Antigen identification and high-throughput interaction mapping by reprogramming viral entry

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Deciphering immune recognition is critical for understanding a broad range of diseases and for the development of effective vaccines and immunotherapies. Efforts to do so are limited by a lack of technologies capable of simultaneously capturing the complexity of adaptive immunoreceptor repertoires and the landscape of potential antigens. To address this, we present receptor-antigen pairing by targeted retroviruses, which combines viral pseudotyping and molecular engineering approaches to enable one-pot library-on-library interaction screens by displaying antigens on the surface of lentiviruses and encoding their identity in the viral genome. Antigen-specific viral infection of cell lines expressing human T or B cell receptors allows readout of both antigen and receptor identities via single-cell sequencing. The resulting system is modular, scalable and compatible with any cell type. These techniques provide a suite of tools for targeted viral entry, molecular engineering and interaction screens with broad potential applications.

The importance of T cell-mediated immunity has led to the development of a series of approaches dedicated to identifying antigens or pMHC–TCR pairs1. Conventional T cell assays, such as ELISPOT or intracellular cytokine staining2, provide direct readouts of T cell function. However, these approaches are highly multiplexed, they do not readily lend themselves to TCR sequencing and are limited in their ability to identify single reactive antigens. More recently, T-Scan and similar approaches have broadened the antigenic scope of functional assays to genome scale3,4, but cannot enable one-pot library-on-library interaction screens by displaying antigens on the surface of lentiviruses and encoding their identity in the viral genome. Antigen-specific viral infection of cell lines expressing human T or B cell receptors allows readout of both antigen and receptor identities via single-cell sequencing. The resulting system is modular, scalable and compatible with any cell type. These techniques provide a suite of tools for targeted viral entry, molecular engineering and interaction screens with broad potential applications.

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Recombinant protein-based screening can broaden either the number of T cell clones or the number of antigens that are feasible to screen. Baculoviral display libraries have enabled screening of ~105 pMHCs by panning of infected Sf9 cells with fluorescently labeled recombinant TCRs. Yeast display of pMHCs enables screening of ~104 unique pMHC antigens, but can only examine a limited number of recombinantly expressed TCRs at a time and can require significant optimization for each MHC allele. Conversely, recent advances in barcoded pMHC multimers enable screens on the order of 103 antigens in bulk or hundreds while maintaining the ability to identify interactions in large pools of multimers; and (3) the relatively small set of MHC molecules that have been recombinantly expressed successfully.

Identifying antigenic targets of B cells poses similar limitations. Recent approaches have shown that antigen–BCR pairs can be identified via the oligonucleotide tagging of recombinantly expressed proteins, which is inherently limited in scale. Thus, our understanding of antigen recognition by both arms of the adaptive immune system is currently constrained by significant hurdles to experimental scale.

Deciphering the full complexity of immune recognition requires the ability to screen for interaction pairs while incorporating diversity of both antigen receptors and their targets at the same time.

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If only the target antigens are known, it is difficult to understand the cellular factors regulating a successful response. On the other hand, if antigen receptors are characterized without knowledge of their specificity, it is difficult to understand which target antigens are important for preventing or eliminating disease. This represents a broader experimental challenge for interaction screens commonly known as ‘library-versus-library screening’. The most well-established approach to this problem is yeast two-hybrid\(^1\), in which intracellular protein pairs are used to drive expression of a reporter gene. Other previously established approaches include automated individual enzyme-linked immunosorbent assay (ELISA) screens of recombinantly expressed protein pairs\(^{22-24}\), as well as mass-spectrometry-based identification of interacting pairs\(^{25,26}\). There have been several recent efforts at designing new systems, including those based on yeast mating or spatial colocalization of DNA-barcoded molecules\(^{27,28}\). However, these approaches can be labor-intensive, are often not suited for complex, extracellular protein complexes such as immune receptors and may be inefficient at low (micromolar) affinities. Together, these limitations have thus far rendered such approaches unsuitable for applications such as identifying TCR–pMHC pairs.

To overcome the above limitations, we have developed a technique that combines lentiviral surface display\(^{29,30}\) with a versatile pseudotyping strategy and viral genome engineering to enable one-pot library-versus-library screening (Fig. 1). We demonstrate that our pseudotyping strategy, an engineered fusogen termed VSVGmut coexpressed with a targeting moiety, is general and versatile for both receptor and ligand usage. We leverage these abilities to present ‘receptor–antigen pairing by targeted retroviruses’ (RAPTR), which matches receptors with their cognate antigens based on specific infection of receptor-expressing cells by antigen-displaying viruses. Putative hits can be identified by bulk or single-cell sequencing, enabling screens of single or polyclonal receptors. We demonstrate the feasibility of this approach for both TCRs and BCRs, including a library-on-library screen consisting of 96 pMHC antigens and receptors enriched from a library of >450,000 TCRs (thousands of potential interactions).

**Results**

**VSVGmut-pseudotyped lentiviruses enable modular tropism.** Developing a scalable pipeline for antigen–receptor screening presents several requirements: (1) a straightforward ability to track both receptor and antigen sequences; (2) mammalian expression systems to maximize the ability to express complex, multimeric proteins; and (3) the ability to generate selection reagents without the need to recombinantly express and characterize each protein for every experiment. To address these needs, we turned to lentiviruses, which have decades of precedent for facile molecular manipulations. They can be created at large scale, are already known to infect T cells and, upon successful infection, leave a permanent record of infection due to their integration into the host genome.

Lentiviruses have been pseudotyped via a number of different strategies to enable their use as biotechnology tools and gene therapy vectors\(^32\). Due to its robustness and efficient infection of many cell types, vesicular stomatitis virus G protein (VSVG, referred to as VSVGwt in this manuscript) is the most common pseudotype for laboratory studies and cell manufacturing for clinical applications\(^{33}\). More recently, approaches to enable cell type-specific targeting, via coexpressing receptor-blinded versions of Sindbis virus\(^{34-36}\) or paramyxovirus envelope proteins\(^ {37-42}\) with targeting moieties have been described. While these approaches show promise, they have been reported to have strict limitations on targeting ligand- and receptor-binding topology due to their mechanism of entry.

As these factors constrain the generality of an interaction screening system, we developed an alternative strategy based upon VSVG. We used the recently described crystal structure of VSVG in complex with its native receptor\(^41\), the low-density lipoprotein receptor (LDLR), to engineer VSVGmut, which incorporates the K47Q and R354A mutations reported to ablate affinity for the LDLR family of receptors (Fig. 1a). To retarget VSVGmut-pseudotyped viruses, we coexpressed a variety of surface-bound molecules during viral production (Fig. 1b). To benchmark against established systems\(^{29}\), we used IL-13 as a viral targeting ligand. After testing several surface architectures (Extended Data Fig. 1), we found that while many distinct constructs conferred specific infection of IL-13Rα1-expressing cells, the dimerized, surface-tethered IL-13 yielded the most efficient infection while retaining excellent specificity (Fig. 1c). We also found that VSVGmut-pseudotyped viruses displaying an anti-CD19 single-chain antibody fragment (scFv) efficiently infected CD19\(^\text{+}\) B cell lines, but not CD19\(^\text{−}\) T cell lines (Fig. 1d).

Next, we sought to exploit the unique modularity of the VSVGmut system to develop synergistic targeting strategies. While display of the anti-CD3 Fab UCHT1 yielded only modest infection of Jurkat T cell lines, CD80 mediated robust infection (Extended Data Fig. 2a). Although the infection rate for the synergistic strategy was not better than for CD80 alone, we reasoned that the ability to incorporate multiple signals during infection could be useful for engineering primary cells by providing user-defined phenotypic inputs. Therefore, we used these viruses to infect primary CD8\(^\text{+}\) T cells and measured infection and CD25 upregulation as a marker of activation (Fig. 1e). We observed that the anti-CD3/CD80 combination viruses both infected and activated cells, but viruses displaying CD80 alone infected but did not activate the cells. We also observed that the anti-CD3/CD80 combination viruses induced robust T cell proliferation, while viruses displaying CD80 alone did not cause proliferation (Fig. 1f). This work presents a viral targeting system capable of simultaneously delivering multiple synergistic signals while infecting cells.

**Antigen-specific T cell targeting by engineered lentiviruses.** Following the observation that anti-CD3 viruses yielded infection of T cells via a component of the TCR complex, we next sought to determine whether we could achieve antigen-specific cell entry via the TCR itself. One potential challenge to repurposing the TCR–pMHC interaction for viral entry is its affinity, which is typically 1-50 μM\(^{44}\). We therefore generated cell lines expressing a range of previously characterized affinity variants for the 1G4 TCR\(^{45,46}\), which recognizes a peptide derived from the NY-ESO-1 cancer testis...
antigen (SLLMWITQV) presented in the context of HLA-A*02:01, with reported affinities ranging from picomolar to micromolar (Fig. 2a). To display these molecules on the virus surface, we expressed them as single-chain trimers47, which consist of covalently linked peptide, β-2-microglobulin and MHC. We found that pMHC-displaying lentiviruses are able to efficiently infect T cells in an antigen-dependent manner. We observed similar infection efficiency across the tested affinity range, with only a modest reduction for the lowest-affinity variant (Fig. 2b). To ensure that TCR-mediated infection was generalizable, we displayed several individual pMHCs as single-chain trimers alongside VSVGmut and used them to infect either Jurkat cells expressing its parental TCR (off-target) or on-target J76 cell lines expressing their cognate TCRs (Fig. 2c). We observed specific infection across three different TCR–pMHC pairs with minimal background infection. As VSVG enables cell entry via an endocytic route48 and because endocytosis is one of the earliest downstream events following T cell activation49–51, we next sought to determine whether viral infection was...
accompanied by TCR signaling. We measured CD69 expression following viral infection (Fig. 2d) and observed robust CD69 upregulation during TCR-mediated viral entry, but not during off-target combinations or VSVGwt infection (Extended Data Fig. 2c). Moreover, TCR-mediated entry was inhibited by dasatinib, a TCR signaling inhibitor, whereas TCR-independent infection via VSVGwt was unaffected (Extended Data Fig. 2b). Thus, we concluded that pMHC-targeted viruses integrate both binding and signaling as a means of infection. Notably, pMHC-based targeting was more efficient than infection by viruses displaying an anti-CD3 Fab. While it is unclear why this is the case, it is possible that the pMHC-based approach is more efficient at inducing TCR signaling when engaged as monomeric proteins or that the orientation of the pMHC is more favorable for viral entry than the CD3 epitope recognized by the anti-CD3 Fab.

We next sought to determine the limitations of our system by characterizing its sensitivity and specificity using the 1G4-NY-ESO-1 system. To determine the ability of our approach to discern on-target interactions in complex mixtures, we labeled 1G4-expressing cells with a cell tracking dye, mixed them at varying ratios with unlabeled Jurkat cells and infected the mixture with NY-ESO-1/HLA-A2-displaying viruses (Fig. 2e). We calculated a selectivity ratio of on-target infection relative to off-target infection by dividing the transduction rate of on-target cells by the
transduction rate of off-target cells and observed on-target selectivity greater than 200:1, down to target cell frequencies of $1 \times 10^{-5}$. Thus, we concluded that our system can detect on-target interactions in the micromolar affinity range, even in complex mixtures containing hundreds of thousands of non-target T cells.

A viral packaging system to maintain protein-barcode linkage. To fully enable interaction screening at library scale, we next turned to the challenge of producing lentiviral libraries. While lentiviral screens have been widely used for functional genomics, standard lentiviral packaging techniques pose inherent limitations that are particularly relevant to interaction screens. The key challenge stems from the fact that at least hundreds of plasmids containing library elements are mixed in each transfected packaging cell (Fig. 3a). In functional genomics, this leads to well-described intermolecular recombination of library elements$^{52,54}$, which is a significant source of noise in combinatorial experiments. To date, solutions to this problem have included plasmid dilution (accompanied by a 100× reduction in viral titer)$^{59}$ or simply restricting libraries to sizes suitable for arrayed screens$^{60-62}$. This problem is compounded in our approach, as multiple plasmids entering the same packaging cell would cause multiple different targeting molecules to be expressed on each virus surface, compromising the link between viral genotype and surface phenotype.

To obviate this issue, we exploited a detail of HIV-1 replication that results in the copying of sequences between the polyuridine tract and the 3′ long terminal repeat (LTR) to the 5′ end of the genome during reverse transcription and integration$^{63}$. There have been several recent reports exploiting this phenomenon to enable genomic screens via copying of CRISPR guide RNA cassettes$^{64,66}$. Here, we insert a promoter to enable repackaging of the viral genome following infection of a cell (Fig. 3b). Conventional third-generation lentiviral packaging approaches are inherently self-inactivating in part due to an inability to transcribe packageable viral genomes following infection; the LTRs, which are weak promoters, are truncated and there is no promoter upstream of the viral genome following successful integration. Our approach, which we term ‘lentiviruses activated by promoter shuffling’ (LeAPS), enables generation of lentiviruses from previously transduced cells upon the re-introduction of helper plasmids by placing a strong promoter 5′ of the integrated viral genome, allowing the integrated vector to produce additional viral genomes at a high copy number. As a result, we can use lentiviral transduction to provide a single library member per packaging cell without incorporating additional viral elements into the genome. We first validated that viruses produced with the LeAPS strategy yielded high-titer viruses ($>10^7$TU·ml$^{-1}$ unconcentrated), whereas conventional constructs (without a LeAPS promoter) yielded very little following transduction (Fig. 3c). By stably expressing pMHC cassettes in LeAPS-transduced packaging cells, we verified that viruses produced in this manner maintain antigen-specific targeting capability (Fig. 3d).

To develop a scalable library assembly strategy, we co-transduced 293T cells with two VSVGwt-pseudotyped viruses: one that drives expression of pMHC and barcode in a single sorting step. This yields a library packaging line that can be used to generate the same lentiviral library indefinitely via single transfections with only helper plasmids. By contrast, conventional approaches require arrayed transfection to produce each library member for each experiment$^{41}$, which poses a substantial impediment to experimental throughput.

To validate the utility of our approach, we employed a two-component system of the HIV SL9 antigen (SYLNTVATL) paired with a LeAPS–mCherry cassette and the NY-ESO-1 antigen paired with a LeAPS–green fluorescent protein (GFP) cassette. By mixing the packaging cells for each at different ratios, we assessed the feasibility of screening pMHC libraries at 100-member and 1,000-member scales. To assess the feasibility of a 1,000-member library, we mixed the SL9 and NY-ESO-1 LeAPS packaging cell lines at a ratio of 1:1,000 by cell number, produced the virus in pools via transfection with helper plasmids and used the resulting viral pools for target infections of TCR knock-in T cell lines (Extended Data Fig. 3). We then used flow cytometry to quantify signal to noise across different frequencies of on-target cells (Fig. 3f). Across both library sizes and extending from target cell frequencies of 10% down to 0.01%, we observed robust enrichment of on-target interactions, but minimal enrichment of off-target interactions (no enrichment of either component on Jurkat cells expressing an irrelevant TCR). These results gave us confidence that the screening step would be feasible at scales of at least 1,000 pMHC variants using the LeAPS library generation approach.

Receptor-antigen pairing by targeted retroviruses. With all the necessary tools for RAPTR validated, we assembled a library of 96 pMHC constructs consisting of known seroprevalent viral or vaccine antigens from the Immune Epitope Database (IEDB)$^{46}$ and further filtered for binding to HLA-A2 using NetMHC4.1 (ref. 44; Supplementary Tables 1 and 2). We co-transduced each into 293T cells alongside a LeAPS vector encoding a barcode, pooled the resulting cells proportionally and sorted for cells expressing both barcode (via mCherry) and pMHC (via GFP). We verified that antigen and barcode expression in our packaging cell library was stable after at least nine passages after sorting (Extended Data Fig. 4a). We next used the library to infect a cell line expressing the C7 TCR$^{44}$, which is known to recognize the cytomegalovirus (CMV) peptide NLVPNVAT (known as NLV). Without any cell-sorting step required, we sequenced genomic DNA from transduced cells to enumerate barcode frequencies (Fig. 4a). We observed a large enrichment of NLV barcode in transduced cells relative to the packaging cell line (Fig. 4b), with NLV representing 15% of all reads in transduced cells and no notable enrichment of other sequences. We further validated this approach in C7 replicates (Fig. 4c and Extended Data Fig. 4b) and in a separate screen of the JM22 TCR$^{44}$, which recognizes the GL9 peptide from influenza A (Fig. 4d and Extended Data Fig. 4c). In all cases, we observed robust enrichment of on-target, but not off-target, barcodes.

RAPTR can be adapted to B cell receptors. Many of the same constraints that limit TCR–pMHC interaction screens are also shared by efforts to identify B cell antigens at scale. Most approaches rely on barcode multimers for very few targets and the largest antigen library reported to date is nine antigens$^{63}$. As BCRs are also rapidly endocytosed upon antigen engagement$^{64,67}$, we reasoned that we could use

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**Fig. 3 | LeAPS-based lentiviral packaging enables scalable viral library construction.** a, Schematic overview of the limitation of conventional lentiviral packaging strategies for surface displayed lentiviral libraries. b, Schematic overview of the mechanism of promoter translocation used in the LeAPS system. c, Packaging cells transduced with viruses containing LeAPS-encoding genomes and then transfected with lentiviral helper plasmids (top) and transduced with viruses containing standard genomes (bottom). d, LeAPS-produced viruses maintain TCR specificity in two independent TCR–pMHC systems. e, Schematic overview of the implementation of LeAPS for viral library assembly. f, On-target enrichment of libraries using defined mixtures of A2-NY-ESO-1 and A2-SL9 packaging cells to represent 100- or 1,000-variant libraries.
Desired lentiviral library
Matched genotype and phenotype: virus surface characteristic also encoded in genome

Conventional lentiviral library
Genotype and phenotype not linked: genome does not represent the virus surface

100-member scale 1,000-member scale
Target cell frequency (%)

On-target enrichment ratio

0.1 1 0.1 0.01

Target cell frequency (%)

293T with known combinations

Co-transduce
Barcode + pMHC

Viral library with strict genotype–phenotype linkage for screening

Reusable packaging cell library maintained as cell line pool

VSVGwt lentivirus produced with pLeAPS barcode and pMHC expression

Barcodes

Conventional

LeAPS Pro

LeAPS Pro

LeAPS Pro

pLeAPS

pLeAPS

LeAPS Pro

Barcode 1

Barcode 2

pMHC 1

pMHC 2

... etc.

Reusability

Virus library with strict genotype–phenotype linkage for screening

1G4wt rep 1

1G4wt rep 2

868 rep 1

868 rep 2

Jurkat rep 1

Jurkat rep 2

Target cell frequency (%)
stabilized viral surface antigens or known immunogens as targeting ligands for viral infection. We thus generated a ‘hybrid’ pseudotype coexpressing either the surface-tethered receptor-binding domain (RBD) or the full-length, prefusion-stabilized spike (S2P) protein from SARS-CoV-2 (ref. 68) alongside VSVGmut (Fig. 5a).

We used each of these to achieve efficient, antigen-specific infection of a Ramos cell line expressing CR3022, a BCR clone that is cross-reactive for the RBDs of SARS-CoV-1 and SARS-CoV-2 (refs. 69, 70; Fig. 5b). Similarly, hybrid pseudotypes incorporating HIV env CD4 binding site (CD4bs) constructs from diverse clades71 each infected cell lines expressing the bNab VRC01 (ref. 72) and a known affinity-reducing mutation (D368R) dramatically reduced, but did not fully abrogate, infection73.

To determine the sensitivity for BCR-mediated cell entry, we mixed Ramos cells expressing either the CR3022 BCR (IgM+), or no BCR (IgM−) and tracked on-target cells via surface IgM expression (Fig. 5c and Extended Data Fig. 5). Using RBD-displaying viruses, we observed specific infection at target cell frequencies as rare as \(1 \times 10^{-5}\), but could not calculate a meaningful selectivity ratio due to a lack of off-target infection. Next, to assess the potential feasible library size, we mixed on-target RBD-displaying viruses encoding GFP with off-target CD4bs-displaying viruses encoding mCherry and used them to infect CR3022 BCR-expressing Ramos cells (Fig. 5d and Extended Data Fig. 5c). Keeping the total amount of virus constant, we were able to robustly detect on-target interactions with minimal off-target infection in mixtures as dilute as 1:1,000 on-target viruses. We observed similar results when displaying SARS-CoV-2 S2P instead of RBD (Extended Data Fig. 5). These results gave us confidence in the feasibility of extracting a meaningful signal from interaction screens using lentiviral infection via the BCR.

Next, we sought to establish the capabilities of RAPTR for BCR profiling in a true library setting. We assembled a library
Fig. 5 | RAPTR enables profiling of B cell reactivity via infection of BCR knock-in cells. a, Overview of BCR-based targeting via the VSVGmut hybrid pseudotype in which a viral protein is coexpressed with VSVGmut. b, Targeting of Ramos cells expressing the SARS-CoV-2 specific CR3022 BCR via VSVGmut lentiviruses displaying either SARS-CoV-2 spike (2P) or surface-bound SARS-CoV-2 S protein RBD or VRC01 BCR via env constructs representing multiple clades of HIV; representative data are from two independent experiments. c, SARS-CoV-2 RBD-displaying viral infection of Ramos cells (on-target, IgM+) mixed into IgM− Ramos cells (off-target, IgM−) at target cell frequencies indicated in red, demonstrating selectivity in a mixed population of cells. d, Infection of CR3022 BCR-expressing Ramos cells with mixtures of SARS-CoV-2 RBD (on-target, expressing mCherry) and HIV env (off-target, expressing GFP) viruses; variant library size indicates the ratio of off-target to on-target virus present. Infection of VRC01-expressing Ramos cells with HIV env hybrid pseudotyped viruses (right) demonstrates that the viral particles remain functional. e, Enrichment of antigen barcodes relative the viral packaging cell line following viral glycoprotein library infection of Ramos cells expressing BCRs for VRC01, mature CR3022 or the germline-reverted version CR3022, demonstrating selectivity even for antibody sequences before affinity maturation.
composed of 21 prefusion-stabilized SARS-CoV-2 spike variants and 22 additional viral surface glycoproteins or immunogens from diverse sources as controls to enable proof-of-concept experiments (Supplementary Tables 3 and 4). When we used this library to infect cells expressing the VRC01 BCR, we observed clear enrichment of only HIV-1 SOSIP (Supplementary Fig. 2c), highlighting the potential for direct receptor de-orphanization. When we infected cells expressing the CR3022 BCR, we observed clear enrichment of SARS-CoV spikes, including multiple SARS-CoV-2 variants. Notably, a version of CR3022 that excluded the mutations acquired via somatic hypermutation yielded enrichment of only SARS-CoV-1 spike and the closely related WIV-1 spike, highlighting the ability of RAPTR to distinguish BCR cross-reactivity even for naive BCRs that have yet to undergo affinity maturation.

**RAPTR enables library-on-library screens.** To fully realize the potential of RAPTR for antigen identification, we sought to apply it to library-versus-library screening by using our 96-member virus library and a polyclonal TCR-expressing T cell line. By mixed infection followed by mCherry+ sorting and single-cell RNA sequencing, we identified enriched TCR clones expressing antigens from both EBV and influenza in two independent infections. The top two clones identified the EBV antigen pMHC and influenza antigen pMHC, with clonal frequencies of 73% and 3% respectively in two independent infections. The antigen barcodes in the top two clones were enriched relative to controls, confirming that RAPTR enables scalable antigen discovery.
pMHC viral library to pan a previously reported library of >450,000 TCRs\(^75\). To simplify the first attempt, we pre-enriched for potentially reactive cells using tetramers for CMV, Epstein–Barr virus (EBV) and influenza (CEF) antigens presented by HLA-A2, resulting in a polyclonal pool of TCRs with greater prevalence of these specificities, similar to the routine process of pre-expanding cells (Fig. 6a). Following transduction and FACS to isolate transduced cells, we used bulk sequencing to identify which antigens were enriched in aggregate. In line with tetramer staining of the untransduced cells, we observed strong enrichment of influenza GL9 and more moderate enrichment of EBV GLCTLVAML (GLC) (Fig. 6b and Extended Data Fig. 6).

With this validation in hand, we performed single-cell RNA sequencing using the 10X Genomics Chromium 5\(^+\) chemistry with V(D)J enrichment. We included a targeted primer at the reverse transcription step and a custom amplification protocol to enable recovery of our pMHC barcodes, akin to ECCITEseq and other similar methods\(^79\). In this experiment, we recovered 1,438 cells, with TCR clonotypes dominated by a few clones (Fig. 6c,e). We grouped cells of common clonotype together and analyzed pMHC barcode expression to determine enrichment relative to packaging library frequency. The most prevalent TCR clone, GL9.1, demonstrated strong enrichment of the GL9 pMHC signal (Fig. 6d). Upon literature search, we noted that GL9.1 represents a well-characterized public clonotype known to recognize the GL9 peptide presented by HLA-A2 (ref. 75).

The next most abundant clone, GLC.1, was identified in a previous screen of this same TCR library by recombinant HLA-A2 GLC dextramer\(^75\). Our screen also identified GLC as the target pMHC (Fig. 6d). Notably, this was the only clone previously identified by GLC dextramer panning that was validated by monoclonal binding and functional tests. Our approach did not identify any of the previously reported false positives as hits, providing further evidence that RAPTR is an efficient means of identifying T cell antigens that integrates both binding and signaling. Finally, while T cells that recognize NLV are common in HLA-A2\(^+\) CMV-seropositive individuals\(^75\), we did not observe any NLV-reactive cells (Fig. 6b). However, this result is consistent with previously reported screening of the same library via orthogonal methods\(^75\) and our tetramer staining data (Extended Data Fig. 6a).

**Discussion**

RAPTR is a high-throughput platform for directly linking immune receptors with their cognate antigens. It is based on the combination of several conceptual and technological advances that enable (1) versatile and efficient targeted viral entry via the VSVGmut pseudotyping system; (2) a method for scalable, reproducible lentiviral library packaging (LeAPs); and (3) the use of viral entry as a means of screening for interactions. We demonstrated a system for efficient, antigen-specific infection via both the TCR and the BCR that integrates both binding and target cell activation. We then exploited this mode of infection to match receptors with target antigens in complex mixtures and in a library-versus-library format.

We demonstrated that the VSVGmut pseudotyping system is efficient, specific and uniquely modular by targeting a variety of cell surface proteins, including cytokine receptors, co-stimulatory molecules, lineage markers and both TCRs and BCRs. We report here as many different targeting approaches for VSVGmut as have been reported for any other single pseudotyping system\(^74\), including antigen-specific infection of lymphocytes and a targeting approach that can simultaneously deliver multiple signals. In addition, the VSVGmut system produces viruses at higher average titers than other systems while maintaining a high degree of specificity. Whereas most pseudotyping systems have been limited to the use of small, stable, high-affinity targeting ligands (such as scFvs or DARPin), VSVGmut is versatile in ligand usage due to the modular design it permits. We report targeting even at micromolar affinities, whereas the efficiency of infection by paramyxoviral pseudotypes was reported to dramatically decrease in the low nanomolar range\(^60\). We observed efficient pMHC-based infection for both antiviral TCRs (C7, JM22 and the TCRs recovered shown in Fig. 6) and Ig4, which recognizes a self-antigen, suggesting that the full range of possible CD8\(^+\) antigens should be able to be profiled.

**Applications of RAPTR.** We have demonstrated the utility of RAPTR for identifying antigens recognized by TCRs and BCRs. For T cell reactivity, our approach could be directly applied to rapidly deorphanize TCRs from existing single-cell studies, which routinely pair TCR sequences with gene expression for cancer, autoimmunity or infectious disease. Existing techniques, including yeast display, SABRs and T-Scan, require individual screens to pair each TCR with its cognate antigen, including at least one cell-sorting step per TCR. In contrast, RAPTR can be performed on pools of TCRs in a single step, with only one cell-sorting step required (and no sorting required for single-TCR experiments). While barcode pMHC multimers can be built into single-cell workflows, large antigen pools are inherently difficult to produce and validate and require an equivalent effort for each new batch. These factors have largely limited their use to only a few specialized labs thus far. RAPTR library production is straightforward and far more scalable in comparison, requiring only a simple transfection per batch following initial library assembly. In principle, these reagents could be made and distributed at a large scale, either via sharing the packaging cell line or by leveraging current industrial infrastructure for production of lentiviruses.

For B cells, we envision applying pools of protein variants, such as the one presented here, to large libraries of BCRs. Such an approach would greatly accelerate isolation and optimization of cross-reactive monoclonal antibodies to a wide range of target classes. Current techniques require the recombinant expression of individual monoclonal antibodies followed by ELISA for each target of interest, which represents a major bottleneck to experimental scale and scope\(^71\). Using RAPTR with BCR knock-in cells, which could themselves be prepared as pooled lentiviral libraries\(^74\), could alleviate this bottleneck. For example, even scales smaller than the one we presented would be sufficient to represent every subtype of influenza hemagglutinin and neuraminidase, multiple representatives of each of the major clades of HIV-1 and more. Larger libraries on the scale of thousands of variants, whose feasibility we demonstrated, could enable detailed antigenic site or epitope mapping for many monoclonal antibodies at once, a process currently limited to structural studies that are difficult to scale.

Finally, the demonstrated modularity of the system opens the door to many potential applications. As our approach only requires HEK cell surface expression of the targeting molecule, we hypothesize that RAPTR can be readily adapted to other MHC class I and class II alleles, as well as non-human systems for vaccine and immunotherapy development. This compares favorably to recombinant MHC expression or yeast display, each of which can require allele-specific optimizations and/or mutations to ensure proper folding\(^73,74,75\). However, our system is not restricted to immunological applications. In principle, it is applicable to any receptor–ligand system in which one binding partner can be displayed on a lentivirus and the other can be expressed as an endocytic receptor on cells. This may enable large-scale interactome and coevolution studies that are not readily achievable by existing techniques. Furthermore, the VSVGmut pseudotyping system may be useful as a new platform for the development of efficient, specific gene therapies.

**Limitations.** As with any technology, RAPTR has several limitations. Our proof-of-concept studies were performed on ~100-antigen scale and while our control experiments indicate that
this screening capability could be readily increased to thousands, these scales will still require preselection of antigen targets for analysis. Further streamlining library assembly and increasing throughput efficiency will be focuses for enabling larger-scale assays. At present, the LEAPS system involves one-time arrayed transfections and transductions on scales and complexity routinely performed for genomics screens55–57, after which lentiviral library generation can be performed in a straightforward manner by standard transfections of the pooled packaging cell library. As a result, library generation is currently the largest bottleneck to throughput. Due to the one-time nature of the bottleneck, we believe that automation will readily enable library sizes of up to 10^4, but adapting our approach for pooled library generation could aid in further scaling.

In addition, the single-cell sequencing step must be performed in sufficient depth to facilitate acceptable signal-to-noise determinations. While we chose to use a commercially available platform to ensure broad applicability, recent and future advances in the scale of single-cell analysis can improve the utility of RAPTR regardless of platform. Like other genetically encoded techniques, RAPTR also requires previous selection of MHC haplotype(s) to generate libraries. The antigen identification studies we presented were on cell lines with receptors knocked in, rather than primary cells. While the decreasing cost and turnaround time of gene synthesis will make such resources increasingly available, future studies can apply RAPTR directly to primary cells. Moreover, immortalized TCR libraries are becoming increasingly common as library generation methods continue to improve and can serve as renewable resources to conduct deep profiling of low-abundance or irreplaceable samples such as patient-derived tumor-infiltrating lymphocytes. Finally, for each of these cases, our data indicate that the ability to enrich for antigen-specific cells at library scale will be beneficial for efficient deorphanization, to ensure that target cells are present at sufficient frequencies for infection and analysis. Existing techniques, including stimulation via peptide pools, should readily achieve this but may require some optimization.

RAPTR is a versatile platform for high-throughput interaction screens that incorporates diversity of both receptors and ligands in a single assay. The resulting tools will enable detailed studies of antigen recognition for understanding and engineering the adaptive immune response, as well as broader interactome studies.

Online content
Any methods, additional references, Nature Research reporting summaries, source data, extended data, supplementary information, acknowledgements, peer review information; details of author contributions and competing interests; and statements of data and code availability are available at https://doi.org/10.1038/s41592-022-01436-z.

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**Methods**

**Ethics statement.** This work was reviewed and approved by the MIT Institutional Review Board (protocol no. 1801190804).

**Media and cells.** HEK293T cells (ATCC CRL-11268) were cultured in DMEM (ATCC) supplemented with 10% fetal bovine serum (FBS; Atlanta Biologics) and penicillin-streptomycin (pen/strep; Gibcco). Jurkat (ATCC TIB-152), J76 cells and Ramos cells (ATCC CRL-1596) were cultured in RPMI-1640 (ATCC) supplemented with 10% FBS and pen/strep. J76 cells were transduced with a gift from M. Heemskerk and M. Davis.

CSS-930 TCR library cells were a gift from D. Johnson and were cultured in RPMI-1640 (Thermo Fisher) supplemented with 10% FBS, pen/strep, 1x non-essential amino acids (Thermo Fisher), 2 mM Glutamax (Thermo Fisher) and 1 mM sodium pyruvate (Thermo Fisher).

**Plasmid construction.** The plasmid pSHIV-EGFP was a gift from R. Welm and Z. Werb (Addgene plasmid 21373) and pMD2.G and psPAX2 were gifts from D. Trono (Addgene plasmid 12259 and 12260). pLentiCRISPR v2 was a gift from E. Zhang (Addgene plasmid 52961). Human CD8+ T cells were isolated using EasySep human CD8+ enrichment kit (StemCell Technologies) as per manufacturer instructions. Primary CD8+ T cells were isolated using EasySep Human CD8+ T Cell Enrichment kit (StemCell Technologies) and cultured in RPMI-1640 (Thermo Fisher) supplemented with 10% FBS, 100 U ml⁻¹ pen/strep (Corning) and 100 μM β-mercaptoethanol (Sigma-Aldrich). After 24 h, an additional 100 μM β-mercaptoethanol was added. Cells were analyzed by flow cytometry 48–72 h after infection. TCR KO J76 cells were transduced as described above and sorted based on TCR expression to establish monoclonal cell lines. Ramos BCR cells were established according to the published protocol.

**Lentiﬁc infections in mixed cell populations.** Cells were labeled with CellTrace dyes (Thermo Fisher) according to the manufacturer’s instructions, counted and mixed at the indicated ratios. After labeling, viral infections were carried out as indicated above. For 48 h, cells were washed in FACS buffer and analyzed via flow cytometry to examine infection in CellTrace− versus CellTrace+ cells. After flow cytometry, the selectivity was calculated as follows:

\[
\text{Selectivity} = \frac{\% \text{ target cells transduced}}{\% \text{ off-target cells transduced}} \times (\text{frequency of target cells})
\]

**Generation of LeAPS libraries.** To generate LeAPS production cell lines, 293T cells were transduced as outlined above while being seeded at 20% confluency on either six-well or 96-well plates (Thermo Fisher). For six-well plates, 500 μl each of unconcentrated, VSVGwt-pseudotyped LeAPS barcode and ligand expression viruses were used for infection. For 96-well plates, 50 μl of each virus was used.

For libraries, cells were transduced in duplicate. One of the duplicates was assessed by flow cytometry to determine the proportion of cells transduced with both ligand (GFP+) and LeAPS barcode (mCherry+). Cells from the remaining duplicate were then pooled, with each library member normalized to include an equivalent number of barcode-expressing (GFP+) and barcode-encoding (mCherry+) cells for each library member. This pool of cells was then sorted based on GFP and mCherry expression to make library packaging cell pools. To produce lentiviruses, library packaging cells were transfected as described above, but using only psPAX2.1 and pMD2-VSVGwt. For library creation, both transfer plasmids and the pMD2 targeting plasmid were omitted, as they were replaced by the LeAPS packaging line. After 48 h, virus was then collected and concentrated as described above.

**Library screening of monoclonal TCR lines.** Concentrated pMHC library virus (10 μl) was used to transduce one million TCR-expressing J76 cells as outlined above. After infection was confirmed by flow cytometry as described above, genomic DNA was isolated using the PureLink Genomic DNA kit (Thermo Fisher). Barcode inserts were then amplified via 25 cycles of PCR and submitted for Amplicon-EZ analysis by Genewiz. Enrichment was calculated for each barcode as the fraction of total barcode-containing reads divided by the barcode frequency in the packaging cells.

**Library screening of monoclonal Ramos BCR lines.** Unconcentrated viral antigen library (500 μl) was used to transduce one million BCR-expressing Ramos cells as outlined above. After infection was confirmed by flow cytometry, genomic DNA was isolated using the PureLink Genomic DNA kit. Barcode inserts were then amplified via 25 cycles of PCR and submitted for Amplicon-EZ analysis by Genewiz or analyzed via MiSeq v3 150 × 150-nt PE Nano kit (MIT BioMicro Center). Enrichment was calculated for each barcode as the fraction of total barcode-containing reads divided by the barcode frequency in the packaging cells.

**Tetramer enrichment of TCR libraries.** To enrich a polyclonal pool of cells from the CSS-930 TCR library for known specificities, 50 million library cells were co-stained with an anti-TRBC antibody (clone IP26) and a pool of HLA-A2 tetramers (National Institutes of Health (NIH) Tetramer Core) presenting the following peptides: NLVPVMATV (NLV), GLIGGFVFTL (GL9) and GLC, all at 1:2000 dilution. Sorting for double-positive cells was performed on a Sony MA900 FACS.
mCherry barcodes were enriched via separate PCRs and sequenced on an Illumina MiSeq (150 x 150-nt paired end reads). CellRanger V(D) was used to assign TCR done identities for each cell. Cell barcodes were used to match TCRs with associated pMHC barcodes counts.

**Antibodies used in flow cytometry.** All antibodies were used at a 1:50 dilution from the stock concentration. Gating strategies shown in Extended Data Fig. 7.

**Software.** Graphs were generated using GraphPad Prism (v.8). Flow cytometry data were analyzed by FlowJo (10.8.0).

**Statistical analysis.** Statistical analyses (calculation of s.d.) were performed using the Prism 8 (GraphPad) software. Sample sizes were not predetermined using statistical methods.

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

**Data availability**
The next generation sequencing datasets generated during and/or analyzed during the current study are available in the National Center for Biotechnology Information Sequence Read Archive (accession no. PRJNA796957). The UniProt database was used to design constructs in this study (https://www.uniprot.org) for determining the peptide sequence of human IL-15 (accession no. P35225), human PDGF (accession no. P09619) and human CD80 (accession no. P33681). All data generated or analyzed during this study are included in this published article, the Sequence Read Archive and its supplementary information files. Source data are provided with this paper.

**Code availability**
A sample R workspace for extracting barcode counts based on CITEseq™ is available in the Birnbaum laboratory code repository (https://github.com/birnbaumlab/Dobson-et-al-2022).

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**Author contributions**
C.S.D. and M.E.B. conceived the RAPTR system. C.S.D., A.R., S.G., B.E.S., E.J.K., J.D., L.R. and V.O. conducted all experiments. C.S.D., A.R., S.G., B.E.S., E.J.K. and J.D. analyzed data. D.L., M.D., S.K.D. and M.E.B. supervised research. C.S.D. and M.E.B. wrote the manuscript. All authors edited the manuscript.

**Competing interests**
The lentiviral targeting approach in this manuscript is the subject of US patent applications with M.E.B., S.G. and C.S.D. as co-inventors. M.E.B. is a founder, consultant and equity holder of Viralogic Therapeutics and Abata Therapeutics, is an equity holder in iST Biosciences, has received consulting fees from Repertoire Immune Medicines and has received research funding from Pfizer that partially funded this work. S.K.D. received unrelated research funding from Novartis Pharmaceuticals, Eli Lilly and Company and Bristol-Myers Squibb and is a founder, science advisory board member and equity holder in Kojin. M.D. is a science advisory board member for Neoleukin, has received research funding from Novartis and Eli Lilly, has received consulting fees from Tillotts Pharma, ORIC Pharmaceuticals, Partner Therapeutics, SQZ Biotech, AzurRx, Mallinckrodt Pharmaceuticals and Moderna. The remaining authors declare no competing interests.

**Additional information**
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Extended Data Fig. 1 | IL-13 Surface Architecture Development. a, Schematic diagram of the different targeting ligand architectures used. b, Summary table with details of each individual construct and titer achieved. c, Flow cytometry plots showing infection of IL-13Rα1-expressing Jurkat cells (top, on-target) and parental Jurkat cells which don’t express the receptor (bottom, off-target).
Extended Data Fig. 2 | See next page for caption.
Extended Data Fig. 2 | Viral entry via the TCR complex and costimulatory receptors, and inhibition by dasatinib. a, 1x10⁵ Jurkat cells were infected with the indicated amount of VSVGmut-pseudotyped virus displaying either anti-CD3 Fab, CD80, or a combination of the two; data shown as mean + SD of three biological replicates. b, 5x10⁴ J76 cells expressing the 868 TCR were pre-incubated with the indicated concentrations of dasatinib for 2 hours to inhibit TCR signaling prior to virus addition (red dots indicate no dasatinib condition). Next, 5 µL of virus displaying either VSVGmut and HLA-A2-SL9 (on-target), VSVGmut and HLA-A2-NY-ESO-1 (off-target), or VSVGwt were added. Infection was measured 48 h post-infection. Results represent the mean and standard deviation of 3 biological replicates. c, The indicated amount of concentrated virus was added to each TCR-expressing J76 cell (100,000 cells/well in a 96-well plate). After 24 h, transduction was measured by fluorescent reporter (left) and activation was measured by CD69 upregulation (right) relative to unstimulated cells. d, 500,000 1G4hi-expressing J.RT3 cells or parental Jurkat cells were infected with 500 µL of unconcentrated VSVGmut-pseudotyped virus displaying HLA-A2 presenting either NYESO-1 or an irrelevant hepatitis B peptide (SLYAVSPSV), or VSVGwt-pseudotyped virus.
Extended Data Fig. 3 | Schematic diagram of LeAPS proof of concept experiments. Packaging cells transduced with (i) SL9 presented on HLA-A2 and LeAPS-mCherry and (ii) NY-ESO-1 presented on HLA-A2 and LeAPS-GFP were mixed at either 1:100 or 1:1,000 for each to represent libraries containing 100 constructs and 1,000 constructs, respectively. These mixtures were then used as packaging lines, transfected with helper plasmids to produce viruses, and used to transduce mixtures of cell tracking dye-labeled J76 cells expressing either 868 TCR or 1G4wt TCR at defined ratios.
Extended Data Fig. 4 | Quality control of LeAPS packaging line and reproducibility of infection. A, LeAPS-barcode expression (mCherry) and pMHC expression (GFP) of the RAPTR 96-member packaging line after 9− passages post-sorting. B, Correlation between infection replicates of the C7 TCR cell line. C, Correlation between infection replicates of the JM22 TCR cell line.
Extended Data Fig. 5 | Signal to noise in virus mixtures containing SARS-CoV-2 full spike and HIV env. a, Infection of CR3022 BCR-expressing Ramos cells with mixtures of SARS-CoV-2 prefusion-stabilized spike (on-target, expressing mCherry) and HIV env (off-target, expressing GFP) viruses; variant library size indicates the ratio of off-target to on-target virus present. Infection of VRC01-expressing Ramos cells with HIV env hybrid pseudotyped viruses (right) demonstrates that the viral particles remain functional. b, Quantification of selectivity ratios for the SARS-CoV-2 RBD infections of CR3022 Ramos presented in Fig. 5c. Selectivity ratio is calculated as the proportion of on-target cells infected divided by the proportion of off-target cells infected times the frequency of on-target cells (see Methods). c, Quantification of on-target enrichment for mixtures of HIV env and either SARS-CoV-2 RBD (presented in Fig. 5d) or full spike (presented in Supplementary Fig. 5a). Enrichment is calculated by fraction of cells expressing on-target signal divided by fraction of cells expressing off-target signal times on-target virus frequency.
Extended Data Fig. 6 | Quality control and reproducibility for the library vs library experiment. 

a) Tetramer staining for each of the pooled specificities used for enrichment. 

b) Correlation between infection replicates in bulk sequencing data for the library vs library experiment.
Extended Data Fig. 7 | Gating strategies used throughout manuscript. Flow cytometry gating strategies for experiments described.
Reporting Summary

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Statistics

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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  - [x] A full description of the statistical parameters including central tendency (e.g. means) or other basic estimates (e.g. regression coefficient) AND variation (e.g. standard deviation) or associated estimates of uncertainty (e.g. confidence intervals)
  - [x] For null hypothesis testing, the test statistic (e.g. F, t, r) with confidence intervals, effect sizes, degrees of freedom and P value noted
    - Give P values as exact values whenever suitable.
  - [x] For Bayesian analysis, information on the choice of priors and Markov chain Monte Carlo settings
  - [x] For hierarchical and complex designs, identification of the appropriate level for tests and full reporting of outcomes
  - [x] 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 | MiSeq 3.1, CellRanger VDJ (3.0) |
|-----------------|----------------------------------|
| Data analysis   | Flowlo (10.8.0), Graphpad Prism (v8). Custom code can be found at https://github.com/birnbaumlab |

For manuscripts utilizing custom algorithms or software that are central to the research but not yet described in published literature, software must be made available to editors and reviewers. We strongly encourage code deposition in a community repository (e.g. GitHub). See the Nature Portfolio guidelines for submitting code & software for further information.

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

- Accession codes, unique identifiers, or web links for publicly available datasets
- A description of any restrictions on data availability
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The next-generation sequencing datasets generated during and/or analyzed during the current study are available in the NCBI sequence read archive (Accession Number PRJNA796957). All data generated or analyzed during this study are included in this published article, the sequence read archive, and its supplementary information files.
Field-specific reporting

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For a reference copy of the document with all sections, see nature.com/documents/rr-reporting-summary-flat.pdf

Life sciences study design

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

| Sample size | No sample-size calculation was performed. For flow cytometry and sequencing experiments, the minimum number of cells analyzed was determined as ten times the inverse of the frequency of the rarest population [e.g. 10,000 cells for a population expected at 0.1%]. |
| Data exclusions | No data were excluded |
| Replication | All attempts at replication were successful with replicates as marked (typically 2-3x) |
| Randomization | This was not relevant to this study as no clinical or animal studies were performed |
| Blinding | This was not relevant to this study as no clinical or animal studies were performed |

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

Antibodies

Antibodies used: All antibodies were obtained from BioLegend unless indicated otherwise: anti-human CD69 (clone FN50, cat. 310910, lot B258744); anti-human TCRbeta (clone JP26, cat. 306708, lot B258839); anti-human EGFR (clone AY13, cat. 352911, lot B321041); anti-human HLA-A2 (clone B87.2, cat. 343308, lot B277594); anti-human IgM (clone MHM-88, cat. 314510, lot B294323)

Validation: Antibodies were validated by commercial vendors using flow cytometry of antibody-stained cells and this information is available on their websites.

Eukaryotic cell lines

Policy information about cell lines:

Cell line source(s): Wild type Jurkat cells [clone: E6-1, from ATCC TIB-152]
TCRbeta deficient Jurkat cells [clone: JRT3-T3.5, from ATCC TIB-153]
HEK293T/17 cells (ATCC CRL-11268)
Ramos Cells (ATCC CRL-1596)
J76 Cells [a gift from Mark Davis]

Authentication: J76 cells were validated by staining for surface TCR beta chain expression. Cell lines have also been authenticated by morphology checking and through use in other experiments.

Mycoplasma contamination: Tested negative for mycoplasma contamination.
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          | Tissue culture samples were resuspended to single cells for staining. |
|----------------------------|---------------------------------------------------------------------|
| Instrument                 | Analysis: data were collected on either the Accuri C6 or the BD Cytoflex S. Sorting: Cells were sorted using the Sony MA900 |
| Software                   | Manual gating was used in Flowjo (10.8.0) for analysis               |
| Cell population abundance  | Sorted cells were directly analyzed by single-cell sequencing. Any cells maintained in culture were subsequently verified for >95% purity by flow cytometry analysis for the marker(s) used to sort originally. |
| Gating strategy            | The gating strategies used are provided in Supplementary Figure 7 and all applicable figures. In brief, gating strategy was: FSC-A vs SSC-A to capture the cell population of interest --> FSC-A vs FSC-H to identify single cells --> Marker(s) of interest |

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