

Structural Basis of Specificity and Cross-Reactivity in T Cell Receptors Specific for Cytochrome c–I-E^k

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T cells specific for the cytochrome *c* Ag are widely used to investigate many aspects of TCR specificity and interactions with peptide-MHC, but structural information has long been elusive. In this study, we present structures for the well-studied 2B4 TCR, as well as a naturally occurring variant of the 5c.c7 TCR, 226, which is cross-reactive with more than half of possible substitutions at all three TCR-sensitive residues on the peptide Ag. These structures alone and in complex with peptide-MHC ligands allow us to reassess many prior mutagenesis results. In addition, the structure of 226 bound to one peptide variant, p5E, shows major changes in the CDR3 contacts compared with wild-type, yet the TCR V-region contacts with MHC are conserved. These and other data illustrate the ability of TCRs to accommodate large variations in CDR3 structure and peptide contacts within the constraints of highly conserved TCR–MHC interactions. *The Journal of Immunology*, 2011, 186: 5823–5832.

In early studies of T cell recognition, a strongly immunodominant peptide from pigeon cytochrome c (PCC) emerged as an informative model Ag for studying the nature of MHC restriction (1–3). This has since been widely employed in studies of TCR genes and proteins to gain important insights into the many aspects of TCR specificity and function, including relating thermodynamic and kinetic properties to T cell responses (4). However, the structural foundations upon which these interactions are based have been elusive, despite efforts spanning decades.

The strength of this experimental system stems from the limited TCR usage of the CD4⁺ T cell population that responds to the single peptide epitope PCC_{88–104}, or the related moth cytochrome c peptide (MCC_{88–103}), presented by I-E^k (PCC/I-E^k or MCC/I-E^k). This population is strongly dominated by cells expressing TCRs

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The structures and coordinates presented in this article have been submitted to the Research Collaboratory for Structural Bioinformatics Protein Data Bank (PDB) (http://www.pdb.org) under PDB ID 3QIB, 3QIW, 3QJF, and 3QJH.

The online version of this article contains supplemental material.

Abbreviations used in this article: APL, altered peptide ligand; H-bond, hydrogen bond; MCC, moth cytochrome c; PCC, pigeon cytochrome c; PDB, Protein Data Bank; pMHC, peptide-MHC; SPR, surface plasmon resonance.

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encoded by the Val1 (TRAV4) and VB3 (TRBV26) gene segments (5–7) that contain a number of reoccurring TCR α and TCRB CDR3 junctional sequence features reported to be critical for clonal recruitment to this oligoclonal response (7-9). The frequency of cells bearing TCRs with the key junctional features that define the dominant PCC reactive clonotypes increases sharply over the course of the primary immune response and continues to focus in response to secondary challenge in a manner that correlates well with the relative affinity for MCC/I- E^{k} (7–9). Interestingly, this relationship between TCR CDR3 usage and affinity has provided important insights into what TCR characteristics facilitate recruitment from the CD4⁺ T cell repertoire to join a polyclonal response to Ag delivered by various adjuvants (10, 11) and those TCR features that promote differentiation to specific Th or CD25⁺ regulatory T cell subsets (12). Finally, variants of the MCC peptide have recently been used to address the relationship between potency and T cell responsiveness or differentiation in vivo (13, 14). Thus, the body of information that has been accumulated with this system has provided important insights into how the diverse CD4⁺ T cell repertoire is accessed and used in response to Ag.

Prior to the availability of TCR and TCR-peptide-MHC (pMHC) crystallographic information, several aspects of the structural basis for the preferred sequence characteristics of MCC/ I-E^k-specific TCRs were deduced from experiments based on peptide (15) and MHC mutagenesis (16) combined with TCR sequence analysis. One such study clearly identified two conserved CDR3 residues (a93 Glu and B100 Asn, henceforth referred to as α 89E and β 97N using the same numbering system in the deposited Protein Data Bank [PDB] files) that appeared to be involved in reciprocal interactions with a critical lysine at the p5 position of the MCC peptide (MCC-99K, p5K) and a critical threonine at the p8 position (MCC-102T, p8T), respectively (17). In some T cells that reacted against MCC peptide with a lysine to glutamate substitution at p5 (MCC-99E), a charge reversal was observed at the CDR3 α 89 position, strongly suggesting that this position would be involved in a charge-charge interaction with the peptide (an example of one such TCR is the 202 TCR; see Supplemental Table II). However, cross-reactive TCRs capable of

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recognizing both versions of MCC (p5K and p5E) were also identified. In subsequent experiments with one of these cross-reactive TCRs, 226, it was found that p5K versus p5E recognition was differentially affected by MHC mutations at several positions on both the α - and β -chains of I-E^k, suggesting that pMHC binding was significantly altered depending on which peptide was being recognized (16). But, it was not clear how the 226 TCR recognizes these two dramatically different peptides (charge reversed at the centrally located p5 position) despite exhibiting the canonical features of a MCC/I-E^k–reactive TCR.

TCR degeneracy and polyspecificity have been the subjects of many studies (18-20). The cross-reactive potential of individual TCRs has been tested by systematically varying one amino acid at a time on the agonist peptide (altered peptide ligands [APLs]) (15, 21-23). Combinatorial approaches have also been taken, including the use of peptide pools fixed at particular residues or higherthroughput display methods, to look for alternate agonist peptides for a given TCR (24-26). The APL studies have revealed that most naturally derived TCRs can tolerate conservative mutations at a few TCR contact residues with occasional strong specificity for a particular residue at a couple key TCR contacts (15, 21, 22). Not surprisingly, certain experimental conditions can give rise to TCRs with drastically reduced specificity for peptide. For instance, TCRs engineered to have higher affinity for given Ag also lose specificity for that Ag and gain the ability to recognize a wide array of APLs (27, 28). Highly degenerate TCRs have also been derived from mice expressing MHC with tethered peptides to restrict the pMHC repertoire available for thymocyte selection (29), and, in some cases, these highly degenerate TCRs are autoreactive. The sum of these data suggests a division of labor for peptide and MHC recognition by TCRs.

From a series of studies on VB8.2-containing TCRs, we previously hypothesized that germline-encoded V-region gene segments have evolved to make conserved contacts on MHC, which we have termed recognition codons, that underlie the structural basis for TCR recognition of pMHC (25, 30, 31). Conserved VB8.2 interactions with a range of I-A allotypes presenting different peptides are also seen in other TCR-pMHC complexes (32), and this germline motif is critical for positive selection (33). However, this notion is complicated by several factors. One is the availability of a large number of MHC alleles, each of which needs to be compatible for a given species' set of TCR V-regions. It is also evident that peptide binding in the groove of MHC can significantly alter the nonpeptide surface of the MHC, which may alter germline-encoded contacts. Lastly, a major technical challenge is that there are a large number of TCR V α and V β pairings available, and each may use a different set of evolved codons to recognize each MHC allele. As of yet, the majority of crystallographic information has come from the mouse system and has focused mostly on comparing TCR-pMHC complex structures of VB8 TCRs. Although much has been learned about how the germline-encoded VB8.2 region contributes to binding various MHC class I and II molecules presenting a range of peptides, it is important to investigate the generality of these findings to other TCR V-regions and MHC molecules.

In this study, we have used the cytochrome *c* system to address the structural basis of TCR cross-reactivity using 226 as a naturally occurring example of a highly degenerate TCR. Previous data suggest that this TCR can use multiple modes of MHC recognition as part of its ability to recognize various ligands (16); therefore, this system serves as a stringent test of the recognition codon hypothesis (30, 31). Furthermore, we analyzed the previously unknown structures of $V\alpha 11^+V\beta 3^+$ TCRs that recognize the mouse MHC allele, I-E^k, which has yet to be crystallized as part of a TCR–pMHC complex. We determined the structures of the 2B4 and 226 TCRs in both the unliganded state as well as bound to an MCC/I-E^k complex and, for 226, bound to the chargereversed p5E-MCC/I-E^k pMHC complex to assess the validity of previous mutational data, suggesting that germline-encoded MHC contacts were significantly altered between these two complexes. We report that all three V α 11/V β 3 TCRs binding to MCC and MCC-p5E/I-E^k had nearly identical germline-encoded contacts with the MHC, despite gross differences in somatically derived CDR3-mediated contacts with the peptide.

Materials and Methods

Preparation and validation of proteins

For all three complexes, the TCR was expressed in bacteria as chimeric proteins made up of mouse 2B4, 5c.c7, or 226 VJ or VDJ regions fused to human constant regions [generously provided by Dr. J. McCluskey and Dr. J. Rossjohn (34) and refolded as described] (32, 35). In addition, the I- E^{k} binding peptides (MCC or MCC-p5E) were tethered to the N terminus of the TCRβ-chain with a 10-residue flexible linker sequence as a tether with sequences MANERADLIAYLKQATKGGGGSGGGGG or MANE-RADLIAYLEQATKGGGGGGGGGGGG. His-tagged I-E^k with a $\beta\text{-chain}$ N-terminally tethered thrombin-cleavable CLIP peptide was expressed in insect cells using a baculovirus expression system and dual promoter construct driving expression of both MHC chains modified from one generously provided by Dr. P. Allen as described (36). After Ni-NTA and size-exclusion purification, the MCC-tethered TCR and CLIP/I-E^k were subjected to an exchange reaction to form a stable tethered TCR-pMHC complex similar to that previously described (37, 38). The MCC-tether-TCR was mixed with CLIP/I-E^k at a 3:1 molar ratio at a final protein concentration of 4 mg/ml in PBS before addition of 3 U/ml thrombin and 1 h incubation at 37°C to remove the tethered CLIP peptide. A total of 30 mM sodium cacodylate (pH 6.2) was then added to adjust pH and facilitate peptide exchange during a 16-h incubation at 37°C. The stable tethered 100-kDa complex was then purified by size exclusion and purity and composition verified by SDS-PAGE electrophoresis in boiled, nonboiled, reducing, and nonreducing conditions.

Exogenous TCR expression in T cell hybridomas for functional studies

Wild-type and mutant 2B4 TCRs were expressed in the $58\alpha^-\beta^-$ TCR⁻ hybridomas for functional studies as described (39). In brief, ecotropic murine stem cell virus retrovirus containing full-length TCR chains as well as CD4 coreceptors were produced in the Phoenix retroviral expression system using FuGene transfection (Roche Molecular Biochemicals). $58\alpha^-\beta^-$ T cell hybridomas were then infected with a combination of viral supernatants from phoenix cells transfected with TCR α , β , and CD4 expression constructs. Each TCR chain is equipped with either puromycin or zeocin antibiotic resistance under control of an internal ribosome entry sequence immediately downstream of the TCR sequence; infected $58\alpha^-\beta^-$ T cell hybridomas were then selected on the basis of drug resistance (2.5 µg/ml puromycin, 50 µg/ml zeocin). To verify TCR (PE-conjugated anti-mouse TCR) and CD4 expression (allophycocyanin-Cy7–conjugated anti-CD4, both Abs obtained from BD Biosciences), the cells were stained with fluorescent Abs and run on a flow cytometer (Beckman Coulter).

To test the function of this exogenously expressed TCR, infected hybridomas were cocultured with various concentrations of peptide-pulsed APCs (CH27 B cells) both at 5×10^5 cells/well in 96-well plates. After ~16 h, supernatants were harvested and subjected to IL-2 ELISA assay. The concentrations of each peptide required to elicit production of either 10 or 50% of the maximal IL-2 produced by wild-type 2B4 transduced cells were determined from plots of IL-2 relative units versus the log of peptide concentration, as described (15).

226 hybridoma functional studies

MCC and altered peptide induced production of IL-2 by 226 hybridomas to assess the extent of 226 cross-reactivity against all possible amino acid substitutions at the three TCR contacts was performed exactly as described (15). For these experiments, irradiated B10.BR splenocytes were used as I-E^k-expressing APCs that were then pulsed with various concentrations of each MCC peptide mutant.

Crystallization and x-ray data collection

Protein crystals were grown via hanging-drop vapor diffusion at 22°C. 2B4-MCC/I-E^k complex crystals were grown in 50% PEG200, 0.2 M sodium chloride, and 0.1 M sodium potassium phosphate (pH 6.2). 226-MCC and 226-MCC–p5K/I- E^k complex crystals were grown in 15–20% PEG5000 MME and Bis Tris (pH 6.5–7). The unliganded 5c.c7 and 2B4 crystals were grown in 20% PEG3350, 0.1 M calcium chloride, and Tris (pH 7.7). Data sets were collected at Stanford Synchrotron Radiation Light Source (Stanford, CA) and Advanced Light Source (Berkeley, CA). Data were indexed, integrated, and scaled using either Mosflm (40) and SCALA from the CCP4 suite (41) or the HKL-2000 program package (42).

Structure determination and refinement

All structures were solved via molecular replacement using the program Phaser (43). The molecular replacement solution for the unliganded 2B4 structure was determined using the variable domains of the 2C TCR (PDB ID 1TCR) and the constant domains of the ELS4 TCR (PDB ID 2NW2). The refined 2B4 coordinates in conjunction with I-E^k (PDB ID 1FNE) were used as search models for the 2B4-MCC/I-E^k complex. The molecular replacement solutions for the 226-I-E^k complex structures were determined using the refined coordinates from the 2B4-MCC/I-E^k complex. To avoid model bias, peptides residues and CDR loops were removed from the search models. Manual model building was preformed in COOT followed by iterative rounds of coordinate and B-factor refinement in either the program CNS (44) or Phenix (45). The program PROCHECK (46) was used to monitor the geometry of the models. The CCP4 suite programs Contact and AreaIMol were used to assess the interface contacts and buried surface areas, respectively. Structural figures were generated using the program Pymol (Delano Scientific, San Carlos, CA).

Surface plasmon resonance

SPR data were collected on a BiaCORE 3000 system or a BiaCORE T100 system (GE Healthcare) as described (47). TCR was expressed as a hybrid receptor consisting of the clonotypic TCR V α and V β domains fused to human LC13 TCR constant domain as described above (32). *Escherichia coli* inclusion bodies were refolded by fast dilution as described (34) and purified on an Äkta system (GE Healthcare) by anion exchange chromatography (MonoQ), and gel filtration (S200). For binding studies, MCC/ $I-E^k$ was expressed, refolded, purified, C-terminally biotinylated, and immobilized to a streptavidin-coated BiaCORE chip (GE Healthcare) as described (47).

Results

Structural overview

We determined the crystal structures of the 2B4 TCR bound to MCC/I-E^k (deposited at PDB [http://www.pdb.org], PDB ID 3QIB), the 226 TCR bound to MCC/I-E^k (PDB ID 3QIU), and the 226 TCR bound to p5E-MCC/I-E^k (PDB ID 3QIW) at a resolution of 2.7, 2.7, and 3.3 Å, respectively (Fig. 1, Supplemental Table I). We also determined structures of the unbound 2B4 (PDB ID 3QJF) and 5c.c7 (PDB ID 3QJH) TCRs (Supplemental Table I). On a gross level, the three bound structures are very similar in that they have buried surface areas of 2605, 2500, and 2494 Å², respectively, and all have a common TCR diagonal docking angle on the pMHC of ~41° (48). The 2B4 and 226 TCRs have very similar V region segments, using Vα11.2 and Vα11.1, respectively (Supplemental Table II). They also use the same V β 3 gene segment; however, they have dramatically different CDR3a sequences and moderately different CDR3ß sequences, partially as a consequence of their different J α and J β gene segments. As a large number of TCRs recognizing MCC/I-E^k have been sequenced and characterized, when comparing these TCRs, it is clear that 226, like 5c.c7, represents an example of the canonical MCC/I-E^k-reactive TCR in that it displays all eight previously identified preferred features of an MCC/I-E^k-reactive TCR (Table I). In contrast, 2B4 displays only three out of the eight possible preferred features. Aside from the use of $V\alpha 11$, the noncanonical nature of this TCR is particularly evident for the α -chain, which has no other previously identified features. Despite the large junctional differences between these TCRs, both 2B4 and 226 TCRs bind to MCC/I-E^k in a similar manner. Furthermore, the recognition of a charge-reversed MCC (MCC-Kp5E) led to very little change in the overall docking orientation of 226 to MCCp5E/I-E^k compared with the structure of this TCR bound to the wild-type $MCC/I-E^k$ complex.

Three TCRs with very different mechanisms of peptide recognition

The CDR3 interactions with the peptides in the three bound structures are illustrated in Fig. 2. Peptide recognition in each of these complexes is achieved by remarkably different chemistries. The main similarities between all three structures are the hydrogen bonds (H-bonds) made by the CDR3_{β3}-N97 side-chain amide with the backbone of the peptide and between the MCC-T8 sidechain hydroxyl with the backbone carbonyl of CDR3ß position 98 (N in 226, W in 2B4). This is not surprising given the conservation of sequences in this region. Contacts made by CDR3a differ the most among all three structures. Only 2B4 makes an H-bond with the side chain of MCC-Y3 using the backbone carbonyl of aG93 (Fig. 2A). Also, only 2B4 makes a β-chain-mediated (CDR3 β D101) salt-bridge with the p5K position of MCC and makes an H-bond with the backbone carbonyl of p5K (via βS99, Fig. 2A). In contrast, 226 uses α E89 to salt-bridge with MCCp5K and makes only one other H-bond using CDR3 α (Fig. 2B), whereas the recognition of MCC-p5E by 226 is completely lacking in polar contacts using CDR3 α (Fig. 2C).

Consistent docking mode despite large changes in CDR3 recognition

As described above, peptide recognition of $MCC/I-E^k$ by the 2B4 and 226 TCR CDR3 loops is accomplished by different strategies. However, superimpositions of the three bound structures solved in this study show nearly identical docking with the MHC and precise superimposition of the germline-encoded HV4 loops (Fig.

FIGURE 1. Structural overview of the three TCR–pMHC extracellular domain complex structures solved in this study. For each, the TCR is a chimeric fusion of the mouse variable and human constant domains (α in light blue, β in dark blue) bound to the mouse MHC protein I-E^k (α in dark yellow, β in light yellow). *A*, The 2B4 TCR bound to MCC/I-E^k. *B*, The 226 TCR bound to MCC/J-E^k. *C*, The 226 TCR bound to MCC-p5E/J-E^k.

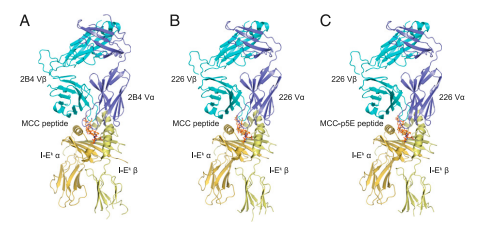


Table I. A list of preferred features of TCRs specific for MCC/I-E^k

No.	Feature	2B4	226	5c.c7
1	$\alpha 89E$ (prev. $\alpha 93E$)	N (L)	Y	Y
2	8 aa CDR3α	N (10)	Y	Y
3	$\alpha 91S$ (prev. $\alpha 95S$)	N (A)	Y	Y
4	Jα 16, 17, 22, 34	N (56)	Y (16)	Y (34)
5	β97N (prev. β100N)	Y	Y	Y
6	9 aa CDR3β	Y	Y	Y
7	β99A/G (prev. β102A/G)	N (S)	Y (A)	Y (A)
8	Jβ1.2, Jβ2.5	Y (2.5)	Y (1.2)	Y (1.2)
	No. of preferred features	3	8	8

A large number of studies on the evolution and properties of in vivo T cell responses to MCC/I- \mathbf{E}^k have identified and subsequently relied on the strong bias for usage of eight common preferred features of MCC/I- \mathbf{E}^k -reactive TCRs. These eight features are listed above. For all TCR residue numbers, we are now using the ImMunoGeneTics numbering system employed for numbering residues in all deposited PDB files. Residue numbers from previous cytochrome *c*-reactive TCRs are provided parenthetically.

3A). Inspection of individual contacting residues within the CDR1 α , HV4 α , CDR2 α (Fig. 3B), and CDR2 β (Fig. 3C) loops shows that, at most, very minor rearrangements are made between these three structures in side-chain conformation. Previous comparisons between several structures using the same V regions, such as the comparisons of many VB8.2-containing structures (30-32), have noted the striking conservation of contacts formed by tyrosine residues, namely Tyr⁴⁸ in V β 8.2 CDR2. For V α 11, it is possible that Tyr⁴⁷ in CDR2 plays a similar role in that it is highly malleable and capable of forming a variety of different type of contacts, from polar to hydrophobic (Fig. 3B). None of the VB3-encoded contacts use aromatic residues, nor do any of these contacts appear to be playing a dominant role in MHC binding, as only a few van der Waals interactions and solvent exposed saltbridge are seen. In contrast to the V α -encoded contacts, we observe a small amount of wobble in the CDR2B contacts when comparing the 226-pMHC structures to the 2B4-pMHC structure (Fig. 3C).

Although it is striking that large differences in CDR3 do not require more significant adjustments to the germline-encoded contacts, it may not be necessarily surprising that these germline-encoded contacts, of which some may be conserved codons of recognition, strictly impose a docking mode for pMHC recognition for this pairing of V α , V β , and MHC allele. It is also possible that different V α and V β pairings may vary in their ability to accommodate multiple docking modes.

Germline-encoded contacts and dominance of $V\alpha$ over $V\beta$

If it is true that this pairing of V α 11 and V β 3 genes imposes a strict docking mode, does one V region predominate? Several metrics point at V α 11's dominant role in contacting the I-E^k allele in this very specific way. For one, all three germline-encoded loops (CDR1 α , CDR2 α and HV α 4 of V α 11) are making a large number of contacts of all types with the MHC (Fig. 3B, Supplemental Fig. 1, burying a total of 1045, 990, and 978 Å² of surface in each of the structures). In contrast, the only V β 3-encoded loop making a few contacts is CDR2β (Fig. 3C, Supplemental Fig. 1A, 1C, burying only 451, 447, and 433 Å² of surface in each of the structures). This skewing toward Va contacts in these structures is further illustrated by comparing the V region contacts made in another mouse MHC class II-restricted (MBP/I-A^u) TCR, 1934.4, which has relatively equal contributions from V α - versus V β encoded contacts (Supplemental Fig. 1B, 1C). Furthermore, the strictness of the requirement for each V region in the mouse response to MCC in the context of I-E^k has been addressed by looking at the V region usages of $5c.c7\alpha$ versus β hemitransgenic mice. In 5c.c7 α transgenic mice, the frequency of non-V β 3expressing T cells responding to MCC is much greater than the frequency of non-Val1-expressing T cells responding to MCC in 5c.c7 β transgenic mice. Thus, one might predict that all V α 11 TCRs selected on the I-E^k MHC are making contact in this same docking mode. It is interesting to note that a TCR composed of 5c. $c7\alpha$ paired with a Vβ8.2 is capable of recognizing p8E-MCC/I- E^k (see Supplemental Table II). One might predict that V β 8.2 is more flexible in this interaction, whereas $V\alpha 11$ is more stringent.

Structural explanation for the preferred features of $MCC/I-E^{k}$ recognition

The 226 TCR displays all of the canonical features of an MCC/ I-E^k-reactive TCR (Table I, Supplemental Table II). Although numerous mutational and sequence comparisons have identified residues critical for recognition, the precise structural basis for each of these features and sequence requirements are not known. For example, MCC peptide charge-reversal experiments leading to a reciprocal charge reversal in CDR3 α of responding TCRs

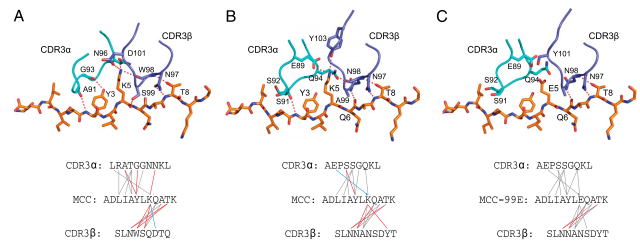


FIGURE 2. A comparison of the CDR3–peptide interactions of each TCR–pMHC complex. Only the CDR3 loops and MHC-bound peptide are considered for 2B4-MCC/I-E^k (A), 226-MCC/I-E^k (B), and 226-MCC-p5E/I-E^k (C). The *top panel* shows the arrangement of the CDR3 loops interacting with the MHC-bound peptide, shown in stick form. Side-chains involved in polar interactions are shown with the H-bonds/salt-bridges shown in red. The *bottom panel* shows the sequence of the CDR3s and the peptide. Lines connecting each residue represent van der Waals interactions (gray), H-bonds (red), and salt-bridges (blue).

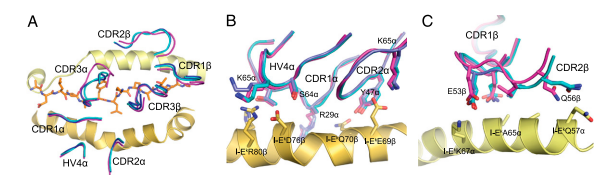


FIGURE 3. TCR docking comparison. In all panels, as in other figures, 2B4 is colored magenta, 226-binding MCC/I-E^k is cyan, and 226-binding MCCp5E/I-E^k is colored blue. *A*, The ends of each CDR and HV4 α loops are overlaid. *B*, Detailed illustration of germline-encoded TCR α contacts with I-E^k β . Relevant side-chains for each structure are overlaid to compare the interface. Note extensive contacts between all three TRAV4 (V α 11) germline-encoded loops, CDR1, 2, and HV4, and no significant difference in conformation between the three structures solved in this study. *C*, TRBV26 (V β 3) germline-encoded residues making contact with I-E^k α are compared. For these chains, only the CDR2 β loop makes contact, and some small differences are seen when comparing the 2B4-bound structure with either 226-bound structure. No differences are seen between the two 226 structures.

strongly suggested that a charge–charge interaction is taking place in the bound structure, which would explain preferred feature 1 in Table I. This is confirmed in the crystal structure, where the central p5K position of MCC forms a salt-bridge with TCR α 89E in the 226-MCC/I-E^k complex (Fig. 4*A*). There is further coordination of the positively charged amine group H-bonds from 226 α 94Q, 226 β 98N, 226 β 100N, and 226 β 103Y.

Preferred feature 2 in Table I is a conserved α 91S that makes a backbone-carbonyl H-bond with the backbone amide of MCCp3Y residue, positioned by side-chain hydroxyl H-bond with I-E^k β -T77 side-chain hydroxyl (Fig. 4*B*). In 2B4, this residue is replaced by an alanine residue, which lacks the H-bond with I-E^k β -T77. However, it does make a van der Waal's contact with this same residue (not shown).

Sequence analysis of canonical MCC (wild-type p8T)-reactive TCR β -chain sequences and comparison with MCC-p8E–reactive TCRs from 5c.c7 α transgenic mice have repeatedly shown a strong conservation of CDR3 β 97N residue usage (preferred feature 5 in Table I) (17) and consistent changes in this residue for p8E-reactive TCRs. These data strongly suggested an interaction between these two residues on the TCR and pMHC. Indeed, all three structures from this study have a consistent mode of interaction between this CDR3 β 97N residue and MCC-p8T (Fig. 2); that is, hydrogen bonding between the TCR β 97N side-chain

amide and the backbone carbonyl oxygen of MCC-p8T. At the same time, there is a precisely aligned H-bond between the backbone amide group of the following CDR3 β 98 position and the hydoxyl oxygen of MCC-p8T (Figs. 2, 4*C*). Slight alterations in this alignment in MCC-p8S peptide mutants are almost certainly the cause of the drastic reductions in binding affinity observed for all MCC/I-E^k-reactive TCRs tested thus far. Any other residue at this position is not tolerated.

In contrast, no salt-bridge is formed between the 226 TCR and the MCC-p5E residue (Fig. 4D). Instead, 226's a89E residue forms an H-bond with the nearby 226a94Q [coded by Ja16 used by the AND, 226, and 202 TCRs, all of which have reduced specificity for MCC-p5K (17, 49)], but absent from 2B4 and 5c. c7, which absolutely require p5K for stimulatory activity (15) (Fig. 4D). It is interesting to note that with the exception of $J\alpha 22$, all four preferred J α gene segments, as well as the J α 56 gene segment used by 2B4, have an N/Q residue at position 94, which can H-bond to MCC-p5K (Fig. 4A) or, in the case of 226 and J α 16's 94Q, make a H-bond with α 98E, when bound to MCC-p5E (Fig. 4D). Thus, we have provided structural data that can be used to explain at least three of the eight preferred features involved MCC peptide recognition. These hypotheses as well as hypotheses for the structural basis for the remaining five related preferred features are summarized in Table II.

FIGURE 4. Illustration of the conserved features of peptide recognition. The structural basis for each of the preferred sequence features of MCC/I-Ek reactive TCRs is illustrated in detail. A, The structural basis for recognition of MCC-p5K is illustrated. Several Hbonds coordinate the positive charge of this central lysine in addition to the predicted salt-bridge between α89E and p5K. B, The interactions made between 226 TCR α 91S and MCC/I-E^k are shown. This residue hydrogen bonds both the peptide at p3Y as well as I-Ek at β77T. C, Reciprocal side-chain and main-chain Hbonds are shown for TCR B98N and the peptide p8T. D, The arrangement of p5K hydrogen-bonding 226 CDR3a residues is shown in the p5E-bound structure. E, The salt-bridge between MCC-p5K and the J β 2.5encoded aspartate of 2B4 is shown together with additional charge-coordinating H-bonds.

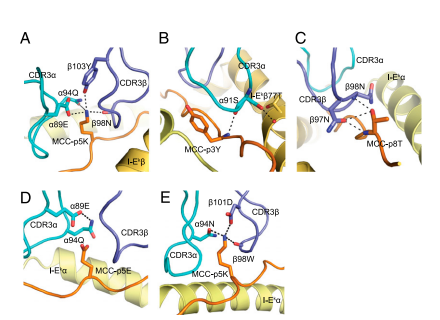


Table II. Structural hypotheses explaining conserved features of MCC/I-E^k reactive TCRs

No.	Feature	Structural Explanation
1	α89E	Forms salt bridge with central MCC-p5K residue (Fig. 4A).
2	8 aa CDR3α	Allows appropriate spacing for germline encoding of Glu at position 89 by TRAV4 and germline encoding of N/Q at position 94 by the preferred J α (feature 4).
3	α91S	Makes a backbone-carbonyl H-bond with the backbone amide of MCC-p3Y residue, positioned by side-chain hydroxyl H-bond with I-E ^k β -T77 hydroxyl side-chain (Fig. 4B).
4	Jα 16, 17, 22, 34	Provide N/Q residue in position for H-bonding central MCC-p5K residue (Jα16 encoded Q appears to be required for MCC-p5K/E cross-reactivity; Fig. 4A, 4D).
5	β97N	An aligned pair of H-bonds of β 96N side-chain amide with MCC-p8T backbone carbonyl and MCC-p8T side-chain hydroxyl with β 95 backbone amide (Fig. 4 <i>C</i>).
6	9 aa CDR3B	
7	β99A/G	H-bond with I-E ^k β70Q and van der Waals interactions with I-E ^k β67F. Small to leave room for MCC-99K to protrude. β98A is at least partially coded by Jβ1.2.
8	Jβ1.2, Jβ2.5	In Jβ1.2 there is a Tyr at the appropriate position (β103Y) for H-bonding MCC-p5K as in 226. In Jβ2.5, there is an Asp residue in place to make a salt-bridge with MCC-p5K as in 2B4 (Fig. 4A).

Based on crystal structure information, we can now hypothesize the structural bases for each of the well-characterized preferred sequences features of MCC/I-E^k-reactive TCRs. These features are described in more detail in the *Results* section.

How then does 2B4 bind to the same positively charged residue, as there is no corresponding negatively charged residue in 2B4? In fact, this salt-bridge is replaced by a salt-bridge between MCC-p5K and 2B4 β D101 coded by J β 2.5 that is further coordinated by H-bonds with 2B4 α N95, 2B4 β W98, and 2B4 β S99 (Fig. 4*E*).

CDR3 flexibility and $MCC/I-E^k$ recognition

The unliganded structures of two MCC/I-E^k–reactive TCRs, 5c.c7 TCR and the 2B4 TCR, were also determined in this study. The CDR loop configurations of these two TCRs are overlaid with the MCC/I-E^k–bound version of 2B4 in Fig. 5A. As had been seen for other TCR structures, we see very similar CDR1 and -2 conformations, but quite different CDR3 conformations. For 5c.c7, the unliganded CDR3 loops would require large conformational changes to assume the structure expected for MCC/I-E^k binding, as is seen for its close relative 226. For 2B4, we see further evidence for CDR3 loop flexibility in that the majority of both CDR3 α and β could not be resolved in the unbound structure. Thus, as has been seen for several other TCRs (31, 50–52), both 5c.c7 and 2B4 TCRs appear to have very flexible CDR3 loops.

To address the role of this CDR3 loop flexibility in T cell recognition, we made a series of proline substitutions in the 2B4 TCR throughout both CDR3 loops. Proline has a cyclic side chain in which a backbone dihedral angle is fixed at approximately -75° , giving it unusual conformational rigidity compared with other amino acids. We postulated that if a given proline substitution did not destroy 2B4's specificity, it might increase its affinity by freezing a CDR3 loop into its optimal conformation for binding MCC/I-E^k, similarly to the way somatic mutation rigidifies CDR loops of Abs (53). To test for the ability of the proline mutant TCRs to bind MCC/I-E^k, we expressed each mutant retrovirally in $58\alpha^{-}\beta^{-}$ hybridomas and tested for the ability of these 2B4 TCR expressing hybridomas to produce IL-2 in response to MCC stimulation (see Materials and Methods). In this manner, 17 CDR3 proline mutants were tested, and 5 were found to maintain $MCC/I-E^{k}$ reactivity (substituted positions are shown in Fig. 5*B*). These positions were not critical peptide contacts in the CDR3, but rather appear to play roles in supporting the backbone conformation of the CDR3s. Three out of the five had higher MCC sensitivity than the wild-type 2B4 TCR (Fig. 5C). We then made rTCR proteins for selected binding mutants. The affinities of these TCRs for MCC/I-E^k correlated well with Ag sensitivity seen for the hybridomas (Fig. 5D). These results indicate that 2B4 can accommodate rigidifying proline mutations and maintain MCC/ $I-E^{k}$ reactivity. In these cases, the affinity of the interaction can be improved, likely due to reduced entropic costs for binding. We did not observe any narrowing of fine specificity for recognition of MCC altered peptide ligands (not shown), which is consistent with the notion that gross CDR3 conformational changes are generally not always required for degenerate recognition of altered peptide ligands (see *Discussion*).

226 is a naturally occurring high-affinity, highly cross-reactive TCR

As described above, the 226 TCR was originally identified for its ability to react with both MCC-p5K/I-E^k and MCC-p5E/I-E^k, accommodating a charge reversal at a well-characterized critical TCR contact residue. To compare the affinities of each of these TCR-pMHC interactions, we subjected the soluble 226 and 2B4 TCR proteins to SPR analysis using SPR. Surprisingly, we found that 226 had an unusually high affinity for MCC/I-E^k at 25°C $(K_{\rm D} = 321 \text{ nM})$ with very slow off-rate $(k_{\rm off} = 0.0029, t_{1/2} = 239 \text{ s};$ Fig. 6A, 6B). In contrast, the values we obtained for 2B4 were similar to those previously described (54) ($K_D = 13.7 \mu M$, $k_{off} =$ 0.032, $t_{1/2} = 21$ s). For 226 binding to MCC-p5E/I-E^k, we were also surprised to find that the binding was detectable (not shown) but extremely weak ($K_D > 500 \mu$ M, Fig. 6B). Thus, our use of tethered peptide strategy to crystallize 226 bound to MCC-p5E/ I-E^k may represent one of the lowest affinity TCR-pMHC cocrystals solved to date and implies that this strategy could be used to crystallize other low-affinity TCR-pMHC interactions such as positive-selecting or null peptides. To assess the extent of 226's degeneracy for recognition of MCC TCR contact variants, IL-2 production by 226 hybridomas was used as readout. As positions 3, 5, and 8 had been previously described as critical for 5c.c7 and 2B4 recognition of $MCC/I-E^k$ (15), we tested all 20 possible aa substitutions at each of these three positions. In contrast to 2B4 or 5c.c7, which have strict requirements at these three positions. 5c. c7 reacts well with F, W, or Y and weakly with P or C at p3, absolutely requires K at p5, and reacts well with only with T and weakly with V, S, W, H, or R at P8. 2B4 reacts well with F or Y and weakly with I, W, or C at p3, absolutely requires K at p5, and only reacts well with T and weakly with W, N, or S at p8. See figures 7 and 8 in Reay et al. (15). 226 reacts with more than half of the possible substitutions at all three of these positions individually. Given 226's very weak recognition of the p5E variant of MCC, it is not surprising that this is an example of a weak agonist for 226 stimulation, despite the fact that 226 was originally cloned for its ability to recognize this peptide. It is also interesting to note that some substitutions such as MCC-p3F,

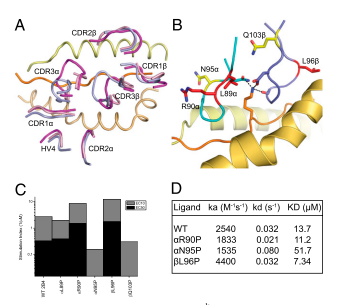


FIGURE 5. CDR3 flexibility and MCC/I-E^k recognition. A, In addition to the three bound structures, the unbound versions of both 2B4 and 5c.c7 TCRs were solved. These two unbound TCRs, 2B4-free in pink and 5c.c7free in blue/purple, are aligned and overlaid with the bound 2B4 TCR in magenta. Note minimal conformational differences in the CDR1, 2, or HV4 loops. In contrast, large differences were seen for the CDR3 loops. In the case of the 2B4-free structure, neither loop could be resolved due to large b-factor probably caused by high mobility in the crystal. B and C, To reduce the mobility of 2B4 CDR3s, proline substitutions were made throughout these loops. These proline-mutant TCRs were then expressed in $58\alpha^{-}\beta^{-}$ T cell hybridomas and tested for their ability to respond to MCC peptide stimulation in the presence of CH27 APCs (see Materials and Methods). The five residues able to accommodate proline substitution and maintain MCC reactivity are highlighted in B. Three out of these five TCR mutants had increased MCC sensitivity and are colored red; the other two had reduced sensitivity and are colored yellow. The sensitivity of each mutant-TCR-expressing hybridoma is shown in C. Stimulatory index is plotted for each hybridoma and represents the inverse peptide concentration required to stimulate 10 (EC10, gray bars) and 50% (EC50, black bars) of maximal IL-2 secretion (see Materials and Methods) (15). D, Three of the five MCC-reactive proline mutant TCRs were expressed as soluble proteins and tested for their ability to bind MCC/I-E^k in solution using SPR. Consistently with the hybridoma stimulation data, the two more sensitive TCRs had a higher affinity (lower K_D) and/or slower offrate (lower k_{off}) than the wild-type 2B4 TCR, whereas the TCR displaying weaker responses to MCC peptide had a much lower affinity and faster offrate than wild-type 2B4.

-p5L, and -p8S were more potent than wild-type MCC (p3Y, p5K, p8T) for stimulating 226 even though 226 already binds MCC with submicromolar affinity, suggesting that in some instances, weaker binders may sometimes provide for a more optimal stimulus [as has been observed for AND (49); see *Discussion*]. Overall, these data indicate that 226's very high affinity for MCC allows it to recognize a large number of MCC variants, such as MCC-p5E, albeit at a much lower affinity.

Discussion

Since Schwartz and colleagues (2, 17) first developed it in the late 1970s, the murine CD4⁺ T cell response to the C-terminal portion of PCC has become one of the best-studied systems in immunology. Many important studies of T cell specificity, MHC restriction (1–3), gene structure (55), thymic selection (56–58), biochemistry (4, 39, 54, 59, 60), thermodynamics (47, 61, 62), and cellular biophysics (63–65) have used it (14, 49). A major problem however, has been the lack of any structural data on the TCRs or

One of these longstanding issues has been the lack of any structural information concerning the findings of Jorgensen et al. (17), who reported that 5c.c7ß TCR transgenic mice immunized with a cytochrome c peptide with a charge reversal (Lys to Glu) at the p5 position elicited a reciprocal charge reversal (Glu to Lys) on CDR3 α at position 89. In this study, we can confirm that, as proposed in Jorgenesen et al. (17), the Glu residue of the closely related 226 TCR is involved in a salt-bridge with the central lysine (p5K) of the MCC peptide. We also found that this same central lysine is further coordinated by a series of H-bonds with several other residues on both CDR3 α and β . Interestingly, we find the situation to be quite different from the 2B4 TCR-MCC/I-E^k complex, where a negatively charged residue (BD101) of the CDR3 β forms a salt-bridge with the p5 lysine of the peptide. This is a further indication of the importance of charge-charge interactions in TCR recognition and also the shows the flexibility of these related TCRs in the way that their CDR3 residues can interact with the peptide.

As several studies from our laboratory and others aimed at understanding the biophysical basis for TCR recognition of pMHC have used the cytochrome system as a model system, there is large amount data that can now be reconciled based on the crystal structure of this interaction. For instance, mutagenesis studies aimed at identifying MHC residues involved in TCR contacts have used IL-2 production as readout for pMHC recognition by 2B4 and 226 T cell hybridomas (16, 17). Purified 2B4 TCR binding to MCC/I-E^k with various MHC and peptide residue substitution has also been investigated (4, 47, 62). In one such study, the kinetics of 2B4 binding to each of these altered ligands was used to determine the relative contribution of various residues to the binding transition state, using φ analysis (47). The residues making contact in the bound structures we have solved in this paper are all consistent with these data. That is, residues previously found to influence either T cell recognition or TCR binding are all making contacts in the appropriate structure (Supplemental Fig. 2 for graphical summaries of these data). It is interesting to note that only some of the residues found to be involved in the transition state are making contacts in the bound structure, consistent with the interpretations of Wu et al. (47) (Supplemental Fig. 2C). One residue (I-E^k α -K67), which has one of the highest φ values (and thus contributes mostly to the transition state), is involved in a solvent-exposed salt-bridge among 2B4-CDR2B-E53. Such a charge-charge interaction may help guide binding, but it is not surprising that it does not contribute significantly to the bound state, as it is solvent exposed. Particularly in the 2B4-MCC/I-E^k structure, these residues had high B factors, indicating that this interaction is not very stable.

Another goal of this study was to follow up the intriguing findings of Ehrich et al. (16), who used a panel of $I-E^k$ mutants to characterize the MHC footprint of the various cytochrome *c*-specific TCRs, including 226. In this study, they found that 226 had a very different sensitivity to the $I-E^k$ mutants when it recognized the wild-type MCC-p5K versus MCC-p5E. These different patterns of sensitivity suggested that 226 was binding in very different orientations depending on the ligand. But when we examine the structure of 226 complexed with either of these two ligands, we find no difference in the orientation of the TCR or in the way it binds $I-E^k$. What we do find is that 226 has a much

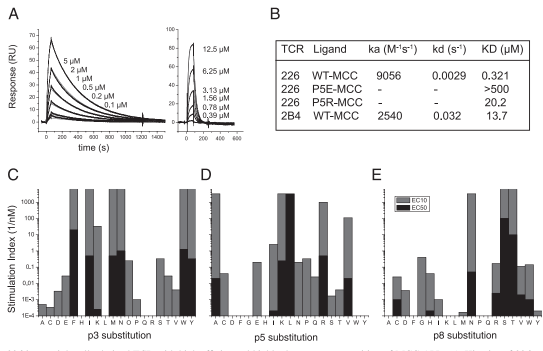


FIGURE 6. 226 is a peripherally derived TCR with high affinity and highly degenerate recognition of MCC-APLs. *A*, Kinetics of 226 versus 2B4 TCR binding are compared. Background subtracted binding SPR sensograms for injections of 226 TCR at 0.1, 0.2, 0.5, 1, 2, and 5 μ M over MCC/I-E^k– immobilized surfaces are shown in comparison with 2B4 TCC applied at 0.39, 0.78, 1.56, 3.13, 6.25, and 12.5 μ M in a different experiment. The 2B4 and 226 responses are plotted with the same scale time-axis. *B*, Kinetic parameters from these sensorgrams as well as those obtained from 226 applied to flow cells containing MCC-p5E and MCC-p5F/I-E^k are tabulated. 226 T cell responsiveness to MCC APLs at peptide positions 3 (*C*), 5 (*D*), and 8 (*E*) were tested using 226 hybridomas and CH27 cells in an IL-2 secretion assay. For each peptide, the inverse peptide concentration required to stimulate 10 (EC10, gray bars) and 50% (EC50, black bars) of maximal IL-2 secretion is plotted (see *Materials and Methods*).

higher affinity, >1000-fold, for MCC/I-E^k than for the p5E variant. Thus, it seems that this very large difference makes the 226 TCR much more tolerant of many of the MHC mutants when MCC is the peptide compared with the p5E variant (16). As to why 226 has such lower affinity for MCC-p5E, this can be explained by the lack of a number of TCR peptide contacts in the complex structure. In particular, the glutamate at p5 does not make any clear contacts with the TCR. Interestingly, the remaining TCR-pMHC interactions are maintained in the same configuration as the 226-MCC-p5K/I-E^k, indicating that the TCR does not use a completely different binding mode for recognition of these two ligands. Thus, even with poor quality peptide contacts, the binding footprint of the 226 TCR is essentially the same with both ligands and the same as the 2B4 TCR, suggesting that, at least with these closely related $V\alpha V\beta$ heterodimers, this particular MHC binding site is highly favored.

Also reported by Ehrich et al. (16), the effect of $I-E^k$ mutations on 226's recognition of MCC-p5K/I-E^k versus MCC-p5E/I-E^k fell into three categories. In one category, I-E^k mutations caused decreased 226 IL-2 production when either peptide was presented (I- $E^{k}Q\beta 64R$, I- $E^{k}T\beta 77Q$). This category of residues represents residues that are involved in 226's binding of both pMHC complexes. It is interesting to note that neither of the I-E^k mutations completely abolished 226's ability to respond to MCC-p5K/I-E^k, consistent with the notion that 226's very high affinity for MCCp5K/I-E^k allows it to accommodate less than ideal ligands. A second category of residues caused an increase in IL-2 production only for MCC-p5E presentation (I-E^kAα65V, I-E^kAα68V). This category could be explained if these two mutants caused an increase in affinity in either case. No change could be detected for MCC-p5K recognition because the affinity for this complex is already extremely high. The final type of mutant was observed for I-E^kHβ81Y, which increased 226's sensitivity for MCC-p5K and decreased it for MCC-p5E. This phenotype can be explained if 226's sensitivity for MCC-p5K goes up when affinity is decreased. We think this is the case, as it can also be seen for several of the MCC peptide APLs (Fig. 6C); p3F, P5L, and p8S all had higher potency for 226 stimulation than p5K, but it is unlikely that these pMHC complexes would be higher affinity for the 226 TCR. Together, we take this as evidence for an incomplete detrimental effect of having TCR–pMHC interactions with too high an affinity. This effect has been seen in the AND system (49) and in vivo for the 5c.c7 system (14). However, it is also clear that there is no absolute ceiling on the functional affinity TCRs have been shown to be functional when expressed in hybridoma T cell lines in vitro (66, 67).

As is clear from the structures presented in this study, the 226 TCR β -chain (same TCR β as 5c.c7) also contributes to the coordination of the central p5 lysine of MCC. Thus, immunization of 5c.c7 β hemitransgenic mice with MCC-p5E selected for TCR α -chains able to overcome a less than ideal TCR β -chain sequence in order recognize this Ag, albeit very weakly ($K_D > 500 \mu$ M). Consequently, the 226 TCR also has very high affinity for MCC-p5K/I-E^k, making it a robust responder to substitutions in contacting residues in the peptide and the MHC. As we have described in this paper, this TCR, which may be highest affinity and the most cross-reactive of the naturally occurring TCRs described to date, appears to accomplish this with very canonical peptide and MHC docking and without significant conformational changes.

Several lines of evidence suggest that cross-reactive T cells may underlie at least some forms of autoimmunity (68, 69). Indeed, there are clear examples of autoreactive TCRs that are also capable of recognizing viral (70) and bacterial peptides (71). Although there is no evidence that 226 is autoreactive in these mice, it is one of the few examples in which we have structural information of a naturally occurring dramatically cross-reative TCR, and so it should help us understand this phenomenon. Like the known autoimmune epitopes, the affinity of 226 for MCC-p5E/I- E^{k} is extremely low (72, 73) and is stimulatory despite suboptimal contacts lacking H-bonds between the TCR and the peptide (as reviewed in Ref. 74). Furthermore, a structure of the MBP-specific Ob.1A12 TCR (a DR2-restricted TCR clone) in complex with a bacterial mimitope shows nearly identical docking as it has with the MBP/HLA-DR2b complex (37, 71), as is seen in comparison of 226's recognition of both p5K- and p5E-MCC. It is also interesting to compare the mechanism of peptide degeneracy for this TCR to structures of previously identified degenerate and autoreactive TCRs. For instance, an extremely cross-reactive TCR isolated from mice expressing 3K-peptide-tethered/I-A^b (29), the Yae62 TCR, was later found to have reduced peptide contacts and a somewhat exceptional docking mode dominated by I-A^b α α -helical contacts (32). In contrast, in an engineered high-affinity version of the 2C TCR, M6, which also has dramatically reduced peptide specificity and endogenous-peptide reactivity (27), docking with the MHC was nearly the same as wild-type 2C, and the affinity gains came from modest rearrangements of CDR3 α , enhancing hydrophobic contacts with the peptide (75). Thus, in the case of the 226 TCR presented in this study, like the M6 TCR, 226's MHC contacts are unchanged compared with its nondegenerate counterpart, 2B4. Similarly modest differences in CDR3a-mediated contacts are capable of dramatically increasing TCR affinity for MCC/I-E^k, allowing the recognition of a wide range of MCC-peptide variants.

In summary, after many years of effort, we have obtained structures of two cytochrome *c*-specific TCRs complexed to two different ligands. These structures have resolved and explained the results of a large number of cellular, molecular, and biochemical studies. They also represent the first structural examples of I-E^k–reactive TCRs, the first structures of the V α 11 and V β 3 TCR gene segments, and, in the case of 226, the highest affinity and most degenerate of the naturally occurring TCRs identified to date. We have found the CDR3 regions of these TCRs can recognize peptide variants of MCC in quite different ways, despite strict constraints imposed by large numbers of V α 11-mediated conserved contacts with the MHC.

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Disclosures

The authors have no financial conflicts of interest.

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