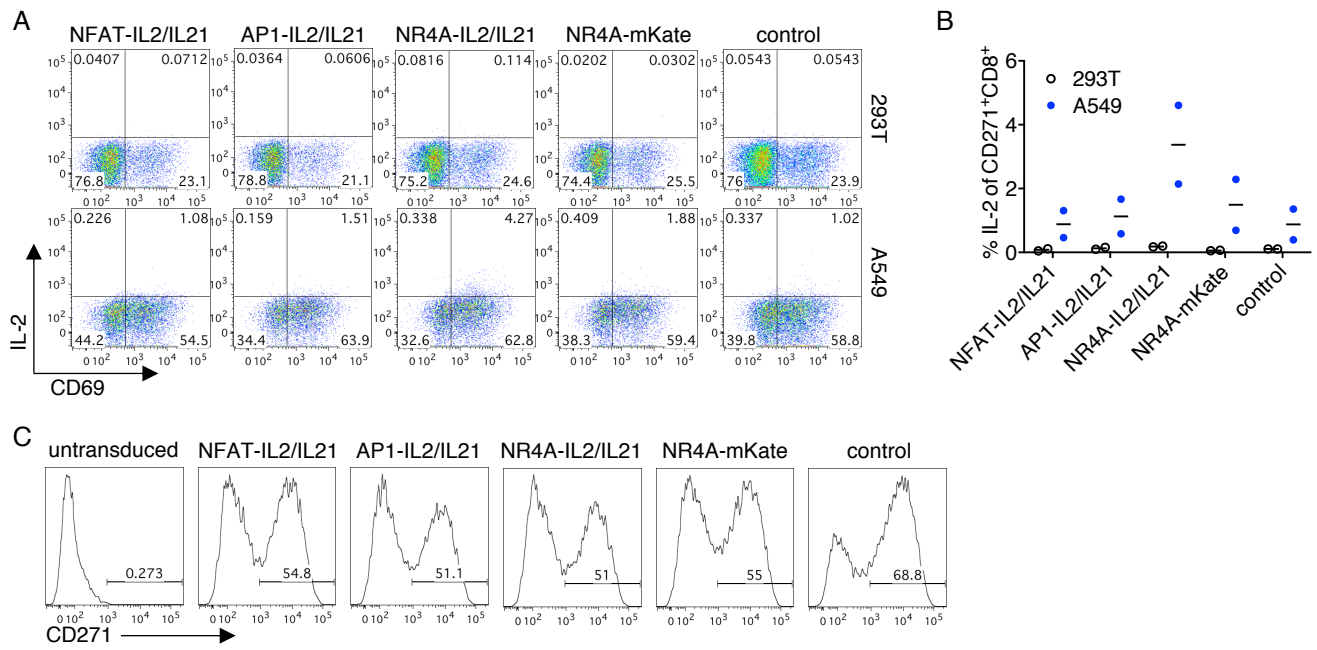
Supplementary Figure 6. Inducible promoter activities in TCR/CAR-T models. **a** Representative flow plots gated on CD271+CD8+ cells for data in Figure 3 are shown. **b** Representative transduction efficiencies of the TCR/CAR vectors. Gated on total live cells. Note that the control M5-28z transduction was unexpectedly low. However, this does not affect the interpretation of the data, as the control M5-28z vector mainly serves as a gating control for mKate encoding vectors.



Supplementary Figure 7. Doubling the number of NFAT binding sites only marginally improves response. **a** Left, vector schematics. Right, promoter sequences and designs. Two copies of the original NFAT promoter with 4 binding sites were cloned to generate 2xNFAT. The NR4A promoter was derived from the SPECS library, where transcription factor binding motifs were spaced by three random nucleotides. **b** Quantification of reporter responses among CD271⁺CD8⁺ or CD4⁺ subsets. **c** Representative flow plots gated on CD271⁺CD8⁺ cells for data shown in **b**. Lines and error bars denote mean \pm standard deviation. n=4 from 2 independent donors tested in 2 technical replicates.



Supplementary Figure 8. Inducible expression of cytokines augments weak antigen-specific response in CAR-T cells. a Representative flow plots gated on CD8⁺ cells for data in Figure 4A-C are shown. **b** Representative flow plots gated on CD8⁺ cells for data in Figure 4D-F are shown.



Supplementary Figure 9. Analyses of inducible payload expression in stimulated CAR-T cells. a CAR-T cells transduced with inducible IL-2/IL-21 or control constructs were stimulated with 293T or A549 for 18 hours. Monesin was then added, and cells were cultured for another 6 hours before staining. Representative flow plots gated on CD271⁺CD8⁺ cells are shown. b Quantification for data shown in a across two independent donors. Lines denote means. c Representative transduction efficiencies of vectors used in this experiment and Figure 4.

ID	Ref.	Full sequence	Length (bp)	Motif
AP1	20	TTGAGTCAAGATTGAGTCATCGTTGAGTCAGACTTGAGTCACTATTGAGTCAACTTTGAGTCATGCTTGAGTCAGTATTGAGTCA	85	TTGAGTCA
AP1_v2		GGTGACTCATGAGAGGTGACTCATGTCCGGTGACTCATGGACGGTGACTCATGCTAGGTGACTCATGACTGGTGACTCATGTGCG GTGACTCATGGTAGGTGACTCATG	109	GGTGACTCATG
ATF	21	CTCTGACGTCAAGACTCTGACGTCACTCGCTCTGACGTACAGCTCTGACGTCACTACTCTGACGTCAACTCTCTGACGTCAAGC TCTGACGTCACTCTGACGTCA	109	CTCTGACGTCA
EGR	27	CCGCCACGCAGACCGCCACGCTCGCCGCCACGCGACCCGCCACGCCTACCGCCACGCACCTCCGCCACGCTGCCCGCCCA CGCGTACCGCCACGC	101	CCGCCACGC
EGR_v2		TGCGTGGGCGAGATGCGTGGGCGTCTGCGTGGGCGGACTGCGTGGGCGCTATGCGTGGGCGACTTGGTGGGCGTCTGCGTGG GCGGTATGCGTGGGCG	101	TGCGTGGGCG
ETS	22	CCGGAAGAGACCCGGAAGTCGCCGGAAGGACCCGGAAGCTACCGGAAGACTCCGGAAGTGCCCGGAAGGTACCGGAAG	77	CCGGAAG
ETS_v2		CCGGAAGTGGCAGACCCGGAAGTGGCTCGCCGGAAGTGGCGACCCGGAAGTGGCTACCGGAAGTGGCACTCCGGAAGTGGCTGCC CGGAAGTGGCGTACCGGAAGTGGC	109	CCGGAAGTGGC
GLI	28	CCTGGGTGGTCCAGACTGGGTGGTCTCGCTGGGTGGTCCGACCTGGGTGGTCCCTACCTGGGTGGTCCACTCTGGGTGGT CCTGCCCTGGGTGGTCCGTACCTGGGTGGTCC	117	CCTGGGTGGTCC
IRF4/BATF	29	GAAATGAGTCAAGAGAAATGAGTCATCGAAATGAGTCAGACGAAATGAGTCACTAGAAATGAGTCAACTGAAATGAGTCATGCG AAATGAGTCAGTAGAAATGAGTCA	109	GAAATGAGTCA
MAF	30	TGCTGACTCAGCAAGATGCTGACTCAGCATCGTGTGACTCAGCAGACTGCTGACTCAGCACTATGCTGACTCAGCAACTTGCTG ACTCAGCATGCTGCTGACTCAGCAGTATGCTGACTCAGCA	125	TGCTGACTCAGCA
MEF2	23	ACTATAAATAGAAGAACTATAAATAGATCGACTATAAATAGAGACACTATAAATAGACTAACTATAAATAGAAGTACTATAAATA GATGCACTATAAATAGAGTAACTATAAATAGA	117	ACTATAAATAGA
NFAT	16	GGAGGAAAAACTGTTTCATACAGAAGGCGTGGAGGAAAAACTGTTTCATACAGAAGGCGTGGAGGAAAAACTGTTTCATACAGAA GGCGTGGAGGAAAAACTGTTTCATACAGAAGGCGT	120	GGAGGAAAAACTGTTTC ATACAGAAGGCGT
NFKB	24	GGGGAAATTTCCCTAGAGGGGAAATTTCCCTTCCGGGGAAATTTCCCTTACGCGGGAAATTTCCCTTACGCGGGAAATTTCCCTACT GGGGAAATTTCCCTTCCGGGGAAATTTCCCTTGT	119	GGGGAAATTTCCCT
NFKB_v2		AGGGGATTTCCAAGGAGAAGGGGATTTCCAAGGTTCGAGGGGATTTCCAAGGGACAGGGGATTTCCAAGGCTAAGGGGATTTCCAA GGACTAGGGGATTTCCAAGGTGCAGGGGATTTCCAAGGGT	126	AGGGGATTTCCAAGG
NR4A	31	AAAGGTCACAGAAAAGGTCACTCGAAAGGTCACGACAAAAGGTCACCTAAAAGGTCACACTAAAGGTCACTGCAAAGGTCACGTAA AAGGTCAC	93	AAAGGTCAC
SP1	32	AAGTGGGCGTGGCCAGAAAGTGGGCGTGGCCTCGAAGTGGGCGTGGCCGACGGCGTGGCCCTAAAGTGGGCGTGGCCACTAATGG GCGTGGCCTGCAAGTGGGCGTGGCCGTA	113	AAGTGGGCGTGGCC
SP1_v2		CCCTCCCCAAGGCTAGACCCTCCCCAAGGCTTCGCCCTCCCCAAGGCTGACCCCTCCCCAAGGCTCTACCCTCCCCAAGGCTACT CCCTCCCCAAGGCTTGGCCCTCCCCAAGCTGTA	118	CCCTCCCCAAGGCT
SRF	25	ATGCCCATATATGGAAGAATGCCCATATATGGATCGATGCCCATATATGGAGACATGCCCATATATGGACTAATGCCCATATATG GAATATGCCCATATATGGATGCATGCCCATATATGGAGTA	126	ATGCCCATATATGGA
TRIM28	33	GGTTTCTCTAGAGGTTTCTCTTCCGGTTTCTCTGACGGTTTCTCTCTAGGTTTCTCTACTGGTTTCTCTTGGGTTTCTCTGTA GGTTTCTCT	94	GGTTTCTCT
USF	26	GGTCACGTGACAGAGGTACAGTACTCGGGTACAGTACGACGGTACAGTACCTAGGTACAGTACACTGGTACAGTACTGCG GTCACGTGACGTAGGTACAGTAC	109	GGTCACGTGAC
VDR	34	GGGTTACCGGGAGAGGGTTACCGGGTTCGGGGTTCACCGGGACGGGTTACCTGGCTAGGGTTACCGGGACTGGGTTACCGG GGTTCGGGTTACCGGGTAGGGTTACCGGG	117	GGGTTACCGGG

Supplementary Table 1. Transcription factor (TF) binding site sequences. Sequences and length for the promoters studied in Supplementary Figure 1 and the conventional NFAT promoter are shown. Bolded promoters were studied in the main figures. References for association of respective TF pathway with TCR signaling are listed.