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High-throughput and single-cell T cell receptor sequencing technologies

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Abstract

T cells express T cell receptors (TCRs) composed of somatically recombined TCRa and TCR β chains, which mediate recognition of major histocompatibility complex (MHC)–antigen complexes and drive the antigen-specific adaptive immune response to pathogens and cancer. The TCR repertoire in each individual is highly diverse, which allows for recognition of a wide array of foreign antigens, but also presents a challenge in analyzing this response using conventional methods. Recent studies have developed high-throughput sequencing technologies to identify TCR sequences, analyze their antigen specificities using experimental and computational tools, and pair TCRs with transcriptional and epigenetic cell state phenotypes in single cells. In this Review, we highlight these technological advances and describe how they have been applied to discover fundamental insights into T cell-mediated immunity.

T cells are critical effector cells of the adaptive immune system, providing protection against pathogens and cancer while maintaining immune self-tolerance. This broad range of functions is enabled by the diversity of T cell phenotypes and antigen specificities that exists in an individual. T cells can adopt a wide spectrum of phenotypes, ranging from highly cytotoxic effector T cells to anti-inflammatory regulatory T cells. This phenotypic specification is largely driven by the specific TCR expressed on each T cell's surface, which mediates antigen recognition and T cell responses and presents another element of diversity within T cell populations. The TCR is composed of two chains, TCR α and TCR β , which are produced through combinatorial somatic rearrangement of multiple variable (V), diversity (D) (for the β -chain only), joining (J) and constant (C) gene segments¹. This recombination process, in conjunction with nontemplated addition or deletion of nucleotides between spliced gene segments, generates a TCR repertoire with a theoretical diversity of approximately 2×10^{19} unique TCRa β pairs¹⁻³, although the diversity in an individual is likely lower owing to selection pressures and the numbers of T cells present^{4–6}. This vast diversity, particularly in the hypervariable complementary-determining region 3 (CDR3) of the TCR^{1,7}, enables the recognition of an extensive array of foreign and self-peptides presented on MHC (pMHC) complexes.

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There are several aspects of the T cell response that can be best understood through analysis of the TCR repertoire. Because of the low probability of somatic recombination producing an exact V(D) J rearrangement twice in an individual³, the TCR sequence can be used as a unique identifier of T cell clones. This feature is a useful proxy for measuring (1) antigen-driven clonal expansion of T cells as an indicator of antigen specificity and response (Fig. 1a,d) and (2) longitudinal clonal dynamics and heterogeneity of the T cell response (Fig. 1b,c). Moreover, joint assessment of the TCR and cellular phenotypes in single cells can provide information on T cell differentiation pathways and the role of TCR sequence and specificity in T cell selection, activation and phenotypic specification (Fig. 1b). This information is useful not only in understanding the etiology and pathology of immune-mediated diseases, but also in designing therapeutic strategies. For example, tracking TCR diversity in patients across time points can reveal patterns of T cell clonal dynamics that correlate with treatment response or other clinically relevant features, and pairing clonal information with cellular phenotype can identify disease-relevant cell types^{8–12}.

In these contexts, meaningful biological interpretation of data is enabled by the use of sensitive and accurate methods to profile the TCR repertoire. Early TCR analysis methods (reviewed previously^{13–15}) revealed fundamental features of the TCR repertoire but were often limited by their ability to capture only partial sequence information. For example, CDR3 spectratyping, which measures the distribution of CDR3 sequence lengths by the separation of PCR amplicons in electrophoresis^{16,17}, was used to provide the first estimates of human and mouse TCR diversity^{4,18}. However, this approach targets only specific V–J rearrangements and does not directly read out sequence information, which is necessary for determining T cell clonality and specificity. A second approach relied on flow cytometry using anti-TRBV monoclonal antibodies, which provided insights into repertoire diversity based on TCR β V gene usage but did not capture sequence information or junctional diversity in the CDR3 (refs. ^{19,20}).

The advent of genomic sequencing enabled early TCR sequencing methods that could capture TCRs at nucleotide resolution by molecular cloning and Sanger sequencing, delivering more precise measurements of CDR3 diversity²¹. These methods were used in pioneering studies of the TCR repertoire in a wide range of contexts, including in immune-mediated diseases, infections and cancer, leading, for example, to the identification of disease-specific expansion of certain TCR β genes²¹. Importantly, these methods also enabled the first estimations of TCR α and TCR β sequence diversity and provided evidence that the naive T cell repertoire is highly diverse while the memory T cell compartment contributes only a small proportion of the total TCR repertoire diversity in an individual^{4,18}. However, these techniques are generally low in throughput (10–1,000 TCRs per assay), which precludes extensive sampling of the TCR diversity present in a given sample or population.

As next-generation sequencing technologies have become commonplace over the past two decades, the sensitivity of TCR detection and ease of applying TCR discovery tools have greatly improved, allowing for the capture of 10^3-10^6 TCRs per assay and analysis of TCR sequences in their cellular contexts. In this Review, we discuss recent advances in high-throughput methods for interrogating the TCR repertoire at bulk and single-cell resolution,

as well as techniques for pairing TCR sequences with antigen specificity, cell state and phenotype. In particular, we highlight how these approaches have enabled deeper insights into T cell clonal dynamics, differentiation and response trajectories, and function in health and disease.

Molecular strategies for TCR sequencing

Profiling the TCR repertoire using high-throughput sequencing technologies has enabled the analysis of millions of T cells in a single experiment³ (Fig. 1a). These methods generally use one of two amplification strategies: (1) multiplex PCR^{22–24} or (2) rapid amplification of 5' complementary ends (5' RACE)^{25–29}.

Multiplex PCR.

Because of the large sequence diversity of TCR V genes, one pair of primers is not sufficient to capture all TCR transcripts. To address this challenge, the first approach developed for targeted sequencing of the TCR used multiplex PCR reactions^{22–24}. Multiplex PCR methods use a set of forward primers complementary to all known V genes and a set of reverse primers for either J or C regions, depending on whether the starting material is genomic DNA (gDNA) or complementary DNA (cDNA), respectively. In this primer amplification strategy, gDNA or RNA is isolated from T cells and subjected to multiple rounds of PCR using these primer sets, which also contain universal primer sequences to enable subsequent sequencing on high-throughput platforms. The use of multiplex PCR methods has led to many early insights into human TCR diversity. For instance, Robins et al. performed deep sequencing of CDR3B and demonstrated that the TCR repertoire at steady state exhibits a strong bias toward certain V-J rearrangements²³. They also observed a significantly greater degree of naive CDR3^β overlap between individuals than could be expected by chance. Taken together, these results suggest that the TCR repertoire is not randomly generated but instead may be biased toward specific rearrangements, perhaps pruned through T cell selection during thymic development or common antigen experiences^{23,30}.

Because this approach relies on multiple rounds of PCR before sequencing, there is potential to introduce sequencing biases and errors that could skew measurements of TCR diversity and/or generate artificial TCR sequences^{31–33}. However, several strategies exist to mitigate such bias. For example, because cDNA transcripts are already spliced, a smaller set of reverse primers can be used to target the C region when using mRNA instead of gDNA as starting material, thereby reducing PCR amplification bias from multiplexed J primers^{34,35}. Conversely, starting from gDNA eliminates the need for reverse transcription, which also minimizes the number of errors introduced during cDNA synthesis. Additionally, the use of synthetic TCR molecules targeting the multiplexed primers allows for the quantification of templates before and after multiplexed PCR, thereby enabling optimization of primer concentrations and correction of amplification biases³⁶.

5' RACE.

A second approach for TCR sequencing is based on 5' RACE^{25,27,28}. In this method, RNA is reverse transcribed by using a reverse transcriptase enzyme with terminal transferase

activity that adds untemplated C nucleotides to the 3' end of the cDNA. A template switch oligonucleotide (TSO) containing a complementary poly(G) stretch then anchors to this untemplated region, enabling the reverse transcriptase to switch templates and continue extending the cDNA to the end of the TSO, which includes a common adaptor sequence³⁷. As a result, one pair of primers targeting the 5' adaptor and the constant region is sufficient to amplify all TCR rearrangements. While 5' RACE strategies require fewer rounds of PCR, thereby reducing amplification biases prevalent in multiplex PCR approaches, these methods are still susceptible to errors from PCR, template switching or sequencing.

Error correction strategies for TCR amplification methods.

While these TCR sequencing strategies aim to provide an accurate representation of the TCR repertoire, they are subject to amplification biases and sequencing errors, as discussed above. The introduction of unique molecular identifiers (UMIs) provided a strategy to correct these errors³⁸. UMIs are random DNA sequences that are appended during cDNA synthesis and uniquely tag individual cDNA molecules³². This strategy has two main advantages: (1) the original distribution of TCR sequences present in the sample can be recovered by counting each UMI once, allowing for more accurate quantification of TCR clonal frequencies^{39,40}, and (2) PCR and sequencing errors can be corrected by grouping reads tagged with the same UMI together, so the true TCR sequence can be inferred by consensus (although errors introduced during reverse transcription will remain unresolved). Because TCRs may differ by only a few nucleotides, this correction step enables the distinction between true TCR variation and error-induced artificial diversity, resulting in more accurate estimations of TCR repertoire diversity. However, this accuracy may come at the cost of lower sensitivity, as low-frequency clones may be filtered out owing to insufficient read coverage per UMI⁴⁰. Error correction algorithms such as those using quality scores for each nucleotide have also been developed to generate more precise measurements of clonotype frequencies within the TCR repertoire⁴¹⁻⁴⁴.

Comparison of TCR targeting methods.

Given the technical differences among TCR sequencing strategies, users should consider several factors when analyzing the TCR repertoire. First, the availability of starting material may dictate which method is applicable. Each method can be used with RNA, allowing for sensitive detection of TCR sequences owing to higher copy numbers of RNA templates as compared to gDNA. However, if the RNA quality is poor, such as in samples obtained from chemically preserved archival tissue, gDNA multiplex PCR methods are preferable, as 5' RACE methods require high-quality RNA as starting material³⁴ (Table 1). Second, the purpose of the TCR analysis may also impact the choice of method. For example, RNA measurements cannot be directly correlated with cell numbers. Therefore, if the goal is to quantify clonal expansion of a T cell population, gDNA methods are preferred, because there is one genomic copy of the TCR per cell. Similarly, because multiplex PCR uses primer sets targeting different regions of the V gene, it is not always possible to capture full-length TCR sequences spanning the entire V(D)J region. This may be sufficient for analyzing the CDR3, which contains the highest sequence variability^{1,7}. However, if answering a particular biological question requires assessing variability in other regions, such as CDR1 and CDR2, then 5' RACE methods are more suitable.

Furthermore, the technical features of each step in these TCR sequencing approaches can have a substantial influence on the resultant repertoire data and, consequently, on their interpretation. In a recent comparative study of multiplex PCR and 5' RACE TCR sequencing methods, Barennes et al. benchmarked nine TCR sequencing protocols on the same bulk T cell sample to assess their reproducibility, replicability and sensitivity⁴⁰. They detected method-specific repertoire profiles that were largely consistent among replicates, suggesting that each protocol imposes unique biases when capturing the TCR repertoire. By comparing TRB usage with flow cytometry frequencies, they observed that RNA-based multiplex PCR methods exhibit stronger skewing toward specific V genes than gDNAbased multiplex PCR or 5' RACE methods, likely reflecting amplification biases due to differences in RNA transcript abundance⁴⁰. The ability to obtain a sequenced repertoire representative of the true biological repertoire is also highly dependent on the amount of starting material, regardless of method, particularly for the detection of rare TCR sequences. If the objective of a study is to capture maximal TCR diversity or detect rare clonotypes in highly diverse samples, it may be optimal to use non-UMI 5' RACE methods, as they are more sensitive than UMI-based methods in capturing rare clones, especially for TCRa chains^{33,40}. Conversely, if the objective is to construct a representative clonal structure of the TCR repertoire or identify TCRs of interest on the basis of expansion levels, then 5'RACE with UMI correction may be most suitable because it more faithfully recapitulates TCR clonotype frequencies in comparison to other approaches, although several replicates or higher sequencing depth may be needed to consistently capture lower-frequency clones⁴⁰.

While these bulk approaches can survey the TCR repertoire at high depth, this comes at the cost of resolving the pairing of TCR α and TCR β chains, because only one chain is captured at a time. This is an important limitation as dimerization of the two chains ultimately determines antigen specificity¹. Additionally, clonotype analyses are more accurately performed with TCR $\alpha\beta$ sequences, as the same TCR β sequence may pair with different TCR α sequences and vice versa⁴⁵. Therefore, considering each chain alone may lead to underestimates of TCR diversity, confounded analyses of intraclonal phenotypes and an inability to accurately identify T cell antigen specificities relevant to immune responses.

High-throughput methods for paired TCRαβ capture

Capturing paired TCR $\alpha\beta$ sequences provides more accurate resolution of the clonal structure present within the TCR repertoire and is necessary for assessing TCR function and antigen specificity. This is generally achieved by (1) computational inference of TCR $\alpha\beta$ pairing using a combinatorics-based strategy^{46,47} or (2) simultaneous TCR α and TCR β sequencing for physically isolated single cells^{10,48–52}.

Combinatorial inference of TCRaß pairing.

One approach for retrieving paired TCR $\alpha\beta$ repertoires relies on combinatorics⁴⁶. In this assay termed pairSEQ, T cells are randomly distributed across a plate such that each well contains a subset of cells tagged with the same well-specific DNA barcode (Fig. 2a). After pooling the samples for sequencing, TCR α and TCR β sequences that have the same set of barcodes are matched computationally and inferred to be a pair, as the likelihood of

two clones sharing the same set of wells is very low owing to the combinatorial nature of TCR rearrangements. In a similar approach developed by Lee et al., CDR3 $\alpha\beta$ pairs are computationally inferred from bulk-sequenced clones sorted into wells by using a repeated sampling strategy⁴⁷. One caveat of these frequency-based pairing methods is that only expanded clones can be resolved, therefore making these assays unsuitable for the discovery of rare TCR $\alpha\beta$ sequences.

Single-cell TCR sequencing techniques.

Recent advances in single-cell isolation techniques have made it possible to capture paired TCR $\alpha\beta$ chain information by using the TCR amplification strategies originally introduced for bulk populations (described above). The earliest methods for paired TCRαβ analysis in single cells relied on isolating individual T cells with micromanipulation techniques such as laser microdissection and micropipetting, followed by multiplex PCR and Sanger sequencing⁵³. While these approaches were instrumental in the first characterizations of the paired TCR $\alpha\beta$ repertoire, they are time-consuming and limited in both efficiency and throughput. Currently, a common method for isolating single cells is by fluorescenceactivated cell sorting (FACS)⁵⁴. An advantage of this approach is the ability to enrich for populations of interest on the basis of surface marker expression rather than sequencing all T cells in a sample, which may be useful when analyzing rare subsets. Additionally, the high read quality and coverage (~1,000 reads per chain) afforded by this method yield high-confidence TCR sequences¹⁰, which is crucial for analyzing clonal diversity. In this method, cells are stained with fluorescent antibodies against surface proteins and individually sorted into microtiter plates, resulting in the isolation of up to several hundred single cells at a time⁵⁵. The TCR chains are then reverse transcribed and amplified via TCR targeting strategies such as multiplex PCR. Initial studies using this strategy yielded the first characterizations of clonal diversity in antigen-specific CD8⁺ T cell responses to viral infection in both humans and mice and revealed a markedly higher frequency of dual-allelic TCR α expression than previously appreciated^{56–58}. More recent adaptations of this FACS-based isolation approach have incorporated high-throughput sequencing through the use of PCR-based barcoding strategies¹⁰. For example, in an approach described by Han et al., oligonucleotide barcodes unique to each well are introduced via a PCR extension step after nested TCR amplification¹⁰ (Fig. 2a). These barcodes preserve the cellular origin of transcripts, thus enabling samples to be pooled and sequenced together on a next-generation sequencer while maintaining TCRaß pairing.

Another strategy for performing paired TCR $\alpha\beta$ sequencing for individual cells is through cell-based emulsion PCR approaches^{48–52}. In these methods, single cells are captured within water-in-oil emulsions containing TCR primers and RT–PCR reagents. After encapsulation and cell lysis, T cells undergo overlap extension RT–PCR (OE-PCR) with primers targeting C regions and a set of V region primers containing a complementary sequence that enables linking of TCR $\alpha\beta$ transcripts within the emulsion, as described by Turchaninova et al.⁴⁸ (Fig. 2b). These fused products containing both chains can then be sequenced while maintaining native TCR $\alpha\beta$ pairing. For example, Munson et al. developed an emulsion OE-PCR method to identify tumor-specific TCR $\alpha\beta$ clones that were shared among patients with breast cancer but absent in healthy samples, providing evidence that antitumor T cell

responses may also be driven by shared antigens and not exclusively by patient-specific neoantigens⁴⁹.

Microfluidic platforms that partition individual cells in microwells or droplets have substantially increased the throughput of single-cell isolation⁵⁹, leading to their adoption in many single-cell TCR sequencing methods developed in the past few years. In well-based approaches, cells are loaded onto an array of nanoliter wells at low densities to confine single cells in nanowells by gravity^{59–62}. Alternatively, in droplet-based approaches, cells are flowed in an aqueous medium through microchannels on a chip filled with carrier oil at a rate and pressure such that the majority of droplets contain a single cell⁶³. A recent iteration of the OE-PCR approach used a high-throughput droplet microfluidic device to capture millions of T cells in a single assay^{50,51} (Fig. 2b), enabling the authors to profile the TCR repertoire of 15 healthy individuals, including six pairs of monozygotic twins⁶. By using the largest dataset of paired TCR $\alpha\beta$ sequences collected thus far, they demonstrated that the extent of TCR repertoire overlap between unrelated individuals is orders of magnitude smaller than previously appreciated by single-chain studies, highlighting the value of paired TCR $\alpha\beta$ data in providing meaningful insights into repertoire diversity. Furthermore, the authors found that V and J gene usage was significantly more correlated in twins than in unrelated individuals, suggesting that genetic influence has a role during thymic selection, likely driven by the presentation of antigens on identical MHC molecules⁶.

Single-cell methods for pairing the TCR with T cell state and function

While capturing TCR sequence information at single-cell resolution has been a powerful technical advance, studying the TCR repertoire in the context of cell state information can provide a more complete view of the functional and physiological role of T cells in immune responses. To this end, several groups have developed techniques to couple TCR sequencing with other modalities^{64–75} (Fig. 1b–d). One class of methods enables the simultaneous measurement of TCR and transcriptional profiles within single cells. This is achieved by either (1) computationally reconstructing TCR chains from single-cell RNA sequencing (scRNA-seq) reads^{64–69} or (2) specifically amplifying the TCR locus in conjunction with gene expression profiles^{70–73} (Fig. 3).

Pairing the TCR with T cell phenotype via computational TCR reconstruction.

Computational methods that extract TCR information from scRNA-seq data commonly rely on a combination of alignment and de novo assembly steps to identify TCR-originating reads and reconstruct TCR chains (Fig. 3c and Table 2). A challenge for TCR reconstruction using prior scRNA-seq methods was that they relied on amplifying the 3' transcript end by using the poly(A) tail as an anchor^{60,76–78}, which does not capture the V(D)J regions of the TCR located at the 5' transcript end. To overcome this, newer modalities use full-length cDNA amplification and sequencing strategies, in which single cells are reverse transcribed by using a template-switching mechanism similar to that of 5' RACE (Fig. 3b)^{25,26,79,80}. After fragmentation and sequencing, individual TCR sequences can be computationally reconstructed and analyzed in the context of their phenotypic profile.

One such TCR reconstruction algorithm, known as scTCRseq, constructs full-length paired TCR sequences by using nucleotide BLAST to align reads against a database of V and C genes and assembling these reads into consensus contigs (Fig. 3c, blue line)⁶⁴. The consensus sequences for the V and C genes are then padded with 'N' nucleotides and concatenated to produce a gapped V-C scaffold against which all reads are realigned and assembled into a final TCR sequence. Stubbington et al. developed a widely used TCR reconstruction tool. TraCeR, which uses a conceptually similar process, except a database of all possible V–J combinations is used for initial read alignment⁶⁵ (Fig. 3c, gray line). Another TCR reconstruction method, VDJPuzzle, includes an additional round of alignment and assembly where the TCR reconstructions from the first round are used as an updated database to capture TCRs that differ sufficiently from the reference genome^{66,67} (Fig. 3c, gold line). In contrast to these methods that require longer reads (>50 bp), the TCR reconstruction algorithm for paired-end single cell (TRAPeS) enables TCR assembly from short (25–30 bp) reads by using mate pairs to identify CDR3-originating reads followed by an iterative dynamic programming process to extend the CDR3 sequence⁶⁸ (Fig. 3c, orange line).

By combining TCR sequence with gene expression, these computational methods have provided some of the first views of T cell clonal phenotype dynamics. For example, Stubbington et al. applied TraCeR to scRNA-seq data generated from CD4⁺ T cells during mouse Salmonella infection by using SMART-seq on the Fluidigm C1 system⁶⁵. By pairing reconstructed TCRs with their cellular transcriptomes, the authors traced T cell clones along their differentiation trajectory from activated to T helper 1 (T_H1) and effector memory T cell states during infection and demonstrated that a single clone could adopt multiple cell fates⁶⁵. This finding confirmed previous adoptive transfer studies performed in the context of viral infections that noted heterogeneity in the cell fates of antigen-reactive T cells, supporting a model of progressive differentiation during T cell responses^{81–83}. Similarly, Ranasinghe et al. identified and characterized unconventional MHC class II-restricted CD8⁺ T cell responses in human immunodeficiency virus (HIV) controllers who are able to maintain undetectable viral loads without antiretroviral therapy⁸⁴. By using TRAPeS, they demonstrated that these antigen-specific CD8⁺ T cells exhibit cytotoxic antiviral phenotypes and are dominated by a single expanded TCR β clonotype within each individual, suggesting that unconventional class II-restricted CD8⁺ T cell responses may have a previously unappreciated role in human viral infection that can be harnessed for future vaccine design.

While these approaches have the advantage of analyzing TCRs by using previously generated scRNA-seq data, successful TCR reconstruction is dependent on sequencing depth and the expression level of the TCR locus, which is variable among cells^{65,85}. Consequently, incomplete capture of the TCR repertoire as a result of poor sequencing depth or limited starting material may skew biological interpretations regarding clonality and diversity.

Pairing the TCR with T cell phenotype via simultaneous RNA and targeted TCR sequencing.

To generate TCR sequencing libraries with higher coverage, several groups have developed techniques that combine RNA profiling with targeted amplification of both TCR chains in

single cells^{10,70–73} (Fig. 3a). One of the first methods to accomplish this used a multiplex PCR strategy to simultaneously target the TCR α and TCR β chains and a panel of 17 phenotyping genes that included cytokines and transcription factors important for T cell function and subset distinction¹⁰. This technique was used to characterize CD4⁺ tumor-infiltrating lymphocytes (TILs) in colorectal cancer and revealed clonal sharing between FOXP3⁺RORC⁺ and FOXP3⁻RORC⁺ T helper 17 (T_H17) subpopulations, suggesting that interleukin (IL)-17-producing cells can originate from a common ancestor that expands in response to antigenic activation and differentiates into distinct phenotypes¹⁰. This finding once again supports the concept that CD4⁺ T cells can exhibit lineage plasticity during an active immune response⁶⁵. Moreover, this study shed light on the ambiguous role of FOXP3⁺ T cells in tumor immunity, especially in conjunction with IL-17 expression, which has been associated with poor prognosis⁸⁶.

Subsequent studies have expanded the throughput of paired analysis by combining targeted TCR sequencing with droplet-based scRNA-seq methods that profile the whole transcriptome. For example, the InDrop method for 3' scRNA-seq was modified by using part of the amplified RNA library to target TCR transcripts with primers specific for the *TRAV* and *TRBV* genes before fragmentation (Fig. 3a, red line)^{70,78}. By applying this technique to regulatory T cells isolated from the mouse intestine, Zemmour et al. showed that cells expressing the same TCR are transcriptionally similar⁷⁰, suggesting that TCR signaling can shape the phenotype of a T cell, an observation that has since been recapitulated by other studies^{11,12}.

A similar method developed by Tu et al. enriches for TCR transcripts within cDNA libraries by using biotinylated oligonucleotide probes mapping to the C regions (Fig. 3a, purple line)⁷¹. Importantly, this approach is compatible with standard 3'-barcoded platforms, including Seq-Well and Drop-seq^{60,77}, thereby bypassing the need for specific kits, which are necessary for InDrop⁷⁸ (Table 1). Another approach for performing targeted TCR capture, known as repertoire and gene expression by sequencing (RAGE-seq), uses a hybridization bait library specific for V and J genes followed by long-read Oxford nanopore sequencing to capture full-length TCR transcripts while the remaining cDNA library is subjected to standard short-read Illumina sequencing for gene expression profiling (Fig. 3a, green line)⁷². Although this method has the advantage of capturing full-length TCR sequences, relative inefficiencies in the TCR recovery rate due to error-prone nanopore sequencing and the requirement for multiple sequencing technologies have perhaps prevented the widespread adoption of this method.

Finally, simultaneous scRNA-seq and TCR sequencing has also been implemented by using 5' RNA amplification⁷³. In this approach, cells are encapsulated within droplets containing barcoded beads in a similar fashion to 3' scRNA-seq methods; however, instead of using cellular barcodes adjacent to the poly(dT) sequence^{61,76–78}, the barcode in this method is adjacent to a TSO that is appended to the 5' end of the cDNA after template extension (Fig. 3a, blue line). Full-length cDNA is then pooled and amplified with PCR primers targeting common sequences added during reverse transcription. An aliquot of the cDNA library is used for targeted enrichment of TCR transcripts via PCR with primers mapping to the TCR constant regions followed by library construction, while the remaining material

is directly taken for construction of the gene expression library. This 5' capture method has been made commercially available by 10x Genomics and enabled several key insights into T cell responses, particularly in cancer. In one of the first applications of this method, Azizi et al. analyzed the TIL compartment in human breast cancer tumors to examine the relationship between TCR repertoire diversity and T cell phenotypic diversity¹¹. Integration of cell state and TCR clonotype data revealed a continuous trajectory of T cell activation that could partially be explained by TCR diversity. Furthermore, the authors observed that T cell clones are largely restricted to subsets of phenotypically related clusters that express similar environmental signatures such as anergy and gluconeogenesis, suggesting that cell state is collectively driven by TCR stimulation and microenvironmental cues¹¹. Yost et al. also used this method to track T cell clonal dynamics before and after anti-PD-1 therapy in site-matched tumors from patients with basal cell carcinoma¹². This study revealed that the post-treatment expanded CD8⁺ T cell population largely comprises exhausted clones that were previously absent from the tumor, suggesting that checkpoint blockade acts by recruiting new T cell clones rather than by reinvigorating existing exhausted clones within the tumor, as previously thought 12,87.

Pairing the TCR with surface protein.

Simultaneous analysis of the TCR and transcriptional profile of single cells can provide meaningful insights into T cell states at high resolution. However, the phenotyping of T cells using protein expression measurements may complement this analysis and more clearly define T cell subsets, as certain cell type markers are poorly detected at the transcript level⁸⁸. Furthermore, single-cell protein measurements may be better able to resolve (1) isoform usage, which is especially important for the identification of effector and memory T cell populations that express isoforms of the PTPRC gene, CD45RO and CD45RA, at varying levels⁸⁹, and (2) intracellular protein activity and modifications, such as the activity of critical T cell transcription factors. To this end, Mimitou et al. recently described a method called expanded CRISPR-compatible cellular indexing of transcriptomes and epitopes by sequencing (ECCITE-seq) that extends the 5' scRNA-seq and TCR sequencing method to capture cell-surface proteins in parallel with gene expression and TCR profiles⁷⁵. This is achieved by detecting proteins with DNA-barcoded antibodies conjugated to oligonucleotides complementary to the TSO. During reverse transcription, the cellular barcode is associated with the antibody-derived tags in addition to cDNA templates within the droplet. After amplification and sequencing, these antibody-derived tags can be read out and used to couple protein expression with TCR clonotype and RNA expression in each cell^{75,88}. Notably, this method can also be adapted to assess TCR antigen specificity by using oligonucleotide-conjugated pMHC multimers, which we discuss below.

ECCITE-seq can also be extended to directly capture single guide RNAs (sgRNA) from pooled CRISPR screens in parallel with RNA, protein and TCR profiling⁷⁵. By using an additional reverse transcription primer that anneals to the invariant 3' region of sgRNAs, the sgRNA sequence can be captured and linked to the cellular barcode via a template-switching mechanism analogous to that used for the RNA fraction. Proof-of-concept experiments have demonstrated the ability for ECCITE-seq to capture changes at both the mRNA and protein level resulting from sgRNA-directed genetic perturbations⁷⁵. This method presents

a promising approach by which future work may investigate how CRISPR-mediated perturbations modulate T cell clonality and phenotype.

Pairing the TCR with chromatin accessibility.

An orthogonal method for analyzing TCR sequences in their cellular context combines single-cell TCR capture with assay for transposase-accessible chromatin with sequencing (ATAC-seq), a technique that measures genome-wide chromatin accessibility^{90,91} (Fig. 1c). Termed transcript-indexed ATAC-seq (T-ATAC-seq)⁷⁴, this method involves the microfluidic isolation of single cells followed by tagmentation of genomic DNA with Tn.5, a transposase that preferentially inserts into open chromatin sites. TCR transcripts are then reverse transcribed by using primers specific for the C region and amplified via multiplex PCR with inner primers targeting the constant and variable regions, while the ATAC-seq fragments undergo 5' extension and PCR amplification. This approach enables the interrogation of how epigenetic factors, including *cis*-acting DNA elements and *trans*-acting transcription factors, may drive the specificity and expansion of T cell clones. For example, this method was used to identify an epigenomic signature of malignant clonal T cells in lymphoma that was previously undetected by standard FACS-based approaches and may potentially be applied to improve current diagnostic abilities to distinguish cancerous from benign T cell proliferation⁷⁴ and inform clinical therapies that target the epigenome⁹².

Strategies for mapping TCR sequence to antigen specificity

Linking antigen specificity to measurements of T cell clonality and phenotype is critical for understanding the drivers of the T cell response (Fig. 1d)^{93–99}. Several aspects of T cell antigen recognition make mapping TCR sequence to specificity a challenging problem. First, the frequency of rare populations of antigen-specific T cells may be as low as one in a million cells⁹⁴, which makes detection of these cells difficult. Second, high levels of variability due to the polymorphic nature of the MHC, the polyspecificity of both the TCR and pMHC, and the wide variety of potential epitopes encoded by a single antigen substantially add to the complexity of resolving TCR antigen specificities^{93,100}. Lastly, the relatively weak affinity and avidity of most TCR–pMHC interactions render selective isolation of antigen-specific T cell populations a nontrivial task^{93,98}.

While many approaches for assessing T cell antigen specificity exist, we focus our discussion here on strategies that also allow for TCR sequencing in parallel and refer interested readers to a recent review on T cell epitope discovery⁹³. A classical approach for studying antigen-specific T cells involves the use of fluorescently labeled pMHC multimers to isolate T cells that recognize the specific peptide in the context of a given MHC molecule via flow cytometry^{101,102}. The antigen-specific T cell populations can then be subjected to any of the methods discussed above for TCR sequencing or paired single-cell TCR and phenotype analysis. Although this approach is limited in throughput, several developments have increased the number of specificities that can be concurrently assessed. Recent advances in DNA and magnetic nanoparticle barcoding have increased the throughput of multimeric TCR specificity assays, allowing for the concurrent detection of more than 1,000 peptide specificities^{95–97,103}. While many of these methods have only

been used to identify the antigen specificities of T cells, several recent developments enable capturing the TCR as well, allowing for the characterization of clonality and TCR sequence variability among antigen-specific T cell populations. One such method is tetramer-associated TCR sequencing (TetTCR-seq), in which T cells are stained with fluorescently labeled DNA-barcoded pMHC tetramers and sorted into single-cell wells by FACS⁹⁶ (Fig. 4a). Following simultaneous RT–PCR amplification of TCR $\alpha\beta$ transcripts and DNA barcodes within each well, cellular barcodes can be used to link TCR sequences with their cognate antigens at the single-cell level. Zhang et al. used this method to screen over 150 cancer neoantigen and wild-type peptide pairs concurrently, demonstrating the utility of TetTCR-seq in identifying functionally reactive neoantigen-specific TCRs in a high-throughput manner⁹⁶. This approach can also be applied to the systematic interrogation of T cell specificity in the contexts of infection, autoimmunity and other immune-mediated diseases and may be pertinent for informing adoptive T cell therapy strategies.

Microfluidic techniques that further increase throughput have also been developed to capture antigen specificity and TCR sequence simultaneously. One method known as microfluidic antigen-TCR engagement sequencing (MATE-seq) uses magnetic nanoparticle-barcoded pMHC tetramers linked to photocleavable TCR-specific primers to capture both the TCR sequence and antigen identity within droplets⁹⁸ (Fig. 4b). Specifically, T cells are incubated with a library of nanoparticle-barcoded pMHCs and purified by magnetic separation⁹⁷. After single cells are encapsulated into droplets and lysed, the nanoparticle-barcoded pMHCs are exposed to UV light, releasing RT–PCR primers targeting TCRαβ C regions. Because these primers are linked to a DNA barcode corresponding to the pMHC, the TCR sequence and antigen specificity can be coupled at the single-cell level even after pooling and sequencing. While this method is currently limited to relatively few pMHC tetramers owing to limitations in library construction (Table 1), future iterations may enable the analysis of T cell populations with a larger array of antigen specificities. Although these advances have greatly improved the throughput and resolution for mapping TCR sequence to specificity, these methods are only applicable if the objective is to characterize T cells with a known antigen specificity and the peptide(s) of interest can be nominated a priori, which in some cases may not be possible.

Similarly to the identification of TCR sequences that bind specific antigens, the converse problem of identifying antigen(s) that are recognized by a particular TCR is also challenging. One approach for linking antigen specificity to TCR sequences that is still relatively nascent is computational epitope prediction^{104–107}. This approach involves training a supervised machine learning model on a collection of known TCR–antigen pairs and using this classifier to predict the antigen specificities of new TCR sequences. Although some models have been able to distinguish TCR specificity for a small number of epitopes^{106,107}, these algorithms are as yet unable to achieve high prediction accuracy on full TCR repertoires, as models must be trained separately for each epitope or set of epitopes. Improvements to these methods will require substantially larger training sets of experimentally verified TCR–pMHC pairs than are currently available through in-house and public databases such as McPAS-TCR, VDJdb and the Immune Epitope Database (IEDB)^{108–111}. This could be facilitated by the continued collection of TCR–pMHC data through the strategies for concurrently capturing TCR sequence and peptide

specificity described above^{94,96,98}, along with studies using TCR-directed methods for epitope discovery⁹³. Additionally, algorithmic advancements that better model the three-dimensional complexity and flexibility of the TCR–pMHC interaction will be instrumental in improving computational epitope prediction approaches. However, even with more sophisticated algorithms that enable improved prediction of TCR epitopes, the issue of identifying T cell clones that are physiologically relevant to a given antigen-specific response is complicated by TCR polyspecificity. This cross-reactivity, whereby one TCR is able to recognize multiple unrelated peptides^{112,113}, makes precise linking of specificity to in vivo biological function difficult. Given these challenges, it is likely that, while these machine learning algorithms may prove helpful in nominating TCRs of interest for further investigation, they may require coupling with additional assays to provide a complete solution for TCR specificity determination.

Standards for TCR sequencing data collection

Importantly, the recent widespread adoption of the high-throughput TCR discovery methods described in this Review has led to the accumulation of large amounts of TCR sequencing data. Thus, there is an impetus for the standardization of TCR sequencing data to facilitate data reproducibility and sharing. To this end, the Adaptive Immune Receptor Repertoire (AIRR) Community has defined the Minimal Information about Adaptive Immune Receptor Repertoire (MiAIRR) standard, a set of guidelines for reporting metadata from antibody and TCR sequencing studies, ranging from details on study design and sample processing to data processing, annotation and submission to NCBI repositories^{114,115}. The establishment of a standardized schema and file format for representing annotated receptor rearrangements has also enabled streamlined access to data from repositories supporting this format¹¹⁶, which are collectively organized into the AIRR Data Commons¹¹⁷. Continued efforts to define data standards and encourage adherence to these guidelines will be instrumental in promoting an ever-improving collaborative ecosystem among the research community that fosters progress through the use of TCR sequencing techniques.

Future perspectives

Although these advances in TCR sequencing have propelled the T cell field forward rapidly, there is a need for improved efficiency, sensitivity and cost, particularly for single-cell methods. Current droplet-based sequencing approaches have singlet capture rates of approximately 65% (ref.¹¹⁸). This low efficiency limits the interpretability of studies, as the cells captured in an assay may not be representative of the starting population, especially when analyzing large samples such as peripheral blood and other immunologically relevant organs. Another limitation of current droplet-based single-cell technologies is the minimum input requirement of approximately 1,000 cells per capture¹¹⁹, which precludes the analysis of rare subsets, such as antigen-specific T cell populations. Additionally, while microfluidic single-cell methods are highly parallelized and carried out in nanoliter reaction volumes, leading to reduced cost in comparison to traditional plate-based methods¹¹⁹, the cost to profile the TCR repertoire using single-cell methods is currently still higher than with bulk methods and may be limiting in some settings⁵⁹. Future work that improves on these challenges will therefore be crucial for generating more faithful representations of the T cell

compartment. For instance, the adaptation of microwell array techniques may provide a way forward in addressing these challenges to TCR sequencing methods, as they are compatible with lower cell inputs and more efficient than droplet-based methods⁶⁰⁻⁶².

Beyond these technical improvements, there is opportunity for combining the technologies discussed here with orthogonal methods to achieve an even finer understanding of how T cell clonality and specificity coordinately influence physiological function. One area with potential for innovation is in resolving T cell specificity and phenotype in situ. While existing single-cell methods can link the TCR repertoire with phenotype, the spatial organization of T cells within a tissue is lost. Recently developed spatial transcriptomic techniques enable high-throughput gene expression profiling while maintaining two-dimensional positional information^{120,121}. Progress on these methods could enable spatially resolved TCR and phenotypic pairing. The ability to study the clonality and diversity of TCRs in their spatial context will be instrumental in dissecting how the anatomical distribution of T cells and their cognate antigens shapes immune responses.

Finally, recent advances in single-cell genetic lineage tracing methods that make use of DNA barcodes to label cells and their progeny have proven to be tremendously valuable in informing paradigms of hematopoiesis and a plethora of other biological processes^{122–124}. Combining lineage tracing with TCR profiling has the potential to refine understanding of T cell developmental trajectories and phenotypic plasticity, specifically in regard to intraclonal phenotypic and functional heterogeneity. Previous efforts to track T cells using fate mapping have relied on mice with transgenic TCRs, which revealed important features of the clonal T cell response, such as the ability for a T cell to give rise to progeny exhibiting heterogeneous patterns of differentiation and growth kinetics^{125,126}. However, these studies did not allow for analysis of the contribution of TCR clonality and specificity in shaping T cell responses. Integration of TCR sequencing and transcriptomic profiling with genetic tracing strategies could open the door for tracking T cell clones in vivo at an unprecedented resolution, enabling the reconstruction of lineage and phenotypic hierarchies within clones. While these approaches may require several iterations before they are available for use in physiological contexts such as human tissues, innovations such as these will nonetheless contribute to a better understanding of how T cell responses are coordinated, which can then be harnessed to develop improved therapies for cancer, infection and other immune-mediated diseases.

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Competing interests

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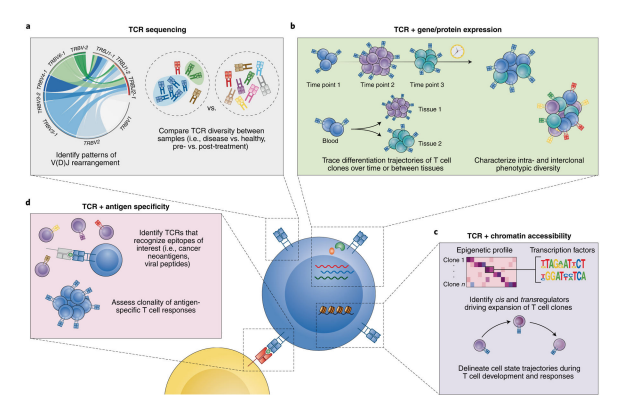


Fig. 1 |. **Characterization of T cell dynamics using multiomic TCR sequencing approaches. a**, TCR sequencing through bulk or single-cell methods enables profiling of TCR repertoire diversity and clonality. **b**, Paired TCR sequencing and profiling of gene or protein expression allows for joint analysis of T cell clonality and phenotype and enables the tracing of clones over time or throughout tissues. **c**, TCR sequencing combined with measuring chromatin accessibility reveals clonal epigenetic signatures and uncovers developmental trajectories that may be obscured at the level of gene or protein expression. **d**, TCR epitope discovery methods identify antigens driving the T cell response.

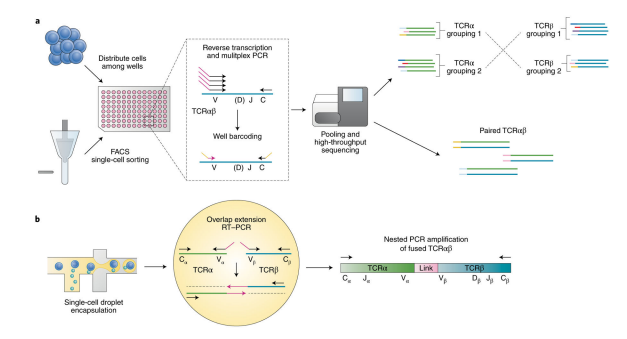


Fig. 2 |. Overview of single-cell TCR sequencing approaches.

a, Left, T cells are distributed among wells on a plate at a density of 10^3-10^5 cells per well in the pairSEQ method (top) or single T cells are sorted by FACS into individual wells (bottom). Middle, within each well, TCR transcripts are reverse transcribed, amplified and attached to well-specific barcodes. Right, after the amplified libraries are pooled and deep sequenced, TCRa β pairings are computationally inferred by analyzing which TCRa and TCR β sequences share the same set of well barcodes (top) or cellular barcodes (bottom). **b**, In emulsion-based methods, single T cells are encapsulated into droplets, where they undergo OE-PCR to enable linking of TCRa and TCR β transcripts. Fused TCRa β transcripts are then pooled, amplified and sequenced.

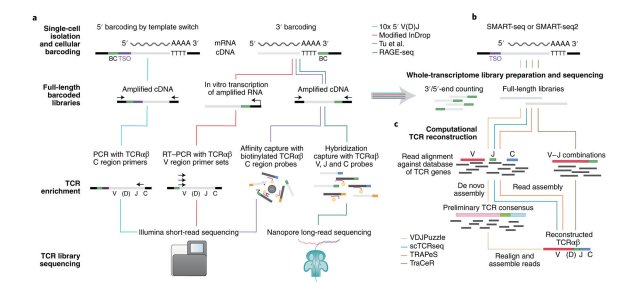


Fig. 3 |. Schematic of paired scRNA-seq and TCR sequencing methods.

a, Whole-transcriptome cDNA single-cell libraries barcoded at either the 3' or 5' end are subjected to gene expression profiling and targeted TCR capture with primers or hybridization probes. TCR and gene expression reads originating from the same cell can be linked on the basis of cellular barcodes. BC, barcode. **b**, Full-length whole-transcriptome cDNA libraries are generated from single cells by the SMART-seq or SMART-seq2 approach and sequenced. **c**, Computational methods for reconstructing TCR sequences from scRNA-seq libraries without targeted TCR amplification involve the alignment of reads to TCR gene reference databases or a 'recombinome' of possible V–J pairings. In **a**–**c**, colored lines indicate the different methods described.

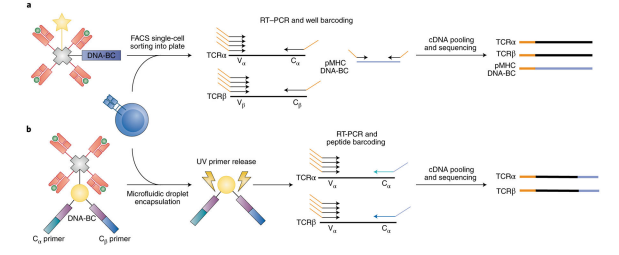


Fig. 4 |. Overview of single-cell methods for linking TCR sequence to antigen specificity. a,b, pMHC tetramers conjugated to a fluorophore (a) or magnetic nanoparticle (b) and a unique DNA barcode (DNA-BC) are incubated with a T cell population and single-cell sorted into wells (a) or encapsulated in droplets (b). In a, TCR transcripts and the pMHC DNA barcode (purple) are simultaneously captured and attached to well barcodes, allowing TCR sequences to be matched to pMHC specificity. In b, TCR transcripts are captured by cell-barcoded V region and C region primers containing the DNA barcode. Final sequenced TCR products contain both the cell barcode and pMHC DNA barcode, enabling pairing of TCR $\alpha\beta$ sequences to antigen specificity.

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Table 1

methods
sequencing
of TCR
Overview

Select advantages	Compatible with either RNA or DNA as starting material	Reduced PCR amplification bias	Can be implemented using standard laboratory reagents and equipment; high read coverage yields high-confidence TCR sequences	Does not require physical isolation of single cells; easily implemented with standard laboratory reagents and equipment	High TCRach pairing efficiency due to physical linkage of transcripts; increased sensitivity and fewer artifacts as compared to standard PCR approaches	Compatible with InDrop amplified RNA libraries	Compatible with most 3' scRNA-seq methods; applicable to preexisting amplified cDNA libraries	Adaptable to any 3' or 5' scRNA-seq method
TCRaß capture efficiency	1	I	~80%	1	NAR	NAR	~35%	~17%
No. of antigen specificities assessed ^a	1	1	1	1	I	I	1	I
Chromatin accessibility	1	I	1	1	1	I	1	1
Protein expression	I	I	1	1	1	1	1	I
Gene expression	1	I	Targeted panel	1	I	Whole transcriptome	Whole transcriptome	Whole transcriptome
TCR targeting	Multiplex PCR	RACE PCR	Multiplex PCR	Multiplex PCR	Overlap extension multiplex PCR	Multiplex PCR	Multiplex PCR	Hybridization capture
TCR regions	CDR3	Full length	CDR3	CDR3	CDR3	CDR3	CDR3	Full length
Paired TCRαβ	No	No	Yes	Yes	Yes	Yes	Yes	Yes
No. of cells assessed	>10 ⁵	>105	10 ² -10 ³	>105	10 ³ -10 ⁶	10^{2} 10 ⁴	10 ³ -10 ⁴	10^{3} - 10^{4}
Platform compatibility	1	I	Plate based	Plate based	Droplet	Droplet	Microwell or droplet	Droplet
Refs.	22- 24	25- 29	10	46	48- 52	70	71	72
Method	Multiplex PCR (bulk)	5' RACE (bulk)	Han et al.	pairSEQ	Emulsion RT-PCR	Modified InDrop	Tu et al.	RAGE-seq

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Method	Refs.	Platform compatibility	No. of cells assessed	Paired TCRαβ	TCR regions	TCR targeting	Gene expression	Protein expression	Chromatin accessibility	No. of antigen specificities assessed ^a	TCRaβ capture efficiency	Select advantages
5' V(D)J with feature barcoding	73	Droplet	10^{3} - 10^{4}	Yes	Full length	RACE PCR	Whole transcriptome	Yes	I	44	~65%	Commercially available (10x Genomics) and easy to use
ECCITE- seq	75	Droplet	10 ³ -10 ⁴	Yes	Full length	RACE PCR	Whole transcriptome	Yes	1	NAR	~65%	Simultaneously captures many modalities, including sgRNA perturbations; highly modular
T-ATAC- seq	74	Microwell	10^{2} 10 ³	Yes	CDR3	Multiplex PCR	I	Ι	Yes	I	~70%	Pairs TCR clonality with epigenomic state
TetTCR- seq	96	Plate based	10 ² -10 ³	Yes	CDR3	Multiplex PCR	I	1	1	315	~70%	Sensitive detection of rare antigen-specific T cell populations; easily scalable to large peptide libraries
MATE-seq	86	Droplet	10 ² -10 ⁴	Yes^b	CDR3	Multiplex PCR	1	1	1	4	NAR	Sensitive detection of rare antigen-specific T cell populations; compatible with lower sample inputs
^a Demonstrated by original paper.	1 by origin	nal paper.										

emonstrated by original paper.

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b Paired TCR $\alpha\beta$ per peptide, not per single cell. NAR, not assessed/reported.

					Table 2			
Single-cel.	l compi	Single-cell computational TCR reconstructic	on methods					
Method	Refs.	Refs. TCR alignment reference	Minimum input read length (bp) ^a	Input data format	Sequencing mode	Language	Code availability	other software dependencies
scTCRseq 64	64	IMGT V and C genes	50	FASTQ	Paired or single end Python and Perl	Python and Perl	https://github.com/ ElementoLab/scTCRseq	SEQTK, Blastall, GapFiller and Vidjil
TraCeR	65	V-J recombinomes	50	FASTQ	Paired or single end Python	Python	https://github.com/ Teichlab/tracer	Bowtie2, Trinity, IgBLAST and kallisto/salmon
VDJPuzzle	66,67	VDJPuzzle 66,67 Reference genome V, D, J and C genes	150	FASTQ or BAM	Paired end	Bash, Python and Perl	https://bitbucket.org/ kirbyvisp/vdjpuzzle	TopHat2 or STAR, Trinity, IgBLAST and MiGMAP
TRAPeS	68	Reference genome V, J and C genes	25	BAM	Paired end	Python and C++	https://github.com/ YosefLab/TRAPeS	Bowtie2 and RSEM
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 $^{\rm a}{\rm For}$ a recovery rate greater than 90%, as demonstrated by the original paper.

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