# Load-based divergence in the dynamic allostery of two TCRs recognizing the same pMHC

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Abstract Increasing evidence suggests that mechanical load on the  $\alpha\beta$  T cell receptor (TCR) is crucial for recognizing the antigenic peptide-loaded major histocompatibility complex (pMHC) molecule. Our recent all-atom molecular dynamics (MD) simulations revealed that the inter-domain motion of the TCR is responsible for the load-induced catch bond behavior of the TCR-pMHC complex and peptide discrimination. To further examine the generality of the mechanism, we perform all-atom MD simulations of the B7 TCR under different conditions for comparison with our previous simulations of the A6 TCR. The two TCRs recognize the same pMHC and have similar interfaces with pMHC in crystal structures. We find that the B7 TCR-pMHC interface stabilizes under ~15-pN load using a conserved dynamic allostery mechanism that involves the asymmetric motion of the TCR chassis. However, despite forming comparable contacts with pMHC as A6 in the crystal structure, B7 has fewer high-occupancy contacts with pMHC and exhibits higher mechanical compliance during the simulation. These results indicate that the dynamic allostery common to the TCR $\alpha\beta$  chassis can amplify slight differences in interfacial contacts into distinctive mechanical responses and nuanced biological outcomes.

#### Introduction

The A6 TCR $\alpha\beta$  and B7 TCR $\alpha\beta$  (herein we call TCR $\alpha\beta$  simply as TCR) are both specific for the same Tax peptide (LLFGYPVYV) of the human T lymphotropic virus 1 (HTLV-1) bound to HLA-A2 (*Garboczi et al., 996a*,b; *Ding et al., 1998*). A6 and B7 derive from T cell clones isolated from two patients with HTLV-1-associated myelopathic/tropical spastic paraparesis (*Utz et al., 1996*; *Ding et al., 1998*). They share the same V $\beta$  germline gene (TRBV6-5) and differ only in the V $\alpha$  germline gene (A6: TRAV12-2; B7:TRAV29DV5), with sequence similarity of 45% for V $\alpha$ , 96% for V $\beta$ , and 100% for C $\alpha$  and C $\beta$  (*Ding et al., 1998*). The only structural differences between the two TCRs are from the

residues of the  $V\alpha$  domain and the highly variable complementarity-determining region 3 (CDR3 $\beta$ ) loop of the  $V_B$  domain that is crucial for peptide recognition (*Ding et al.*, 1998; Bourcier et al., 2001; Rudolph et al., 2006). In crystal structures, both TCRs bind in a diagonal orientation to the Tax peptide-bound major histocompatibility complex (pMHC), such that both  $V\alpha$  and  $V\beta$  contact the MHC  $\alpha$ 1 and  $\alpha$ 2 helices (*Garboczi et al.*, 996a). The similar diagonal binding modes are achieved by interactions involving different CDR residues of A6 and B7 contacting largely the same sets of pMHC residues (Ding et al., 1998). Only one out of 17 residues contacting pMHC in B7 is also found in the A6 pMHC interaction (Ding et al., 1998). T cell response assays demonstrated that single-residue mutations to the Tax peptide elicit different responses in the two TCRs (Ding et al., 1998: Hausmann et al., 1999). Interfacial interactions may play a role in this TCR-specific response. as residue charges at the surface of the A6 and B7 variable domains show different electrostatic profiles, where the pocket for the Tax peptide Y5 residue is positively charged in A6 but negatively charged in B7 (*Ding et al.*, 1998). Overall, the B7  $V\alpha$  surface has fewer charged residues exposed than A6 V $\alpha$ . While A6 and B7 recognize Tax-MHC with similar affinities and kinetics, it has been suggested that they achieve binding via different thermodynamic pathways (Davis-Harrison et al., 2005).

However, since  $\alpha\beta$ TCR is a mechanosensor (*Reinherz et al., 2023*), the TCR-pMHC bond lifetime under physiological piconewton (pN) level load should be more functionally relevant than the equilibrium binding pathway. In this regard, we have previously used all-atom MD simulations to show that contacts with pMHC of A6 (*Chang-Gonzalez et al., 2024*) and JM22 (*Hwang et al., 2020*) TCRs are stabilized when an adequate 15–20 pN force is applied. The force-induced stabilization occurs as the asymmetric domain motion of the TCR chassis leads to weakening of the interface with pMHC either in the absence of an adequate level of force or if the sequence of the bound peptide is incompatible with maintaining contacts in the loaded state. The goal of this study is to determine whether the load-dependent control of the binding with pMHC is also present in B7 TCR, and identify any differences with A6 that may impact the response of the T-cell while responding to the same pMHC.

We find that the mechanism of dynamic allostery is largely conserved in B7, yet the loaded state does not stabilize contacts with pMHC as robustly as in the A6. Thus, while both A6 and B7 possess comparable equilibrium binding affinity for pMHC, A6 appears to exhibit a stronger catch bond behavior under load. Given the nuanced and dynamic nature of TCRs against the same pMHC, including the effect of ligand abundance (*Akitsu et al., 2024*), difference in mechanical response between A6 and B7 suggests T-cell clones bearing those TCRs may operate differently *in vivo*. Our study also underscores the importance of dynamics when comparing between TCRs that have similar crystal structures and equilibrium binding affinities.

#### Results

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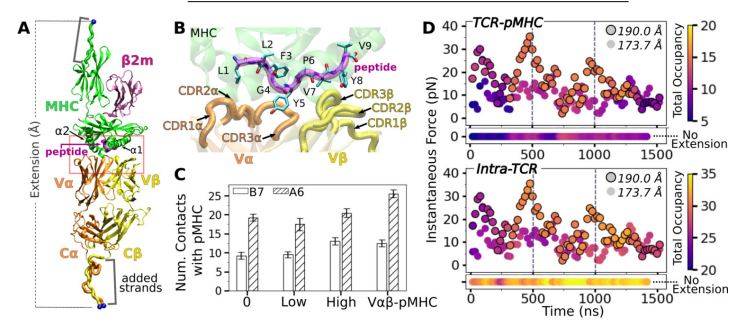
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Our simulation systems include the B7 TCR bound to Tax pMHC (*Figure 1*A,B) with no, low, and high extensions to apply different loads, and an isolated B7 TCR (*Table 1*). We also used systems without the C-module ( $V\alpha\beta$  and  $V\alpha\beta$ -pMHC) to study the role of the C-module. To apply a load, we held the terminal  $C_{\alpha}$  atoms of the added strands at different extensions (*Figure 1*A). This reflects the constant spacing between a T cell and an APC maintained by adhesion molecules such as CD2 and CD58 (*Reinherz et al., 2023*). In *in vitro* single-molecule experiments, pulling to a fixed separation and holding is also commonly done. On the other hand, simulation for B7 $^{0}$  was performed by lightly holding the MHC  $\alpha$ 3 domain (*Figure 1*A) in a construct without added strands (Methods). It thus does not have any extension nor applied load.

We analyzed TCR-pMHC intermolecular and intra-TCR (intramolecular) interactions and domain motion to test whether the TCR-pMHC interface is stabilized by load, and to find the underlying allostery mechanism that involves the motion of the TCR chassis. We analyze dynamics during the full trajectory and average behavior during 500-1000 ns.

**Table 1.** Simulation systems constructed based on PDB 1BD2 (*Ding et al., 1998*). Extension is the distance between the harmonic potentials on the terminal restrained atoms (blue spheres in *Figure 1*A), which was selected for B7<sup>low</sup> and B7<sup>high</sup> to yield average low and high loads among simulations scanning different extensions (see Methods). Average load is calculated between 500-1000 ns. The standard deviation (std) in load as measured in 40-ns intervals between 500-1000 ns is shown in parentheses. Their values are consistent with those for A6 (*Figure 1—figure Supplement 2*). The extension and force (average±std) for A6 corresponding to B7<sup>low</sup> and B7<sup>high</sup> are 182.6 Å and 13.2±5.65 pN, and 187.7 Å and 18.2±9.17 pN, respectively (*Chang-Gonzalez et al., 2024*), which indicates that B7 is more compliant compared to A6. For B7<sup>0</sup>, B7<sup>low</sup>, and B7<sup>high</sup>, simulations were further extended for additional time-dependent stability analysis. However, averaging was done for the 500-1000 ns interval for consistency.

Label	Structure	Time (ns)	Extension (Å)	Load (pN)
Vαβ	$V\alpha$ - $V\beta$ only (no pMHC)	1000	-	_
$T\alpha\beta$	$TCR\alpha\beta$ only (no pMHC)	1000	-	-
$V\alpha\beta$ -pMHC	$V\alpha\beta$ with pMHC (no C-module)	1000	-	-
B7 <sup>0</sup>	TCRlphaeta-pMHC	1450	-	-
B7 <sup>low</sup>	TCRlphaeta-pMHC	1550	173.7	9.01 (3.96)
B7 <sup>high</sup>	TCRlphaeta-pMHC	1550	190.0	14.5 (7.20)



**Figure 1.** B7 TCR-pMHC interface. (A) Overview of the base complex used in simulations. The 4 subdomains of TCR $\alpha\beta$  are V $\alpha$ , V $\beta$ , C $\alpha$ , and C $\beta$ . Load was applied by holding the C $_\alpha$  atoms of terminal residues (blue spheres at the ends of "added strands") at a given distance from each other.  $\beta$ 2m:  $\beta$ 2 microglobulin. (B) Magnified view of red box in panel A showing labeled CDR loops and side chains of peptide residues in stick representation. (C) Number of contacts with greater than 50% average occupancy and 80% maximum instantaneous occupancy from 500-1000 ns. Bars: std. Criteria for counting contacts and values for A6 are from **Chang-Gonzalez et al.** (2024). (D) Total contact occupancy measured in 40-ns overlapping intervals. TCR-pMHC (top; intermolecular) and intra-TCR (bottom; intramolecular) contacts are shown separately. Intra-TCR contacts exclude C $\alpha$ -C $\beta$  contacts (Methods). Circles with outline: B7<sup>high</sup>; without outline: B7<sup>low</sup>. Horizontal bar below each panel: B7<sup>0</sup>.

**Figure 1—figure supplement 1.** Model of  $TCR\alpha\beta$  with leucine zipper domain fused to the C-terminal added strands.

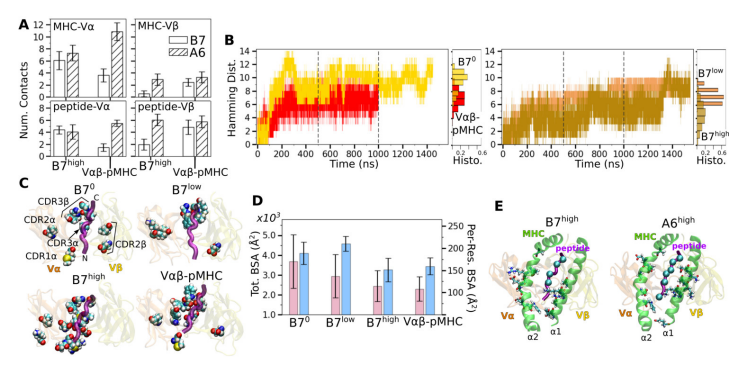
Figure 1—figure supplement 2. Standard deviation vs. average of the applied force.

# Stabilization of the TCRlphaeta-pMHC interface with load

- <sup>91</sup> Compared to the no- (B7<sup>0</sup>) and low-load (B7<sup>low</sup>) cases, the number of high-occupancy contacts were
- $_{92}$  more numerous for the high-load case (B7 $^{\text{high}}$ ), indicative of a catch bond behavior. Absence of the
- <sup>93</sup> C-module ( $V\alpha\beta$ -pMHC) also promoted more contacts with pMHC, suggesting the allosteric role of
- the C-module for binding with pMHC (Figure 1C). These results agree well with the behaviors seen
- 95 in A6 and JM22 TCRs in our previous studies (Chang-Gonzalez et al., 2024; Hwang et al., 2020).
- <sub>96</sub> However, the number of pMHC contacts with B7 was reduced compared to A6 (Figure 1C). This is

despite their comparable number of contacts with the pMHC in crystal structures and comparable equilibrium binding affinity in solution (*Ding et al., 1998; Davis-Harrison et al., 2005*). In fact, we had to modify our simulation protocol to avoid premature breakage of contacts between B7 and pMHC when preparing the system for production run under load (see Methods).

Time-dependent behavior of the TCR-pMHC interface further supports the load-mediated enhancement of binding. For this, we calculated instantaneous force and contact occupancy in 40-ns overlapping intervals (*Figure 1*D). B7<sup>high</sup> had overall more inter- and intramolecular contacts (more orange-yellow circles) than B7<sup>low</sup> (more purple), suggesting that the increased load stiffens the TCR-pMHC complex. B7<sup>0</sup> had fewer TCR-pMHC contacts but more intra-TCR contacts (horizontal bars in *Figure 1*D), indicating decoupling of the TCR from pMHC in the absence of load. However, after about 1300 ns, B7<sup>high</sup> started to lose the initial high-occupancy contacts (*Figure 1*D). This is likely because the fixed extension (end-to-end distance) imposed in simulation can adversely impact the interfacial stability of the B7 TCR-pMHC complex that has a higher compliance than A6, as discussed further below. For analysis of average properties, we thus used the 500-1000 ns interval, to be consistent with other systems.



**Figure 2.** Load dependent behavior of the B7 TCR-pMHC interface. (A) Number of MHC-V $\alpha$ , MHC-V $\beta$ , peptide-V $\alpha$ , and peptide-V $\beta$  contacts in B7 and A6 TCRs between 500-1000 ns. Data for A6 are from *Chang-Gonzalez et al.* (2024). (B) Hamming distance  $\mathcal{H}$ . Histograms were calculated using data between 500-1000 ns (marked by vertical dashed lines), the interval between the dashed lines. (C) Location of V-module residues forming contacts with pMHC with greater than 50% average occupancy. The frame at 1000 ns is used for visualization. The backbone of the Tax peptide is shown as a purple tube. CDRs are labeled in the first panel. (D) Total (pink) and per-residue (blue) BSA for interfacial residues between 500-1000 ns. (E) pMHC residues forming contacts with the V-module with average occupancy greater than 50% in the high-load case. Left: B7<sup>high</sup>, right: A6<sup>high</sup>. MHC residues are shown as sticks and C $_{\alpha}$  atoms of the peptide residues are shown as spheres. Viewing direction is the same as in panel C.

**Figure 2—figure supplement 1.** Additional characterization of the TCR-pMHC interface.

Comparing average counts of high-occupancy pMHC contacts for both A6 and B7 TCRs indicates that interfacial contacts were dominated by MHC-V $\alpha$  (*Figure 2*A). Also, V $\alpha$  formed more contacts with MHC than V $\beta$ , while V $\beta$  formed more contacts with the peptide than it does with MHC (*Figure 2*A). Similar to A6 (*Chang-Gonzalez et al., 2024*), occupancy heat maps for individual contact residue pairs show reduced or fragmented contacts for B7 $^0$  and B7 $^{low}$  (*Figure 2-figure Supplement 1*A,B, more red compared to blue) while B7 $^{high}$  and V $\alpha\beta$ -pMHC exhibit more persistent contact

profiles (Figure 2-figure Supplement 1C,D, more blue compared to red).

Temporal progression of the number of contacts was measured via the Hamming distance  $\mathcal{H}$ , the number of the initial high-occupancy contacts lost over time (*Figure 2*B; see Methods). For B7<sup>0</sup>,  $\mathcal{H}$  rapidly increased and by 200 ns, most of the initial high-occupancy contacts were lost. While the increase in  $\mathcal{H}$  for Vαβ-pMHC was comparable to that of B7<sup>low</sup> (*Figure 2*B), the contact occupancy heat maps reveal that Vαβ-pMHC maintained contacts after a brief initial adjustment (arrow in *Figure 2-figure Supplement 1*D) while contacts were lost in B7<sup>low</sup> (red in *Figure 2-figure Supplement 1*B). On the other hand, B7<sup>high</sup> maintained  $\mathcal{H}$  comparable to Vαβ-pMHC until about 1300 ns, after which it approached that of B7<sup>0</sup>. This occurred as some of the high-occupancy contacts broke (*Figure 2-figure Supplement 1*C). Importantly, the breakage is not due to a high force, but instead it happened when instantaneous force was low. Low-force states are reached at around 750 ns and 1300 ns (*Figure 1*D), followed by stepwise increase in  $\mathcal{H}$  (*Figure 2*B). The difference in extension between B7<sup>low</sup> and B7<sup>high</sup> is 16.3 Å whereas it is 5.1 Å for A6 (*Table 1*), for similar ~5-pN difference between low- and high-load cases. Being more compliant, the interface of B7 can reach a low-force state more easily, hence it is more prone to destabilization.

Location of V-module residues forming contacts with pMHC with greater than 50% average occupancy were concentrated along the peptide for B7<sup>high</sup> and V $\alpha\beta$ -pMHC, but scattered in B7<sup>0</sup> and B7<sup>low</sup> (*Figure 2*C). This trend was also observed in A6 and JM22. Such concentration of high-occupancy contacts may protect TCR-pMHC interactions from breakage by water. In A6 and JM22 TCRs, the greater number of contacts with pMHC in the high-load cases and V $\alpha\beta$ -pMHC correlated with larger buried surface area (BSA) of the residues forming contacts (*Chang-Gonzalez et al., 2024*; *Hwang et al., 2020*). B7 did not follow this trend, as B7<sup>high</sup> and V $\alpha\beta$ -pMHC had reduced total and per-residue BSA than B7<sup>0</sup> and B7<sup>low</sup> (*Figure 2*D). This is likely because the fewer high-occupancy contacts in B7 (*Figure 1*C) tend to be more exposed, making the relationship between the BSA and load less direct. Consistent with this, the total BSA of B7 was 67.4 % (B7<sup>0</sup>) to 44.2% (B7<sup>high</sup>) of the corresponding values of A6.

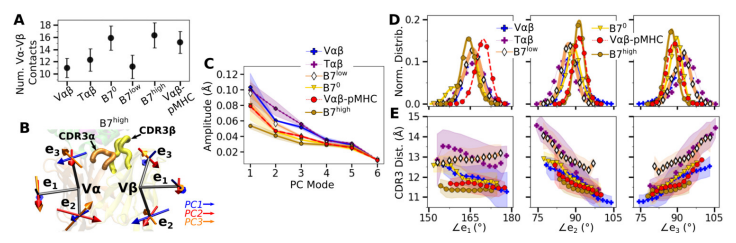
The residues of pMHC forming greater than 50% average occupancy under high load differed between A6 and B7 (*Figure 2*E), as did the location of V-module residues forming respective contacts with the pMHC residues (*Figure 2*C), further highlighting their divergence in the interfacial footprint under load. Despite this, the root-mean square fluctuation (RMSF) of  $C_{\alpha}$  atoms of the Tax peptide measured after 500 ns was reduced in B7<sup>high</sup> and V $\alpha\beta$ -pMHC compared to B7<sup>0</sup> and B7<sup>low</sup> (*Figure 2-figure Supplement 1*E), as observed for A6 (*Chang-Gonzalez et al., 2024*), which supports load-induced stabilization of the complex.

We calculated the distance between the V-module and pMHC as another measure of the interfacial stability (*Figure 2-figure Supplement 1*F; Methods). The distance was stably maintained in  $V\alpha\beta$ -pMHC and B7<sup>high</sup> (before 1300 ns) whereas it fluctuated more in B7<sup>0</sup> and B7<sup>low</sup>. Of note, the former two systems maintained the distance greater than that in the crystal structure by 0.3–0.9 Å. Thus, a slight separation engendered by force or in the absence of constraint imposed by the C-module provides room for adjusting residues to form more stable contacts. The more stable maintenance of the distance between V-module and pMHC in  $V\alpha\beta$ -pMHC and B7<sup>high</sup> is consistent with other measures of their stability explained above.

# CDR3 positions are controlled by load-dependent $V\alpha$ - $V\beta$ motion

The greater number of  $V\alpha$ - $V\beta$  contacts in B7<sup>0</sup> (*Figure 3A*) is consistent with the increase in total intra-TCR contact occupancy (horizontal bar in *Figure 1*D bottom panel). Without load this does not translate to a stronger TCR-pMHC interface explained above. B7 in general had fewer  $V\alpha$ - $V\beta$  contacts (11.0–16.3) than A6 (15.9–23.1) (*Chang-Gonzalez et al., 2024*). The ~70% reduction in  $V\alpha$ - $V\beta$  contacts for B7 is comparable to the ~50% reduction in contacts with pMHC between the two TCRs (*Figure 1*C).

 $V\alpha$ - $V\beta$  motion was measured via triads (orientational markers) assigned to respective domains and by performing principal component analysis (PCA) (*Figure 3*B; Methods). PC amplitude was



**Figure 3.** V $\alpha$ -V $\beta$  motion of B7. (A) Number of V $\alpha$ -V $\beta$  contacts with greater than 50% average occupancy and 80% maximum instantaneous occupancy between 500-1000 ns. Bars: std. (B) V-module triads {e<sub>1</sub>, e<sub>2</sub>, e<sub>3</sub>}. Arrows denote directions of the first 3 PC modes for B7<sup>high</sup> as an example. CDR3s are labeled. (C) Amplitudes for the first 6 PCs. PCA was performed between 500-1000 ns. Transparent bands: std for PCA performed in three overlapping intervals (500–800 ns, 600–900 ns, and 700–1000 ns). (D) Histograms of the V-module triad angles. (E) CDR3 distance vs. triad angles. Transparent bands: std. Both D and E were determined from data between 500-1000 ns.

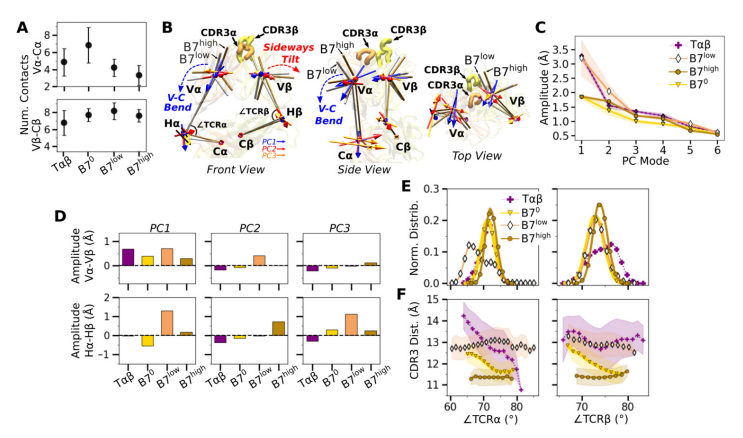
the lowest for B7<sup>high</sup> and V $\alpha\beta$ -pMHC (*Figure 3*C), which is consistent with the greater number of V $\alpha$ -V $\beta$  contacts. Regarding the direction of motion, the mutually orthogonal PC directions can be difficult to interpret (arrows in *Figure 3*B). We instead measured angles between the matching arms of the two triads named  $\angle e_i$  (i=1,2,3), to examine the V $\alpha$ -V $\beta$  motion in structurally interpretable directions (*Figure 3*D) (*Chang-Gonzalez et al., 2024*). For example,  $\angle e_1$  is the angle between each  $e_1$  arm from V $\alpha$  and V $\beta$ , which describes a 'flapping' or 'twisting' motion of the two domains. Since  $e_2$  and  $e_3$  lie approximately parallel to the V $\alpha$ -V $\beta$  interface, they vary reciprocally, corresponding to a 'scissoring' motion (*Hwang et al., 2020*).

Measuring the distance between CDR3 $\alpha$  and CDR3 $\beta$  ("CDR3 distance") revealed that this distance is the shortest for B7<sup>high</sup> followed by V $\alpha\beta$ -pMHC. Comparing CDR3 distance versus triad angles (*Figure 3*E) shows that CDR3 distance varied in opposite directions with  $\angle e_2$  and  $\angle e_3$ , which reflects their reciprocal relation (opposite slopes in *Figure 3*E). In comparison, the CDR3 distance of the B7 crystal structure is 12.0 Å, which is larger than those of B7<sup>high</sup> and V $\alpha\beta$ -pMHC (*Figure 3*E). The slight separation between the V-module and pMHC (*Figure 2-figure Supplement 1*F) in V $\alpha\beta$ -pMHC and B7<sup>high</sup> allows CDR3 loops to come closer together compared to the crystal structure, akin to pinching the central protrusion of the peptide.

## Asymmetric V-C bending in the B7 TCR is suppressed with applied load

Next we considered the motion between the V- and C-modules ("V-C motion"). The number of high-occupancy contacts for the  $C\alpha$ - $C\beta$  interface (26.2–27.5) was considerably greater than those for  $V\alpha$ - $V\beta$  (11.2–16.3), indicating that the C-module acts as a single base for the V-C motion, as noted for A6 and JM22 TCRs (*Hwang et al., 2020*; *Chang-Gonzalez et al., 2024*). Continuing this general feature, there were fewer high-occupancy contacts for the  $V\alpha$ - $C\alpha$  interface compared to the  $V\beta$ - $C\beta$  interface (*Figure 4*A).

The V-C motion was analyzed by using the bead-on-chain (BOC) model that tracks individual domains and the hinge between them (*Figure 4*B). PC motion directions were compared by calculating the dot products between the corresponding PC vectors (*Figure 4-figure Supplement 1*A). PC1 corresponding to the V-C bending in B7<sup>high</sup> (*Figure 4*B) was similar in other systems, whereas B7<sup>low</sup> differed the most (darker color for B7<sup>low</sup> in *Figure 4-figure Supplement 1*A, PC1). Amplitudes of PC1 shows a clear distinction where the unliganded  $T\alpha\beta$  and B7<sup>low</sup> were more mobile than B7<sup>0</sup> and B7<sup>high</sup> (*Figure 4*C). Comparing PC amplitudes of the elements of the BOC between  $\alpha$  and  $\beta$  chains revealed that  $V\alpha$  moves more relative to the C-module than  $V\beta$  (*Figure 4*D), similar to A6



**Figure 4.** V-C motion of B7. (A) Number of V-C contacts per chain measured with the same criteria as *Figure 3*A. (B) Average BOC for B7<sup>low</sup> and B7<sup>high</sup>. The V-module of B7<sup>high</sup> is less bent compared to B7<sup>low</sup>. The arrows for the first 3 V-C PC modes are shown, where PC1 corresponds to the V-C bending motion. (C) V-C PC amplitudes for the first 6 PC modes. Transparent bands: std measured in the same way as in *Figure 3*C. (D) Differences in amplitudes for the first 3 PCs between matching V- and H-beads of *α* and *β* chains. (E) Histograms of hinge angles defined in panel B. (F) CDR3 distance versus hinge angles. Transparent bands: std. All panels are generated with data between 500-1000 ns.

Figure 4—figure supplement 1. Additional characterization of the V-C motion.

(*Chang-Gonzalez et al., 2024*). Amplitude of the hinge motion in the two chains varied, where  $H\alpha$  had greater amplitude in PC1 for B7<sup>low</sup> compared to B7<sup>high</sup> (*Figure 4*D, PC1 in bottom row). This suggests a more pronounced asymmetric motion in B7<sup>low</sup>. The no-load B7<sup>0</sup>  $H\beta$  amplitude was larger compared to  $H\alpha$  (*Figure 4*D, bottom, negative value for PC1 of B7<sup>0</sup>). For B7<sup>0</sup>, the small PC1 amplitude of the overall V-C motion without load (*Figure 4*C) does not suppress the motional asymmetry between  $\alpha$  and  $\beta$  chains, while in B7<sup>high</sup>, the chassis becomes less mobile under load.

The V-C angles (*Figure 4*B,  $\angle$ TCR $\alpha$  and  $\angle$ TCR $\beta$ ) reveal the motional asymmetry in addition to PCA. As in the case of low-load A6 (*Chang-Gonzalez et al., 2024*),  $\angle$ TCR $\alpha$  of B7<sup>low</sup> shows a wide bimodal distribution (*Figure 4*E).  $\angle$ TCR $\alpha$  decreased at around 700 ns as B7<sup>low</sup> bent more (*Figure 4-figure Supplement 1*B). This would put the V-module in an unfavorable orientation to bind pMHC (*Hwang et al., 2020*). The dependence of CDR3 distance on V-C angles was most steady in B7<sup>high</sup> (*Figure 4*F). This supports the allosteric mechanism by which the asymmetric V-C motion of the whole TCR controls the V $\alpha$ -V $\beta$  motion and the V-C orientation that in turn affects the stability of the TCR-pMHC interface.

## **Concluding Discussion**

As a protein-protein complex, TCR-pMHC forms a weak interface. A typical heterodimeric protein-protein interface with BSA comparable to that of TCR-pMHC ( $\sim$ 1700 Å<sup>2</sup>) has sub- $\mu$ M binding affinity (*Chen et al., 2013*) while the affinity of the TCR-pMHC complex ranges between  $\mu$ M to hundreds of  $\mu$ M (*Rudolph and Wilson, 2002*; *Rudolph et al., 2006*). Given the low equilibrium affinity, recent find-

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ings highlight the importance of mechanosensing, where force generated during immune surveillance of  $\alpha\beta$  T-cells is utilized to discriminate cognate versus non-cognate pMHCs (reviewed in **Zhu et al.** (2019); **Liu et al.** (2021); **Reinherz et al.** (2023)). Our earlier simulation studies of the JM22 and A6 TCRs showed that the dynamical motion of the TCR chassis is responsible for the force-driven stabilization of the contacts with pMHC, *i.e.*, catch bond formation (**Hwang et al.**, 2020; **Chang-Gonzalez et al.**, 2024). Examining additional TCRs via all-atom MD simulation can inform how the mechanism is adopted. By finding similarities and differences between A6 and B7 TCRs, which recognize the same Tax pMHC, we can elucidate how mechanical force is utilized by each to impact the differential function of T cells.

Catch bond mechanisms generally differ among proteins (*Thomas et al., 2008*). For example, FimH and vinculin have relatively well-defined weak- and strong-binding states where there are corresponding crystal structures (*Bakolitsa et al., 1999*; *Le Trong et al., 2010*; *Sauer et al., 2016*; *Mei et al., 2020*). Availability of the end-state structures enable using simulation approaches such as enhanced sampling of individual states and studying the transition between the two states (*Languin-Cattoën et al., 2023*; *Ccoa et al., 2024*). In contrast, TCRs do not have any structurally well-defined weak- or strong-binding states, which requires a different approach. As demonstrated in our present work as well as in our previous papers (*Hwang et al., 2020*; *Chang-Gonzalez et al., 2024*), our microsecond-long simulations of the complex under realistic pN-level loads and a combination of analysis methods are effective for elucidating the catch bond mechanism of TCRs, which shows that the load-dependent stabilization of the TCR-pMHC complex differs from simple relaxation of the structure in the initial crystallographic confinement.

Recently, *Choi et al.* (2023) proposed a mathematical model of TCR catch bond formation. Their assumptions are based mainly on the steered MD (SMD) simulation (*Wu et al.*, 2019) where partial unfolding of MHC and tilting of the TCR-pMHC interface were considered as required. Our mechanism does not conflict with their assumptions since the complex in the fully folded state should first bear load in a ligand-dependent manner in order to allow any subsequent larger-scale changes.

We found that the dynamic allostery of the TCR chassis observed in A6 (Chang-Gonzalez et al., 2024) and IM22 (Hwang et al., 2020) is largely conserved in B7.  $C\alpha$  and  $C\beta$  domains form extensive contacts to construct a base, and the  $V\beta$ - $C\beta$  contacts are more extensive than  $V\alpha$ - $C\alpha$  contacts. This leads to an asymmetric V-C motion where  $V\alpha$  is more mobile relative to the C-module than  $V\beta$ . In turn, the motional asymmetry affects the relative positioning of CDR loops, as measured by the CDR3 distance as well as the orientation of the TCR-pMHC interface relative to the loading direction. Unless an adequate load is applied to the complex to suppress the motion, destabilization of the interface occurs. As noted in A6 and IM22, the C domain renders a disadvantage to binding except under force. This is indicated by the greater number of interfacial contacts and decreased  $V\alpha$ - $V\beta$ motion of  $V\alpha\beta$ -pMHC (no C-module) simulations. The increased interfacial stability of  $V\alpha\beta$ -pMHC is also consistent with our discovery that the C-module likely undergoes a partial unfolding to an extended state, where the bond lifetime increases (Das et al., 2015; Akitsu et al., 2024). Therefore, while the variable domain dictates TCR fit with pMHC, a logical evolutionary pressure would be for the constant domain to maximize discriminatory power by adding instability to the TCR chassis. Furthermore, considering single-chain versions of an antibody lacking the C-module (scFv) are in widespread use (Ahmad et al., 2012) including CAR T cells (Reinherz et al., 2023), a better understanding of a TCR lacking the C-module may help with developing a novel TCR-based immunotherapy. Given the conservation of the asymmetric motion, we also predict the C $\beta$  FG-loop deletion mutant of B7 TCR to possess a diminished catch-bond response, as our previous experiment (Das et al., 2015) and simulation (Hwang et al., 2020; Chang-Gonzalez et al., 2024) demonstrated for other TCRs.

While dynamic allostery is overall conserved, A6 and B7 differ in the behavior of the TCR-pMHC interface under load. The crystal structures of the two complexes have a comparable number of TCR-pMHC contacts, with a total of 33 for A6 and 29 for B7 in terms of residue pairs, while the number of distinct atom pairs forming contacts are 46 for A6 (*Garboczi et al.*, 996a) and 63 for B7

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(Ding et al., 1998). During MD simulations, many of these contacts become transient, resulting in fewer high-occupancy contacts. In the thermally fluctuating state, there is a 4-fold increase in the difference in number of contacts between A6 and B7 over the difference for static crystal structures (Figure 1C). Previously, contacts found in crystal structures have been used to explain differences in responses to point mutations to the Tax peptide. For example, the sensitivity of B7 to the mutation of Y5 of Tax was explained based on the Y5-D30α hydrogen bond and stacking of Y5 with Y1018 of B7 (Y1018 in the present study was numbered Y1048 in Ding et al. (1998) and Hausmann et al. (1999)). In comparison, the more amenable pocket for Y5 on A6 was suggested to be responsible for the greater tolerance of A6 under mutations of Y5. In MD simulations of B7, the Y5-D30 $\alpha$ hydrogen bond is formed only in B7high, after about 300 ns (Figure 2-figure Supplement 1C), and the Y5-Y101\(\theta\) nonpolar contact breaks during simulation of B7\(^0\) and is not present in B7\(^1\) (Figure 2-figure Supplement 1A.B) while it persists with 62% occupancy in B7<sup>high</sup> and 52% occupancy in  $V\alpha\beta$ -pMHC (*Figure 2C*,D; these occupancies are for the whole simulation period). Changes to T cell specificity by point mutations on Y5 are thus unlikely to arise solely from the size or the polarity of its binding pocket as a static structure. Instead, point mutations affect the organization and dynamics of the surrounding contacts in addition to fit (Chang-Gonzalez et al., 2024). In a related vein, intermolecular motional network has been suggested to be a controlling factor for allostery in a peptide-SH3 domain complex (Gomez et al., 2024), which aligns with the nonlocal and dynamic role of contacts at the TCR-pMHC interface. The inter-domain motion of TCR enables long-range allosteric discrimination of pMHCs, integrating the motion of the two entities such that they influence each other.

The 14.5-pN force on B7<sup>high</sup> was applied under 190.0-Å extension (*Table 1*), whereas the corresponding high-load simulation for A6 was with 18.2 pN at a slightly lower 187.7-Å extension (*Chang-Gonzalez et al., 2024*). The average total intra-TCR contact occupancy for B7<sup>high</sup> between 500-1000 ns is 30.4±0.49 contacts (*Figure 1*C), compared to 38.7±0.87 for A6 under high load (after 500 ns), which further supports a higher compliance of the former. With its higher compliance, the B7 TCR-pMHC complex needs to be under a greater extension than A6 to apply comparable levels of force, and it would be more difficult to achieve load-induced stabilization of the TCR-pMHC interface. This leads us to predict that B7 will exhibit a weaker catch bond where the peak of the TCR-pMHC bond lifetime is at a lower force. And the destabilization of B7<sup>high</sup> at 1300 ns (*Figure 1*D and *Figure 2*B) may have been a result of our constant-extension approach to apply load, which allows reaching low-force states by the compliant B7. Although this approach was motivated by the constant spacing between a T cell and an APC (*Reinherz et al., 2023*), the actin cytoskeleton also adjusts the load applied to the TCR-pMHC complex (*Feng et al., 2017*; *Hu et al., 2024*). How fluctuation in the applied load affects the TCR-pMHC dynamics is a subject of a future study.

As a related example to A6 and B7 TCRs, PA59 and PA25 TCRs share the same *TRBV* and *TRBJ* genes, and they share 11-aa CDR3 $\beta$  loops that differ only at a single position (Trp vs. Leu) (*Akitsu et al.*, *2024*). Recognizing the same pMHC (PA<sub>224-233</sub>/D<sup>b</sup>), their maximum bond lifetimes and the peak catch bond forces differ considerably, 75 s at 21 pN for PA59, and 13 s at 15 pN for PA25 (*Akitsu et al.*, *2024*). These differences may facilitate actions of the corresponding T-cells in tissues that present diverse mechanical environments provided by cell movement, adhesion molecules, tight junction, and non-compliant inflammation.

Likewise, T cells bearing A6 and B7 may also perform differently *in vivo* depending on tissue localization (*Akitsu et al.*, *2024*) even though they behave similarly *in vitro* in terms of cytotoxicity and secretion of select cytokines ( $\gamma$ -IFN, MIP-1 $\alpha$ , and TNF $\alpha$ ) (*Ding et al.*, *1998*; *Hausmann et al.*, *1999*). We also predict the different catch bond profiles of A6 and B7 will render the two types of T cells to respond differently when the pMHC copy number on the APC is limited (*Akitsu et al.*, *2024*). Differences in the catch-bond profiles may also affect the energy input for downstream signaling events that involve reversible conformational transitions in TCR $\alpha\beta$  within a certain range of forces and extends the TCR-pMHC bond lifetime by more than an order of magnitude (*Akitsu et al.*, *2024*). Elucidating details of those differences is beyond the scope of the present work.

Disparate biological outcomes between structurally similar TCRs recognizing the same pMHC are mechanically possible since slight changes in interfacial contacts can result in altered distribution of loads across the TCR chassis so that changes in its dynamics affect interaction with CD3 signaling subunits (*Reinherz et al.*, 2023). Structural details of the dynamic amplification and propagation of the recognition signal warrant further investigation.

# **Computational Methods**

# Structure preparation

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B7 TCR $\alpha\beta$ -pMHC was built from PDB 1BD2 (*Ding et al., 1998*) using CHARMM (*Brooks et al., 2009*; *Hwang et al., 2024*). Non-numeral residue IDs in the PDB were renumbered to follow sequential numbering used in the present study. We used MODELLER (*Šali and Blundell, 1993*) to generate coordinates for missing loops in the C $\alpha$  domain (S133-K136 and S170-D172 in the PDB numbering scheme) followed by a brief energy minimization. We visually verified MODELLER results, comparing generated loops to those of the related A6 TCR (*Garboczi et al., 996a*; *Ding et al., 1998*). The constant domain of TCR $\alpha$  (C $\alpha$ ) was also missing coordinates for F204-S210 (F198-S204 after renumbering), which were added with the TCR $\alpha$  linker as detailed below. Disulfide bonds were assigned between cysteine residues as defined in the PDB file. Crystal waters within 2.8 Å from the protein were kept for the truncated structures, and all waters were kept for the full structure.

Histidine protonation state was determined to promote hydrogen bond formation with neighboring residues. The histidine  $N^{\delta}$  atom was protonated as follows: MHC residues 3, 93, 114, 145, 151, 188, 260;  $\beta$ 2m residues 13, 51; TCR $\alpha$  all histidine residues; and TCR $\beta$  residues 29, 47, 154. For the remaining histidine residues, the  $N^{\epsilon}$  atom was protonated.

As done in *Hwang et al.* (2020) and *Chang-Gonzalez et al.* (2024), we extended the MHC and  $TCR\alpha\beta$  termini as handles for applying positional restraints. For MHC, we used UniProt P01892 to add <sup>276</sup>LSSQPTIPI<sup>284</sup>. For  $TCR\alpha$  we used GenBank AAA60627.1 to add <sup>205</sup>CDVKLVEKSFETDT<sup>218</sup>. For  $TCR\beta$  we used GenBank AAC08953.1 to add <sup>245</sup>CGFTSESYQQGVLSA<sup>259</sup>. We placed an interchain disulfide bond between  $\alpha$ C205- $\beta$ C245. Added strands in the initially straight conformations were relaxed to a state similar to that in *Figure 1*A by performing a series of brief energy minimization and MD simulation with the FACTS implicit solvent model (*Haberthür and Caflisch, 2008*).

Truncated structures were built based on the prepared B7 TCR-pMHC complex as:

- $V\alpha\beta$ : The last residues were  $\alpha$ P110 and  $\beta$ V113.
- $T\alpha\beta$ : The last residues were  $\alpha$ D206 and  $\beta$ G246 (no C-terminal strands).
- $V\alpha\beta$ -pMHC: includes  $V\alpha\beta$ , pMHC, and  $\beta$ 2m. The last residue of MHC was L276.
- B70: includes  $T\alpha\beta$ , pMHC, and  $\beta$ 2m. The last residue of MHC was L276.

## 353 MD simulation protocol

Solvation, energy minimization, heating, and equilibration of the B7 complexes followed the protocol in *Chang-Gonzalez et al.* (2024), except for systems which include the pMHC, where we modified the preparation protocol prior to production runs as detailed below.

## 57 Laddered extensions

Applying the same protocol as done for A6 to achieve the laddered extensions in B7 resulted in substantial breakage of the TCR-pMHC contacts within the first 50 ns in several production runs. To mitigate this, we introduced distance restraints to selected atom pairs forming contacts between the TCR and pMHC to prevent them from breaking during preparatory simulations. This ensured that the complex could structurally adapt as we modified the extension distance, yet all laddered extensions maintained a core set of initial TCR-pMHC contacts. Atom pair distance restraints were removed in production runs.

Twelve atom pairs between TCR and pMHC were selected that were within 5 Å of each other in the equilibration restart file of the TCR-pMHC complex with added linkers. A 2-kcal/[mol·Å²]

flat-bottom harmonic restraint potential was applied to keep the atom pair distance within the value at the end of the equilibration run. Then a 2-ns CPT simulation was carried out while also applying a 1-kcal/[mol·Å²] harmonic potential to the  $C_{\alpha}$  atoms of the C-terminal MHC (I284), TCR $\alpha$  (T218), and TCR $\beta$  (A259) residues. Production run followed upon removing the atom pair distance restraints . During the production run, the 1-kcal/[mol·Å²] harmonic potential to the TCR and MHC end-residues and a 10 Å flat-bottom harmonic distance restraint between  $\alpha$ T218 and  $\beta$ A259 were applied, the latter to prevent the ends of the C-terminal added strands from separating (*Figure 1—figure Supplement 1*). Throughout the production run, we intermittently measured average and rolling force in 40 ns intervals and found these to be around 10 pN, an ideal target for the low-load simulation. This simulation is the 173.7 Å B7low system reported in *Table 1*.

Using the structure at the end of the 2-ns simulation with the TCR-pMHC atom-pair distance restraints, we increased the extension by 8 Å by moving the center of the 1-kcal/[mol·Å²] harmonic potential on the end-residue  $C_{\alpha}$  atoms by 4 Å at each end. While keeping the TCR-pMHC atom-pair distance restraints applied, we launched another 2-ns simulation under the increased extension. Atom pair distance restraints were then removed and the production run was launched. The extension averaged after 500 ns of this simulation was 181.7 Å. Following the same way, we increased the extension by another 8 Å, which led to the 190.0 Å B7<sup>high</sup> system reported in *Table 1*. We also decreased the extension by 8 Å from the initial 173.7-Å extension, which led to a 165.7-Å extension.

Among the 4 extensions tested, the 181.7-Å extension was not selected primarily because the average force of the simulation from 500 ns to 850 ns (the total length of the simulation), was 9.26 pN, only barely higher than the reported load for B7<sup>low</sup>. For the 165.7-Å simulation, the average force from 500 ns to 900 ns (total length of the simulation) was 15.7 pN. We had observed this high force at low extension for A6 TCR (*Chang-Gonzalez et al., 2024*) and attribute this to folding of the flexible added strands leading to contacts between the stands and the TCR constant domains.

# $V\alpha\beta$ -pMHC and B70

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We also applied a 2-kcal/[mol·Å²] flat-bottom harmonic distance restraint during preparatory simulations of  $V\alpha\beta$ -pMHC and B7°. We attempted to use the same set of atom pairs as in the laddered extension simulations, but considerable interface breakage occurred, likely due to changes in interfacial contacts after equilibration in these systems. We thus selected different atom pairs for  $V\alpha\beta$ -pMHC and B7°. For consistency, we selected 12 atom pairs, the same number as in the laddered extensions, and distributed in the same way between the Tax peptide or MHC residues to  $V\alpha$  or  $V\beta$  residues. The distance restraint was applied to the atom pairs for a 2-ns CPT simulation, then released for production runs.

# Systems without load

The following additional restraints were used for systems without load.

- $V\alpha\beta$ ,  $T\alpha\beta$ : no positional restraints were applied.
- $V\alpha\beta$ -pMHC: we applied a weak 0.01-kcal/[mol·Å²] harmonic positional restraint to the backbone  $C_{\alpha}$  atoms of MHC  $\alpha$ 3 (P185-L276) to prevent large transverse rotation of the whole molecule in the orthorhombic box.
- B7°: we applied a 0.2-kcal/[mol·Å²] harmonic positional restraint to the backbone  $C_{\alpha}$  atoms of MHC  $\alpha$ 3 with RMSF less than 0.5 Å calculated from the simulations of B7low. These residues were: L201-Y209, T240-O242, T259-H263.

## Production runs

Production runs were performed similar to *Chang-Gonzalez et al.* (2024). We used OpenMM (*East-man et al.*, 2017) with the CHARMM param36 all-atom force field (*MacKerell Jr et al.*, 2004) and the particle-mesh Ewald method to calculate long-range electrostatic interactions. We used an Ewald error tolerance of  $10^{-4}$  which is 1/5 of the default value in OpenMM and a 12 Å cutoff distance for

- nonbonded interactions. The complexes were simulated at 300 K with a 2-fs time step using the
- Nose-Hoover integrator in OpenMM. Production run lengths are in *Table 1*.

# 416 Trajectory analyses

- 417 Analysis methods are detailed in Chang-Gonzalez et al. (2024). Below we mainly explain B7-specific
- residue selections. We used the frames for 500-1000 ns of the production runs to calculate the
- <sub>419</sub> average and standard deviation of the number of contacts, BSA, CDR3 distance, PCA, and triad and
- V-C angles. With a coordinate saving rate of 20 ps, this leaves at least 25,000 frames for analysis.

## V-module to pMHC distance

- The distance from TCR V-module to pMHC (Figure 2-figure Supplement 1F) was measured between
- the center of mass of the  $C_{\alpha}$  atoms of the same residues used to build the V-module triads (de-
- scribed below) to the center of mass of five C<sub>n</sub> atoms from each of the central 4 strands forming
- the  $\beta$ -sheet floor located above the  $\alpha 1$  and  $\alpha 2$  helices of MHC (20 MHC atoms in total; *Figure 1*A).
- <sup>426</sup> These were R6-T10, I23-Y27, Q96-G100, and Y113-A117. RMSF of these residues after 500 ns was
- below 1.4 Å in all B7 systems, so the measured distance is minimally affected by the intra-domain
- conformational motion.

## 429 CDR3 distance

- Distance between CDR3 $\alpha$  and CDR3 $\beta$  (Figure 3E, Figure 4F) was measured using the midpoint be-
- tween backbone  $C_{\alpha}$  atoms of two residues at the base of each CDR3, which are E93 and K97 for
- 432 CDR3 $\alpha$  and S94 and E102 for CDR3 $\beta$ .

#### 433 V-module triads

- We assigned triads (*Figure 3B*) based on the backbone  $C_{\alpha}$  atoms of the stably folded  $\beta$ -sheet core of
- each variable domain (Hwang et al., 2020; Chang-Gonzalez et al., 2024). Selected residues for triad
- assignment of the B7 systems were as follows. For  $V\alpha$ , I19-Y24, F32-K37, H71-I76, and Y87-M92. For
- Vβ, T20-Q25, M32-Q37, D73-L78, and Y89-S94. Prior to triad assignment we aligned all complexes
- to the first frame of B7<sup>low</sup> using the selected residues to monitor the relative motion between the
- two triads without global translation nor rotation.

#### 440 V-C BOC

- We assigned BOCs (Figure 4B) as detailed in Hwang et al. (2020); Chang-Gonzalez et al. (2024). To
- place beads for the C-module, we used the following residues. For  $C\alpha$ , A118-R123, V132-D137, Y153-
- T158, and S171-S176. For C<sub>B</sub>, L143-T148, L157-N162, L190-R195, and F208-O213. For the hinges,
- we used:  $\alpha$ N114 for H $\alpha$ , and  $\beta$ D116 and  $\beta$ L117 for H $\beta$ . We aligned all complexes to the backbone
- 445 C<sub>x</sub> atoms of the selected residues of B7<sup>low</sup> then built BOCs to monitor the motion of the V-module
- relative to the C-module.

# 447 Acknowledgments

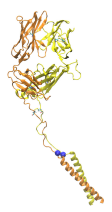
- This work was funded by US National Institutes of Health Grants P01AI143565 and R01AI136301.
- 449 Simulations were performed by using computers at the Texas A&M High Performance Research
- 450 Computing facility.

## References

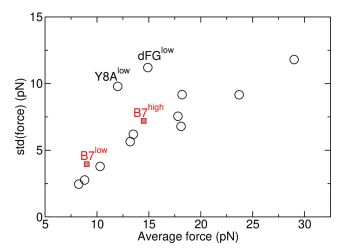
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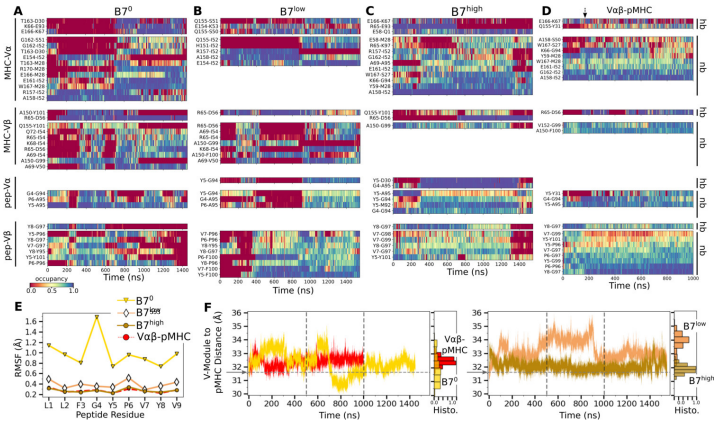


**Figure 1—figure supplement 1.** Model of TCR $\alpha\beta$  with leucine zipper domain fused to the C-terminal added strands. Model was built based on PDB 1NFD (*Wang et al., 1998*). The construct was used in single-molecule optical tweezers experiments (*Das et al., 2015*). Blue spheres denote  $C_{\alpha}$  atoms of residues corresponding to  $\alpha$ T218 and  $\beta$ A259 of B7 TCR (*Figure 1*A, bottom). Their distance is 6.7 Å. The 10-Å flat-bottom harmonic distance restraint on those atoms (see Methods) mimic the presence of the leucine zipper that prevents large separation of the added strands.

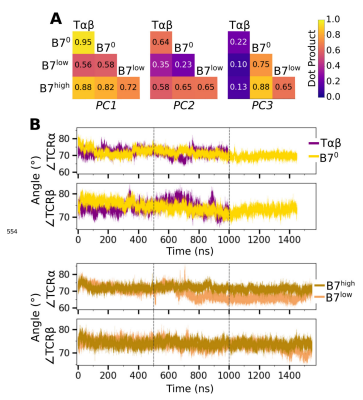


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**Figure 1—figure supplement 2.** Standard deviation vs. the average of the applied force. All data are measured between 500-1000 ns. Open circles are for A6 (*Chang-Gonzalez et al., 2024*). Red squares for B7<sup>low</sup> and B7<sup>high</sup> are from Table 1. Except for Y8A<sup>low</sup> (an antagonist) and dFG<sup>low</sup> (C $\beta$  FG-loop deletion mutant) for A6, the near-linear relation between the std and average in force is a consequence of the force being applied to the restraining potential via random conformational fluctuation of the complex (*Burgess, 1973*). See Appendix 3 of *Chang-Gonzalez et al.* (*2024*) for further explanation. The fact that data for both B7 and A6 lie on approximately the same line also reflects consistency in our force measurement.



**Figure 2—figure supplement 1.** Additional characterization of the TCR-pMHC interface. (A–D) Contact occupancy heat maps. (A) B7<sup>0</sup>, (B) B7<sup>low</sup>, (C) B7<sup>high</sup>, and (D) Vαβ-pMHC. Contacts with overall occupancy greater than 30% and instantaneous occupancy greater than 80% are shown. hb: hydrogen bond, np: nonpolar contact. Arrow in panel D denotes the approximate time when the initial adjustment of contacts in Vαβ-pMHC happens (**Figure 2**B). (E) RMSF of peptide backbone C<sub>α</sub> atoms between 500-1000 ns. C<sub>α</sub> atoms were aligned to those at the beginning of the production run for RMSF calculation. (F) Distance between the V-module and pMHC. Histograms were calculated using data between 500-1000 ns, the interval between the vertical dashed lines. For Vαβ-pMHC and B7<sup>high</sup>, avg±std between 500-1000 ns are 32.5±0.31 Å and 31.9±0.30 Å, respectively. V-module to pMHC distance measured from the B7 crystal structure (1BD2) is denoted by a dashed horizontal line indicated by an arrow at 31.6 Å.



**Figure 4—figure supplement 1.** Additional characterization of the V-C motion. (A) Dot products computed between the BOC PC vectors of listed systems. Values closer to 1.0 denote similar V-C BOC direction of motion. (B) V-C hinge angle trajectories over time.