Title: Previously Hidden Dynamics at the TCR-pMHC Interface Revealed

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**Keywords:** MHC | TCR | conformational dynamics | molecular dynamics

Running Title: TCR-pMHC dynamics

### **Abstract**

A structural characterization of the interaction between  $\alpha\beta$  T cell receptors (TCR) and cognate peptide-MHC (pMHC) is central to understanding adaptive T cell mediated immunity. X-ray crystallography, although the source of much structural data, traditionally provides only a static snapshot of the protein. Given the emerging evidence for the important role of conformational dynamics in protein function, we interrogated 309 crystallographic structures of pMHC complexes using ensemble refinement, a technique that can extract dynamic information from the X-ray data. We found that a large number of systems contain significant conformational plasticity that confounds, and in some cases overturns, previous functional interpretations that were based on a single static structure. Notable examples include the interpretation of differences in the disease association of HLA subtypes, the relationship between peptide prominence and TCR recognition, the role of conformational flexibility in vaccine design, and discriminating between induced fit and conformational selection models of TCR binding. We show that the currently widespread practise of analyzing pMHC interactions via the study of a single crystallographic structure fails to properly distinguish real results from noise, and argue that biological observations should be explained through the properties of ensembles rather than isolated structures, as these are less prone to observer bias. This new analysis therefore not only raises significant questions regarding the interpretation of decades of structural data, but also provides previously missing information concerning the dynamics of existing characterized TCR-pMHC interactions.

Introduction

αβ T cell receptors (TCR) expressed on the surface of cytotoxic T cells engage endogenous peptide antigens from infected or malignant cells that are displayed by major histocompatibility complex (MHC) proteins. Productive engagement of the TCR with the peptide-MHC (pMHC) leads to T cell activation and intracellular signal transduction via the CD3 signaling subunits. The TCR-pMHC molecular recognition event is thus key to the cell-mediated adaptive immune response and has been widely studied using X-ray crystallography. These studies have focused on the conformation of the peptide when bound to the MHC and the specific interactions between peptide and MHC side-chains, with or without bound TCR. While providing insights into the key features of TCRpMHC recognition, the detailed examination of a single static structure by X-ray crystallography reveals little about the conformational dynamics and degree of flexibility of the system in solution<sup>1</sup>-<sup>3</sup>. Given that protein flexibility is inextricably linked to protein function, consideration of protein dynamics is essential in order to achieve a full understanding of pMHC and TCR-pMHC binding. Recent years have seen an increasing use of atomistic molecular dynamics (MD) simulations, which can probe the conformational flexibility of pMHC systems by computing iterative numerical solutions of Newton's equations of motion as the system evolves over time<sup>4</sup>. Unfortunately, the heavy computational demands of MD limits the technique to examination of relatively short time spans (usually less than a microsecond), and thereby excludes many larger motions that occur over longer, more biologically relevant timescales. Results of MD are also somewhat difficult to validate given the known imperfections of existing force fields<sup>5-7</sup>

To circumvent some of these limitations, ensemble X-ray structure refinement can be employed<sup>8</sup>. Instead of the usual practice of refining against a set of crystallographic reflections to produce a single protein structure, ensemble refinement combines the experimentally-derived reflection data with the results of a short, steered MD simulation to produce an ensemble of similar but distinct conformations of the protein. The structural diversity observed within the resulting ensemble thus

provides information about the conformational flexibility of the protein, and in many cases improves the agreement between the structure and data as measured by the crystallographic free R-factor ( $R_{\text{free}}$ ). Notable recent applications of this method include an investigation of the conformational landscape of an insect carboxylesterase protein over the course of its catalytic cycle<sup>9</sup>, and the discovery of a self-inhibition mechanism mediating the transition between active and inactive protein of human factor  $D^{10}$ . Therefore, this approach may offer significant new insights into protein function where crystallographic data already exists. Here we apply ensemble refinement to investigate protein dynamics at the TCR-pMHC interface.

## Results

To acquire a sample of appropriate pMHC systems whose dynamics could be investigated using ensemble refinement, we first searched the Protein Data Bank (PDB) for pMHC complexes with a resolution of better than 3 Ångströms. Some of the resulting entries were rejected owing to lack of the necessary crystallographic reflection files, leaving 284 pMHC structures which were subjected to ensemble refinement as described in Methods. Each of these systems consisted of the heavy and light chains of the MHC molecule, the bound peptide, and also the  $\alpha$  and  $\beta$  chains of the TCR when this was present. The ensemble refinement process takes as input a single crystallographic structure and returns as output an ensemble of several dozen structures that collectively fit the X-ray diffraction data roughly as well as the original single structure. Of all the systems we subjected to this process, we informally estimated that about half contained conformations noticeably different from the original PDB structures. It was therefore necessary to reduce this to a more manageable number of systems that we could subject to detailed structural analysis. First, in order to eliminate possible confounds we discarded all systems with bound ligands and missing peptide residues, leaving a total of 194 ensembles. Next, we ranked these systems according to the degree of conformational flexibility displayed by the peptide and MHC peptide-binding helices, as measured by the root mean square fluctuation (RMSF) of these residues calculated over the ensemble structures. We then focused our analysis only on published structures of human MHCs (human leukocyte antigens or HLA), and selected the top 30 most flexible MHC class I ensembles from the RMSF ranking. For each of these systems, we compared our results in detail to those reported in each original publication.

In comparing our ensemble results with the analyses reported in the original papers, we found it necessary to run an additional 25 ensembles to properly interpret our ensemble results in relation to structural and functional claims made in the literature. Of the 30 systems we examined in detail, we selected 12 illustrative cases in which our ensemble results provided the greatest functional insights

pertinent to the claims made in the original literature. Given that in some cases the original study compared multiple pMHC systems, these 12 examples included a total of 20 distinct PDB structures, all of which are summarized in Table 1. In our analysis of these systems, we highlight several new insights into TCR-pMHC engagement yielded by ensemble analysis compared to traditional approaches that consider only a single static structure. To supplement our ensemble results, five of the systems were selected for analysis in standard all-atom MD simulations, and the results compared to the corresponding ensembles.

**Table 1.** pMHC systems investigated by ensemble refinement.

			R <sub>free</sub>						
PDB	Res (Å)	PDB	Phenix refine <sup>†</sup>	Ensemble	$\Delta R_{\rm free}^{\&}$	Ensemble size	Class I MHC (HLA-)	Peptide Sequence	Peptide Name
1TVH	1.80	0.247	0.196	0.197	0.001	75	A*02:01	IMDQVPFSV	gp100 <sub>209</sub>
2BNQ	1.70	0.253	0.228	0.202	-0.026	80	A*02:01	SLLMWITQV	ESO 9V
3H9H*	2.00	0.226	0.200	0.205	0.005	42	A*02:01	MLWGYLQYV	Tel1p
3KLA	1.65	0.239	0.224	0.223	-0.001	56	A*02:01	SLLMWITQL	ESO 9L
3GSO	1.60	0.229	0.217	0.208	-0.009	60	A*02:01	NLVPMVATV	NLV
3MRC	1.80	0.223	0.210	0.207	-0.003	43	A*02:01	NLVPMCATV	NLV <sub>V6C</sub>
3MRD*	1.70	0.226	0.210	0.217	0.007	63	A*02:01	NLVPMGATV	NLV <sub>V6G</sub>
3MRG	1.30	0.202	0.160	0.180	0.020	89	A*02:01	CINGVCWTV	CIN
3MRH	2.40	0.308	0.264	0.269	0.005	34	A*02:01	CISGVCWTV	CIN <sub>N3S</sub>
3MRL	2.41	0.279	0.250	0.252	0.002	30	A*02:01	CINGVVWTV	CIN <sub>C6V</sub>
2A83*	1.40	0.150	0.133	0.163	0.030	125	B*27:05	RRRWHRWRL	pGR
1UXS*	1.55	0.177	0.192	0.204	0.012	89	B*27:05	RRRWRRLTV	pLMP2
1K5N	1.09	0.148	0.149	0.168	0.019	131	B*27:09	GRFAAAIAK	m9
3LKN*	2.00	0.252	0.244	0.223	-0.021	56	B*35:01	LPFERATIM	1918-NP <sub>418</sub>
3LKP	1.80	0.244	0.243	0.226	-0.017	50	B*35:01	LPFDKSTIM	1972-NP <sub>418</sub>
3LKR	2.00	0.266	0.254	0.236	-0.018	25	B*35:01	LPFERATVM	2009-NP <sub>418</sub>
2FZ3	1.90	0.275	0.251	0.248	-0.003	60	B*35:08	HPVGEADYFEY	HPVG
3KPN	2.00	0.244	0.231	0.229	-0.002	28	B*44:03	EEYLQAFTY	allotope
3KPP	1.90	0.230	0.225	0.216	-0.009	38	B*44:05	EEYLQAFTY	allotope
3KPQ	1.84	0.269	0.263	0.242	-0.021	43	B*44:05	EEYLKAWTF	mimotope

<sup>\*</sup> Systems for which MD was also run.

### Conformational polymorphism and peptide variants

A common goal in studying HLA-A\*02 systems is to determine the structural basis for differential TCR recognition of peptide or MHC variants, and the effect of such variations on T cell activation. For example, Chen et al<sup>11</sup> compared two peptide analogs of the NY-ESO-1 tumor antigen peptide, ESO 9L (PDB ID 3KLA) and ESO 9V (PDB ID 2BNQ), both bound to HLA-A\*02:01. The peptides are generally very similar, except that HLA-A\*02-ESO 9V has a six-fold greater binding affinity for the 1G4 TCR compared to HLA-A\*02-ESO 9L. The authors attribute this affinity difference to the movement of peptide position 9 (P9) and a reverberating shift in the main chain of

<sup>†</sup> Using the structure sourced from PDB\_REDO

 $<sup>^{\&</sup>amp;}\Delta R_{\text{free}}$  is derived from subtracting the ensemble refinement  $R_{\text{free}}$  from the Phenix refinement  $R_{\text{free}}$ .

residues P4-P9 of the ESO 9L peptide (Figure 1A). This difference is caused first by the greater length of the P9-Leu residue compared to P9-Val, and second by the I6 residue blocking the Tyr116-Arg97 conformational switch, which can produce an alternate conformation<sup>12</sup> (Figure 1A, coral/steel blue) that would allow the longer P9-Leu residue to fit in the groove. Our ensemble results (Figure 1B), however, indicate that the ESO 9L peptide is in fact easily flexible enough for the 'forbidden' Tyr116-Arg97 flip to occur, thereby enabling the peptide to adopt a wide range of conformations. The ESO 9V ensemble, by contrast, does not show such flexibility (data not shown), indicating that the affinity differences the authors attributed to structural differences may in fact be a result of varying levels of peptide flexibility. Similar bond flips have been observed in other ensembles of HLA-A\*02:01 systems. Borbulevych et al<sup>13</sup>, for instance, found that a melanoma antigen (gp100<sub>209</sub>) bound to HLA-A\*02:01 (PDB ID 1TVH) in two distinct conformations, differing in the bond flip of one key side chain (P7-Phe) (Figure 1C). Our ensemble results support the existence of both of these alternate conformations, though the large number of additional conformations observed highlight that for these systems it may be more accurate to speak of conformational polymorphism than conformational dimorphism (Figure 1D).

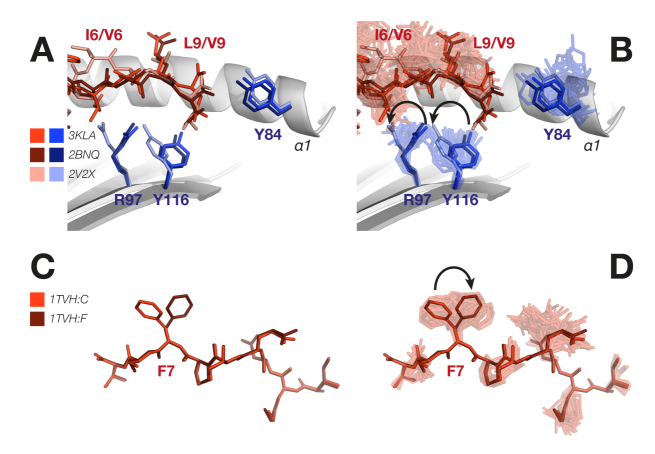


Figure 1: Conformational polymorphism and peptide variants

**A.** An overlay of HLA-A\*02:01-ESO 9L (red/blue) (PDB ID 3KLA) and HLA-A\*02:01-ESO 9V (brown/navy) (PDB ID 2BNQ), and an alternate 'forbidden' conformation of a similar peptide (SLFNTVATL, pale red/pale blue) bound to HLA-A\*02:01 (PDB ID 2V2X), compared to **B.** ensemble results of HLA-A\*02:01-ESO 9L showing high flexibility of the peptide, and MHC side chains undergoing the 'forbidden' bond flips. The MHC α2-helix is omitted. **C.** Two conformations of a melanoma antigen (gp100<sub>209</sub>) observed in different molecules of the crystallographic asymmetric unit (PDB ID 1TVH, chains C and F), and **D.** ensemble results from a single molecule showing interconversion between these conformations, in particular the side chain rotation observed in P7-Phe. MHC α-helices have been omitted.

## Conformational populations of pMHC

As in the case of HLA-A\*02 systems, analyses of HLA-B\*27 pMHC systems which fail to incorporate the effect of conformational flexibility are liable to make inaccurate ascriptions of the basis of functional differences. Fiorillo et al<sup>11</sup>, for example, compared the structure of HLA-B\*27:05 (PDB ID 1UXW) and HLA-B\*27:09 (PDB ID 1UXS) bound to the Epstein-Barr virus (EBV)-latent membrane protein 2 (LMP2, termed pLMP2). These two alleles differ only by a single residue, with Asp116 in HLA-B\*27:05 substituted with His in HLA-B\*27:09. The authors observed a substantial difference in the conformation of the peptide between the two alleles (Figure 2A), which they argue may contribute to the observed differential TCR recognition. Ensemble results for these systems, however, show that while the two peptide conformations are not observed to fully interconvert, the peptide side chains are highly flexible (Figure 2B), especially around the P4-P7 residues where the largest conformational difference is observed. These results indicate that the true structural differences between the two MHC alleles may not be quite so drastic as claimed, particularly when it is considered that peptide flexibility at physiological temperatures will almost certainly exceed that observed at cryogenic temperatures used in crystallographic data collection. Further support for this is observed in our MD results for the HLA-B\*27:09 system (PDB ID 1UXS), which show a similar pattern of flexibility to that observed in the ensembles, with even clearer evidence of interconversion between the two conformations (Figure S1A and S1B). Taken together, our ensemble and MD results indicate that at physiological temperatures, the two distinct conformations observed by Fiorillo et al. are most likely simply two members of a large ensemble of conformations that the peptide adopts when bound to the MHC. It is thus unlikely that static structural differences in peptide binding are solely responsible for the differential immunogenicity between the two pMHC. Rather, it is possible that the peptide occupies particular conformations more frequently when bound to one MHC allele compared to the other, which in turn influences their relative immunogenicities. It may therefore be more realistic to examine how conformational

populations in the pMHC are perturbed by polymorphisms, and how this altered energy landscape affects TCR engagement.

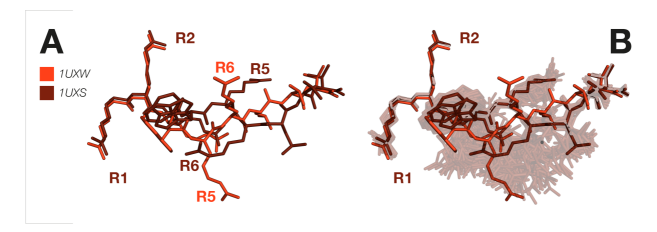


Figure 2: Conformational populations in MHC alleles

**A.** Superposition of original structures of Epstein-Barr virus pLMP2 peptide bound to HLA-B\*27:05 (red) (PDB ID 1UXW) and HLA-B\*27:09 (brown) (PDB ID 1UXS) showing the different conformations. **B.** Ensemble results of the HLA-B\*27:09 system, showing the degree of flexibility and possibility of interconversion of the two binding modes. MHC α-helices have been omitted.

### **Overinterpretation of static interactions**

A differential T cell response is observed when HLA-B\*27:05 and HLA-B\*27:09 present the same pLMP2 EBV peptide. By inference, a differential T cell response to arthritogenic peptides may be the reason why HLA-B\*27:05, but not HLA-B\*27:09, is associated with spondyloarthropathies<sup>15</sup>. To determine whether structural differences could account for the allele-specific disease association, Hülsmeyer et al.<sup>16</sup> compared the structures of HLA-B\*27:05 and HLA-B\*27:09, each bound to a token peptide (m9). In their analysis they identify several subtle structural differences between the F-pocket regions of the two alleles, related to the exact position and orientation of P9-Lys, several of the surrounding MHC side-chains, and the network of water molecules mediating salt bridges between these residues (Figure 3A). They suggest such differences alter the dynamics of the HLA-

B\*27:09 F-pocket region and accordingly could affect TCR recognition. Our ensemble analysis, however, suggests a greater degree of flexibility of the F-pocket region of HLA-B\*27:09 (PDB ID 1K5N), with all of the indicated side-chains displaying considerable variation in position and orientation (Figure 3B). Given this substantial intrinsic flexibility, the network of hydrogen bonds identified by the authors most likely exists only transiently. Our ensemble results therefore indicate that the difference in disease association of the two HLA-B\*27 alleles is unlikely to be merely the result of the static structural differences reported in the study. Further work is necessary to determine whether the differences can instead be explained in terms of differing patterns of side-chain and peptide conformational variability.

A study by Rückert et al<sup>17</sup> also used X-ray crystallography to examine the association of HLA-B\*27:05 with spondyloarthropathies, and hence investigate the potential role of molecular mimicry and T cell cross-reactivity in pathogenesis more generally. They determined the structure of HLA-B\*27:05 bound to a glucagon-derived self-peptide (pGR) (PDB ID 2A83). pGR was selected based on its sequence similarity to pVIPR and pLMP2, which are respectively self and foreign peptides for which a proportion of T cells cross-react in ankylosing spondylitis patients compared to healthy controls. The aim was to determine if pGR could adopt the structurally similar and unusual conformation observed of both pVIPR and pLMP2 when bound to HLA-B\*27:05, thereby indicating that cytotoxic T cell cross-reactivity may be due to molecular mimicry. Two pGR conformations were observed in the structure, and a network of contacts supporting those conformations reported (Figure 3C). Both pGR conformations were also observed in our ensemble, as well as a wide range of intermediate conformations and further variants (Figure 3D). We conducted MD on this system and found a very similar overall pattern of peptide flexibility to that seen in the ensembles (Figure S1C and S1D), further emphasizing that many hydrogen-bonding networks in pMHC systems are not stable at physiological temperatures, but are in a constant state of dynamic flux.

In order to investigate the influence of MHC polymorphism on peptide conformation and the T cell response, Miles et al<sup>18</sup> determined the structures of HLA-B\*35:01 (PDB ID 2FYY) and HLA-B\*35:08 (PDB ID 2FZ3) bound to the same EBV peptide (HPVG) (Figure 3E). These two alleles differ only at position 156, with Leu156 found in HLA-B\*35:01 compared with Arg156 in HLA-B\*35:08. The authors identified a set of hydrogen bonds between the P5-Glu residue and two MHC side chains of HLA-B\*35:08, which were not found in HLA-B\*35:01. The authors argued that these hydrogen bonds were likely responsible for the 'relatively rigid' nature of the peptide when bound to HLA-B\*35:08 (as determined by B factor analysis), compared to its very high flexibility when bound to HLA-B\*35:01, and show that peptide mobility influences the T cell response. Our ensemble results, however, indicate that HPVG has considerable flexibility when bound to HLA-B\*35:08 (Figure 3F), with the stabilizing interactions identified by the authors present only some of the time. As such, even if these interactions are involved in stabilizing the peptide, such stabilization clearly comes in varying degrees, and may still result in an overall highly flexible system.

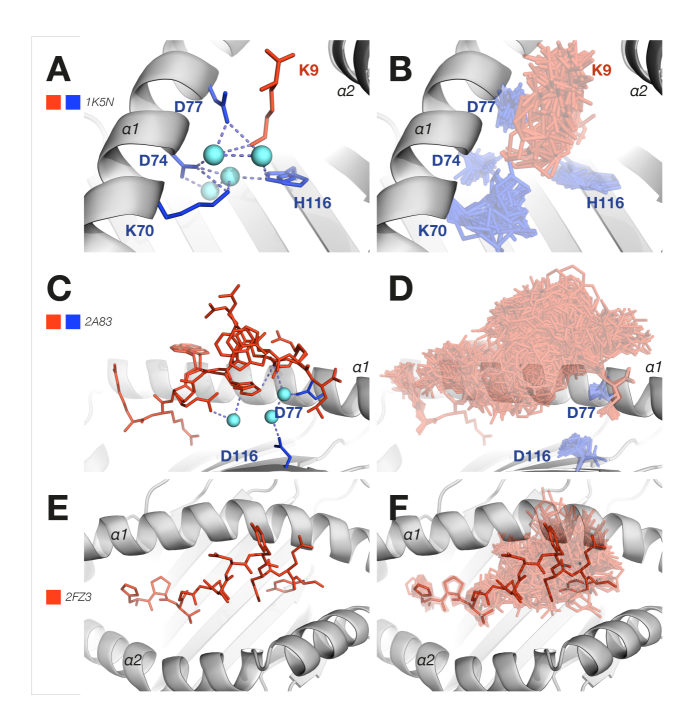


Figure 3: Overinterpretation of static contacts

**A.** Original structure of HLA-B\*27:09-m9 (PDB ID 1K5N) with proposed stabilizing bonding network of the F-pocket region, compared to **B.** ensemble results, illustrating how many of the hydrogen bonds (shown in dashed lines) identified in the original structure are not stable over the ensemble. Water molecules are shown as light blue spheres. MHC  $\alpha$ -helices have been omitted. **C.** In the original structure two conformations of the glucagon-derived self-peptide were observed when bound to HLA-B\*27:05 (PDB ID 2A83) showing the proposed conserved hydrogen-bonding network. Hydrogen bonds are depicted as dashed lines, and water molecules are shown as light blue spheres. **D.** Ensemble results for the same system showing the hydrogen-bonding network is not conserved over the ensemble. The MHC  $\alpha$ 2-helices have been omitted. **E.** Original structure (PDB ID 2FZ3) of Epstein-Barr virus peptide HPVG bound to HLA-B\*35:08 which the authors describe as 'relatively rigid', compared to **F.** the ensemble results of the same system showing considerable conformational variability of this peptide.

## Prominence of MHC-bound peptides in TCR recognition

Reiser et al<sup>19</sup> sought to identify pMHC structural features that correlate with pMHC immunogenicity by examining eight different viral and tumor peptides bound to HLA-A\*02:01. While no linear relationship was found between peptide solvent exposure and the frequency of pMHC-specific T cells, several structural differences between the peptides were identified as likely causes of varying pMHC antigenicity, suggesting an optimal intermediate degree of prominence<sup>19</sup>. Specifically, structural differences between the peptides of  $NLV_{V6C}$  (PDB ID 3MRC) and  $NLV_{V6G}$ (PDB ID 3MRD) variants compared to the wild type NLV (PDB ID 3GSO), were attributed to the absence of the buried stabilizing P6-Val side chains of the modified peptides (Figure 4A). Our ensemble analysis, however, shows that both modified NLV peptides (especially the V6G variant) exhibit considerable flexibility around the P5 and P6 residues (Figure 4A-D), which suggests that when intrinsic structural flexibility is properly considered these variants are not as structurally dissimilar as suggested by an analysis of static structures. This finding is further supported by our MD simulation results of the NLV<sub>V6G</sub> peptide system, which also showed considerable variation around these residues (Figure S1E and S1F). In a similar analysis of static structures of HLA-A\*02:01 bound to CIN (PDB ID 3MRG), CIN<sub>N3S</sub> (PDB ID 3MRH) or CIN<sub>C6V</sub> (PDB ID 3MRL) variants (Figure 4E), the authors found that the frequency of pMHC-specific T cells is influenced by peptide prominence from the cleft, peptide mobility, and by the altered conformation of the MHC Gln155 side chain between the systems. Our ensembles show that both the peptide prominence and the position of the Gln155 residue vary considerably within each system (Figure 4F-H), thereby highlighting the fact that peptide prominence cannot be determined independently of peptide mobility, but must be considered over a range of conformations in order to accurately characterize their potential relationship with TCR recognition.

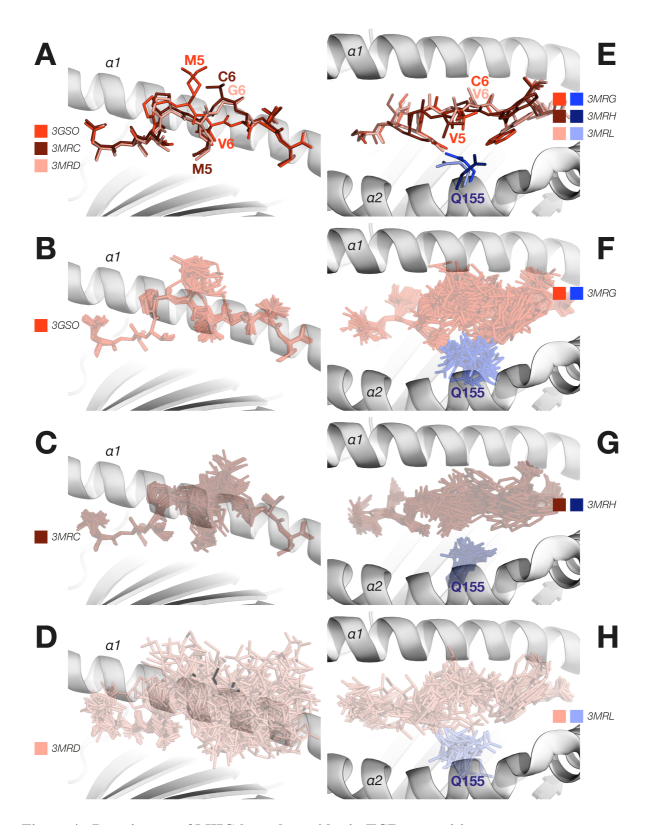


Figure 4: Prominence of MHC-bound peptides in TCR recognition

**A.** Original peptide conformations of HLA-A\*02:01-NLV (red) (PDB ID 3GSO), HLA-A\*02:01-NLV<sub>V6C</sub> (brown) (PDB ID 3MRC), and HLA-A\*02:01-NLV<sub>V6G</sub> (pale red) (PDB ID 3MRD) in the original structures, compared to ensemble results for **B.** HLA-A\*02:01-NLV, **C.** HLA-A\*02:01-NLV<sub>V6C</sub>, and **D.** HLA-A\*02:01-NLV<sub>V6G</sub>, showing that the variability within the NLV<sub>V6G</sub> variant ensemble substantially exceeds the variation between any of the systems. The MHC  $\alpha$ 2 helix is omitted in all left-hand side panels. **E.** An overlay of the original peptide conformations of HLA-A\*02:01-CIN (red/blue) (PDB ID 3MRG), HLA-A\*02:01-CIN<sub>N3S</sub> (brown/navy) (PDB ID 3MRH),

and HLA-A\*02:01-CIN<sub>C6V</sub> (pale red/pale blue) (PDB ID 3MRL), compared to ensemble results for **F.** HLA-A\*02:01-CIN, **G.** HLA-A\*02:01-CIN<sub>N3S</sub> and **H.** HLA-A\*02:01-CIN<sub>C6V</sub>, showing that the variability within each ensemble is similar in magnitude to the variability between systems. The highly flexible P5 and P6 residues are found around the center of the peptide.

### Implications of conformational flexibility for vaccine design

Gras et al<sup>20</sup> sought to identify the structural basis for why the 2009 pandemic influenza virus, A/Auckland/1/2009(H1N1), exhibits a relatively high rate of cross-reactive CD8+ T cells with the 1918 strain A/Brevig Mission/1/1918(H1N1), but much lower cross-reactivity with seasonal strains from intermediate years (e.g. 1947 and 1972). They examined the structures of HLA-B\*35:01 complexed with six nucleoprotein (NP)<sub>418</sub> peptides derived from influenza virus strains from six different years spanning the period 1918 to 2009. Their interpretation of the crystallographic data is that the solvent exposed P5-Arg residue common to both the 1918-NP<sub>418</sub> (PDB ID 3LKN, Figure 5A) and 2009-NP<sub>418</sub> (PDB ID 3LKR, Figure 5C) peptides produces a different surface recognition profile compared to the Asp-Lys (DK) motif found in NP<sub>418</sub> peptides from intermediate years (eg 1972, Figure 5E), and that this may partly explain the differential CD8+ T cell cross-reactivity (Figure 5A for 1918 strain, 5E for 1972 strain, and 5C for 2009 strain). In particular, they suggest the P5-Lys found in the NP<sub>418</sub> of intermediate years (1947 and 1972) moves further away from the neighboring MHC residue Gln155, breaking the hydrogen bond found between the P5-Arg and Gln155 residues in 1918-NP<sub>418</sub> and 2009-NP<sub>418</sub>. Examination of the ensemble results, however, show that the P5 residues for all three systems exhibit considerable flexibility, with hydrogen bonds formed with the Gln155 residue in some of the ensemble structures but not others (Figure 5B for 1918 strain, 5F for 1972 strain, and 5D for 2009 strain).

We further explored the dynamics of HLA-B\*35:01-1918-NP<sub>418</sub> through MD, which showed even greater flexibility of the MHC Arg62 and Gln155 side chains than observed in the ensemble (Figure S1G and S1H). Both the ensemble and the MD results clearly indicate that the mobility of the 1918-NP<sub>418</sub> peptide when bound to HLA-B\*35:01 is greater than any differences found between the

systems of different years, thereby casting significant doubt on many of the fine structure interpretations of the singular crystallographic model. The authors also identify two distinct conformations of the Arg62 residue found in different years, but once again, both conformations are found in all three of our ensembles. While the original study concluded that identifying key solvent-exposed residues might be important for vaccine design, our results indicate that a more careful comparison of the conformational dynamics of the peptide variants is required in the development of new vaccines, since conformational flexibility is a key characteristic feature of these pMHC systems.

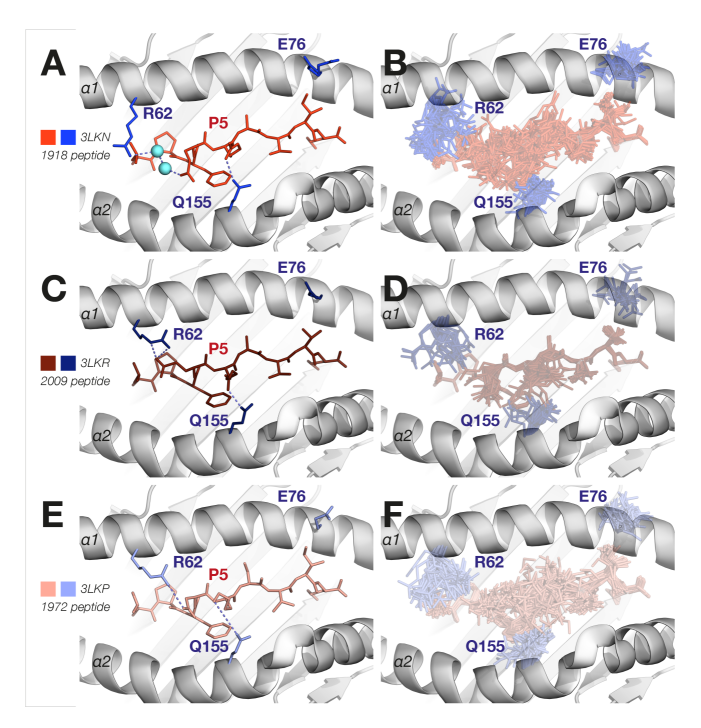


Figure 5: Implications of conformational flexibility for vaccine design

**A.** Original structure of 1918 influenza NP peptide bound to HLA-B\*35:01 (PDB ID 3LKN) compared with **B.** ensemble results from the same system. The MHC residues Arg62, Glu76, and Gln155 are shown in blue sticks. **C.** Original structure of 1972 influenza NP peptide bound to HLA-B\*35:01 (PDB ID 3LKP) compared with **D.** ensemble results from the same system. The MHC residues Arg62, Glu76, and Gln155 are shown in navy sticks. **E.** Original structure of 2009 influenza peptide bound to HLA-B\*35:01 (PDB ID 3LKR) compared with **F.** ensemble results from the same system. The degree of flexibility of the 1918 and 1972 systems is seen to be similar in magnitude to the difference between the systems. The MHC residues Arg62, Glu76, and Gln155 are shown in pale blue sticks. Hydrogen bonds are shown as dashed lines.

## TCR recognition models

By providing a means of examining the conformational diversity of pMHC systems, ensemble refinement has the potential to provide much richer information for discerning between induced fit and conformational selection models of TCR binding than is possible with traditional structurebased methods. For example, Borbulevych et al<sup>21</sup> compared the structures of HLA-A\*02:01 bound to the human T-lymphotropic virus type 1 (HTLV-1) Tax peptide (PDB ID 3IXA) or bound to the similar Tellp peptide (PDB ID 3H9H) from Saccharomyces cerevisiae (Figure 6A). Despite very similar binding conformations when bound to the MHC alone, the two peptides adopted substantially different conformations upon A6 TCR recognition (Figure 6A), distinguishable by a  $\sim$ 180° flip of the peptide backbone  $\phi$  angle at P6, and different orientations of P4 and P5 side chains. The authors complement their structural observations with experimental measurements and MD, and conclude that these conformational differences are the result of the greater inherent flexibility of the Tellp peptide relative to the Tax peptide, which enables the former to adopt a conformational binding pattern inaccessible to the latter. Our ensemble results provide strong support for this hypothesis, as both conformations of the Tellp peptide, in particular the bond flip of P5, are present in the ensembles (Figure 6B). The structural variation found within the Tellp ensemble is more than sufficient to account for the alternate peptide conformation associated with the A6 TCR-bound form, thereby supporting the contention of Borbulevych et al. that the Tellp conformation observed following A6 TCR binding is due to a conformational selection mechanism, rather than to induced fit. This finding is further corroborated by our MD results, which show very similar bond-flipping behavior to that observed in the ensembles (Figure S1I and S1J).

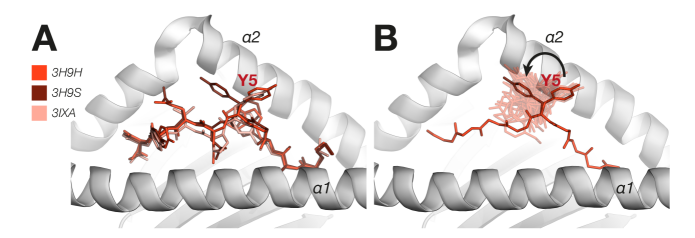


Figure 6: TCR recognition models

**A.** An overlay of HLA-A\*02:01-Tel1p without A6 TCR (red) (PDB ID 3H9H), HLA-A\*02:01-Tel1p with A6 TCR (brown) (PDB ID 3H9S), and HLA-A\*02:01-Tax without the TCR (pale red) (PDB ID 3IXA) compared to **B.** the original HLA-A\*02:01-Tel1p without A6 TCR (red) with overlaid ensemble results (transparent red). The bond flip of P5 distinguishing bound and unbound peptide forms is observed within the ensemble of the original Tel1p structure without TCR.

Another study examined the basis of LC13 TCR alloreactivity by comparing the structures and binding modes of two peptides (which are referred to as 'allotope' and 'mimotope') bound to three different HLA-B\*44 alleles (HLA-B\*44:02, HLA-B\*44:03 and HLA-B\*44:05)<sup>22</sup>. The purpose of the study was to understand the mechanism underpinning LC13 alloreactivity with peptide-bound HLA-B\*44:02 and HLA-B\*4405, but not HLA-B\*44:03, which differs from the other alleles by 1 and 2 polymorphisms respectively. The authors argue that their data show that the HLA-B\*44:05-bound mimotope (PDB ID 3KPQ) and allotope (PDB ID 3KPP) peptides only adopt a similar conformation after LC13 TCR ligation, indicating an induced-fit 'forcing' by the TCR. In particular, for the allotope they identify a downward shift of the P3 residue upon binding to the TCR (PDB ID 3KPS) (Figure 7A), while the mimotope exhibits an even more dramatic rearrangement of the P3, P5, and P7 side chains (PDB ID 3KPR) following TCR binding (Figure 7C). They further argue that the reason the HLA-B\*44:05 and HLA-B\*44:02 complexes bind with the LC13 TCR, while the HLA-B\*44:03 complex (PDB ID 3KPN) does not, is because the latter does not accommodate the conformational change to the peptide necessary for TCR recognition. Specifically, they identify P3 of the allotope complex as being prevented from rotating into the TCR-binding position by the

unfavorable location of Leu156 in HLA-B\*44:03 (Figure 7E). Our ensemble results, however, are

not consistent with these conclusions. All HLA-B\*44:05 allotope and mimotope complexes

(Figures 7B and 7D) exhibit considerable flexibility within each ensemble, with both the unbound

and TCR-bound peptide conformations clearly present within the ensembles. These results indicate

that, rather than being induced by the TCR, the conformational plasticity the authors describe may

be due to pre-existing conformational variability with each system. The ensemble results for the

HLA-B\*44:03 complex likewise exhibits considerable flexibility (Figure 7F), with the P5 residue

clearly adopting a range of conformations including the supposed 'inaccessible' conformation seen

in the TCR-bound forms of the HLA-B\*44:05 system. This result renders implausible the

hypothesis that HLA-B\*44:03 is unable to accommodate peptide conformational change necessary

to bind to the TCR. This highlights the necessity of more detailed analysis of differences in

structural dynamics to address differential TCR recognition.

Overall our data highlights the danger in assuming that an observed difference between static

pMHC and TCR-pMHC complexes is necessarily due to TCR binding, rather than the natural

flexibility of each system. The traditional focus on static structures may have inadvertently had the

effect of biasing analyses in favour of an induced fit mechanism, since the underlying

conformational heterogeneity of the initial system which forms the basis for a conformational

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selection mechanism is overlooked.

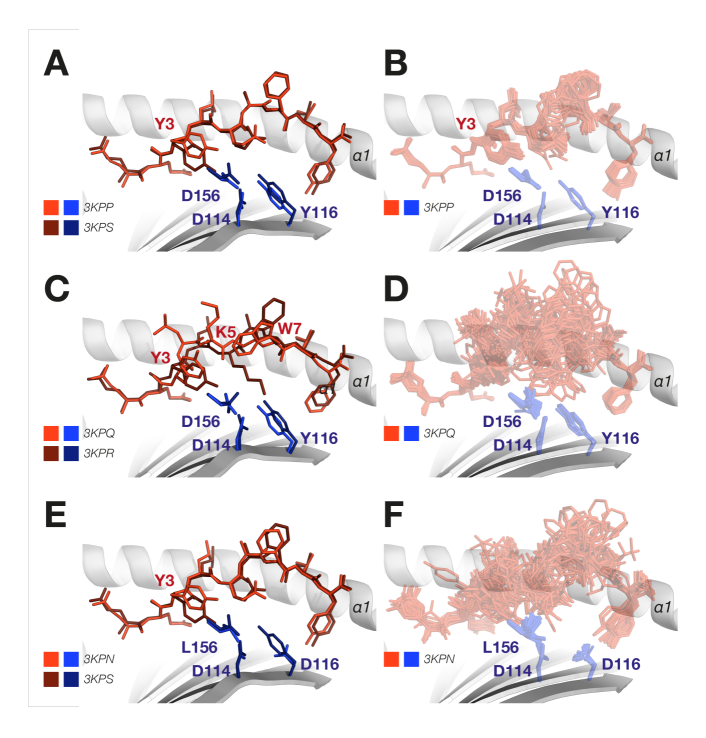


Figure 7: Conformational Plasticity and Molecular Mimicry

**A.** Overlay of the original structures of the HLA-B\*44:05-allotope unbound (red/blue) (PDB ID 3KPP) and bound to TCR (brown/navy) (PDB ID 3KPS), compared to **B.** the ensemble results for HLA-B\*44:05-allotope, showing that structural variability within the ensemble is greater than the difference between TCR-bound and unbound structures. The MHC  $\alpha$ 2-helices have been omitted. **C.** Overlay of the original structures of the HLA-B\*44:05-mimotope unbound (red/blue) (PDB ID 3KPQ) and bound to TCR (brown/navy) (PDB ID 3KPR), compared to **D.** the ensemble results for the HLA-B\*44:05-mimotope ensemble, showing that structural variability within the ensemble is greater than the difference between TCR-bound and unbound structures. The MHC  $\alpha$ 2-helices have been omitted. **E.** Overlay of the original structures of TCR-ligated HLA-B\*44:05-allotope (brown/navy) (PDB ID 3KPS) with the unbound HLA-B\*44:03-allotope (red/blue) (PDB ID 3KPN), showing the different position of P5 which is alleged to be the result of the unfavourable position of Leu156. **F.** The ensemble results for the HLA-B\*44:03-allotope complex, however,

show that the P5 residue is highly flexible, adopting a range of conformations overlapping with that found in the TCR-bound form of the HLA-B\*44:05 complex. The MHC  $\alpha$ 2-helices have been omitted.

### **Discussion**

X-ray crystallography has played a considerable role in increasing our understanding of the structural basis of TCR-pMHC engagement, but is limited in that it provides only a static snapshot. Therefore the role of conformational dynamics in TCR recognition of the pMHC has been largely overlooked, despite its important role in determining biological outcomes. Our study strongly emphasizes that a sizeable number of pMHC complexes exhibit considerable conformational flexibility that is not apparent in the published single-structure crystallographic model. Such dynamic variability, shown in both our ensemble and MD results, extends well beyond mere alternate side-chain rotamers or bond flips, and encompasses a vast range of motions including large-scale backbone movements and interconversion between substantively different peptide conformations. This result initially seems at odds with the relatively low B-factors reported in some of these systems. However, it is likely that B-factors substantially underestimate atomic motions, such that much of information on the flexibility and motions present in the molecule is lost in the refinement process<sup>23,24</sup>. Our results corroborate these findings, and highlight the importance of using ensemble refinement techniques to more accurately gauge protein conformational flexibility.

Many contemporary TCR-pMHC structural studies attempt to determine the factors that influence TCR binding specificity or cross-reactivity by comparing the structures of two or more complexes and identifying similarities and differences. Thus, for instance, if a peptide adopts one conformation when bound to MHC alone and a different conformation when bound to MHC complexed with TCR, it is often inferred that this represents a change in conformation brought about by the binding of the TCR. However, as we have emphasized in the case studies discussed above, a baseline degree of structural heterogeneity must be expected in any protein system, resulting from the intrinsic flexibility of protein molecules. The precise position of backbone atoms, orientation of side chains,

extent of hydrogen bonding networks, etc., is therefore expected to vary over time and between samples. Crystallographic structures are thus properly regarded as the result of random draws from the underlying distribution of protein conformational states. Observed structural differences between systems should therefore be compared in magnitude to the intrinsic structural variation within each system, to determine whether there is sufficient reason to justify claiming a significant difference between the systems. In principle this is much the same as statistical tests routinely applied in disciplines outside of structural biology, where a difference between two groups is only deemed significant if this inter-group difference is sufficiently greater than the intra-group variation. Although it is possible to estimate coordinate error from the crystallographic data, no such method yet exists for statistical comparison of protein ensembles. The resulting failure to consider this baseline of structural heterogeneity has resulted in numerous false positive identifications of putative differences, and thereby hampered the growth in our understanding of MHC and T-cell function.

In addition to the question of whether two protein systems exhibit a structural difference that is significantly greater than background heterogeneity, there is the further question as to what is the most likely cause of this difference. In many structural studies there is a tendency to readily infer that, if an MHC-bound peptide changes conformation upon TCR ligation, then the TCR has induced this change in conformation. Though this is undoubtedly the case for some systems, in other systems it is likely that causation runs in the opposite direction, meaning that the reason the TCR has bound to the peptide is because the peptide has adopted that particular conformation. This distinction corresponds to the dispute between the 'induced fit' and 'conformational selection from existing equilibrium' models of protein interaction. As we noted previously, ensemble refinement provides a method uniquely suited to examine this question, since if the TCR-bound peptide conformation is already observed in the unbound pMHC ensemble, this constitutes at least *prima facie* evidence for a conformational selection mechanism<sup>25</sup>. Our ensemble results show that, at least for some peptide-MHC systems, there is a considerable degree of intrinsic flexibility, from which

the TCR has considerable 'choice' in selecting a favorable form with which to bind. Recent work in

support of these ideas has shown that conformational variability may facilitate several rounds of

association and disassociation of peptides with MHC complexes, with low affinity peptides rapidly

disassociating and thereby gradually establishing an equilibrium ensemble where a given MHC is

only bound to its corresponding high affinity peptides. Our results build upon these findings in

providing further evidence of the importance of a structural ensemble-based approach in the study

of MHC dynamics and affinity.

A third, related and important consideration when examining TCR-pMHC systems is that the

intrinsic structural heterogeneity resulting from peptide and MHC flexibility need not always be

simply random fluctuations about a single functional state. Rather, it is likely that in at least some

cases different conformations of the system fulfil different functional purposes. As discussed in a

number of the cases we considered, the flexibility of the system itself can play a role in determining

binding specificity, for example through bond flips that alternate between bound and unbound

peptide conformations. Such behaviors indicate that one cannot simply infer that because a peptide

is observed in one particular conformation when crystallized bound to the TCR, this was the same

conformation that the peptide adopted during the process of TCR recognition — it is possible that

conformational changes may occur after the initial binding event. There may be one or more

intermediates and/or transition states with subtly or considerably different conformations to that

seen in the final complex, with the conformational variability an intrinsic part of the binding

process. Peptide flexibility should therefore be considered from the outset as an essential functional

aspect of the system when analyzing peptide, MHC and TCR binding dynamics.

**Conclusions** 

Our analysis highlights the importance of analyzing the properties of protein ensembles rather than

isolated structures. Examination of a variety of TCR-pMHC complexes suggests that reliance on

single structures entails significant pitfalls for understanding the rules of productive TCR ligation,

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particularly for static interpretations involving fine details such as hydrogen-bonding networks and side chain orientation. Furthermore, our findings indicate that structural differences between the pMHC and TCR-pMHC conformations may be due to intrinsic flexibility rather than any change elicited by binding itself, and highlights the risk of false-positives in assuming that an observed difference between static pMHC and TCR-pMHC complexes is necessarily due to the binding process, rather than a reflection of the natural flexibility of each system. Broadly, our results imply that biological observations should be explained through the properties of ensembles rather than isolated structures, as these are less prone to observer bias. We propose that extracting dynamic information from X-ray data through the use of ensemble methods will not only mitigate these concerns but lead to a richer, dynamic understanding of TCR-pMHC interactions.

**Materials and Methods** 

Computational resources

Calculations, modeling and simulations were performed on a range of computing resources:

ORCHARD 800 core x86 cluster (Monash University; X-ray ensemble refinement); Multi-modal

Australian ScienceS Imaging and Visualisation Environment (MASSIVE; atomistic MD).

Ensemble refinement

All atomic coordinates (.pdb) and crystallographic reflections (.mtz) files were sourced from the

PDB REDO server<sup>27</sup>. Ensembles were calculated with PHENIX 1.9-1692<sup>28</sup> by first passing each

system through the ReadySet tool, and then using the Ensemble refinement tool with default

parameters<sup>8</sup>. Ensemble analysis was performed using PyMOL version 1.3<sup>29</sup> and python scripts.

Atomic coordinates, modeling and graphics

In MD simulations, atomic coordinates were obtained from the following PDB entries: 1UXS,

2A83, 3H9H, 2LKN, and 3MRD. Structural representations were produced using PvMOL version

1.3<sup>29</sup> and VMD 1.9.1<sup>30</sup>. Trajectory manipulation and analysis was performed using MDTraj<sup>31</sup> and

VMD 1.9.1<sup>30</sup>.

Molecular dynamics (MD) systems setup and simulation

Each protein, with protonation states appropriate for pH 7.0<sup>32,32</sup>, was placed in a rectangular box

with a border of at least 12 Å, explicitly solvated with TIP3P water34, counter-ions added, and

parameterized using the AMBER ff14SB all-atom force field<sup>35,36,37</sup>. After an energy minimization

stage and an equilibration stage, production simulations were performed in the NPT ensemble. Four

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independent replicates of each system were simulated for 500 ns each using NAMD 2.9<sup>38</sup>.

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## Acknowledgements

This work was supported by grants from the National Health and Medical Research Council (NHMRC) and the Australian Research Council (ARC). NAB is funded by an ARC Future Fellowship (110100223). This work was supported by the Multi-modal Australian ScienceS Imaging and Visualisation Environment (MASSIVE) (www.massive.org.au).

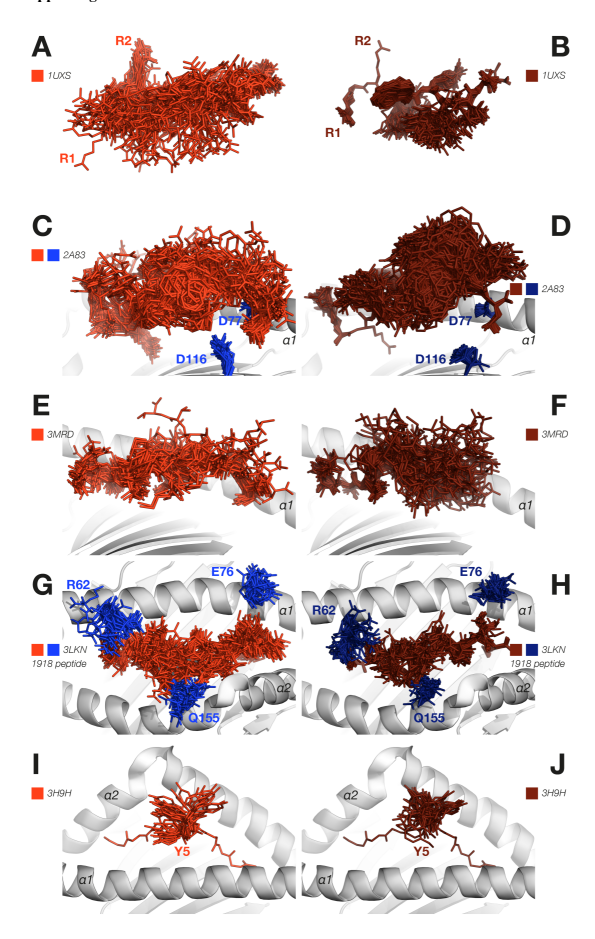
### **Author Contributions**

AMB designed the study. JF performed ensemble refinement and molecular dynamics simulations. BTR prepared the figures. JF, BTR, NAB and AMB wrote the manuscript.

## **Competing financial interests**

The authors declare no competing financial interests.

# **Supporting Information**



## Figure S1: Molecular Dynamics and Ensemble Refinement Comparison

A. Overlay of 50 frames from 2 us MD simulation of an Epstein-Barr virus peptide pLMP2 bound to HLA-B\*27:09 (red sticks) (PDB ID 1UXS) compared to **B.** ensemble results of the same system (brown sticks), showing very similar patterns of dynamics. MHC  $\alpha$ -helices have been omitted. C. Overlay of 50 frames from 2 µs MD simulation of a glucagon-derived self-peptide (pGR) (red sticks) bound to HLA-B\*27:09 (blue sticks, cartoon) (PDB ID 2A83) compared to **D.** ensemble results of the same system (peptide as brown sticks; MHC as navy sticks, cartoon), with Asp77 and Asp116 shown as sticks, showing very similar patterns of dynamics. The MHC  $\alpha$ 2-helix has been omitted. E. Overlay of 50 frames from 2 µs MD simulation of NLV<sub>V6G</sub> peptide (red sticks) bound to HLA-A\*02:01 (cartoon) (PDB ID 3MRD) compared to F. ensemble results of the same system (peptide as brown sticks; MHC as cartoon), showing comparable overall degrees of flexibility. The MHC  $\alpha$ 2 helix has been omitted. G. Overlay of 50 frames from 2  $\mu$ s MD simulation of 1918-NP<sub>418</sub> (red sticks) bound to HLA-B\*35:01 (blue sticks, cartoon) (PDB ID 3LKN) compared to H. ensemble results of the same system (peptide as brown sticks; MHC as navy sticks, cartoon), with very similar patterns of flexibility evident in each. The MHC residues Arg62, Glu76, and Gln155 are shown as sticks. I. Overlay of 50 frames from 2 µs MD simulation of Tellp (red sticks) bound to HLA-A\*02:01 (cartoon) (PDB ID 3H9H) compared to J. ensemble results of the same system (peptide as brown sticks; MHC as cartoon), showing the very similar bond flips of P5-Tyr in both cases.