The first step of peptide selection in antigen presentation by MHC class I molecules

Malgorzata A. Garstka^a, Alexander Fish^b, Patrick H. N. Celie^b, Robbie P. Joosten^b, George M. C. Janssen^c, Ilana Berlin^a, Rieuwert Hoppes^a, Magda Stadnik^b, Lennert Janssen^a, Huib Ovaa^a, Peter A. van Veelen^c, Anastassis Perrakis^b, and Jacques Neefjes^{a,1}

Departments of ^aCell Biology and ^bBiochemistry, Netherlands Cancer Institute, 1066 CX Amsterdam, The Netherlands; and ^cDepartment of Immunohematology and Blood Transfusion, Leiden University Medical Center, 2333 ZA Leiden, The Netherlands

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Results

MHC class I molecules present a variable but limited repertoire of antigenic peptides for T-cell recognition. Understanding how peptide selection is achieved requires mechanistic insights into the interactions between the MHC I and candidate peptides. We find that, at first encounter, MHC I H-2K b considers a wide range of</sup> peptides, including those with expanded N termini and unfitting anchor residues. Discrimination occurs in the second step, when noncanonical peptides dissociate with faster exchange rates. This second step exhibits remarkable temperature sensitivity, as illustrated by numerous noncanonical peptides presented by H-2K^b in cells cultured at 26 °C relative to 37 °C. Crystallographic analyses of H-2K^b-peptide complexes suggest that a conformational adaptation of H-2K^b drives the decisive step in peptide selection. We propose that MHC class I molecules consider initially a large peptide pool, subsequently refined by a temperature-sensitive induced-fit mechanism to retain the canonical peptide repertoire.

antigen presentation | peptide binding | anchor residues | dynamics | entropy

MHC class I molecules present a wide array of peptides to cytotoxic T lymphocytes, allowing the immune system to scan for intracellular pathogens and mutated proteins. These peptides are not chosen at random, but rather are selected for their ability to bind to the polymorphic MHC class I peptidebinding groove. Antigenic peptide precursors are produced by the proteasome and further trimmed by cytosolic aminopeptidases. They are translocated into the endoplasmic reticulum (ER) lumen by the peptide transporter TAP that has broad peptide specificity. Peptides can be further trimmed in the ER lumen by ER aminopeptidases before selection by a defined MHC class I allele (reviewed in ref. 1). Only few peptides from a broad peptidome are presented by a given MHC class I allele. How a defined MHC I allele selects the correct peptides for presentation out of a large and diverse peptide pool is unclear.

In principle, MHC class I molecules could consider only "optimal" peptides and ignore the remainder of the TAPtranslocated peptidome. Alternatively, MHC class I could bind all available peptides followed by a selection step for the optimal candidates for presentation. To discriminate between these scenarios, we studied peptide association (on-rates) and dissociation (off-rates) from the mouse MHC class I molecule $H-2K^b$. To characterize the biophysical basis of discrimination between candidate peptides we complemented the kinetic data with five crystal structures of $H-2K^b$ with peptide variants. A comprehensive analysis of our in vitro data indicated that discrimination against suboptimal peptides by MHC class I may exhibit a strong temperature dependency, as illustrated by the H-2K^b peptidome from cells cultured at 26 °C versus 37 °C. We thus arrive at a two-step model for peptide selection by MHC class I molecules, which explains how not so "empty MHC class I molecules come out in the cold" (2).

Expanding the H-2K^b–Associated Peptidome Under Experimental Conditions. Peptide loading is a limiting factor for MHC class I complex assembly in cells (3). To study the initial steps of this process, we aimed to alleviate such limitation experimentally. First, the effect of temperature on the production of $H-2K^b$ -peptide complexes in vivo was assessed, using TAP-deficient (RMA-S) and control (RMA) mouse leukemia cells. The levels of peptidebound $H-2K^b$ increased significantly in RMA-S and RMA cells when cultured at 26 \degree C compared with 37 \degree C. (Fig. 1B), as observed previously (2, 4, 5).

To explore whether the increase in peptide-bound $H-2K^b$ is attributable to an altered peptide repertoire we evaluated binding of suboptimal peptides to $H-2K^6$. Optimal MHC class I peptides are typically nine residues in length with defined residues at particular positions that dock into specialized pockets within the MHC I peptide-binding groove (6, 7) (Fig. 1A). As a model peptide, we used FAPGNYPAL (Sendai virus nucleoprotein peptide 324–332) (5) and designed variants with anchor residues Tyr at position P(Ω –3) (third amino acid from the C terminus) and Leu at the C-terminal position $P(\Omega)$ substituted by either smaller or larger residues ([Table S1](http://www.pnas.org/lookup/suppl/doi:10.1073/pnas.1416543112/-/DCSupplemental/pnas.201416543SI.pdf?targetid=nameddest=ST1)). In addition, we replaced Gly with Lys (yielding FAPKNYPAL) to attach a biotin for the isolation of peptide complexes (Fig. 1A). Biotinylated peptide variants were incubated with lysates of RMA-S cells at

Significance

MHC class I molecules select and present a limited set of peptides from a broad repertoire provided by TAP. How MHC class I makes this selection is unclear. We show that MHC class I H-2K^b molecules initially bind many peptides because of highly flexible binding pockets. Peptide binding is followed by a selection step wherein a large fraction of these peptides is released, leaving the canonical peptides for presentation. The peptide presentation has a remarkable temperature dependency and explains the lowaffinity peptides found associated to MHC class I molecules in cells cultured at low temperature. Our data suggest that MHC class I goes through rounds of considering and rejecting peptides until peptides with high affinity are acquired for presentation.

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¹To whom correspondence should be addressed. Email: j.neefjes@nki.nl.

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Fig. 1. Binding and stability of H-2K^b in association with suboptimal peptides. (A) Schematic representation of H-2K b binding groove with 9-mer</sup> peptide. Anchor residues at position P($Ω$ –3) and P($Ω$) (red) bind into the C and F pockets, respectively. Residue P4 was exchanged to Lys (cyan) to attach biotin/TAMRA for isolation/detection purposes. (B) RMA and RMA-S cells were cultured at 26 °C or 37 °C and lysed and peptide-bound forms of H-2K^b were isolated using Y3 antibody and examined by immunoblotting (IB) with anti-exon8 antibody (recognizing all forms of H-2K^b). Efficiency of peptide binding is expressed as Y3-reactive H-2K^b as percentage of total H-2K^b. A representative experiment is shown; quantification represents the average \pm SD of two independent experiments. (C) Peptides with nonclassical anchor residues bind to H-2K^b in lysates. RMA-S cells cultured at 37 °C were lysed and incubated with biotinylated peptide. Peptides used were FAPKNYPAL or its derivatives: FAPKNXPAL [where X is 3,5-diiodo–Tyr (¥), Trp, Phe, Leu, or Ala]; FAPKNYPAX (where X is Trp, Phe, or Ala); or FAPKNXPAX (where X is Trp or Ala) [\(Table S1\)](http://www.pnas.org/lookup/suppl/doi:10.1073/pnas.1416543112/-/DCSupplemental/pnas.201416543SI.pdf?targetid=nameddest=ST1). H-2K^b–peptide complexes were isolated with streptavidin beads and examined by IB with anti-exon8 antibody. Binding is represented as the amount of isolated H-2K^b relative to total H-2K^b. A representative experiment is shown; quantification represents the average \pm SD of four independent experiments. (D) Thermal denaturation of various H-2K^b-peptide complexes was determined using the barycentric mean fluorescence (BCM) [\(Fig. S1\)](http://www.pnas.org/lookup/suppl/doi:10.1073/pnas.1416543112/-/DCSupplemental/pnas.201416543SI.pdf?targetid=nameddest=SF1). Fit of BCM to the first derivative yields melting temperature T_m [\(Fig. S1](http://www.pnas.org/lookup/suppl/doi:10.1073/pnas.1416543112/-/DCSupplemental/pnas.201416543SI.pdf?targetid=nameddest=SF1)D). The average T_m \pm SD of four independent experiments for all H-2K^b-peptide complexes is shown.

 4° C and isolated with streptavidin beads, and bound H-2K^b was examined by immunoblotting with exon8 (Fig. 1C). Exchanging the Tyr anchor at $P(\Omega-3)$ for a similar or larger amino acid reduced (Tyr→Phe/Trp) or prevented [Tyr→3,5-diiodo-L-Tyr (¥)] peptide binding to $\overline{H^22K^b}$. Replacement of Tyr by smaller amino acids (Leu/Ala) did not yield detectable $H-2K^b$ complexes. Substitution of Leu at $P(\Omega)$ with Phe was allowed (Fig. 1C). Other replacements (Leu→Trp/Ala) significantly decreased binding to $H-2K^b$. This suggests that under steady-state conditions binding of peptides to $H-2K^b$ is not strictly limited to the presence of canonical anchor residues, as would be expected from the binding rules extracted from the H-2K^b-associated peptidome (6).

Because steady-state association of modified peptides with H-2K^b in cell lysates was assayed at 4° C, no information was obtained regarding stability of the resulting complexes. Therefore, $H-2K^b$ complexes with altered peptides were produced [\(Table S2\)](http://www.pnas.org/lookup/suppl/doi:10.1073/pnas.1416543112/-/DCSupplemental/pnas.201416543SI.pdf?targetid=nameddest=ST2) for thermal stability experiments, where intrinsic Trp fluorescence and light scattering at 473 nm were monitored over a temperature range to measure protein unfolding and aggre-gation, respectively ([Fig. S1](http://www.pnas.org/lookup/suppl/doi:10.1073/pnas.1416543112/-/DCSupplemental/pnas.201416543SI.pdf?targetid=nameddest=SF1)A). H-2K^b-FAPGNYPAL complex showed two transitions, the first at ∼25–26 °C and the second at ∼46–52 °C, which corresponds to its reported melting temperature (T_m) (8). Aggregation was not observed before reaching the latter temperature range [\(Fig. S1](http://www.pnas.org/lookup/suppl/doi:10.1073/pnas.1416543112/-/DCSupplemental/pnas.201416543SI.pdf?targetid=nameddest=SF1) $A-C$ $A-C$). Two distinct folding events were also observed for $H-2K^b$ in complex with suboptimal peptides, with the first conformational change at ∼26 °C, and T_m ranged between 30 °C and 40 °C, depending on the peptide (Fig. 1D and [Fig. S1](http://www.pnas.org/lookup/suppl/doi:10.1073/pnas.1416543112/-/DCSupplemental/pnas.201416543SI.pdf?targetid=nameddest=SF1)D). There was one exception, H-2K^b-FAPGNYPAF, with a T_m of ~53 °C (similar to the original peptide FAPGNYPAL), consistent with the efficient binding to $H-2K^b$ in lysates (Fig. 1C).

These biophysical experiments suggest that $H-2K^b$ molecules can associate with a broader set of peptides than defined strictly on the basis of canonical anchor residues. Most of these peptides do not stabilize $H-2K^b$ sufficiently to survive at physiological temperatures.

Association and Disassociation Kinetics of 9-Mer Peptide Variants Binding to H-2K^b. To better understand the peptide selection process by MHC I we explored the kinetic association (k_{on}) and dissociation (k_{off}) rates of our noncanonical peptides. Because empty MHC class I molecules are unstable (9) , thermolabile H-2K^b–FAPGNAPAL complex (T_m ~30.5 °C, Fig. 1D) was used as a source of peptide-receptive MHC I (Fig. 2A). We monitored the binding in time of TAMRA-labeled peptides to $H-2K^b$ at 26 °C or 32 °C using fluorescence polarization (FP) (Fig. 2B) and calculated k_{on} , k_{off} , and the equilibrium dissociation constant $(K_d = k_{off}/k_{on})$ ([Table S3](http://www.pnas.org/lookup/suppl/doi:10.1073/pnas.1416543112/-/DCSupplemental/pnas.201416543SI.pdf?targetid=nameddest=ST3)). FAPKNYPAL is characterized by a high on-rate and a low off-rate, resulting in stable binding to H-2K^b (K_d ~16 nM, [Table S3\)](http://www.pnas.org/lookup/suppl/doi:10.1073/pnas.1416543112/-/DCSupplemental/pnas.201416543SI.pdf?targetid=nameddest=ST3) (10).

We first considered the effect of anchor residue size at $P(\Omega-3)$ and $P(\Omega)$ (Fig. 2 C and D and [Table S3\)](http://www.pnas.org/lookup/suppl/doi:10.1073/pnas.1416543112/-/DCSupplemental/pnas.201416543SI.pdf?targetid=nameddest=ST3). As in the cell lysate experiments (Fig. 1C), replacing Tyr for a small Ala residue

Fig. 2. Kinetics of modified peptide binding to H-2K b as a function of</sup> temperature. (A) Schematic representation of peptide exchange by H-2K^b. FAPGNAPAL from thermolabile H-2K^b-FAPGNAPAL complex is exchanged for TAMRA-labeled peptide. (B) Primary data of FP measurements of FAPGN**A**PAL in H-2K^b complex exchanged for FAPK(TAMRA)NYPAL. H-2K^{b_} FAPGNAPAL was used at concentrations indicated and peptide FAPK (TAMRA)NYPAL at 2 nM. From the binding curves, association rates (k_{eq}) , dissociation rates (k_{off}), and K_d were calculated (see [SI Experimental Proce](http://www.pnas.org/lookup/suppl/doi:10.1073/pnas.1416543112/-/DCSupplemental/pnas.201416543SI.pdf?targetid=nameddest=STXT)[dures](http://www.pnas.org/lookup/suppl/doi:10.1073/pnas.1416543112/-/DCSupplemental/pnas.201416543SI.pdf?targetid=nameddest=STXT)). (C–F) k_{on} and k_{off} values of peptides with variable anchor residues (C and D) or altered or extended N or C termini (E and F; for sequence details see [Fig. S2](http://www.pnas.org/lookup/suppl/doi:10.1073/pnas.1416543112/-/DCSupplemental/pnas.201416543SI.pdf?targetid=nameddest=SF2)C). FP measurements were performed at 26 °C and 32 °C, as in-dicated (also see [Table S3\)](http://www.pnas.org/lookup/suppl/doi:10.1073/pnas.1416543112/-/DCSupplemental/pnas.201416543SI.pdf?targetid=nameddest=ST3). Average k_{on} , and k_{off} values \pm SD from at least two independent experiments are shown. ND, no binding resolved at the highest concentration of H-2K^b (4 μ M).

abolished association with $H-2K^b$. Expanding the anchor residue with Trp or 3,5-diiodo-L-Tyr (¥) allowed efficient association, whereas dissociation increased with size of the anchor, resulting in low affinity. Importantly, exchange of the C-terminal Leu to a smaller (Ala) or larger (Phe/Trp) residue incurred minor effects on binding kinetics to $H-2K^b$, suggesting that the peptide-binding groove is receptive to more peptides than it ultimately presents. However, its flexibility is not unlimited, because $H-2K^b$ ignored the FAPKNWPAW peptide with two expanded anchors. These data indicate that $H-2K^b$ molecules can sample a variety of noncanonical peptides, as long as those alterations are compatible with productive association.

Comparison of peptide on- and off-rates (Fig. 2 C and D, respectively) revealed that although most peptides bind with similar on-rates both at 26 °C and 32 °C off-rates can differ significantly between these temperatures. This implies that despite some peptide selection by MHC class I at the level of initial binding (on-rate) the most significant filtering step is dictated by peptide dissociation (off-rate).

We next varied anchor residue charge (Tyr→Arg/Asp, Leu→ Arg/Asp, Fig. $S2A$ and B). The F pocket (with net neutral charge) responsible for accommodation of the C-terminal anchor residue (11) did not allow any charged amino acid at the P(Ω). The net negatively charged C pocket (12) accommodating residue at $P(\Omega-3)$ accepted positively but not negatively charged residues. Interestingly, although FAPKNRPAL has an on-rate comparable to that of FAPKNYPAL it displays a considerably higher off-rate, reducing overall binding affinity (Fig. $S2 \, A$ and B and [Table S3](http://www.pnas.org/lookup/suppl/doi:10.1073/pnas.1416543112/-/DCSupplemental/pnas.201416543SI.pdf?targetid=nameddest=ST3)), which may explain the underrepresentation of such peptides in the $H-2K^b$ peptidome under physiological conditions (6).

Finally, we considered the role of secondary anchors by replacing most residues in FAPKNYPAL for Ala. Such peptides still associated with $H-2K^b$, given the presence of canonical anchors, but showed higher off-rates. Rather surprisingly, although the single $P(\Omega-3)$ Tyr \rightarrow Ala mutant peptide FAPKNAPAL was ignored by H-2K^b (Fig. 2 C and D), poly-Ala peptides (FAPKAAAAL and AAPKAAAAL) had association rates similar to that of FAPKNYPAL. Their dissociation rates, however, were markedly elevated, resulting in compromised binding to $H-2K^b$ (Fig. $S2A$ and B and [Table S3\)](http://www.pnas.org/lookup/suppl/doi:10.1073/pnas.1416543112/-/DCSupplemental/pnas.201416543SI.pdf?targetid=nameddest=ST3). This suggests that peptides with a small anchor residue at P(Ω –3) are considered by H-2K^b if placed within an otherwise favorable sequence context. Furthermore, selection of peptides for presentation is determined mostly by higher dissociation rates of suboptimal peptides and consequent retention of those that manage to stay bound.

Peptide Size Selection During the First Binding Steps to H-2K $^{\rm b}$. ${\rm MHC}$ class I typically harbors peptides of nine amino acids in length, restricted by the hydrogen bonds formed between MHC class I and the α NH₂- and α COOH-terminal ends of the peptide. Longer peptides then bulge outward to favorably accommodate their N and C termini within the binding groove (13–16). To establish how modification of peptide length impinges on the individual binding parameters to $H-2K^b$ we reduced or increased peptide length (Fig. 2 E and F and [Table S3](http://www.pnas.org/lookup/suppl/doi:10.1073/pnas.1416543112/-/DCSupplemental/pnas.201416543SI.pdf?targetid=nameddest=ST3)). Removal of N-terminal Phe or N-terminal extension (AAA/HGE) resulted in low but detectable on-rates and high off-rates. Removal of C-terminal Leu abolished binding to $H-2K^b$, whereas C-terminal extensions affected k_{on} and k_{off} values in a sequence-dependent manner (compare extensions AAA and WSY). This suggests a role for C-terminal anchor docking for the first step of peptide binding to MHC class I, although both termini need to be properly positioned to produce a stable complex with $H-2K^b$. This could be due to the peptide $NH₂-$ and COOH-terminal groups that interact with the peptide-binding groove. We modified these groups to reduce the number of potential hydrogen donors or acceptors ([Fig. S2](http://www.pnas.org/lookup/suppl/doi:10.1073/pnas.1416543112/-/DCSupplemental/pnas.201416543SI.pdf?targetid=nameddest=SF2)C). Acetylation of the N terminus (*FAPKNYPAL) reduced on- and increased off-rates similarly

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Fig. 3. Binding of peptides with expanded anchor residues to H-2K^b. (A) The electron density maps of various peptides crystallized in complex with H-2K^b. The peptides are represented as sticks and the electron density maps are contoured at 1σ level, with a carve of 1.5 Å. The N-terminal region of FAPGN¥PAL is disordered as observed by poor electron density. Trp in the FAPGNWPAL peptide was modeled with two discrete conformers. K_d values are shown for peptide binding to H-2 K^b as determined by FP ([Table S3\)](http://www.pnas.org/lookup/suppl/doi:10.1073/pnas.1416543112/-/DCSupplemental/pnas.201416543SI.pdf?targetid=nameddest=ST3). (B) Surface representation of the space-filling model of H-2K^b with the peptide represented as sticks in top view. Expanded anchor residues exert only minimal effects on the peptide conformation in H-2K^b.

to removal of the N-terminal Phe residue. Amidation of the C-terminal COOH group (FAPKNYPAL#) or its conversion into an alcohol (FAPKNYPA&) did not significantly affect peptide on- and off-rates. Replacement of Leu by isopentylamine (FAPKNYPA^) to eliminate all C-terminal hydrogen donors reduced peptide on-rates twofold and enhanced off-rates 70-fold. Only when both N- and C-terminal ends were blocked (*FAPKNYPAL#) was the peptide was completely ignored (Fig. $2 E$ and F and [Table S3](http://www.pnas.org/lookup/suppl/doi:10.1073/pnas.1416543112/-/DCSupplemental/pnas.201416543SI.pdf?targetid=nameddest=ST3)). Our data suggest that the C-terminal amino acid (for all noncharged amino acids tested) has to fill the F pocket in order for the peptide to be considered by $H-2K^b$. The free N and C termini of the peptide are of lesser importance for initial binding unless both groups are prevented from making hydrogen bonds. This then suggests that both termini have to dock into the MHC class I peptide-binding groove for detection by our FP assays.

Collectively, these data suggest that MHC class I considers peptides of different sequences and sizes during the first binding step, as evidenced by measurable association constants. This, however, is followed by a second step, wherein peptides of canonical sequence and size remain due to low off-rates.

Structural Flexibility in the Peptide-Binding Groove of H-2K $^{\sf b}$. $\rm The$ peptide-binding groove should be flexible to accommodate expanded anchors $(17–22)$. To visualize this, we crystallized H-2K^b molecules in complex with the original FAPGNYPAL peptide and its variants (FAPGNWPAL, FAPGN¥PAL, FAPGNYPAF, and FAPGNYPAW). We failed to obtain crystals of $H-2K^b$ with FAPGNAPAL or FAPGNYPAA. The crystal structures were determined to resolutions from 2.0 to 2.8 Å, all with excellent R_{free} values in the 23.4–28.1 range, no Ramachandran outliers, and all belonging to the 99th or 100th percentile of the Mol-Probity score [\(Table S4](http://www.pnas.org/lookup/suppl/doi:10.1073/pnas.1416543112/-/DCSupplemental/pnas.201416543SI.pdf?targetid=nameddest=ST4)). In all structures the peptides could be resolved and fit into the electron density (Fig. 3A). No major changes in the $H-2K^b$ structures were observed as a function of different peptides, indicating that the pockets in the peptidebinding groove of $H-2K^b$ can accommodate amino acid side chains that are considerably larger than the canonical ones [\(Fig.](http://www.pnas.org/lookup/suppl/doi:10.1073/pnas.1416543112/-/DCSupplemental/pnas.201416543SI.pdf?targetid=nameddest=SF3) [S3](http://www.pnas.org/lookup/suppl/doi:10.1073/pnas.1416543112/-/DCSupplemental/pnas.201416543SI.pdf?targetid=nameddest=SF3)). A closer examination of the $H-2K^b$ peptide structures reveals differences that may explain the nonselective first peptide binding step and a second more selective step required for the selection of stable peptides.

An expanded anchor at $P(\Omega-3)$ hampers H-2K^b interactions with the peptide. Significant structural changes are observed within the $H-2K^b$ -FAPGN¥PAL complex, containing 3,5-diiodo-L-Tyr (¥). The Pro ring at P3 rotates ∼90° in comparison with the other structures, and three N-terminal residues (Phe, Ala, and Pro) become significantly disordered, as evidenced by lack of clear electron density. Additionally, \angle anchor slightly changes the width of the peptide-binding groove, especially around the N terminus and the middle part of the peptide ([Fig. S3](http://www.pnas.org/lookup/suppl/doi:10.1073/pnas.1416543112/-/DCSupplemental/pnas.201416543SI.pdf?targetid=nameddest=SF3) C and D). Fewer hydrogen bonds and hydrophobic interactions between FAPGN¥PAL and the H-2 K^b are observed than for other complexes [\(Table S5](http://www.pnas.org/lookup/suppl/doi:10.1073/pnas.1416543112/-/DCSupplemental/pnas.201416543SI.pdf?targetid=nameddest=ST5)). The diiodo-Tyr substitution has a relatively small twofold effect on the on-rate, indicating that the expanded anchor residue can easily access $H-2K^b$ during the initial binding step. The off-rate, however, is 40-fold higher (Fig. 2 C and D and Table $\overline{S3}$), implying that the peptide cannot adapt well to the $H-2K^b$ pocket.

An expanded anchor at $P(\Omega)$ alters the F pocket. Additional structural changes are observed within the H-2K^b-FAPGNYPAW complex, where the large side chain of Trp in position $P(\Omega)$ "pushes" away" Tyr116 of the H-2K^b peptide-binding groove to create space in the F pocket (Fig. $4A$). In the H-2K^b-FAPGNYPAL complex, Tyr116 interacts with Gln114 via a water molecule and thus shapes the C pocket, occupied by the Tyr anchor at $P(Ω-3)$ of the peptide. In addition, Tyr116 with Ile124 and Tyr123 comprise the F pocket (Fig. 4A), occupied by the Leu anchor at P($Ω$). Binding of FAPGNYPAW induces interactions between Tyr116 and Gln114. This plasticity in the $H-2K^b$ peptide-binding groove may allow binding of unusual anchor residues without a significant decrease in the association rate.

Dynamics and entropy are factors controlling stable interactions. Although FAPGNYPAW and other suboptimal peptides associate to $H-2K^b$ they are also swiftly released from $H-2K^b$ (Fig. 2 and [Table S3](http://www.pnas.org/lookup/suppl/doi:10.1073/pnas.1416543112/-/DCSupplemental/pnas.201416543SI.pdf?targetid=nameddest=ST3)). We considered the dynamics within the $\angle H^2 K^b$ peptide structures by exploring the X-ray data and generated a representative structural ensemble for each of the $H-2K^b$ peptide structures by time-averaged refinement, in which local molecular vibrations are sampled with molecular dynamics (23). Here, the dynamics are calculated from the X-ray data rather than being based entirely on force field calculations. Zooming into the H-2K^b peptide-binding groove with bound FAPGNYPAL revealed considerable flexibility of Tyr116. FAPGNYPAX peptide, where X is Phe or Trp, restricted the dynamics of Tyr-116 by increased filling of the F pocket (Fig. $4B$ and Fig. $S4B$). This implies that the F pocket can accommodate larger anchor residues owing to intrinsic flexibility of Tyr116—a conserved residue found in many MHC class I alleles (11). Tyr116 dynamics contributes to accommodation of the larger than canonical anchors also in C pocket as observed for FAPGNWPAL peptide. Beyond a certain

Fig. 4. Dynamics and the gatekeeper's role of Tyr116 in peptide binding. (A) Zoom-in into the C-terminal end of the peptide-binding groove of H-2K b </sup> in complex with FAPGNYPAL (in yellow) and FAPGNYPAW (in magenta). Model shows the peptide and H-2K b residues Gln114 as well as Tyr116,</sup> Tyr123, and Ile124 that form part of the C and F pockets, respectively. Trp of FAPGNYPAW repositions Tyr116 of H-2K^b. (B) Representative ensemble of peptides (Left) and H-2K^b amino acids 114–117 (Right). Note the selective restrained effect of expanded anchor residues at P(Ω–3) and P(Ω) on flexibility of H-2K^b amino acids Gln114 and Tyr116. The 3,5-diiodo-Tyr-containing peptide is largely disordered (quantification in [Fig. S4\)](http://www.pnas.org/lookup/suppl/doi:10.1073/pnas.1416543112/-/DCSupplemental/pnas.201416543SI.pdf?targetid=nameddest=SF4).

size of the anchor residue—as shown here for diiodo-Tyr (¥) long-range effects on the $H-2K^b$ peptide-binding groove and increased peptide flexibility are observed ([Fig.](http://www.pnas.org/lookup/suppl/doi:10.1073/pnas.1416543112/-/DCSupplemental/pnas.201416543SI.pdf?targetid=nameddest=SF4) $4B$ and Fig. [S4](http://www.pnas.org/lookup/suppl/doi:10.1073/pnas.1416543112/-/DCSupplemental/pnas.201416543SI.pdf?targetid=nameddest=SF4)B). This may drive the release of FAPGN¥PAL from H-2 K^b (Fig. 2D).

Peptides with Noncanonical Motifs Join the H-2K^b Peptidome at 26 °C. Because suboptimal peptides bind to $H-2K^b$ at 26 °C [\(Table S3\)](http://www.pnas.org/lookup/suppl/doi:10.1073/pnas.1416543112/-/DCSupplemental/pnas.201416543SI.pdf?targetid=nameddest=ST3), and because $H-2K^b$ -peptide complexes are more abundant at 26 °C than 37 °C, even in the absence of TAP (Fig. 1B) (2), $H-2K^b$ could present a significant fraction of suboptimal peptides at 26 °C. RMA cells were cultured for 3 days at either 26 °C or 37 °C, and H-2 K^b –associated peptidomes were analyzed by mass spectrometry, as described (24) . We identified 1,381 and 553 peptides bound to $H-2K^b$ molecules isolated from cells cultured at 26 °C and 37 °C, respectively, with 341 peptides in common (Fig. 5A and [Dataset S1\)](http://www.pnas.org/lookup/suppl/doi:10.1073/pnas.1416543112/-/DCSupplemental/pnas.1416543112.sd01.xlsx). NetMHC algorithm (25–27) predicted peptides presented by $H-2K^b$ at 26 °C to be worse binders (higher IC₅₀) relative to those presented at 37 °C (Fig. 5B), harboring differences in sequence and length. As expected, 8-mer, 9-mer, and 10-mer peptides constituted the largest pool at both temperatures. However, a substantial fraction of longer peptides $(25\%$ of total) was isolated from H-2K^b molecules from cells cultured at 26 °C, compared with only 3% at 37 °C (Fig. 5C).

Fig. 5. Characterization of H-2K^b-associated peptidome in cells as a function of temperature. Peptides associated with H-2K^b were isolated from RMA cells cultured at 26 °C or 37 °C and identified by mass spectrometry. (A) Venn diagrams shows the number of unique and shared peptides between H-2K^b peptidomes of RMA cells cultured at 26 °C and 37 °C (for peptide sequences see [Dataset S1](http://www.pnas.org/lookup/suppl/doi:10.1073/pnas.1416543112/-/DCSupplemental/pnas.1416543112.sd01.xlsx)). (B) Distribution and median IC_{50} values of peptides in the peptidome of H-2K^b from cells cultured at 26 °C or 37 °C predicted using the NetMHC algorithm (median $IC_{50} = 33$ for 26 °C and 20 for 37 °C). Distribution of predicted IC₅₀ values for the peptides identified at 26 °C and 37 °C were significantly different ($P = 0.0009$) according to the nonparametric Mann–-Whitney test. (C) Distribution according to size of the H-2K^b-associated peptidome of RMA cells cultured at 26 °C or 37 °C. The total pool at given temperature is set at 100%. (D and E) Sequence conservation of anchor residue at P(Ω–3) (D) and P(Ω) (E) in the H-2K^b–associated peptidome of RMA cells cultured at 26 °C or 37 °C. Pie charts illustrate the proportion of nonpolar aliphatic (Ala, Val, Leu, Ile, and Met, white), nonpolar aromatic (Phe, Trp, and Tyr, magenta), nonpolar special (Pro, Gly, and Cys, yellow), polar positive (His, Lys, and Arg, blue), polar negative (Asp and Glu, red), and polar neutral (Ser, Thr, Asn, and Gln, green) amino acids at P(Ω–3). Amino acids are represented in the following order: Ala, Val, Leu, Ile, Met, Phe, Trp, Tyr, Pro, Gly, Cys, His, Lys, Arg, Asp, Glu, Ser, Thr, Asn, and Gln starting from the 12:00 position on the pie chart.

Moreover, the presence of canonical anchor residues Phe or Tyr at P(Ω –3) was reduced from 57% (at 37 °C) to 35% (at 26 °C) (Fig. 5D). The presence of canonical aliphatic amino acids at P(Ω) was reduced from 93% (at 37 °C) to 76% (at 26 °C) (Fig. 5E).

Thus, the H-2 K^b peptidome at 26 °C has many aspects in common with the suboptimal peptides that bound efficiently to $H-2K^b$ but were also swiftly released in our biophysical experiments (Fig. 2 and [Table S3\)](http://www.pnas.org/lookup/suppl/doi:10.1073/pnas.1416543112/-/DCSupplemental/pnas.201416543SI.pdf?targetid=nameddest=ST3). These observations also indicate that at lower temperature the peptide repertoire of MHC class I can be expanded beyond the consensus sequences and lengths, resulting in presentation of a broader peptide pool.

Discussion

It has long been appreciated that one peptide transporter, TAP, feeds a wide variety of MHC class I alleles. Because these peptides may greatly differ in sequence, a central question therefore arises as to how a defined MHC class I allele $(H-2K^b)$ in our study) is able to select the correct peptides for presentation. One possible option could be that binding of only those peptides that are ultimately presented is considered. This should then result from an intrinsic quality of the MHC class I molecule. Alternatively, $H-2K^b$ may in fact first consider a range of available peptides, followed by a selection step for presentation

In an effort to advance our understanding of the events occurring during the initial confrontation of MHC class I molecules with candidate peptides we engineered systematic variations within a model peptide, FAPGNYPAL, to study the effects of altered sequence, length, and termini accessibility on peptide selection by $H - 2K^b$. Evaluation of association and dissociation parameters of a battery of peptides, containing targeted departures from the model, revealed that peptide acquisition by $H-2K^b$ takes place in two distinct phases. During the first phase, the peptide-binding groove of \hat{H} -2K^b accepts a wide range of candidates, including even amino acid side chains much larger than their optimal size, which implies significant flexibility in the anchor residue-accepting pockets (17, 19–22). Following the initial association event, ill-suited peptides exhibit greater propensity for dissociation and are thus selected against.

There are several reasons for selecting peptides for presentation. The pockets for anchor residues show considerable flexibility. This flexibility is in turn associated with higher entropic costs (Fig. 4), which may yield elevated dissociation rates and a compromised overall stability of the complex. This point is particularly apparent for Tyr116, an amino acid conserved in many MHC class I alleles, essential for tapasin-independent peptide loading (28, 29) and influencing MHC I–T-cell receptor interactions (30). Ensemble refinements calculated on the basis of X-ray diffraction data, where local low resolution essentially corresponds to higher local dynamics, indicate that mobility of Tyr116 is markedly restricted by larger than typical peptide anchors (Fig. 4B). These observations indicate how MHC class I molecules can accommodate a great variety of peptides in the first step of binding and subsequently release suboptimal candidates.

It stands to reason that if the entropic constraints within the peptide-binding groove contribute to higher dissociation rates of peptides harboring expanded anchors, then alleviating the necessity to maintain such flexibility by lowering the temperature during complex assembly could shift the peptidome equilibrium away from the cannon. Indeed, we observed a considerable reduction in dissociation rates for peptides acquired at 26 °C, relative to those at 32 °C, with on-rates remaining only marginally affected in this temperature range (Fig. 2). Moreover, comparing the peptidomes of $H-2K^b$ molecules from cells cultured at 26 °C and 37 °C also showed more elongated peptides and the loss of conservation of anchor residues (Fig. 5). This pattern is similar to that observed in cells where peptide loading is obstructed by tapasin deficiency or where peptide trimming is reduced by inactivation of endoplasmic reticulum aminopeptidase associated with antigen processing (ERAAP) (31, 32).

These observations hint at temperature-dependent conformational adaptations in the $H-2K^b$ peptide-binding groove, following the initial binding event. In fact, we detect a conformational change in H-2K^b at around 26 °C by Trp fluorescence [\(Fig. S1](http://www.pnas.org/lookup/suppl/doi:10.1073/pnas.1416543112/-/DCSupplemental/pnas.201416543SI.pdf?targetid=nameddest=SF1)). This may constitute a step wherein $H-2\tilde{K}^b$ molecules are able to stably acquire noncanonical peptides at 26 °C that would otherwise be entropically disfavored in the second phase of peptide consideration. Importantly, a shift of these MHC class I complexes to higher temperatures results in a swift release of suboptimal peptides, producing a high yield of peptide-receptive MHC class I molecules (4), an observation advantageous in vaccination studies (33, 34).

On the way to understanding the nature of peptide selectivity by MHC class I our work defines a number of simple rules: (i) noncanonical peptides with expanded anchor residues associate with rates similar to those of their canonical counterparts, owing to accommodation by the flexible pockets within the peptide-binding groove of MHC class I; (ii) peptides harboring incompatible charges are ignored, likely by simple repulsion; and (*iii*) stable complex formation between MHC class I and peptide requires each peptide terminus to contribute at least two hydrogen bonds, as well as the F pocket to be properly occupied by an amino acid side chain. The composition of the peptide–MHC class I repertoire is determined primarily by peptide-binding properties of MHC class I itself, but tapasin and ERAAP further shape the peptide repertoire presented (Fig. 5) (31, 32). Of note, Duan et al. (35) showed that cancer neopeptides that elicit CD8+ T-cell immunity may reflect the peptide pool detected at 26 °C

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and then expand the MHC class I-associated peptidome considered by the immune system. Collectively, this suggests that, at physiological temperature, MHC class I undergoes rounds of consideration and rejection until high-affinity peptides are acquired. This mechanism simultaneously allows for a wide range of sampling while ensuring that only the best available options are chosen for presentation.

Materials and Methods

Purification, Characterization, and Crystallization of H-2K^b Peptide Complexes in Vitro. H-2K^b was produced in *Escherichia coli*, refolded, and purified. For kinetic measurements, H-2K^b-FAPGNAPAL complexes were incubated at 26 °C or 32 °C with various TAMRA-labeled peptides for FP experiments. Proteins were crystallized and diffraction data solved by molecular replacement to 2.0–2.8 Å resolution. Protein dynamics were done by ensemble refinement modeling (23).

H-2K^b-Associated Peptidome. H-2K^b molecules were isolated from 10¹⁰ RMA cells, cultured at 26 °C or 37 °C for 72 h. Peptides were analyzed by a conventional Agilent 1100 gradient HPLC system coupled to a Q Exactive mass spectrometer and matched against the International Protein Index database using the Mascot search engine.

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