Promiscuous binding of extracellular peptides to cell surface class I MHC protein

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Promiscuous binding of extracellular peptides to cell surface class I MHC protein

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Algorithms derived from measurements of short-peptide (8–10 mers) binding to class I MHC proteins suggest that the binding groove of a class I MHC protein, such as Kb, can bind well over 1 million different peptides with significant affinity (<500 nM), a level of ligand-binding promiscuity approaching the level of heat shock protein binding of unfolded proteins. MHC proteins can, nevertheless, discriminate between similar peptides and bind many of them with high (nanomolar) affinity. Some insights into this high-promiscuity/high-affinity behavior and its impact on immunodominant peptides in T-cell responses to some infections and vaccination are suggested by results obtained here from testing a model developed to predict the number of cell surface peptide–MHC complexes that form on cells exposed to extracellular (exogenous) peptides.

The heterodimeric αβ antigen-binding receptors on T cells (TCR) and the antigen-binding Fab fragments of antibodies are similar structurally and in the great diversity of their ligand-binding site. Their diversity is also generated by similar stochastic gene segment rearrangements in developing B and T cells. But antibodies can be raised against virtually any existing or imagiable organic structures (antigens), if sufficiently stable, while the antigens recognized by T cells’ TCRs are remarkably limited to short peptides and (iii) they are encoded by genes in the major histocompatibility complex (SI Text, R1 and R2).

Although the peptides are bound with 1:1 stoichiometry, well over 1 million different peptides are estimated to bind with significant affinity to an MHC protein’s binding site (see below). This extreme binding promiscuity, or degeneracy, is probably matched only by the binding of amino acid sequences in unfolded proteins by some heat shock proteins (1, 2) (SI Text, R3). It is not unlimited, however, because each MHC type (encoded by an MHC allele) usually exercises preferences for peptides having particular motifs (e.g., one or two strategically placed amino acid residues) (3). The MHC binding site can also distinguish between peptides that differ by single amino acid mutations or substitutions, apparently binding them with different affinities (4, 5).

Moreover, in the in vivo responses to foreign proteins such as to those proteins of an invading virus, only one (immunodominant) peptide is recognized, thereby focusing of T-cell responses to some pathogens and vaccines on only a few immunodominant peptides.

Model and the Strategy for Testing

The model (Fig. 1) stems from evidence that extracellular (exogenous) peptides can bind to those cell surface MHC molecules that have vacant peptide-binding sites (16) (SI Text, R10). These empty MHC molecules (M) might have been loaded with an endogenous peptide that subsequently dissociated to leave a vacant groove, or they might have appeared on the cell surface as newly synthesized MHC molecules that escaped being loaded in the endoplasmic reticulum with an endogenous peptide. Whatever their origin, the empty MHCs are unstable: they either bind the extracellular peptide (P), forming an MP complex, or undergo denaturation (17).

A third way to generate cell surface MHC, the way with which we are here concerned, is to simply expose cells to extracellular peptides. Although generally thought not to occur in vivo, this last process (often termed peptide pulsing or loading) is critically important for efforts to identify peptides that are recognized by particular TCR and stimulate T cells with peptide-pulsed dendritic cells, such as in some peptide-based vaccines (SI Text, R7 and R8). That it can also occur under physiological conditions is suggested by the binding of peptides in lymphatic fluid (8) and by evidence for extracellular loading of cells in the pancreas with an insulin-derived peptide (9).

The dynamics of intracellular (endogenous) peptide production and their association with MHC have been extensively examined (10, 11, 12) (SI Text, R9), and the thermodynamics and kinetics of some pMHC interactions have been characterized in considerable detail (13–15). Still, although peptide pulsing is indispensable for efforts to identify peptides recognized by T cells, the binding of extracellular peptides to cell surface MHC is not sufficiently well-understood to predict the number of pMHCs formed on cells after exposing them to an extracellular peptide. Our aims here are to evaluate a model that may serve as the basis for making such estimates and predictions and to see if the model leads to any insights into promiscuous peptide binding by MHC proteins and some of its ramifications. Our results show good agreement between predicted and measured numbers of cell surface complexes formed by extracellular peptides, and they support evidence that promiscuous binding arises from conformational flexibility of MHC proteins and peptides. They also support and extend evidence that (i) a class I MHC protein can bind millions of different short peptides and (ii) the very slow dissociation of some pMHC complexes accounts for their high (nanomolar) affinity; (iii) they also suggest how MHC binding promiscuity might support the focusing of T-cell responses to some pathogens and vaccines on only a few immunodominant peptides.


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Materials and Methods

The model focuses on changes over time in the number of cell surface empty MHC that bind a P and form MP complexes. As is shown below, the number of these complexes is determined by the rates at which empty MHC appear on the cell surface (dM/dt) and undergo denaturation (k\text{den}) and the rates at which they associate with the peptide (k\text{on}) and the MP complexes dissociate (k\text{off}) (Materials and Methods). To determine k\text{off}, cells were pulsed with a peptide, and loss of the resulting MP complexes was then followed over time. Values for the three parameters were obtained by measuring the numbers of MP on cells that had been incubated with a peptide at diverse concentrations for various times. All of the measured MP values, together with the independently derived k\text{off}, were then subjected to a multidimensional search program (MATLAB fminsearch) to obtain k\text{on}, k\text{den}, and dM/dt values that best fit the measured MP.

We examined the binding of two synthetic octapeptides to K\text{b}, a mouse class I MHC molecule. The K\text{b}-expressing cells were two cell lines, RMA-S and DC2.4, and bone marrow-derived dendritic cells (1° DCs) from C57BL/6 mice. Most experiments were performed with RMA-S cells, which lack the transporter (TAP) activity that translocates proteasome-generated peptides into the endoplasmic reticulum; such TAP-deficient cells are thought to have (at steady state) a relatively large number of empty class I MHCs on the cell surface. DC2.4, a dendritic cell line, and the 1° DCs are TAP-competent.

The peptides tested were SIINFEKL (often referred to as OVA257-264 but here referred to simply as OVA) and SIYR-YYGL (termed SIY). OVA is produced by cells that express chicken ovalbumin; the OVA-K\text{b} complex is a potent agonist for the widely studied TCR on CD8+ T cells from OT-1 TCR transgenic mice (5). The SIY peptide was identified in a combinatorial library (4); the SIY-K\text{b} complex is a potent agonist for another widely studied TCR on cloned CD8+ T cells (2C) that are maintained as cultured cell lines or obtained from 2C TCR transgenic mice (18).

Results

Peptide–MHC Dissociation Rate Constants (k\text{off}). When fluorescein-labeled OVA was previously added to live cells under conditions similar to those used in the present work, some peptide was endocytosed and bound to K\text{b} in the endoplasmic reticulum (19). In another earlier study, some cell surface pHMC complexes formed from an exogenous peptide with L5 were rapidly endocytosed and later displayed on the cell surface after a lag of many hours (20). To minimize endocytosis, cytochalasin D was added to cells before and during loading of the exogenous peptides. Under these conditions, peptide dissociation from MP followed exponential kinetics (Fig. S1). For OVA, k\text{off} was 0.0495/h (t1/2 ~ 14 h), and for SIY, k\text{off} was 0.119/h (t1/2 ~ 5 h) (Table 1). Previously reported k\text{off} values for dissociation of various peptides from pHMC complexes have been found to span a wide range, from a few minutes (21) to days, for a class II MHC (SI Text, R12).

For OVA-K\text{b}, the finding of a t1/2 of 14 h at 37 °C (Table 1) is consistent with the 21-h t1/2 value found at 25 °C by surface plasmon resonance with recombinant K\text{b}, and the OVA peptide modified covalently for immobilization on a chip (22).

Table 1. Affinity and rate constants for SIINFEKL (OVA) and SIYRYYGL (SIY) binding to cell surface K\text{b}

<table>
<thead>
<tr>
<th>Rate constants</th>
<th>SIINFEKL</th>
<th>SIYRYYGL</th>
</tr>
</thead>
<tbody>
<tr>
<td>k\text{on} (M⁻¹h⁻¹)</td>
<td>1.627 × 10⁷</td>
<td>7.889 × 10⁷</td>
</tr>
<tr>
<td>k\text{off} (h⁻¹)</td>
<td>0.0495</td>
<td>0.1191</td>
</tr>
<tr>
<td>K⁰ = k\text{on}/k\text{off} (nM)</td>
<td>3.042</td>
<td>151</td>
</tr>
<tr>
<td>k\text{den} (h⁻¹)</td>
<td>1.0872</td>
<td>0.4683</td>
</tr>
<tr>
<td>dM/dt (h⁻¹)</td>
<td>[1.00 – 3.19] × 10⁴</td>
<td>[1.36 – 1.88] × 10⁴</td>
</tr>
<tr>
<td>fi(0)</td>
<td>[2.44 – 5.69] × 10⁴</td>
<td>1.23 × 10⁵</td>
</tr>
</tbody>
</table>

*In other studies of OVA binding to K\text{b}, K\text{b} values ranged from 1.5 to 7 nM (21, 22) (SI Text, R25).

RMA-S cells.

DC2.4 cells.

Primary dendritic cells.

Binding of OVA and SIY to K\text{b} (k\text{on}, k\text{den}, dM/dt). The numbers of OVA–K\text{b} complexes that formed on RMA-S and DC2.4 cells after incubating them for 20 min to 4 h with OVA at concentrations that ranged from 0.01 to 10 μM were measured with the 25D1.16 antibody. The data (Fig. 2A) were all combined and analyzed according to the equations shown in Materials and Methods using the fminsearch program (MATLAB) and the independently determined k\text{off} value (Fig. S1). The best-fitting values for k\text{on}, k\text{den}, and dM/dt are shown in Table 1.

Antibodies that are highly specific for a particular pHMC complex—like mAb 25D1.16 for the OVA–K\text{b} complex—are rare, perhaps because they are more stable than most other pHMCs and thus exceptionally immunogenic. It was of interest, therefore, to test the model with more commonplace anti-MHC antibodies, which react with allele-specific elements of MHC proteins regardless of the associated peptide. mAb Y3 recognizes such elements in the α1 and α2 domains of K\text{b} (23). Hence, fluorescein-labeled Y3 was used to follow the binding of the SIY peptide to K\text{b} on RMA-S cells. Under these conditions, it was necessary to subtract from the amount of Y3 that bound to SIY-pulsed cells the amount of Y3 bound to cells that were not peptide-pulsed. After incubating the cells with SIY at various concentrations and different times, the numbers of SIY–K\text{b} complexes formed were measured (Fig. 2B). The SIY–K\text{b} dissociation constant (k\text{off}) was determined as shown in Fig. S1, Lower, and the values found for k\text{on}, k\text{den}, and dM/dt for SIY binding to K\text{b}, shown in Table 1, were obtained as described above for the OVA–K\text{b} interaction.

Unlike k\text{on} and k\text{den}, which reflect inherent properties of OVA and SIY interactions with empty K\text{b} and of empty K\text{b} alone, it was expected that dM/dt might vary considerably from experiment to experiment, because the experiments were carried out over a period of many months with cells maintained more or less continuously in culture. The results of each experiment were, therefore, analyzed separately for dM/dt using the k\text{off}, k\text{on}, and k\text{den} values from Table 1. dM/dt was seen to vary only from about 10,000 to 20,000 cells⁻¹ h⁻¹ for RMA-S cells and from about 20,000 to 60,000 cells⁻¹ h⁻¹ for DC2.4 cells.

The experimentally determined and predicted numbers of MP per cell are compared in Fig. 3. Agreement was evidently obtained over short and long pulsing times (20 min to 4 h) with low- or high-peptide concentrations and both the TAP− RMA-S and the TAP+ DC2.4 cells.

Sensitivity of Calculated Numbers of MP per Cell to Changes in Parameter Values. To examine the impact of parameter value variations on calculated MP values, each parameter was allowed...
Fig. 2. Binding of peptides to cell surface $K^b$ (37°C). The curves are based on Eq. 4 and values for $k_{on}$, $k_{off}$, and $dM_d/dt$ in Table 1. (A) Binding of OVA peptide to $K^b$ on RMA-S (a–f), DC2.4 (g and h), and primary dendritic cells (i). Peptide concentrations are indicated by color in order of orange, green, purple, and (when applicable) blue. (a) 7.85E-009, 7.85E-008, 7.85E-007; (b) 6.30E-008, 6.30E-007, 6.30E-006; (c) 3.00E-007, 3.00E-006, 3.00E-005; (d) 3.00E-007, 3.00E-006, 3.00E-005; (e) 1.00E-008, 1.00E-007, 1.00E-006; (f) 1.00E-008, 1.00E-007, 1.00E-006; (g) 1.00E-006, 1.00E-005; (h) 1.00E-006, 1.00E-005; (i) 1.00E-007, 1.00E-006, 1.00E-005. (B) Binding of SIY peptide to $K^b$ on RMA-S cells. SIY concentrations are indicated by color in the order of orange, green, purple, and (only for a) blue. (a) 1.00E-008, 1.00E-007, 1.00E-006, 1.00E-005; (b) 8.30E-008, 8.30E-007, 8.30E-006.

Fig. 3. Scatter plot comparing predicted and measured number of MP complexes per cell (y and x axes, respectively) from data in Fig. 2. For OVA peptide binding, red dots are from RMA-S cells, blue dots are from DC2.4 cells, and green dots are from primary bone marrow-derived dendritic cells. For SIY peptide binding, purple dots are from RMA-S cells. Linear regression values (95% confidence intervals in parentheses) are slope = 0.9757 (0.9291–1.022), y-intercept = −405, and $r^2 = 0.922$. Dashed line corresponds to perfect match between predicted and measured values.

Frequency of Short Peptides That Bind to Class I MHC ($K^b$). The affinities of various class I MHC proteins for thousands of peptides, measured as IC$_{50}$ values, have served as the basis for predictive algorithms that sort peptides with considerable accuracy into strong and moderate binders (IC$_{50} < 500$ nM) and weak or nonbinders (IC$_{50} > 500$ nM) to various MHC-I proteins (25). Using the 500-nM affinity cutoff, it has recently been estimated that about 1% of the ~10 million unique nonamers in the human proteome bind to human MHC-I: 0.7% bind to what may be the least promiscuous class I MHC protein, HLA-B57, and 1.8% bind to a more conventional protein, HLA-B7 (26). Using a more stringent 50-nM cutoff to identify strong binders, Istrail et al. (27) found a lower frequency of short peptides that bind (0.2–0.5% depending on the MHC-I allele), but importantly, the frequency was the same for predicted proteins from a wide variety of proteomes: human, mouse, Drosophila, Caenorhabditis elegans, many viruses, bacterial species (including archeobacteria from the ocean floor), and even the randomly permuted (shuffled) amino acid sequences of various proteomes.

To further examine this issue, we considered the frequency of $K^b$-binding octamers from two proteins that are evolutionarily remote from mammals and their pathogens. The selected proteins were from the set of putatively plant-specific proteins that have been identified in a plant genome (Arabidopsis thaliana; i.e., proteins with similar sequences are found only in genomes of plant species and not in genomes from Eukaryota, Bacteria, and Archea) (28). In one of these plant proteins (a β-amylase, 77 kDa; accession no. AT2G45880, UniProt 90/Swiss KB database) with 684 unique 8-mers, nine (1.3%) octamers are predicted by NetMHCpan (version 2.4) (25) to bind to $K^b$ (with higher affinity than IC$_{50} < 500$ nM). In
another plant-specific protein (a methyltransferase, 77 kDa; accession no. AT1G78240, UniProt 90/Swiss KB database) with 677 unique 8-mers, seven (1.0%) octamers are predicted to bind to K\(^b\) with affinity above the cutoff (one of them, SSFAYSRIL, with a predicted 2-nM affinity). Thus, it seems that, in general, a representative MHC-I protein, such as K\(^b\), can bind with significant affinity about 1% of all potential octamers. It can be estimated that the number of unique octamers in all 1.3 million proteins in the UniProt 90/Swiss KB database (as of 2005) is about 354 million. Thus, although K\(^b\), a typical MHC-I protein, binds only around 1% of octamers, this small fraction amounts to over 3 million different peptides. And this number is an underestimate because (i) the 500-nM affinity cutoff is somewhat arbitrary (Table 2), (ii) there are limitations to the algorithm’s accuracy—estimated in the work by Kosmrly et al. (26) for a similar algorithm to be about 80% accurate, and (iii) excluded from the total are peptides with covalent modifications, such as phosphoryl groups, and other posttranslational modifications, haptenated peptides [like 2,4-dinitrophenyl (DNP) and the great many other low molecular-mass substances that cause allergic contact dermatitis and drug hypersensitivities], and peptides generated in combinatorial libraries, like the one in which the SIY peptide was identified (4). Indeed, the SIY peptide does not exist in known proteomes (as of 2005).

Table 2 shows that, of several extensively studied peptides, NetMHCPan correctly identified three binders to K\(^b\); it also shows, correctly, that peptide #3 (dEV8) is a better binder than peptide #4 (p2Ca) to K\(^b\), consistent with K\(^b\) having been successfully crystallized in association with dEV8 but not with p2Ca (SI Text, R14). Similarly, p2Ca (LSFPFPEQL), with affinity for K\(^b\) that is below the 500-nM cutoff, actually forms complexes with K\(^b\) (SI Text, R15) that are recognized by a TCR (2C), albeit very weakly (29), and elicits a specific cytolytic response by cloned CD8\(^+\) 2C T cells (30). Likewise, the two peptides (#4 and #5) that are predicted not to be bind significantly to L\(^d\) actually form complexes with L\(^d\) that may be the most potent natural T-cell agonists known (29, 31). It may be that predicted values for peptide-L\(^d\) complexes are erratic because the training set is limited.

**Discussion**

According to the model illustrated in Fig. 1, four parameters (\(k_{on}, k_{off}, k_{on},\) and \(dM/dt\)) determine how many pMHC complexes are formed on cells that are exposed to extracellular peptides. How well do the parameters values found here match the corresponding values reported previously, and what bearing do they have on binding promiscuity? The \(k_{off}\) and \(K_D\) values found here are consistent with those reported previously for various pMHC complexes, including OVA–K\(^b\) (21, 22) (Table 1, SI Text, R25, and Fig. S1).

The association (\(k_{on}\) and denaturation rate constants (\(k_{on}\), however, are of particular interest for promiscuity. The procedures used to measure \(k_{on}\) for peptide binding to ostensibly empty MHC molecules have varied widely, and the reported association rate constants have spanned a wide range, roughly from 10\(^4\) to 10\(^9\) M\(^−1\) s\(^−1\). Many reported values are at the low end of this range (21) (SI Text, R16 and R17), similar to those values found here and also for peptide binding to a class II MHC (32). When expressed in conventional units (M\(^−1\) s\(^−1\)) rather than in the units (M\(^−1\) h\(^−1\)) used in Table 1, we found 4,900 M\(^−1\) s\(^−1\) for OVA–K\(^b\) and 217 M\(^−1\) s\(^−1\) for SIY–K\(^b\) (Table 1). In contrast, the highest \(k_{on}\) reported was about 10\(^8\) M\(^−1\) s\(^−1\) for a nonamer binding to D\(^\alpha\) (33). The latter, however, is still 10–100 times slower than the value found for some antibody–hapten reactions; for instance, the \(k_{on}\) for binding an e,2,4-DNP-lysine nonamer [DNP-(lysine)] by a homogeneous antibody (myeloma protein) was 10\(^9\) M\(^−1\) s\(^−1\), and the same protein’s binding of the smaller hapten, e,2,4-DNP-lysine, was 10\(^8\) M\(^−1\) s\(^−1\), which was almost diffusion limited (SI Text, R18 and R19).

For these disparate \(k_{on}\) values is suggested by studies in which the binding of dansyl- or fluorescein-labeled peptides to recombinant MHC proteins was monitored continuously (13–15, 33). Under these conditions, the association kinetics were biphasic: one rate, reflecting a slow unimolecular step, was attributed to changes of MHC conformation from peptide-unreceptive to -receptive conformations, and the other rate was attributed to peptide binding to the MHC’s receptive conformation. Because there is also evidence that peptides are flexible and can adopt different configurations (9, 34, 35) (SI Text, R26), the very slow overall on-rate values could reflect the time required to achieve sufficient mutual configurational complementarity for flexible MHC and flexible peptides to form an initial complex, which might then undergo an induced fit process. The resulting very slow dissociation of some pMHC could account for their high affinity (Table 1, OVA–K\(^b\)). The binding of flexible MHC and peptides is expected to have an entropic cost. To assess the cost, the entropy change for a peptide–MHC interaction was determined and actually found to be favorable, probably because of the hydrophobicity of the peptide and the MHC’s binding site (13). Whether other peptide–MHC reactions are also entropy-driven remains to be seen.

Although conformational variability underlies an MHC molecule’s ability to bind many different peptides (13), a structural basis for this promiscuity is evident in X-ray crystallographic findings. Most of the hydrogen bonds between bound peptide and an MHC’s binding site residues involve the peptide’s backbone main chain atoms, a common feature of peptides bound in extended conformation (SI Text, R20–R22). In contrast, the preferential binding of some peptides (determinant selection) arises from interactions of side chains of peptide residues at the anchor and some other positions of the peptide with MHC residues in depressions or pockets in the binding site.

**Table 2.** Predicted relative affinity (IC\(_{50}\)) of some class I-MHC proteins for the indicated peptides

<table>
<thead>
<tr>
<th>Peptide</th>
<th>MHC-I</th>
<th>Affinity (IC(_{50}) nM)*</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. SIVRYGGL</td>
<td>K(^b)</td>
<td>13.27</td>
</tr>
<tr>
<td>2. SIINFKEK</td>
<td>K(^b)</td>
<td>215.07</td>
</tr>
<tr>
<td>3. EQYKFYSV</td>
<td>K(^b)</td>
<td>236.63</td>
</tr>
<tr>
<td>4. LSFPFPEQL</td>
<td>K(^b)</td>
<td>716.55</td>
</tr>
<tr>
<td>5. LSFPFPEQL</td>
<td>L(^d)</td>
<td>17,349</td>
</tr>
<tr>
<td>6. QLSFPFPEQL</td>
<td>L(^d)</td>
<td>28,392</td>
</tr>
</tbody>
</table>

*Values are from the netMHCPan predictive algorithm, v2.4 (23).
If slow association rates and MHC conformational variability are the keys to promiscuous peptide binding, the lifetime of empty MHC molecules (indicated by \( k_{\text{den}} \)) is significant. The \( k_{\text{den}} \) found here correspond to \( t_{1/2} \)-values of 38–88 min (at 37 °C). These values (~50 min on average) (Table S1) are similar to the other \( t_{1/2} \) (55 min) for an empty MHC of which we are aware; the latter was found for a human class II MHC, HLA-DR [calculated by Kevin Fowler from data in the work by Grotenbreg et al. (32)] using a model that differed considerably from the model used here]. Whether these unexpectedly long-lived empty MHCs found for one mouse class I and one human class II MHC are representative of MHC proteins in general remains to be seen. For effective peptide pulsing of cells, the lifetimes of empty MHCs should exceed peptide–MHC association rates (\( k_{\text{on}} > k_{\text{den}} \)). This inference is supported by the simulated effect of reducing \( k_{\text{den}} \) on the number of predicted pMHC complexes formed (Fig. 4).

In contrast to promiscuous peptide binding to MHC, only one or very few of the short peptides that can be potentially derived from a virus’s proteins are actually recognized in vivo by the most of the T cells that respond to some virus infections (6). An extreme example of such immunodominant peptides is seen in C57/BL6 (B6) mice, which have two MHC-I proteins, \( K^{\alpha} \) and \( D^{\alpha} \). In these mice, the majority of CD8+ T cells produced in response to influenza virus infection of the respiratory tract recognize only 1 octamer of viral origin (NPfl, in association with \( D^{\alpha} \)) of the over 4,000 potential octamers in the virus’s proteome (about 4,400 aa residues encoded in 11 proteins). Similar immunodominance of a few peptides is also evident in humans infected with influenza virus, but the dominant peptides differ among various individuals, even those individuals sharing the same restricting MHC (36). Altering the route of vaccinia virus infection in mice also results in changes in the vaccinia virus-derived dominant peptides that elicit T-cell responses (37, 38).

Various mechanisms could contribute to peptide immunodominance (6). It could also be favored by promiscuous peptide binding if, as is likely, binding encounters between unselected pMHC and naive, unselected T cells only rarely leads to activation of a T cell. MHC binding promiscuity allows the generation of large pMHC libraries from antigenic proteins, and the larger and more diverse the library, the greater the chance that one or a few of its pMHC will be bound strongly enough to the TCR on a T cell to stimulate the cell to proliferate and thereby identify an immunodominant peptide. Because somewhat different pMHC libraries may be generated by differences in antigen-presenting cells at various anatomic sites, the immunodominant peptides arising from a pathogen may well vary and depend on the route of infection (37, 38).

It is of interest to note similarities between promiscuous peptide binding by MHC proteins and by some heat shock proteins (Hsp70 family). DnaK, the bacterial homolog of ubiquitous Hsp70 proteins, binds with 5-nM to 5-pM affinity many short amino acid sequences in unfolded proteins, the consensus sequence of bound peptides consisting of four to five residues enriched in hydrophobic amino acids flanked by cationic residues. These sequences are typically separated in unfolded proteins by around 50–100 intervening residues (1, 2), a frequency not unlike that of octamers that bind to class I MHC proteins.

The enormous peptide binding promiscuity of MHC is to be expected, of course, because the proteins encoded by the few class I MHC genes in each individual (e.g., HLA-A, -B, and -C in humans and HLA-A, -B, -C, and -D in mice) have to be able to effectively present to an individual’s T cells short peptides generated from a great multitude of proteins, including those from exotic microbial pathogens never encountered in a species’ evolutionary history. Although many of the T cells produced in immune responses to an antigenic peptide may be elicited by only one or a few of the many peptides that are potentially generated from that protein, it is ironic that the adaptive immune system, which is noted for its great specificity, should depend on an initiating ligand–protein reaction that is among the most promiscuous known, ranking with the binding of unfolded proteins by some heat shock proteins.

**Materials and Methods**

**Model.** The processes shown in Fig. 1 can be represented by (Eq. 1)

\[
\frac{dM}{dt} = \frac{dM_0}{dt} - k_{\text{on}} \cdot P \cdot M + k_{\text{off}} \cdot MP - k_{\text{den}} \cdot M. \tag{1}
\]

where \( M \) refers to empty MHC molecules and \( dM_0/dt \) refers to the rate at which empty MHC molecules appear on the cell surface. \( k_{\text{on}} \) is the rate constant for \( P \) binding to empty MHC to form \( MP \), \( k_{\text{off}} \) is the rate constant for \( P \) dissociation from \( MP \), and \( k_{\text{den}} \) is the rate constant for denaturation of the empty MHC molecules (i.e., the rate at which they lose, essentially irreversibly, the ability to bind peptides).

The rate at which the pMHC complexes of interest (\( MP \)) are formed is (Eq. 2)

\[
\frac{dMP}{dt} = k_{\text{on}} \cdot P \cdot M - k_{\text{off}} \cdot MP. \tag{2}
\]

Eqs. 1 and 2 combine to (Eq. 3)

\[
\frac{dM}{dt} = \frac{dM_0}{dt} - \frac{dMP}{dt} - k_{\text{den}} \cdot M. \tag{3}
\]

The solution to Eq. 3, given in SI Text, yields the following formula (Eq. 4) for \( MP(t) \), the number of cell surface MP per cell formed by incubating cells for a specified time with the peptide at a known concentration. The derivation of Eq. 4 assumes there are no significant changes in peptide concentration or rate of appearance of empty MHC during the period of observation (Eq. 4):

\[
MP(t) = A \left( \frac{k_{\text{den}}}{a_1} - 1 \right) \left( 1 - e^{-t/a_2} \right) - \left( \frac{k_{\text{den}}}{a_1} - 1 \right) \left( 1 - e^{-t/a_2} \right). \tag{4}
\]

where \( A = k_{\text{on}} \times a_2 \), \( a_1 = 2.2 \times (e^{\gamma_1/2} - 1) \), \( a_2 = (k_{\text{on}}P + k_{\text{off}} + k_{\text{den}}) \), and \( h = k_{\text{off}}k_{\text{den}} \).

**Cells.** RMA-S cells were from American Type Cell Culture. DC2.4 cells, a gift from K. L. Rock (University of Massachusetts, Worcester, MA), are c-myc-transfected (immortalized) dendritic cells from B6 mice. Primary dendritic cells were derived from bone marrow of B6 mice by standard procedures (SI Text, R23).

**Monoclonal Antibodies.** Hybridoma cells producing the 25D1.16 antibody (39) were a gift from Ron Germain (National Institutes of Health, Bethesda, MD). The hybridoma that produces antibody Y3 (SI Text, R24) was from American Type Cell Culture. Both antibodies were purified from hybridoma cell culture supernatants on a protein A column. Antibodies were labeled as described in SI Text with \( ^{125}I \), AlexaFluor 680 (AF), or fluorescein (F), or they were doubly labeled with both \( ^{125}I \) and AF. The number of AF or F groups per antibody molecule was determined by UV absorption using a molar extinction coefficient for fluorescein = 77,000 (pH 7.3, 494 nm) and a molar extinction coefficient for AF = 187,400 cm−1 (679 nm). Protein concentrations were determined by bichromonic acid (BCA) assay or UV absorption at 280 nm and corrected for chromophore absorption at 280 nm by subtracting 0.2× absorption at 679 nm for AF or subtracting 0.2× absorption at 494 nm for fluorescein. In various preparations, there were 1.3–1.5 AF groups per antibody molecule and 1.7–3.38 F groups per antibody molecule.

**Exogenous Peptide Binding to Cells.** Typically, 200,000 cells in 180 μL RPMI 1640-based medium (10% heat-inactivated FCS) were mixed with 20 μL peptide at various concentrations in PBS or with PBS alone (control) in a total volume of 200 μL (in round-bottomed 96-well plates. After incubating the plates at 37 °C (5% CO2) for various times, they were centrifuged, and the cells were washed one time (with cold PBS) and stained on ice by first adding an Fc blocker and then after 10 min, F- or AF-labeled antibody. After 45 min on ice, the cells were washed and analyzed using a flow cytometer (FACSCaliber; BD Sciences) for F antibody-stained cells or a Liorc plate reader for AF antibody-stained cells (Liorc).

**Measurement of Number of Cognate pMHC per Cell (MP).** Standardized beads, with specified numbers of fluorescein equivalents per bead (Bang Laboratory), were run through the flow cytometer immediately before (and/or after) the stained cells. Calibration curves based on the beads’ fluorescence allowed...
conversion of fluorescence intensity of stained cells to moles of fluorescein per cell and hence (given the number of chromophore groups per antibody molecule), moles of antibody bound per cell. The latter value, multiplied by two to correct for bivalent antibody binding, yielded the number of antibody-bound MP per cell. The validity of this approach was indicated from the agreement found for MP values when T2–Kb cells were stained with fluorescein-labeled Y3 antibody or 125I-labeled Y3 (340,000 peptide–Kb complexes per cell vs. 320,000 complexes per cell). For cells stained with F-25D1.16, background staining of nonpulsed cells was negligible (<10%). For cells stained with F-Y3, however, it was necessary to subtract from their fluorescence values found for cells that had been treated identically but not peptide-pulsed. To test the validity of this procedure, RMA-S cells were loaded with OVA, and the resulting OVA–Kb complexes were measured with both antibodies; with F-25D1.16, there were 194,864 complexes per cell, and with F-Y3, there were 176,016 complexes per cell.

For experiments in which the cells were stained with AF antibody or antibody doubly labeled with 125I and AF, cell fluorescence was read in the infrared (700 nm) using the Odyssey system on a Licor plate reader. Cells stained with 125I-labeled antibody were read in a γ-detector (Packard), with the known specific radioactivity allowing determination of the number of antibody molecules bound per cell.

Peptides. These were synthesized and purified by HPLC at the Koch Institute Biopolymer Facility.

Dissociation of Peptides from MP on Cells. Cells were incubated with cytochalasin D (final concentration = 1–10 μg/mL) before and while they were being loaded with exogenous peptide. After the cytochalasin D-treated cells were peptide-pulsed, they were washed to remove unbound peptide, and the loss of cell surface MP was followed beginning about 0.5–1 h after the washed cells were resuspended in peptide-free medium. When pMHC dissociation was measured with Y3 antibody, Brefeldin A was added to a final concentration of 1 μg/mL to prevent the background cell surface level of Kb from increasing during the period of observation.

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