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Can we predict T cell specificity with digital biology and machine learning?

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Abstract

Recent advances in machine learning and experimental biology have offered breakthrough solutions to problems such as protein structure prediction that were long thought to be intractable. However, despite the pivotal role of the T cell receptor (TCR) in orchestrating cellular immunity in health and disease, computational reconstruction of a reliable map from a TCR to its cognate antigens remains a holy grail of systems immunology. Current data sets are limited to a negligible fraction of the universe of possible TCR-ligand pairs, and performance of state-of-the-art predictive models wanes when applied beyond these known binders. In this Perspective article, we make the case for renewed and coordinated interdisciplinary effort to tackle the problem of predicting TCR-antigen specificity. We set out the general requirements of predictive models of antigen binding, highlight critical challenges and discuss how recent advances in digital biology such as single-cell technology and machine learning may provide possible solutions. Finally, we describe how predicting TCR specificity might contribute to our understanding of the broader puzzle of antigen immunogenicity.

Sections

Introduction

State of the art

Key challenges

Conclusions and call to action

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Introduction

T cells typically recognize antigens presented on members of the MHC protein family via highly diverse heterodimeric T cell receptors (TCRs) expressed at their surface (Fig. 1). These antigens are commonly short peptide fragments of eight or more residues, the presentation of which is dictated in large part by the structural preferences of the MHC allele¹. Lipid, metabolite and oligosaccharide T cell antigens have also been reported²⁻⁴. TCRs typically engage antigen–MHC complexes via one or more of their six complementarity-determining loops (CDRs), three contributed by each chain of the TCR dimer.

The pivotal role of the TCR in surveillance and response to disease, and in the development of new vaccines and therapies, has driven concerted efforts to decode the rules by which T cells recognize cognate antigen–MHC complexes. However, cost and experimental limitations have restricted the available databases to just a minute fraction of the possible sample space of TCR–antigen binding pairs (Box 1). As we discuss later, these data sets^{5–8} are also poorly representative of the universe of self and pathogenic epitopes and of the varied MHC contexts in which they may be presented (Fig. 2).

The research community has therefore turned to machine learning models as a means of predicting the antigen specificity of the so-called orphan TCRs having no known experimentally validated cognate antigen. Accurate prediction of TCR-antigen specificity can be described as deriving computational solutions to two related problems: first, given a TCR of unknown antigen specificity, which antigen–MHC complexes is it most likely to bind; and second, given an antigen–MHC complex, which are the most likely cognate TCRs?

A critical requirement of models attempting to answer these questions is that they should be able to make accurate predictions for any combination of TCR and antigen-MHC complex. These should cover both 'seen' pairs included in the data on which the model was trained and novel or 'unseen' TCR-epitope pairs to which the model has not been exposed⁹. Impressive advances have been made for specificity inference of seen epitopes in particular disease contexts. For example, clusters of TCRs having common antigen specificity have been identified for Mycobacterium tuberculosis¹⁰ and SARS-CoV-2 (ref.¹¹), providing possible avenues for new vaccine and pharmaceutical development. However, as discussed later, performance for seen epitopes wanes beyond a small number of immunodominant viral epitopes and is generally poor for unseen epitopes^{9,12}. This matters because many epitopes encountered in nature will not have an experimentally validated cognate TCR, particularly those of human or non-viral origin (Fig. 2). In the text to follow, we refer to the case for generalizable TCRantigen specificity inference, meaning prediction of binding for both seen and unseen antigens in any MHC context.

We must also make an important distinction between the related tasks of predicting TCR specificity and antigen immunogenicity. The former, and the focus of this article, is the prediction of binding between sets of TCRs and antigen–MHC complexes. The latter can be described as predicting whether a given antigen will induce a functional T cell immune response: a complex chain of events spanning antigen expression, processing and presentation, TCR binding, T cell activation, expansion and effector differentiation. Although great strides have been made in improving prediction of antigen processing and

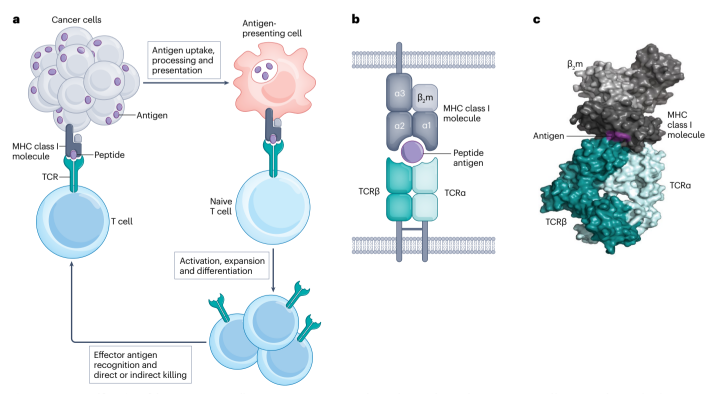


Fig. 1 | Structure and function of the TCR. a, Cartoon illustrating cancer cell antigen presentation to a naive T cell; T cell activation and expansion and effector T cell engagement of the cancer cell. b, Antigen recognition by conventional T cells through the interaction of the $\alpha\beta$ T cell receptor (TCR)

heterodimer with peptide antigen presented by an MHC class I molecule. **c**, Crystal structure of the affinity-enhanced A3A TCR engaging with melanomaassociated antigen 3 (MAGE-A3)-derived peptide presented by HLA-A*01 (ref.¹⁰¹) (generated with data from ref.¹⁰¹ and visualized with PyMOL (see Related links)).

presentation for common HLA alleles, the nature and extent to which presented peptides trigger a T cell response are yet to be elucidated¹³. A significant gap also remains for the prediction of T cell activation for a given peptide^{14,15}, and the parameters that influence pathological peptide or neoantigen immunogenicity remain under intense investigation¹⁶. We believe that only by integrating knowledge of antigen presentation, TCR recognition, context-dependent activation and effector function at the cell and tissue level will we fully realize the benefits to fundamental and translational science (Box 2).

State of the art

From deepening our mechanistic understanding of disease to providing routes for accelerated development of safer, personalized vaccines and therapies, the case for constructing a complete map of TCR-antigen interactions is compelling. We now explore some of the experimental and computational progress made to date, highlighting possible explanations for why generalizable prediction of TCR binding specificity remains a daunting task.

Experimental methods

The development of recombinant antigen–MHC multimer assays¹⁷ has proved transformative in the analysis of TCR–antigen specificity, enabling researchers to track and study T cell populations under various conditions and disease settings^{18–20}. Nonetheless, critical limitations remain that hamper high-throughput determination of TCR–antigen specificity. We direct the interested reader to a recent review²¹ for a thorough comparison of these technologies and summarize some of the principal issues subsequently.

Antigen-MHC multimers may be used to determine TCR specificity using bulk (pooled) T cell populations, or newer single-cell methods. Bulk methods are widely used and relatively inexpensive, but do not provide information on $\alpha\beta$ TCR chain pairing or function. As a result, single chain TCR sequences predominate in public data sets (Fig. 2). However, both α -chains and β -chains contribute to antigen recognition and specificity^{22,23}. We shall discuss the implications of this for modelling approaches later. Multimodal single-cell technologies provide insight into chain pairing and transcriptomic and phenotypic profiles at cellular resolution, but remain prohibitively expensive, return fewer TCR sequences per run than bulk experiments and show significant bias towards TCRs with high specificity²⁴⁻²⁶. The appropriate experimental protocol for the reduction of nonspecific multimer binding, validation of correct folding and computational improvement of signal-to-noise ratios remain active fields of debate^{25,26}. Indeed, concerns over nonspecific binding have led recent computational studies to exclude data derived from a 10× study of four healthy donors²⁷.

Although bulk and single-cell methods are limited to a modest number of antigen–MHC complexes per run, the advent of technologies such as lentiviral transfection assays^{28,29} provides scalability to up to 96 antigen–MHC complexes through library-on-library screens. However, previous knowledge of the antigen–MHC complexes of interest is still required. This precludes epitope discovery in unknown, rare, sequestered, non-canonical and/or non-protein antigens³⁰.

The advent of synthetic peptide display libraries (Fig. 3a) permits the extension of binding analysis to hundreds of thousands of peptides per TCR³⁰⁻³³. Using transgenic yeast expressing synthetic peptide–MHC constructs from a library of 2×10^8 peptides, Birnbaum et al.³¹ dissected the binding preferences of autoreactive mouse and human TCRs, providing clues as to the mechanisms underlying autoimmune targeting in multiple sclerosis. High-throughput library screens such

Box 1

The extraordinary diversity of TCR-antigen pairs

At a conservative estimate of 5 million unique T cell receptors (TCRs) per individual at a given time¹⁰², a global population of 8 billion sharing 11% of their TCRs¹⁰² would represent a unique TCR pool of 3.6×10^{15} . This figure excludes recognition of antigens from over 1,400 pathogens known to be capable of infecting humans¹⁰³, binding to self and neoantigens and presentation of antigens in over 34,000 HLA contexts¹⁰⁴. The universe of feasible TCR-antigen-MHC combinations is, therefore, likely to be orders of magnitude higher, especially when accounting for degeneracy in TCR-antigen recognition.

as these provide opportunities for improved screening of the antigen-MHC space, but limit analysis to individual TCRs and rely on TCR-MHC binding instead of function. There remains a need for high-throughput linkage of antigen specificity and T cell function, for example, through mammalian or bead display³⁴⁻³⁷.

As a result of these barriers to scalability, only a minuscule fraction of the total possible sample space of TCR-antigen pairs (Box 1) has been validated experimentally. At the time of writing, fewer than 1 million unique TCR-epitope pairs are available from VDJdb, McPas-TCR, the Immune Epitope Database and the MIRA data set⁵⁻⁸ (Fig. 2). Just 4% of these instances contain complete chain pairing information (Fig. 2a). About 97% of all antigens reported as binding a TCR are of viral origin, and a group of just 100 antigens makes up 70% of TCR-antigen pairs (Fig. 2b,c). Where the HLA context of a given antigen is known, the training data are dominated by antigens presented by a handful of common alleles (Fig. 2d). Many antigens have only one known cognate TCR (Fig. 2e). These limitations have simultaneously provided the motivation for and the greatest barrier to computational methods for the prediction of TCR-antigen specificity.

Computational methods

A comprehensive survey of computational models for TCR specificity inference is beyond the scope intended here but can be found in the following helpful reviews^{15,38-42}. Broadly speaking, current models can be divided into two categories, which we dub supervised predictive models (SPMs) (Fig. 3b) and unsupervised clustering models (UCMs) (Fig. 3c) on account of their respective use of supervised learning and unsupervised learning. A non-exhaustive summary of recent open-source SPMs and UCMs can be found in Table 1.

Supervised predictive models. SPMs are those which attempt to learn a function that will correctly predict the cognate epitope for a given input TCR of unknown specificity, given some training data set of known TCR-peptide pairs. The past 2 years have seen an acceleration of publications aiming to address this challenge with deep neural networks (DNNs). Although there are many possible approaches to comparing SPM performance, among the most consistently used is the area under the receiver-operating characteristic curve (ROC-AUC). One would expect to observe 50% ROC-AUC from a random guess in a

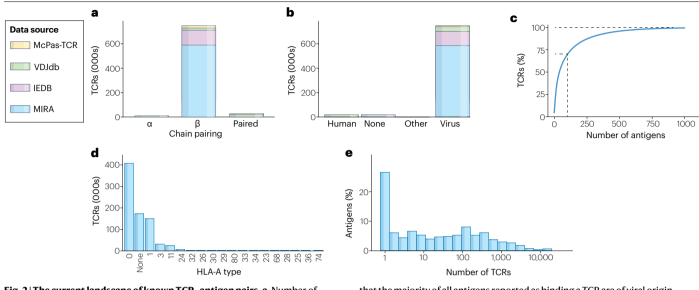


Fig. 2 | The current landscape of known TCR-antigen pairs. a, Number of T cell receptors (TCRs) containing α -chains, β -chains or paired chains, showing variation in numbers according to the data set (manually curated catalogue of pathology-associated TCR sequences (McPas-TCR), VDJ database (VDJdb), Immune Epitope Database (IEDB) and multiplex identification of TCR antigen specificity (MIRA)). **b**, Number of TCRs per antigen species of origin, showing

that the majority of all antigens reported as binding a TCR are of viral origin. **c**, Cumulative frequency of antigens, showing that a group of 100 antigens makes up 70% of TCR-antigen pairs. **d**, Number of TCRs by HLA-A type, showing that known antigens are reported in complex with only a few common HLA alleles. **e**, Frequency histogram showing that most antigens have only one known cognate TCR in the combined data set⁵⁻⁸.

binary (binding or non-binding) task, assuming a balanced proportion of negative and positive pairs.

Performance by this measure surpasses 80% ROC-AUC for a handful of 'seen' immunodominant viral epitopes presented by MHC class I^{9,43}. However, representation is not a guarantee of performance: 60% ROC-AUC has been reported for HLA-A2*01-CMV-NLVPMVATV⁴⁴, possibly owing to the recognition of this immunodominant antigen by diverse TCRs. Critically, few models explicitly evaluate the performance of trained predictors on unseen epitopes using comparable data sets. Weber et al.¹² achieved an average of $62 \pm 6\%$ ROC-AUC for TITAN, compared with 50% for ImRex on a reference data set of unseen epitopes from VDJdb and COVID-19 data sets. Values of 56 ± 5% and 55 ± 3% were reported for TITAN and ImRex, respectively, in a subsequent paper from the Meysman group⁴⁵. Other groups have published unseen epitope ROC-AUC values ranging from 47% to 97%; however, many of these values are reported on different data sets (Table 1), lack confidence estimates following validation⁴⁶⁻⁴⁹ and have not been consistently reproducible in independent evaluations⁵⁰.

Together, these results highlight a critical need for a thorough, independent benchmarking study conducted across models on data sets prepared and analysed in a consistent manner^{27,50}. Until then, newer models may be applied with reasonable confidence to the prediction of binding to immunodominant viral epitopes by common HLA alleles. However, SPMs should be used with caution when generalizing to prediction of any epitope, as performance is likely to drop the further the epitope is in sequence from those in the training set⁹.

Unsupervised clustering models. Unlike SPMs, UCMs do not depend on the availability of labelled data, learning instead to produce groupings of the TCR, antigen or HLA input that reflect the underlying statistical variations of the data^{19,51} (Fig. 3c). Applied to TCR repertoires, UCMs take as their input single or paired TCR CDR3 amino acid sequences, with or without gene usage information, and return a mapping of sequences to unique clusters. Clustering is achieved by determining the similarity between input sequences, using either 'hand-crafted' features such as sequence distance or enrichment of short sub-sequences, or by comparing abstract features learnt by DNNs (Table 1).

Clustering provides multiple paths to specificity inference for orphan TCRs³⁹⁻⁴¹. Epitope specificity can be predicted by assuming that if an unlabelled TCR is similar to a receptor of known specificity, it will bind the same epitope⁵². One may also co-cluster unlabelled and labelled TCRs and assign the modal or most enriched epitope to all sequences that cluster together⁵¹. Finally, DNNs can be used to generate 'protein fingerprints', simple fixed-length numerical representations of complex variable input sequences that may serve as a direct input for a second supervised model^{25,53}.

As for SPMs, quantitative assessment of the relative merits of hand-crafted and neural network-based UCMs for TCR specificity inference remains limited to the proponents of each new model. Although some DNN-UCMs allow for the integration of paired chain sequences and even transcriptomic profiles⁴⁸, they are susceptible to the same training biases as SPMs and are notably less easy to implement than established clustering models such as GLIPH and TCRdist^{19,54}. However, these established clustering models scale relatively poorly to large data sets compared with newer releases^{51,55}. Recent analyses^{27,53} suggest that there is little to differentiate commonly used UCMs from simple sequence distance measures. Here again, independent benchmarking analyses would be valuable, work towards which our group is dedicating significant time and effort.

Key challenges

Despite the exponential growth of unlabelled immune repertoire data and the recent unprecedented breakthroughs in the fields of data science and artificial intelligence, quantitative immunology still lacks a framework for the systematic and generalizable inference of T cell antigen specificity of orphan TCRs. Among the most plausible explanations for these failures are limitations in the data, methodological gaps and incomplete modelling of the underlying immunology.

Data

As we have set out earlier, the single most significant limitation to model development is the availability of high-quality TCR and antigen-MHC pairs. The need is most acute for under-represented antigens, for those presented by less frequent HLA alleles, and for linkage of epitope specificity and T cell function. Meanwhile, single-cell multimodal technologies have given rise to hundreds of millions of unlabelled TCR sequences^{8,56}, linked to transcriptomics, phenotypic and functional information. However, these unlabelled data are not without significant limitations. Notably, biological factors such as age, sex, ethnicity and disease setting vary between studies and are likely to influence immune repertoires. Differences in experimental protocol, sequence pre-processing, total variation filtering (denoising) and normalization between laboratory groups are also likely to have an impact: batch correction may well need to be applied⁵⁷. Therefore, thoughtful approaches to data consolidation, noise correction, processing and annotation are likely to be crucial in advancing state-of-the-art predictive models.

Modelling

The exponential growth of orphan TCR data from single-cell technologies, and cutting-edge advances in artificial intelligence and machine learning, has firmly placed TCR-antigen specificity inference in the spotlight. However, we believe that several critical gaps must be addressed before a solution to generalized epitope specificity inference can be realized.

First, models whose TCR sequence input is limited to the use of β-chain CDR3 loops and VDJ gene codes are only ever likely to tell part of the story of antigen recognition, and the extent to which single chain pairing is sufficient to describe TCR-antigen specificity remains an open question. Structural⁵⁸ and statistical⁵⁹ analyses suggest that α -chains and β -chains contribute equally to specificity, and incorporating both chains has improved predictive performance⁴⁴. However, chain pairing information is largely absent (Fig. 2a), and many state-ofthe-art SPMs and UCMs rely on single chain information alone (Table 1). Although CDR3 loops may be primarily responsible for antigen recognition, residues from CDR1, CDR2 and even the framework region of both α -chains and β -chains may be involved⁵⁸. Subtle compensatory changes in interaction networks between peptide-MHC and TCR, altered binding modes and conformational flexibility in both TCR and MHC may underpin TCR cross-reactivity^{60,61}. Explicit encoding of structural information for specificity inference has until recently been limited to studies of a limited set of crystal structures^{19,62}. However, the advent of automated protein structure prediction with software programs such as RoseTTaFold, ESMFold and AlphaFold-Multimer provide potential opportunities for large-scale sequence and structure interpretations of TCR epitope specificity⁶³⁻⁶⁵. This has been illustrated in a recent preprint in which a modified version of AlphaFold-Multimer has been used to identify the most likely binder to a given TCR, achieving a mean ROC-AUC of 82% on a small pool of eight seen epitopes⁶⁶.

To train models, balanced sets of negative and positive samples are required. In the absence of experimental negatives, negative instances may be produced by shuffling or drawing randomly from healthy donor repertoires⁹. However, these approaches assume, on the one hand,

Box 2

Implications of accurate TCR specificity prediction

The ability to accurately predict the cognate ligand of a given T cell receptor (TCR) or antigen–MHC complex has important implications for the design of new therapies and vaccines and our understanding of the biological role of T cells in health and disease³⁸.

In oncology, T cell antigen recognition has become the focus of new drug development efforts, including checkpoint inhibitors, chimeric antigen receptor (CAR) T cells, endogenous or affinityenhanced TCRs, and cancer vaccines¹⁰⁵. Cross-reactivity in TCRbased T cell therapies has presented a major roadblock to the development of safe interventions, and gaps in preclinical screening have led to tragedies in the clinic¹⁰⁶. Accurate and generalizable specificity inference could provide an additional safety net to robust experimental screens, predicting likely autoreactivity for a given patient population in oncology and beyond^{107/08}.

Beyond the implications for new medicines development, there is significant potential to use predictive tools to dissect the fundamental role of T cells in the surveillance of malignancy. For example, there are reports of the accumulation of clones with driver mutations in sun-exposed skin¹⁰⁹, but the extent to which mutational burden is reflected in TCR repertoires is not well understood¹¹⁰. Exhausted cytotoxic CD8⁺ T cells have long been known to be a hallmark of an inefficient antitumour immune response^{111–113}. However, although early data are emerging^{114,115}, we do not yet fully know whether T cells with particular antigen specificity are more likely to be exhausted.

For infectious diseases such as SARS-CoV-2, predictors of T cell specificity could be of great use in understanding the magnitude and dynamics of antigen-specific T cell responses to the disease¹¹⁶ and vaccination¹¹⁷. However, there remains a significant opportunity to improve open-source systems immunology tools for confident linkage of T cell antigen specificity to differential vaccine-induced response.

Linkage of expanded effector T cell populations to their cognate self-antigen will also provide vital diagnostic clues as to disease aetiology of autoimmune conditions. This is exemplified by a recent longitudinal study that demonstrated an association between Epstein–Barr virus infection and the incidence of multiple sclerosis, supportive of new vaccine development¹¹⁸.

a Multiplex analysis of pMHC-TCR binding

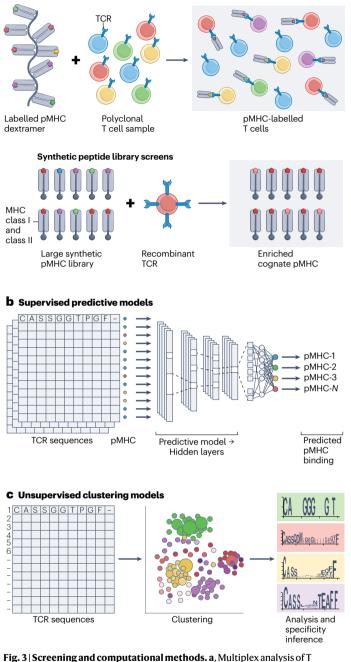


Fig. 3 | **Screening and computational methods. a**, Multiplex analysis of T cell receptor (TCR)–peptide–MHC (pMHC) antigen specificity and synthetic peptide library screens for interrogation of peptide specificity of a single TCR. **b**, Representation of a deep neural network for supervised prediction of TCR–antigen specificity. **c**, Unsupervised clustering analysis of TCR–antigen specificity showing (centre) an example clustering visualization and (right) complementarity-determining loop 3 sequence logos.

that TCRs do not cross-react and, on the other hand, that the healthy donor repertoires do not include sequences reactive to the epitopes of interest. A recent study from Jiang et al.⁶⁷ provides interesting strategies to address this challenge.

Finally, developers should use the increasing volume of functionally annotated orphan TCR data to boost performance through transfer learning: a technique in which models are trained on a large volume of unlabelled or partially labelled data, and the patterns learnt from those data sets are used to inform a second predictive task. This technique has been widely adopted in computational biology, including in predictive tasks for T and B cell receptors^{49,66,68}. Indeed, the best-performing configuration of TITAN made used a TCR module that had been pretrained on a <u>BindingDB</u> database (see Related links) of 471,017 protein–ligand pairs¹². Incorporating evolutionary and structural information through sequence and structure-aware representations of the TCR and of the antigen–MHC complex^{69,70} may yield further benefits.

Immunology

It is now evident that the underlying immunological correlates of T cell interaction with their cognate ligands are highly variable and only partially understood, with critical consequences for model design. Importantly, TCR-antigen specificity inference is just one part of the larger puzzle of antigen immunogenicity prediction^{16,18}, which we condense into three phases: antigen processing and presentation by MHC, TCR recognition and T cell response.

Antigen processing and presentation pathways have been extensively studied, and computational models for predicting peptide binding affinity to some MHC alleles, especially class I HLAs, have achieved near perfect ROC-AUC^{15,71} for common alleles. However, this problem is far from solved, particularly for less-frequent MHC class I alleles and for MHC class II alleles⁷.

A key challenge to generalizable TCR specificity inference is that TCRs are at once specific for antigens bearing particular motifs and capable of considerable promiscuity^{72,73}. This contradiction might be explained through specific interaction of conserved 'hotspot' residues in the TCR CDR loops with corresponding two to three residue clusters in the antigen, balanced by a greater tolerance of variations in amino acids at other positions⁶⁰. TCRs may also bind different antigen-MHC complexes using alternative docking topologies⁵⁸. Despite the known potential for promiscuity in the TCR, the pre-processing stages of many models assume that a given TCR has only one cognate epitope. Another under-explored yet highly relevant factor of T cell recognition is the impact of positive and negative thymic selection and more specifically the effect of self-peptide presentation in formation of the naive immune repertoire⁷⁴.

Many groups have attempted to bypass this complexity by predicting antigen immunogenicity independent of the TCR¹⁴, as a direct mapping from peptide sequence to T cell activation. However, similar limitations have been encountered for those models as we have described for specificity inference. Many predictors are trained using epitopes from the Immune Epitope Database labelled with readouts from single time points⁷. However, Achar et al.⁷⁵ illustrated that integrating cytokine responses over time improved prediction of quality. Antigen load and affinity can also play important roles^{74,76}. Thus, models capable of predicting functional T cell responses will likely need to bridge from antigen presentation to TCR-antigen recognition, T cell activation and effector differentiation and to integrate complex tissue-specific cytokine, cell phenotype and spatiotemporal data sets. Our view is that, although T cell-independent predictors of immunogenicity have clear translational benefits, only after we can dissect the relative contribution of the three stages described earlier will we understand what determines antigen immunogenicity.

Glossary

Area under the receiver-

operating characteristic curve (ROC-AUC). ROC-AUC and the area under the precision-recall curve (PR-AUC) are measures of model tendency to different classes of error. These plots are produced for classification tasks by changing the threshold at which a model prediction falling between zero and one is assigned to the positive label class, for example, predicted binding of a given T cell receptor-antigen pair. ROC-AUC is the area under the line described by a plot of the true positive rate and false positive rate. ROC-AUC is typically more appropriate for problems where positive and negative labels are proportionally represented in the input data. PR-AUC is the area under the line described by a plot of model precision against model recall. PR-AUC is typically more appropriate for problems in which the positive label is less frequently observed than the negative label.

Library-on-library screens

Experimental screens that permit analysis of the binding between large libraries of (for example) peptide-MHC complexes and various T cell receptors.

Machine learning models

A broad family of computational and statistical methods that aim to identify statistically conserved patterns within a data set without being explicitly programmed to do so. Machine learning models may broadly be described as supervised or unsupervised based on the manner in which the model is trained. Many recent models make use of both approaches.

Neural networks

A family of machine learning models inspired by the synaptic connections of the brain that are made up of stacked layers of simple interconnected models. Although each component of the network may learn a relatively simple predictive function, the combination of many predictors allows neural networks to perform arbitrarily complex tasks from millions or billions of instances. Neural networks may be trained using supervised or unsupervised learning and may deploy a wide variety of different model architectures. Deep neural networks refer to those with more than one intermediate layer.

Shuffling

In the absence of experimental negative (non-binding) data, shuffling is the act of assigning a given T cell receptor drawn from the set of known T cell receptorantigen pairs to an epitope other than its cognate ligand, and labelling the randomly generated pair as a negative instance.

Supervised learning

Models that learn a mathematical function mapping from an input to a predicted label, given some data set containing both input data and associated labels. Common supervised tasks include regression, where the label is a continuous variable, and classification, where the label is a discrete variable.

Synthetic peptide display libraries

Experimental systems that make use of large libraries of recombinant synthetic peptide–MHC complexes displayed by yeast³⁰, baculovirus³² or bacteriophage³³ or beads³⁵ for profiling the sequence determinants of immune receptor binding. Peptide diversity can reach 10⁹ unique peptides for yeast-based libraries.

Training data

The training data set serves as an input to the model from which it learns some predictive or analytical function.

Unsupervised learning

Models that learn to assign input data to clusters having similar features, or otherwise to learn the underlying statistical patterns of the data. Unlike supervised models, unsupervised models do not require labels. Common unsupervised techniques include clustering algorithms such as *K*-means; anomaly detection models and dimensionality reduction techniques such as principal component analysis⁸⁰ and uniform manifold approximation and projection.

Validation

Analysis done using a validation data set to evaluate model performance during and after training. A given set of training data is typically subdivided into training and validation data, for example, in an 80%:20% ratio. Models may then be trained on the training data, and their performance evaluated on the validation data set.

New experimental and computational techniques that permit the integration of sequence, phenotypic, spatial and functional information and the multimodal analyses described earlier provide promising opportunities in this direction^{75,77}. Integrating TCR sequence and cell-specific covariates from single-cell data has been shown to improve performance in the inference of T cell antigen specificity⁴⁸. By taking a graph theoretical approach, Schattgen et al.⁷⁸ reported an association between clonotype clustering with the cellular phenotypes derived from gene expression and surface marker expression. We believe that such integrative approaches will be instrumental in unlocking the secrets of T cell antigen recognition.

Conclusions and call to action

Together, the limitations of data availability, methodology and immunological context leave a significant gap in the field of T cell immunology in the era of machine learning and digital biology. We believe that by harnessing the massive volume of unlabelled TCR sequences emerging from single-cell data, applying data augmentation techniques to counteract epitope and HLA imbalances in labelled data, incorporating sequence and structure-aware features and applying cuttingedge computational techniques based on rich functional and binding data, improvements in generalizable TCR-antigen specificity inference are within our collective grasp. To aid in this effort, we encourage the following efforts from the community.

First, a consolidated and validated library of labelled and unlabelled TCR data should be made available to facilitate model pretraining and systematic comparisons. Second, a coordinated effort should be made to improve the coverage of TCR-antigen pairs presented by less common HLA alleles and non-viral epitopes. We encourage the continued publication of negative and positive TCR-epitope binding data to produce balanced data sets. Third, an independent, unbiased and systematic evaluation of model performance across SPMs, UCMs and combinations of the two (Table 1) would be of great use to the community. Such a comparison should account for performance on common and infrequent HLA subtypes, seen and unseen TCRs and epitopes, using consistent evaluation metrics including but not limited to ROC-AUC and area under the precision-recall curve. We encourage validation strategies such as those used in the assessment of ImRex and TITAN^{9,12} to substantiate model performance comparisons. In the future, TCR specificity inference data should be extended to include multimodal contextual information as a means of bridging from TCR binding to immunogenicity prediction.

The scale and complexity of this task imply a need for an interdisciplinary consortium approach for systematic incorporation of the latest immunological understandings of cellular immunity at the tissue level

Table 1 | A non-exhaustive list of supervised and unsupervised models for inference of TCR epitope specificity published since 2020

| Model | Date | TCR chain input | Training data | Method | Availability |
|--------------------------------|-----------------------------|------------------|--|-----------------------------|---|
| Supervised predi | ctive models | | | | |
| ATM-TCR ⁸¹ | 07/2022 | Single | IEDB ⁷ McPas-TCR ⁶ VDJdb ⁵ | DNN-SPM | https://github.com/Lee-CBG/ATM-TCR |
| ImmuneML ⁸² | 11/2021 | Paired | Heikkila ⁸³ VDJdb ⁵ | DNN-SPM | https://immuneml.uio.no/ |
| NetTCR2 (ref. 44) | 10/09/2021 | Single or paired | IEDB ⁷ VDJdb ⁵ 10× ⁸⁴ | DNN-SPM | https://services.healthtech.dtu.dk/service.php?NetTCR-2.0 |
| SwarmTCR ⁸⁵ | 07/09/2021 | Single or paired | IEDB ⁷ VDJdb⁵ Private | SPM | https://github.com/thecodingdoc/SwarmTCR |
| ImRex ⁹ | 07/2021 | Single | VDJdb ⁵ Dean ⁸⁶ | DNN-SPM | https://github.com/pmoris/ImRex |
| Luu et al. ⁴³ | 04/2021 | Single | IEDB ⁷ McPas-TCR ⁶ PIRD ⁸⁷ VDJdb ⁵ | DNN-SPM | https://github.com/jssong-lab/TCR-Epitope-Binding |
| TCRGP ⁸⁸ | 03/2021 | Single or paired | Dash et al. ⁵⁴ VDJdb ⁵ | SPM | https://github.com/emmijokinen/TCRGP |
| TcellMatch ⁴⁸ | 08/2020 | Paired | IEDB ⁷ VDJdb ⁵ 10× ⁸⁴ | DNN-SPM | https://github.com/theislab/tcellmatch |
| SETE ⁸⁹ | 06/2020 | Single | Dash et al. ⁵⁴ VDJdb ⁵ | SPM | https://github.com/wonanut/SETE |
| Unsupervised clu | stering models ^a | I | | | |
| ClusTCR ⁵⁵ | 12/2021 | Single or paired | VDJdb ⁵ Emerson ²³ | UCM | https://github.com/svalkiers/clusTCR |
| TCRdist3 (ref. ¹¹) | 11/2021 | Single or paired | Dash et al. ⁵⁴ Nolan ⁸ Snyder ⁹⁰ VDJdb ⁵ | UCM | https://github.com/kmayerb/tcrdist3/ |
| GIANA ⁵¹ | 08/2021 | Single | Dash et al. ⁵⁴ Glanville et al. ¹⁹ IEDB ⁷ VDJdb ⁵ Zhang et al. ⁹¹ | UCM | https://github.com/s175573/GIANA |
| GLIPH2 (ref. ¹⁰) | 04/2020 | Single or paired | Private VDJdb⁵ | UCM | http://50.255.35.37:8080/ |
| iSMART ⁹² | 03/2020 | Single | Emerson ²³ VDJdb ⁵ TCGA | UCM | https://github.com/s175573/iSMART |
| Other | | | | | |
| TCRDock ⁶⁶ | 08/2022 | Paired | BFD ⁷⁰ PDB ⁹³ | Pre-trained DNN and DNN-SPM | https://github.com/phbradley/TCRdock |
| TCR-BERT ⁴⁹ | 11/2021 | Single | PIRD ⁸⁷ VDJdb ⁵ TCRdb ⁹⁴ | Pre-trained DNN and DNN-SPM | https://huggingface.co/wukevin/tcr-bert |
| TITAN ¹² | 07/2021 | Single | BindingDB ⁹⁵ VDJdb ⁵ ImmuneCODE ⁹⁶ | Pre-trained DNN and DNN-SPM | https://github.com/PaccMann/TITAN |

Table 1 (continued) | A non-exhaustive list of supervised and unsupervised models for inference of TCR epitope specificity published since 2020

| Model | Date | TCR chain input | Training data | Method | Availability |
|--------------------------|---------|------------------|--|------------------------------------|--|
| Other (continue | d) | | | | |
| pMTNet ⁴⁷ | 09/2021 | Single | Chen et al. ⁹⁷ Huth et al. ⁹⁸ Joglekar et al. ³⁷ PIRD ⁸⁷ McPas-TCR ⁶ VDJdb ⁵ Zhang et al. ⁹¹ 10× ⁸⁴ | Pre-trained DNN and DNN-SPM | https://github.com/tianshilu/pMTnet |
| ERGO-II ⁹⁹ | 04/2021 | Single or paired | Kanakry et al. ¹⁰⁰ McPas-TCR ⁶ VDJdb ⁵ Zhang et al. ⁹¹ | Pre-trained DNN-UCM and DNN-SPM | https://github.com/IdoSpringer/ERGO-II |
| ICON-TCRAI ²⁵ | 05/2021 | Single or paired | McPas-TCR ⁶ VDJdb ⁵ 10× ⁸⁴ | Pre-trained DNN and DNN-SPM | https://github.com/regeneron-mpds/ICON |
| TCRMatch ⁵² | 03/2021 | Single | IEDB ^{7,a} 10× ⁸⁴ | Unsupervised: nearest neighbour | https://www.github.com/IEDB/TCRMatch |
| DeepTCR ⁵³ | 03/2021 | Single or paired | Dash et al. ⁵⁴ Glanville et al. ¹⁹ Sidhom et al. ⁵³ 10× ⁸⁴ Multiple public sources: see DeepTCR GitHub repository | DNN-UCM or DNN-SPM | https://github.com/sidhomj/DeepTCR |

ATM-TCR, multi-head self-attention model-TCR; BFD, Big Fantastic Database; DNN, deep neural network; ERGO, peptide TCR matching prediction; GIANA, geometric isometry-based TCR alignment algorithm; GLIPH2, grouping of lymphocytes by paratope hotspots 2; ICON-TCRA, integrative context-specific normalisation TCR-artificial intelligence; IEDB, Immune Epitope Database; ImRex, interaction map recognition; McPas-TCR, manually curated catalogue of pathology-associated TCR sequences; PIRD, pan-immune repertoire database; pMTNet, pMHC-TCR binding prediction network; SETE, sequence-based ensemble learning; SPM, supervised predictive model; TCGA, The Cancer Genome Atlas; TCR, T cell receptor; TCR-BERT, TCR-bidirectional encoder representations from transformers; TCRGP; TCR Gaussian process; TITAN, TCR epitope bimodal attention networks; UCM, unsupervised clusteringmodel; VDJdb, VDJ database. *Not all UCMs are explicitly trained, and so data sets reported for non-DNN UCMs are those on which the models were evaluated.

and cutting-edge developments in the field of artificial intelligence and data science. This should include experimental and computational immunologists, machine-learning experts and translational and industrial partners. Considering the success of the critical assessment of protein structure prediction series⁷⁹, we encourage a similar approach to address the grand challenge of TCR specificity inference in the short term and ultimately to the prediction of integrated T and B cell immunogenicity. Competing models should be made freely available for research use, following the commendable example set in protein structure prediction^{65,70}.

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Author contributions

H.K. and D.H. researched and wrote the article. R.A.F., M.B. and G.O. reviewed and edited the manuscript before submission.

Competing interests

G.O. is a co-founder of T-Cypher Bio. D.H. and R.A.F provide consultancy services to companies active in T cell antigen discovery and vaccine development. The other authors declare no competing interests.

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