# 01 open and split

October 1, 2024

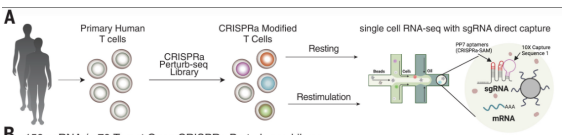
#### 0.0.1 Paper

# 1 CRISPR activation and interference screens decode stimulation responses in primary human T cells

https://www.science.org/doi/10.1126/science.abj4008#body-ref-R56-1

### 2 Data

- https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE190604
- https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE174292
- https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE190846
- https://zenodo.org/records/5784651 (R Repository with results)



B ~150 sgRNA / ~70 Target Gene CRISPRa Perturb-seq Library

We performed CRISPRa Perturb-seq characterization of regulators of stimulation responses in ~56,000 primary human T cells, targeting 70 hits and controls from our genome-wide CRISPRa cytokine screens (Fig. 4, A and B, and fig. S17, A to C). First, we confirmed that sgRNAs led to significant increases in the expression of their target genes (fig. S17D). Next, uniform manifold approximation and projection (UMAP) dimensionality reduction revealed discrete separation of the resting and restimulated cells (fig. S17E) and showed relatively even distribution of cells from two donors (Fig. 4C and fig. S17F). Gene signatures allowed us to resolve most T cells as either CD4+ or CD8+ (Fig. 4D and fig. S17, G and H). Thus, we generated a high-quality CRISPRa Perturb-seq dataset.

#### 2.0.1 Data experimento:

- ~150 sgRNA
- $\sim 70$  Target Gene
- ~56,000 primary human T cells

## 3 Data Loading and preprocess

#### 3.0.1 Raw Data (scRNA-seq)

```
Empty DataFrame
Columns: []
Index: [AAACCCACAACAAGAT-1, AAACCCACAACGGCTC-1, AAACCCACACAGAAGC-1,
AAACCCACACCTGTT-1, AAACCCACTATGTG-1, AAACCCAGTACAGGTG-1, AAACCCAGTATGAGAT-1,
AAACCCAGTTACGATC-1, AAACCCATCGGCATTA-1, AAACCCATCTACCTTA-1, AAACCCATCTGTGCAA-1,
AAACGAAAGAACTCCT-1, AAACGAAAGCATGCAG-1, AAACGAAAGCTATCCA-1, AAACGAAAGGAGTATT-1,
AAACGAAAGTCGCCCA-1, AAACGAAAGTGGACTG-1, AAACGAACAACTGCCG-1, AAACGAACAAGTTTGC-1,
AAACGAACAGAAATCA-1, AAACGAACAGCTTCGG-1, AAACGAACAGTCGGTC-1, AAACGAACAGTTAAAG-1,
AAACGAAGTCACTTAG-1, AAACGAAGTCTCGCGA-1, AAACGAAGTGTCCTAA-1, AAACGAAGTTCATCTT-1,
AAACGAATCAAGCCAT-1, AAACGCTAGAAGTCTA-1, AAACGCTAGAGCATTA-1, AAACGCTAGCCACAAG-1,
AAACGCTAGTACAGCG-1, AAACGCTAGTACGAGC-1, AAACGCTAGTGGCGAT-1, AAACGCTAGTTGCATC-1,
AAACGCTCAGCGTACC-1, AAACGCTCATCAGCTA-1, AAACGCTGTCACGACC-1, AAACGCTGTGTAACGG-1,
AAACGCTGTTGATGTC-1, AAACGCTTCCAAGCAT-1, AAACGCTTCGGTAGGA-1, AAAGAACAGCCTCTTC-1,
AAAGAACAGCCTGTGC-1, AAAGAACAGCTCCGAC-1, AAAGAACAGGACTATA-1, AAAGAACAGTGCAACG-1,
AAAGAACAGTTGTACC-1, AAAGAACCAGTCTCTC-1, AAAGAACCATATCTCT-1, AAAGAACGTCGTGGTC-1,
AAAGAACGTTACGATC-1, AAAGAACTCAACACCA-1, AAAGAACTCAACGTGT-1, AAAGAACTCAGTAGGG-1,
AAAGAACTCCACAGCG-1, AAAGAACTCTATCGTT-1, AAAGGATAGAAACCC-1, AAAGGATAGATGAAGG-1,
AAAGGATAGCATCAAA-1, AAAGGATAGGTTGCCC-1, AAAGGATCACGCACCA-1, AAAGGATCAGTTGTCA-1,
AAAGGATCAGTTGTTG-1, AAAGGATGTAGGACCA-1, AAAGGATGTATGACAA-1, AAAGGATGTGGACCAA-1,
AAAGGATGTTACGTAC-1, AAAGGATTCAGCGCAC-1, AAAGGATTCTAACGGT-1, AAAGGATTCTAGTGTG-1,
AAAGGGCAGATTGGGC-1, AAAGGGCCAACCACAT-1, AAAGGGCCAAGACTGG-1, AAAGGGCCAGAGTGTG-1,
AAAGGGCCAGGACTTT-1, AAAGGGCCAGGGTCTC-1, AAAGGGCCAGTGACCC-1, AAAGGGCCATCAACCA-1,
AAAGGGCCATGCAGCC-1, AAAGGGCGTCCGAAGA-1, AAAGGGCGTCTCGCGA-1, AAAGGGCGTGAACGGT-1,
AAAGGGCTCGCTTGCT-1, AAAGGGCTCGGTAGAG-1, AAAGGGCTCTGCATGA-1, AAAGGTAAGGGCCCTT-1,
AAAGGTACATACAGAA-1, AAAGGTACATTGTCGA-1, AAAGGTAGTAACATAG-1, AAAGGTAGTACGACTT-1,
AAAGGTAGTCACCTTC-1, AAAGGTAGTCACTGAT-1, AAAGGTAGTGACTGAG-1, AAAGTCCAGAACTGAT-1,
AAAGTCCAGGGCAAGG-1, AAAGTCCAGGGCTGAT-1, AAAGTCCCACGACCTG-1, AAAGTCCCACGATTCA-1,
AAAGTCCCAGACCTAT-1, ...]
```

[103805 rows x 0 columns]

Raw data has 36755 genes and 103805 cells

The number -1, -2, etc, represent the chromium well

A chromium well refers to the microfluidic chambers in the 10x Genomics Chromium Controller that are used to encapsulate single cells and barcoded beads into individual droplets, enabling high-throughput single-cell genomics. These wells play a key role in isolating single cells, capturing their RNA, and associating it with unique barcodes for sequencing.

Gene Expression: Represent the transcriptomic profile of the cells (the genes that are being expressed)

CRISPR Guide Capture: Which sgRNAs (and therefore which genes) were targeted in each cell.

	gene_ids	feature_types
MTD 1000 OIIG	0 -	- v-
MIR1302-2HG	ENSG00000243485	Gene Expression
FAM138A	ENSG00000237613	Gene Expression
OR4F5	ENSG00000186092	Gene Expression
AL627309.1	ENSG00000238009	Gene Expression
AL627309.3	ENSG00000239945	Gene Expression
•••	•••	•••
TRIM21-2	TRIM21-2	CRISPR Guide Capture
VAV1-1	VAV1-1	CRISPR Guide Capture
VAV1-2	VAV1-2	CRISPR Guide Capture
WT1-1	WT1-1	CRISPR Guide Capture
WT1-2	WT1-2	CRISPR Guide Capture
[36755 rows	x 2 columns]	
fo	ature tunes coun	+

feature\_types count

Gene Expression 36601

CRISPR Guide Capture 154

#### 3.0.2 Cell Metadata

From the CRISPR Guide Capture columns we can get the cell metada reference: 01\_build\_metadata\_table\_for\_guide\_calls

	condition	crispr	guide_id	gene	well
cell_barcode					
GGGAGATAGACCGTTT-1	Nostim	perturbed	ABCB10-1	ABCB10	1
GACGCTGCATTGTCGA-1	Nostim	perturbed	ABCB10-1	ABCB10	1
TTAATCCTCGTGCACG-1	Nostim	perturbed	ABCB10-1	ABCB10	1
ACACGCGTCGACCTAA-1	Nostim	perturbed	ABCB10-1	ABCB10	1
CATCCACCATCGATGT-1	Nostim	perturbed	ABCB10-1	ABCB10	1
•••	•••	•••			
GTTGTCCGTGGTTTAC-8	Stim	perturbed	WT1-2	WT1	8
GTCTAGAAGGCACTCC-8	Stim	perturbed	WT1-2	WT1	8
TCCTAATCATACACCA-8	Stim	perturbed	WT1-2	WT1	8
AGACCCGGTATTGACC-8	Stim	perturbed	WT1-2	WT1	8
AGTGTTGTCATTTACC-8	Stim	perturbed	WT1-2	WT1	8

[60657 rows x 5 columns]

#### Perturbed vs No TARGET

crispr count
0 perturbed 56774
1 NT 3883

# $4 \sim 56,000 \text{ cels}$

### Total cells per guide id

	guide_id	count
0	TRAF3IP2-1	1056
1	LAT2-2	983
2	EMP3-1	931
3	CD27-1	849
4	TNFRSF1B-2	797
	•••	•••
 149	 PRDM1-2	<b></b> 20
 149 150	 PRDM1-2 IRX4-2	 20 15
150	IRX4-2	15
150 151	IRX4-2 DEF6-2	15 11

[154 rows x 2 columns]

# $5 \quad {\sim}150{,}000 \ sgRNAs$

## Total cells per gene

	gene	count
0	NO-TARGET	3883
1	EMP3	1588
2	TRAF3IP2	1564
3	CD27	1455
4	TNFRSF1B	1344
		•••
 69	 EOMES	<b></b> 262
 69 70	 EOMES HELZ2	 262 128
••		
70	HELZ2	128
70 71	HELZ2 TCF7	128 115

[74 rows x 2 columns]

# $6 \sim 70 \text{ genes}$

## Total cells per gene per guide\_id

ABCB10 AKAP12 ALX4 APOBEC3C guide\_id count guide\_id count guide\_id count guide\_id count

0	ABCB10-2	208	AKAP12-1	255	ALX4-1	380	APOBEC3C-1	486
1	ABCB10-1	189	AKAP12-2	104	AT.X4-2	132	APOREC3C-2	168

# Total cells per well

	well	count
0	2	7838
1	3	7750
2	1	7742
3	5	7642
4	7	7502
5	4	7491
6	6	7430
7	8	7262

	ABCB10-1	ABCB10-2	AKAP12-1	AKAP12-2	ALX4-1	ALX4-2	APOBEC3C-1	\
well								
0	29	37	45	19	62	26	71	
1	28	34	39	18	51	20	70	
2	24	33	34	14	48	19	62	
3	24	27	33	12	47	16	62	
4	24	26	30	12	47	15	59	
5	20	22	26	12	44	14	57	
6	20	15	24	10	41	12	54	
7	20	14	24	7	40	10	51	

	APOBEC3C-2	APOBEC3D-1	APOBEC3D-2
well			
0	27	40	57
1	26	34	50
2	25	31	47
3	21	30	45
4	19	28	43
5	18	28	42
6	18	27	41
7	14	25	34

## 6.0.1 Merge and clean data

	condition	crispr	guide_id	gene	well
AAACCCACAACAAGAT-1	Nostim	perturbed	PLCG2-2	PLCG2	1
AAACCCACAACGGCTC-1	Nostim	perturbed	HELZ2-1	HELZ2	1
AAACCCACACAGAAGC-1	NaN	NaN	NaN	NaN	${\tt NaN}$
AAACCCACACCCTGTT-1	Nostim	perturbed	OTUD7B-1	OTUD7B	1
AAACCCACACTATGTG-1	Nostim	perturbed	CD247-1	CD247	1
•••	•••	•••			
TTTGTTGGTCCAGCCA-8	NaN	NaN	NaN	NaN	${\tt NaN}$
TTTGTTGGTCCCACGA-8	NaN	NaN	NaN	NaN	NaN

```
TTTGTTGGTGAGTGAC-8
                          Stim perturbed
                                               IL2RB-1
                                                          IL2RB
                                                                    8
TTTGTTGGTGCAAGAC-8
                            {\tt NaN}
                                                   {\tt NaN}
                                        NaN
                                                            {\tt NaN}
                                                                 {\tt NaN}
TTTGTTGTCTTCGCTG-8
                            NaN
                                        NaN
                                                   NaN
                                                            NaN NaN
[103805 rows x 5 columns]
{'NO-TARGET'}
['NO-TARGET-1',
 'NO-TARGET-2',
 'NO-TARGET-3',
 'NO-TARGET-4',
 'NO-TARGET-5',
 'NO-TARGET-6',
 'NO-TARGET-7',
 'NO-TARGET-8']
```

#### Data merged

```
View of AnnData object with n_obs × n_vars = 60657 × 73
   obs: 'condition', 'crispr', 'guide_id', 'gene', 'well'
   var: 'gene_ids', 'feature_types'
```

### Around 4000 cells have crispr NT

3883