Structural Basis of Cytochrome c Presentation by IE^k

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Abstract

The COOH-terminal peptides of pigeon and moth cytochrome c, bound to mouse IE^k , are two of the most thoroughly studied T cell antigens. We have solved the crystal structures of the moth peptide and a weak agonist–antagonist variant of the pigeon peptide bound to IE^k . The moth peptide and all other peptides whose structures have been solved bound to IE^k , have a lysine filling the p9 pocket of IE^k . However, the pigeon peptide has an alanine at p9 shifting the lysine to p10. Rather than kinking to place the lysine in the anchor pocket, the pigeon peptide takes the extended course through the binding groove, which is characteristic of all other peptides bound to major histocompatibility complex (MHC) class II. Thus, unlike MHC class I, in which peptides often kink to place optimally anchoring side chains, MHC class II imposes an extended peptide conformation even at the cost of a highly conserved anchor residue. The substitution of Ser for Thr at p8 in the variant pigeon peptide induces no detectable surface change other than the loss of the side chain methyl group, despite the dramatic change in recognition by T cells. Finally, these structures can be used to interpret the many published mutational studies of these ligands and the T cell receptors that recognize them.

Key words: T cell receptor • X-ray crystallography • antigen presentation • peptide • cytochrome

Introduction

One of the most heavily studied antigenic peptides in mice comes from the COOH-terminal end of pigeon cytochrome c (PCC),* encompassing amino acids 89–103 (peptide derived from the COOH terminus of PCC [pPCC]). This peptide contains the dominant pigeon cytochrome c epitope for CD4⁺ T cells in many strains of mice (1). Over the years, many immunological principles have been defined and studied using this peptide and its homologues from the cytochrome c's of other species, especially the equivalent peptide from moth cytochrome c (pMCC).

In H-2^k or H-2^a mice, the IE^k MHC class II molecule (MHCII) presents pPCC and pMCC to T cells. The phenomenology associated with the response to these peptides led to the discovery of two complementing "immune response" genes that later turned out to be the structural genes for the α and β chains of the IE^k molecule (2). Mutational studies have predicted which of the pPCC and pMCC amino acid side chains are important for the binding of the peptides to the IE^k molecule (3, 4). An important difference between pMCC and pPCC is the fact that pPCC has an Ala inserted at amino acid 103. This insertion causes pPCC to raise a "heteroclitic" response to the pMCC, which means that although pPCC and pMCC are virtually completely cross reactive, essentially all T cell clones raised by immunization with pPCC respond better to pMCC (5). The data have suggested that this phenomenon is linked to the interaction of the peptide with the MHC molecule rather than the $\alpha\beta$ TCR, but the structural basis of this difference has remained unknown (6).

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^{*}Abbreviations used in this paper: CPK, Corey, Pauling, Koltan; MHCI, MHC class I; MHCII, MHC class II; pMCC, peptide derived from the c-terminus of moth cytochrome c; PCC, pigeon cytochrome c; pPCC, peptide derived from the COOH terminus of pigeon cytochrome c.

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The repertoire of $\alpha\beta$ TCR expressed by T cells responding to pPCC and pMCC is dominated by receptors bearing AV11S1 and AV11S2 (Va11.1 and Va11.2, respectively) and BV3S1 (V β 3) (7). Many V α 11/V β 3⁺ T cell clones/hybridomas specific for these peptides are in widespread use (8–10) and the genes for the $\alpha\beta$ TCRs of a number of these clones have been used to produce $\alpha\beta$ TCR-transgenic mice (11–14). These mice, as well as those transgenic for various versions of the IE^k molecule (15, 16), have been used over the years to study the principles of T cell activation as well as positive and negative selection in the thymus. The interaction of $\alpha\beta$ TCRs with IE^k/pPCC or IE^k/pMCC has been extensively studied both in vitro and in vivo. These studies have predicted which of the $\alpha\beta$ TCR, IE^k, and peptide amino acids are important in the interaction (4, 6, 17–21).

pPCC and pMCC have also been used to study the phenomenon of antagonism. Amino acid variants of a number of antigenic peptides have been identified, which not only fail to stimulate particular T cell clones or hybridomas that had been raised to the wild-type peptide, but also render the T cell unresponsive to a subsequent or simultaneous challenge with the wild-type peptide (22). Although the mechanism of this phenomenon is not fully understood, it has been shown to involve some type of aberrant or incomplete receptor signaling that effectively desensitizes the receptor to subsequent ligation. In the case of pPCC and pMCC, several clones were antagonized by variant peptides bearing only slight differences from the wild-type peptides (23, 24)

In the current study, we have solved the crystal structure of the IE^k molecule in complex with either pMCC or pPCC bearing a single amino acid change (Thr to Ser at amino acid 102) that renders it either a poor agonist or an antagonist for many T cell clones. The structures reveal the basis for the differences between pMCC and pPCC in binding to IE^k. They also show that the Thr to Ser change results in only a very minor surface difference between the two molecules. Finally, the structures support many of the predictions made in previous studies concerning contacts between the peptides and either the IE^k molecule or the T cell receptor.

Materials and Methods

IE^k–pMCC and IE^k–pPCC(T102S). The expression, production, and purification of soluble IE^k with covalently attached peptides have been previously described (25–27). In this study two different peptides were used: pMCC, RDLIAYLKQATK, corresponding to amino acids 92–103 of moth cytochrome c, and pPCC(T102S), RDLIAYLKQASAK, corresponding to amino acids 92–104 of pigeon cytochrome c, with the exception that the Thr at amino acid 102 was changed to Ser. In discussing the structure of individual peptide amino acids when bound to IE^k, we use the convention of designating the first peptide anchor amino acid as p1 and number the other amino acids accordingly, e.g., p9Lys or p9K. IE^k amino acids are designated by chain and number, e.g., β9Glu or α66Asp. Also, for simplicity, MHCII covalent peptide complexes are referred to with a (–) connecting

the MHC and peptide name, e.g., IE^k -pMCC, whereas nonco-valent complexes use a (/), e.g., IE^k /pMCC.

Crystallization. Both IE^k–pMCC and IE^k–pPCC(T102S) were crystallized by the hanging drop method. IE^k–pMCC crystals were grown in 1- μ l drops containing ~11 mg/ml of protein, 12% PEG-4000, 12% 2-propanol, and 60 mM of sodium acetate buffer at pH 5.5. IE^k–pPCC(T102S) crystals were grown in 1- μ l drops containing ~10 mg/ml of protein, 14% PEG-4000, 14% 2-propanol, 1.4% ethylene glycol, 70 mM of sodium acetate buffer at pH 5.0, and 320 mM of ammonium sulfate.

Data Collection. IE^k-pMCC and IE^k-pPCC(T102S) crystals were cryopreserved by briefly soaking in mother liquor with the addition of 20% ethylene glycol and 10% glycerol followed by rapid cooling in a gaseous stream of nitrogen at 100°K (Oxford Cryosystems). Diffraction data for a single IE^k-pMCC crystal were collected in the X4A beamline at the National Synchrotron Light Source at Brookhaven National Laboratory using a wavelength of 0.9795 Å and PhosphorImager plates with 2° oscillations. Data were processed with Denzo and scaled/merged with SCALEPACK (HKL Research, Inc.; reference 28). Diffraction data for a single IE^k-pPCC(T102S) crystal were collected in the crystallography facility at the Howard Hughes Medical Institute using a wavelength of 1.54 Å, a Rigaku rotating anode generator, Yale mirrors, and a RAXIS IV detector with 1° oscillations. Data were processed, scaled, and merged using the BIOTEX program (Molecular Structure Corp.). Both of the crystals were of space group C2 with similar dimensions and packing, and both had two molecules in the asymmetric unit. Details of the collection and processing statistics are in Table I.

Structure Determination. The structures of IEk-pMCC and IE^k-pPCC(T102S) were solved by molecular replacement using the program AMoRe (29, 30). The search model was the IE^kpHB structure with the peptide and linker removed (31). The very low resolution data (>10 Å) from the IE^k-pPCC(T102S) dataset were omitted due to problems with saturation. The initial models were improved with rigid body refinement using the program Xplor/CNS (32). Multiple rounds of refinement with Xplor/CNS and model building with oxygen (33) were used to improve the model when the peptides were added. The final model of IE^k-pMCC had 792 amino acids and that of IE^kpPCC(T102S) had 794 amino acids. As in the case of the other IE^k structures that we have solved, relatively weak electron density was seen for most of the linker attaching the peptide to the β chain NH₂ terminus, and for the loop connecting the first and second β strands of the β 2 domain. In the later stages of refinement, the following N-acetyl glucosamines were added to some asparagines based on positive electron density in Fo-Fc maps: six to the IE^k-pMCC structure and four to the IE^k-pPCC(T102s) structure. In addition, the following water molecules were added: 215 to the IE^k-pMCC structure and 317 to the IE^kpPCC(T102s) structure. Refinement and final model statistics are listed in Table I. For figures, average structures were calculated for both molecules from the two molecules in the asymmetric units using the programs MOLEMAN and AVEPDB (34, 35). Coordinates have been submitted to the Protein Data Bank (available at http://www.rcsb.org/pdb/ under accession numbers 1KT2 and 1KTD).

Results and Discussion

The Peptide Motif for Binding to IE^k . In previous studies (31, 36), we reported that the structure of the IE^k molecule

Table I. Data Collection	and Refinement Statistics
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Data collection	рМСС	pPCC(T102S)
Space group	C2	C2
Unit cell dimensions (Å)	a = 147.8, b = 56.9, c = 116.0	a = 145.5, b = 57.3, c = 116.6
Unit cell angles (°)	$\beta = 92.1$	$\beta = 94.2$
No. of molecules in AU	2	2
Resolution limits (Å)	20-2.8 (2.95-2.80) ^a	100-2.4 (2.56-2.40)
Unique reflections	22,827 (3,185)	36,433 (5,970)
Completeness (%)	95.2 (93.4)	95.2 (90.2)
Averave redundancy	2.5	3.1
Average I/ σ	11.1 (2.4)	16.8 (2.8)
R _{merge} (%) ^b	9.7 (31.7)	4.7 (26.4)
Refinement		
Resolution (Å)	20-2.8 (2.89-2.80)	10-2.4 (2.5-2.4)
Rejection criterion	$F \leq 0$	$F \le 0$
Total reflections	22,378 (1,845)	35,965 (3,773)
Reflections used for Rfree	1,088 (85)	1,785 (191)
Rworking ^c (%)	22.1 (32.2)	22.0 (31.3)
Rfree ^c (%)	29.3 (42.2)	27.9 (36.4)
Average B factors (Å ²)	47.9	48.8
Ramachandran data		
Percentage of residues in:		
Favored regions	85.7	88.0
Allowed regions	13.3	10.9
Generously allowed regions	0.8	0.9
Disallowed regions	0.3	0.1
Rmsd		
Bonds (Å)	0.0069	0.0064
Angles (°)	1.397	1.288
Bfactor main chain (1.5) ^e (Å ²)	1.95	1.31
Bfactor side chain (2.0) ^e (Å ²)	1.45	1.39
Cross-validated coord. error (Å)	0.47	0.41

^aAll data (outer shell).

^bRmerge = $\Sigma(|I - \langle I \rangle)/\Sigma(I)$.

^cRworking/free = $\Sigma ||F_o| - |F_c||/\Sigma |F_o|$. ^dExcluding Gly and Pro.

"Excluding Gly al

^eTarget value.

bound to two different peptides: pHB, a highly immunogenic peptide derived from an allele of the β chain of mouse hemoglobin and pHSP, a peptide derived from HSP70, which is a major self-peptide that occupies natural IE^k. These structures revealed four anchor residues in each peptide interacting with four pockets in the IE^k peptide binding groove (Fig. 1 A). The very hydrophobic p1 pocket accepted amino acids with aliphatic side chains (Ile in pHB and Val in pHSP). The large, somewhat hydrophobic p4 pocket was filled with a Phe in both cases. The very hydrophilic p6 pocket contained a Glu in the case of pHB. The carboxylate of this amino acid participated in an extensive hydrogen bonding network that included two IE^k acidic amino acids (α 11Glu and α 66Asp). In the case of pHSP, the Ala at this position only filled the very top of the pocket, with an extra water molecule now maintaining the hydrogen bonding network. Finally, the p9 pocket of IE^k consisted of a long tunnel culminating in a negatively charged β 9Glu that formed a salt bridge with the ϵ amino group of a p9Lys at that position in both peptides.

This structural information, as well as other published data (1, 4, 37–45), were used to align and predict the anchor residues of pMCC and pPCC(T102S), as well as other peptides that are the dominant IE^k epitopes in for-

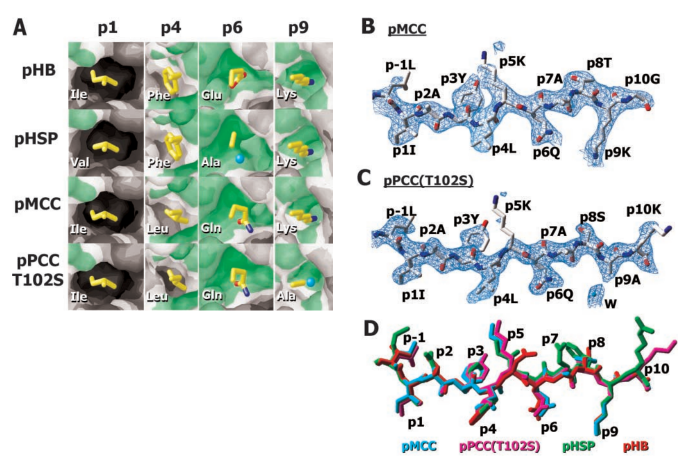


Figure 1. Structure of pMCC and pPCC(T102S) bound to IE^k . (A) Solvent-accessible surface (probe radius of 1.4 Å) representation of the four peptide amino acid binding pockets (p1, p4, p6, and p9) of IE^k with the side chains of four bound peptides are shown. IE^k surface colored by relative hydrophobicity (Gilliard scale). Gray, hydrophobic; white, neutral; green, hydrophilic. Peptide side chains are shown as wire frame. Yellow, carbon; blue, nitrogen; red, oxygen; cyan, water. The figure was made with Protein Explorer (http://proteinexplorer.org). (B) Peptide/water omit electron density map (Fo-Fc, 2.5 σ) surrounding pMCC from p1 to p11. The figure was made with the Swiss PDBViewer. (C) This shows the same as panel B for pPCC(T102S), and also shows the electron density for an extra water molecule (W) in the p9 pocket. (D) Overlay of four peptides from IE^k structures. Cyan, pMCC; magenta, pPCC(T102S); green, pHSP; red, pHB. The structures were superimposed via the backbones of the $\alpha 1$ and $\beta 1$ domains using Swiss PDBViewer. The figure was made with the Swiss PDB Viewer.

eign protein antigens, or major self-peptides isolated from IE^k naturally present on the surface of B cells, and thymic epithelium in unimmunized mice (Fig. 2). This alignment predicts an aliphatic amino acid at the p1 position. The p4 pocket is predicted to be more forgiving, preferring hydrophobic amino acids such as Phe and Leu, but also allowing other amino acids. The p6 pocket appears to have a preference for amino acids that can participate in the hydrogen bonding network underlying the peptide (Glu, Gln, and Asn), but small side chains such as the Ala in pHSP are also allowed. The p9 pocket is predicted almost invariably to be Lys.

Using this alignment, the p1, p4, and p6 pockets of IE^k were predicted to contain Ile, Leu, and Gln, respectively, for both pMCC and pPCC(T102S). Although the p9 amino acid was predicted to be Lys for pMCC, two possibilities existed for pPCC(T102S). pPCC(T102S) has an Ala insertion at p9 that shifts the Lys to p10. Therefore, either the Ala must now occupy the p9 pocket or the peptide backbone must kink to allow the shifted Lys side chain to remain in

the p9 pocket. In either case, the adjustments to the Ala insertion might contribute to poorer binding by pPCC to IE^k .

Interactions of pMCC and pPCCT102S with IE^k. То test these predictions, we solved the crystal structures of IE^k bound to pMCC and pPCC(T102S) by molecular replacement to resolutions of 2.8 and 2.4 Å, respectively, using the high resolution IE^k-pHB structure with the peptide removed as a search model. In the final models, the backbones of the IE^k portions of the structures were virtually identical to that of the IE^k in the IE^k-pHB structure. For example, using the program Swiss PDBViewer (46) to compare the combined $\alpha 1/\beta 1$ domains, the RMSD's were 0.28 Å for backbone atoms and 0.54 Å for all atoms comparing IE^k-pMCC to IE^k-pHB. Comparing IE^kpPCC(T102S) to IE^k-pHB, the values were 0.32 and 0.51 Å, respectively. Peptide omit electron density maps clearly showed the peptides in the binding groove (Fig. 1, B and C) and in the final models, the paths of the peptides through the binding grooves were nearly identical to those seen with IE^k-pHB and IE^k-pHSP (Fig. 1 D).

Source	aa Nos.	Sequence									
НВ	(68-76)	I	T	A	F	N	E	G	L	K	
HSP	(238-246)	v	N	H	F	I	A	E	F	ĸ	
MCC	(95-103)	I	A	Y	L	ĸ	Q	A	т	ĸ	
PCC	(95-104)	I	A	Y	L	ĸ	Q	A	т	A	ĸ
PCC	(T102S)	I	A	Y	L	ĸ	Q	A	s	A	ĸ
HEL	(88-96)	I	т	A	s	v	N	c	A	ĸ	
SWMb	(69-77)	L	т	A	L	G	A	I	L	K	
λrep	(18-26)	L	ĸ	A	I	Y	E	ĸ	к	ĸ	
OVA	(255-263)	L	Ε	s	I	I	N	F	Ε	K	
Snase	(89-97)	L	A	Y	1	Y	A	D	G	ĸ	
LLO	(218-226)	I	A	ĸ	F	G	T	A	F	K	
FluHA	(246-254)	I	s	1	Y	W	Т	I	۷	K	
HIV	(424-432)	I	N	Μ	W	Q	E	v	G	ĸ	
Moß2m	(37-45)	I	Q	M	L	к	N	G	ĸ	к	
MoCCinh	(45-53)	I	Q	N	A	v	Q	G	۷	ĸ	
MoSA	(351-359)	٧	E	A	A	R	N	L	G	R	
MoER60	(448-461)	I	Y	F	S	P	A	N	ĸ	ĸ	I
Relative	Position (p)	1	2	3	4	5	6	7	8	9	10

Figure 2. Peptide motif for binding to IEk. The core sequences of pHB and pHSP are aligned based on their solved structures bound to IE^k, with the positions of the four anchor residues (p1, p4, p6, and p9) highlighted. A series of other IEk binding peptides that were either identified as the major epitopes from foreign antigens or selfpeptides isolated from IE^k, were aligned based on the predicted occupancy of the four binding pockets with particular weight on p1 and p9. Only the central regions of the peptides are shown. Abbreviations and refer-

ences: HB, mouse hemoglobin (37); HSP, mouse heat shock protein 70, (38); MCC, moth cytochrome c (1); PCC, pigeon cytochrome c (1); HEL, hen egg lysozyme (39); SWMb, sperm whale myoglobin (40); λ rep, λ repressor (39); OVA (255–263), chicken ovalbumin (4); Snase, staphylococcal nuclease (41); LLO, listeriolysin O (42); FluHA, influenza hemagglutinin (43); HIV, HIV gp120 (44); Mo β 2m, mouse β 2 microglobulin (38); MoCCinh, mouse cytochrome c inhibitor (38); MSA, mouse serum albumin (38); ER60, mouse ER-60 protease (45).

The structure of pMCC bound to IE^k confirmed the MHCII binding motif prediction (Fig. 1, A, B, and D). Side chains from four amino acids, Ile at p1, Leu at p4, Gln at p6, and Lys at p9, filled the four IE^k pockets. Amino acids at positions p1, p6, and p9, but not at p4, were predicted by mutagenesis experiments to be important in the interaction of pMCC with IE^k (4). In these experiments virtually any amino acid could be substituted at the p4 position without penalty. These results suggest great flexibility in this large pocket in accepting amino acid side chains and relatively little contribution from the side chain to the overall energy of pMCC binding.

As expected, the pPCC(T102S) peptide bound to IE^k (Fig. 1, A, C, and D) also has the Ile at p1, Leu at p4, and Gln at p6, all in conformations very similar to those in pMCC. The structure also clearly shows the adjustment to the inserted Ala at the p9 position of the peptide. Although the side chain of the Lys at p9 of pMCC points down into the pocket, this position is occupied by the inserted Ala whose methyl side chain only fills the very top of the pocket, whereas the p10 Lys is shifted and rotated so that the side chain is now completely solvent-exposed. An extra water molecule is now found in the p9 pocket. This suggests that the loss of the buried Lys side chain is most likely a major reason for the weaker binding of pPCC compared with pMCC.

Details of the peptide interactions with IE^k in the region of the p6 and p9 pockets are shown in Fig. 3. The side chain of the p9Lys of pMCC extends through a hydrophobic tunnel to a salt bridge with β 9Glu and interacts with a water molecule (W2). Other interactions at this end of the peptide include the highly conserved hydrogen bonds between β 61Trp and α 69Asn, the peptide backbone seen in nearly all MHCII structures, and an extensive hydrogen bond network that includes p6Gln, α 66Glu,

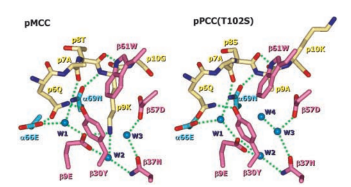


Figure 3. Details of the p6 and p9 pockets of IE^k–pMCC and IE^k– pPCC(T102S). Shown are wire frame representations of residues p6-p10 of pMCC (left) and pPCC(T102S) (right). Also shown for both molecules are side chains of α 66Asn, α 69Asn, β 9Glu, β 30Tyr, β 37Asn, β 57Asp, and β 61Trp. Light yellow, peptide carbon; cyan, α chain carbon; magenta, β chain carbon; blue, nitrogen; red, oxygen. Three water molecules (W1–W3) common to both molecules are shown as well as a water molecule (W4) that is unique to the IE^k–pPCC(T102S) structure (light blue). Some of the predicted hydrogen bonds among the atoms are shown as green dotted lines. Figures were made with WebLab ViewerPro (Accelrys c.).

 β 30Tyr, and several water molecules. In the IE^k–pPCC (T102S) structure, the interactions at p9 are lost and the pocket is partially filled with an extra water molecule (W4). However, the other details of the hydrogen bonding network are preserved.

The main chain of pPCC(T102S) has not kinked to maintain a Lys in the p9 pocket, but rather its path through the binding groove is identical to that of pMCC. Kinks in peptides bound to MHC class I (MHCI) are common (47, 48) and function to maintain the conserved binding of MHCI to the peptide NH₂ and COOH termini, and allow the insertion of favorable amino acid side chains in the MHCI binding pockets. However, in all solved structures of MHCII, including those presented here, the peptide takes a very similar, nearly linear course through the binding groove, held in place by hydrogen bonding between its backbone and the side chains of conserved amino acids on the MHC α helices (31, 49–53).

It is difficult to predict the net changes in the IE^k– pPCC(T102S) structure if the peptide were to be kinked and rotated to preserve a Lys in the p9 pocket. The van der Waals interactions with the Lys side chain as well as the salt bridge and the hydrogen bonds to its ϵ -amino group, would be gained. However, it is likely that the conserved peptide backbone interactions with β 61Trp and α 69Asn would be lost. Also, depending on the precise nature of the kink, the hydrogen bonding network involving the p6Gln might be disturbed. Apparently, the energy gained by the new interactions would not compensate for the loss of these other interactions.

Peptide Amino Acids Interacting with $\alpha\beta TCRs$. The interaction of IE^k/pMCC and IE^k/pPCC with the $\alpha\beta$ TCRs of many different T cell clones has been extensively studied. Various mutagenesis experiments have shown that the amino acids at p2, p3, p5, p7, and p8 are involved in recog-

nition by various T cell clones (4, 6). These represent all of the surface-exposed amino acids in the center of the bound peptides (Fig. 4 A). Of these, the p5Lys has been shown to be the most important. For most T cell clones raised by immunization with pMCC or pPCC, effective stimulation is critically dependent on a Lys at this position. This Lys is directly in the center of the binding pocket with its side chain pointing up and out of the groove into the solvent.

T cell recognition of these peptides is also usually very dependent on the p8Thr. Substitution of any other amino acid at this position, even the conserved Thr to Ser substitution shown here, often compromises T cell activation. For the Thr to Ser change, the peptide is converted to a weaker agonist in that equivalent T cell activation now re-

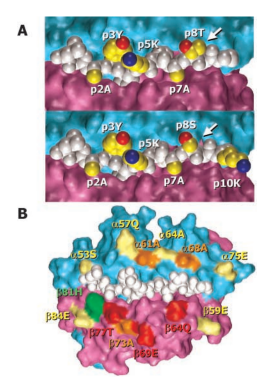


Figure 4. Surface amino acids of IE^k-pMCC and IE^k-pPCC(T102S) that are important for $\alpha\beta$ TCR recognition. (A) A portion of the solventaccessible surfaces of the $\alpha 1$ and $\beta 1$ domains of IE^k-pMCC (top) and IE^k-pPCC(T102S) (bottom) are shown. Cyan, α chain; magenta, β chain. A van der Waal's Corey, Pauling, Koltan (CPK) representation of the peptide is shown from p1 to p10. The peptide atoms are white except for the side chains of the five central surface exposed amino acids and the Lys at p10 of pPCC(T102S). These are colored by atom type. Yellow, carbon; blue, nitrogen; red, oxygen. The methyl group of the p8Thr of pMCC and its absence on the p8Ser of pPCC(T102S) are indicated with white arrows. The figure was made with WebLab ViewerPro. (B) The solvent-accessible surfaces of the $\alpha 1$ and $\beta 1$ domains of IE^k-pMCC are shown. A van der Waal's CPK representation of the peptide is shown from p1 to p10 in white. Except for the 13 amino acids mutated in the study by Jorgensen et al. (20), the IE^k α chain is colored cyan and the β chain is colored magenta. The other 13 amino acids are colored according to the effects of their mutation. Red, the response of at least three out of four T cell hybridomas compromised by mutation; orange, the response of only one out of four T cell hybridomas compromised by mutation; green, the response of one hybridoma improved by mutation; yellow, no effect of mutation. The figure was made with WebLab ViewerPro.

quires more peptide (4, 24). For one T cell clone, partial blocking of the function of CD4 with a monoclonal antibody converted this mutant peptide to an antagonist, i.e., exposure of the clone to the p8Ser peptide under these conditions inhibited its response to the wild-type peptide (54). Another mutation at this position (Thr to Gly) has also been shown to produce an even more antagonistic peptide (24).

In the region of p8, the only detectable surface structure alteration induced by the Thr to Ser substitution was the loss of the Thr side chain methyl group (Fig. 3 and Fig. 4 A). The C β carbons and hydroxyl oxygens appear to be in identical positions in both structures. The backbone of the peptide and the position of the other amino acid side chains in this region did not change significantly. Therefore, either this methyl group is an essential contact point for the $\alpha\beta$ TCR or, less likely, its loss confers some kinetic instability to this region of the molecule that is not revealed by this static structure. As previously described, the insertion of an Ala at p9 of pPCC(T102S) has forced the Lys at p10 out of its binding pocket and onto the molecule surface. This alteration does not seem to be detected by most T cell clones, which suggests that the $\alpha\beta$ TCR does not usually contact this area of the surface.

The requirements for some of the other solvent-exposed peptide amino acids are not quite as stringent for recognition by T cell clones. For example, several receptors accept an Ala to Gly substitution at p2 or p7, but for the most part larger amino acids are not tolerated, perhaps because of steric problems rather than essential receptor–Ala contacts. Phe, or sometimes Trp but not Leu or Met, is accepted in place of the p3Tyr, which suggests that an aromatic amino acid may be essential at this position. In both of our structures, the close proximity of the Tyr aromatic ring to the side chain of the p5Lys may indicate a role for this interaction in positioning this crucial Lys for $\alpha\beta$ TCR recognition. This interaction has been suggested by nuclear magnetic resonance studies (55).

 IE^k Amino Acids Interacting with $\alpha\beta TCRs$. A mutational study of IE^k revealed a number of amino acids on the α and β chain α helices that were important for the recognition of the IE^{k} -pMCC complex by T cells (21). In these experiments, single mutations were introduced into the IE^k molecule at six positions of the α chain (α 53Ser to Asn, α 57Gln to Arg, α 61Ala to Val, α 64Ala to Val, α 68Ala to Val, and α 75Glu to Lys) and seven positions of the β chain (β 59Glu to Lys, β 64Gln to Arg, β 69Glu to Lys, β 73Ala to Val, β 77Thr to Gln, β 81His to Tyr, and β 84Glu to Lys). Although no IE^k structure was available at the time, the authors predicted that these amino acids would lie on the top of the α helices based on sequence homology to MHCI (56). The IE^k structures show that this prediction was indeed correct (Fig. 4 B).

Each mutated IE^k molecule was expressed at the cell surface and tested for binding and presentation of the pMCC to four different T cell hybridomas. In general, the α chain mutations had less effect than those on the β chain, although several of the α chain substitutions were quite conservative (Ala to Val). These amino acids are colored on the surface of the IE^k molecule (Fig. 4 B) according to the effect of these mutations. Several of the mutations (B64Gln to Arg, β69Glu to Lys, and β77Thr to Gln) severely reduced the response of at least three of the hybridomas (red). Others (α 61Ala to Val, α 68Ala to Val, and β 73Ala to Val) compromised the response of only one of the four hybridomas (orange). Mutation of B81His to Tyr (green) had no effect on three of the hybridomas, but greatly enhanced the response of the fourth. Mutations at the other positions, which were mainly at the extremities of the α helices, had no effect on T cell recognition (yellow). In total, these data suggest that the main sites of $\alpha\beta$ TCR interaction with this MHCII peptide ligand are clustered on the upward face of the complex and focused on central peptide residues and the tops of the arched α helices.

Predicted αβ*TCR Interactions with* pMCC and pPCC. Several studies have investigated αβ*TCR* amino acids important in IE^k–pMCC and IE^k–pPCC recognition. For example, experiments have been performed using pMCC and variants at position p5 and p8 to immunize mice bearing a transgene encoding either the α or β chain of an αβ*TCR* from an IE^k/pMCC–reactive T cell (20). Thus, the responding T cells had one of the αβ*TCR* chains fixed by the transgene, whereas the other was allowed to vary. From the sequences of the CDR3s of the varying αβ*TCR* chain, the authors argued convincingly that the p5Lys interacted with a conserved Glu at position 94 of VαCDR3, whereas the p8Thr interacted with a conserved Asn at position 97 of VβCDR3.

These two alleles of V β 3 have been identified in mice: V β 3a and V β 3b (BV3S1A1 and BV3S1A2, respectively). These differ only at amino acid 31 in what is predicted to be the V β 3 CDR1. This amino acid is Val in V β 3a and Phe in V β 3b. V β 3 dominates the response to pPCC or pMCC in mouse strains that carry V β 3a, but not in those with V β 3b (57). Furthermore, direct mutation of amino acid 31 from Val to Phe in the receptor of a V β 3a bearing IE^k/pMCC-reactive T cell hybridoma eliminated IE^k/pMCC recognition (10). These findings suggest a critical interaction between the V β 3 CDR1 loop and the IE^k-pMCC and IE^k-pPCC ligand, which is disrupted by the Val to Phe substitution.

There are no crystal structures of $\alpha\beta$ TCRs bound to IE^k peptide. However, it is reasonable to hypothesize that the orientation of an $\alpha\beta$ TCR bound to IE^k will be similar to those seen in crystal structures of other $\alpha\beta$ TCR complexes with MHCI and MHCII ligands (58–63). In these structures, the $\alpha\beta$ TCR sits diagonally on the top of the MHC molecule, wedged between the arches of the MHC α helices with the V α , and V β CDR3s are centered over the middle of the peptide. So far, in all cases the orientation of the $\alpha\beta$ TCR has been similar with V α on the peptide NH₂-terminal side and the V β on the peptide COOH-terminal side. Within these constraints, however, the various $\alpha\beta$ TCRs swivel by up to 35°.

Therefore, in order to interpret some of these findings we superimposed IE^k -pMCC on the MHCII portion of

two solved $\alpha\beta$ TCR-MHCII peptide complexes (61, 62) to predict how the CDRs of an anti–IE^k/pMCC $\alpha\beta$ TCR might sit on the ligand (Fig. 5 A and B). These models strongly support the results of the mutational studies. The V α and V β CDR3 loops (cyan) are predicted to lie directly over the center of the pMCC peptide. The side chain of p5Lys is in close proximity to the predicted positions of the V α 94, and the side chain of p8Thr is in close proximity to the predicted positions of the V β 97. It is easy to see how mutations in either one of these peptide amino acids might alter the $\alpha\beta$ TCR binding kinetics and affinity. The predicted position of V β 31 in V β CDR1 is such that a substitution of Phe for Val at this position would probably interfere with the interaction of V β CDR1 with the α helix of the IE^k α chain. The seven amino acids of IE^k that affect $\alpha\beta$ TCR interaction when mutated all lie in positions close to the $\alpha\beta$ TCR CDR loops. Overall, this model agrees remarkably with the published biochemical and mutational studies.

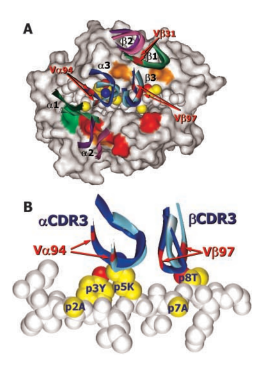


Figure 5. Predicted T cell receptor interactions with IE^k/pMCC. (A) Top view of the solvent-accessible surface of the $\alpha 1$ and $\beta 1$ domains of IE^k-pMCC are colored white except for $\alpha 61$, $\alpha 68$, $\beta 64$, $\beta 69$, $\beta 73$, $\beta 77$, and β81, which are colored as in Fig. 4 B. A van der Waal's CPK representation of the peptide is shown from p1 to p10 and is colored as in Fig. 4 A. Two $\alpha\beta$ TCRs, one from HA1.7 (specific for DR1 plus a peptide from influenza hemagglutinin) (64) and the second from D10 (specific for IA^k plus a peptide from conalbumin) (65), whose structures were solved bound to their MHCII peptide ligands (61, 62), were aligned on the IE^k-pMCC structure by superimposing the backbones of the MHC molecule $\alpha 1$ and $\beta 1$ domains. Ribbons, the V α and V β CDR loops ($\alpha 1$, $\alpha 2$, α 3, β 1, β 2, and β 3) of the two $\alpha\beta$ TCRs; dark green, HA1.7 $\alpha\beta$ TCR $V\alpha/\beta$ CDR1; purple, $V\alpha/\beta$ CDR2; blue, $V\alpha/\beta$ CDR3; light green, D10 $\alpha\beta$ TCR V α/β CDR1; magenta, V α/β CDR2; cyan, V α/β CDR3. The positions of three $\alpha\beta$ TCR amino acids, V α 94, V β 31, and VB97, are labeled and colored red. (B) Side view of the peptide and CDR3 portions of Fig. 4 A with other parts of the molecules removed. The figures were made with WebLab ViewerPro.

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