NMR-directed design of pre-TCRβ and pMHC molecules implies a distinct geometry for pre-TCR relative to αβTCR recognition of pMHC

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The pre-T cell receptor (pre-TCR) guides early thymocytes through maturation processes within the thymus via interaction with self- ligands displayed on thymic epithelial cells. The pre-TCR is a disulfide-linked heterodimer composed of an invariant pre-TCR α (pTα) subunit and a variable β subunit, the latter of which is incorporated into the mature TCR in subsequent developmental progression. This interaction of pre-TCR with peptide-major histocompatibility complex (pMHC) molecules has recently been shown to drive robust pre-TCR signaling and thymocyte maturation. Although the native sequences of β are properly folded and suitable for NMR studies in isolation, a tendency to self-associate renders binding studies with physiological ligands difficult to interpret. Consequently, to structurally define this critical interaction, we have re-engineered the extracellular regions of β, designated as β-c1, for prokaryotic production to be used in NMR spectroscopy. Given the large size of the full extracellular domain of class I MHC molecules such as H-Kb, we produced a truncated form termed Kβ-t harboring properties favorable for NMR measurements. This system has enabled robust measurement of a pre-TCR-pMHC interaction directly analogous to that of TCRαβ-pMHC. Binding surface analysis identified a contact surface comparable in size to that of the TCRαβ-pMHC but potentially with a rather distinct binding orientation. A tilting of the pre-TCRβ when bound to the pMHC ligand recognition surface versus the upright orientation of TCRαβ would alter the direction of force application between pre-TCR and TCR mechanosensors, impacting signal initiation.

αβT cell–mediated immunity serves to combat cellular-based pathogens that afflict the vertebrate host, including those caused by viruses and cancers. At the core of T cell recognition and activation function is the surface T cell receptor (αβTCR), consisting of a clone-specific TCRαβ heterodimer and invariant dimeric CD3 signaling subunits (CD3ζγ, CD3ζδ, and CD3ζζζζ) that are non-covalently associated to form an eight-transmembrane-component complex. TCRαβ recognizes “foreign” peptides derived from proteins unique to a pathogenic process such as a viral protein or mutated host gene product within a transformed cell. A key feature of this recognition is that a peptide is presented in the context of peptide-major histocompatibility complex (pMHC) molecules on the surface of the abnormal cell or cross-presented on dedicated antigen-presenting cells that acquire the protein and surface-array the peptide. The pathogenically derived foreign pMHCs must be recognized in the context of tens of thousands of self-pMHCs generated as a result of the breakdown of endogenous proteins that are displayed on the same host cell. This discrimination is tuned through the harnessing of external cellular forces generated as a T cell scans its environment for foreign pMHC during immune surveillance as well as through the activation of intracellular cytoskeletal forces that dynamically tune immune recognition (1–5). Indeed, it appears that the T cell utilizes piconewton-level forces to initiate signaling-linked structural transitions that extend TCR–pMHC bond lifetime (1, 4, 6, 7), although the atomistic details of this are yet to be fully resolved.

The requirement for exquisite ligand discrimination, to selectively sense foreign pMHC over self-pMHC, drives selection processes in thymocyte development to eliminate αβTCRs with excess self-reactivity without removing those harboring the potential to bind foreign peptides. This is no simple task as a protective αβTCR should recognize self-MHC molecules while distinguishing foreign peptides from self-peptides in the groove of the self-MHC molecule. This is more difficult consid-

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NMR resonance assignments for VSV8/Kb and VSV8/Kb-t were deposited in the Biological Magnetic Resonance Data Bank (BMRB) with accession numbers 27285 and 27286, respectively.

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The abbreviations used are: TCR, T cell receptor; pTα, pre-TCR α; pre-TCR, pre-T cell receptor; pMHC, peptide-major histocompatibility complex; CDR, complementarity-determining region; VSV8, vesicular stomatitis virus nucleoprotein octapeptide, RGYVYQGL; TROSY, transverse relaxation optimized spectroscopy; CCSC, combined 1H-15N chemical shift change; CST, cross-saturation transfer; TRACT, transverse relaxation correlation time; βm, β2-microglobulin; HSQC, heteronuclear single quantum coherence; τc, rotational correlation time; ib, inclusion body.

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ering that at the TCR–pMHC interaction surface the TCR makes many more atomic contacts with the MHC than with the peptide.

The pre-TCR pTα-β heterodimer is an obligate precursor to TCRαβ in the development of αβ T cells (8). Pre-TCRs express the somatically rearranged TCRβ gene product, the β subunit, paired with an invariant pTα subunit, itself comprising a single Cα-like constant domain and lacking a variable domain and associated complementarity-determining regions (CDRs), which drive ligand recognition within TCRαβ. Nevertheless, the pre-TCR is itself a fully functional receptor that includes the same CD3 subunits to initiate signaling (6, 9). Both the CDR loops of Vβ and a hydrophobic patch on Vβ, exposed as a consequence of being in an unpaired V module topology, mediate ligand binding to pMHC (9). The self-pMHC interaction specificity of the pre-TCR drives expansion of the CD4–CD8 double-negative thymocytes (6, 9) and appears to foster preferential expansion of those thymocytes that recognize peptides in the double-positive stage and/or further development of those thymocytes that recognize peptides in the double-positive stage (8, 12–14).

Because the pre-TCR plays a major role in this two-step T cell repertoire refinement, we sought to characterize the binding event between β and pMHC to decipher the parameters of this recognition relative to those of the TCRαβ–pMHC interaction (9). More importantly, we would like to delineate differences between the pre-TCR and the αβTCR recognition events to better understand the biological function of the pre-TCR while providing new insights into TCR function (3, 4, 6). Due to inherent self-association of component molecules (15) and the large size of the complex, attempts to fully define the interactions have been difficult (9, 15). We therefore designed a model system to explore the pre-TCR–pMHC interaction in atomic detail. Experiments presented here establish a working model of the pre-TCR–pMHC interaction through the design of a modified β chain (9) and a truncated mutant of pMHC (16) to describe the pre-TCR–pMHC interaction surfaces.

Results

Interaction of pMHC with the β subunit of the pre-TCR is hampered by self-association of β through the constant domain

In an attempt to characterize an interaction between a β chain and a pMHC, we utilized two distinct β chains with only 34% sequence identity within their Vβ regions, N30β and N15β, for which the αβTCR of each recognizes the same pMHC, a vesicular stomatitis virus nucleoprotein octapeptide bound to H-2Kb (VSV8/Kb) (9, 17, 18). Each β subunit promotes β selection in thymic culture systems (9) and could potentially provide complementary information on the pre-TCR–pMHC complex. We first added unlabeled VSV8/Kb to a purified 15N-labeled sample of N30β and acquired 1H-15N HSQC spectra with transverse relaxation optimized spectroscopy (TROSY) selection of the mixture and compared it with a spectrum of N30β. Utilizing the backbone assignments of N30β (18), we were able to determine the interaction site by mapping the combined 1H-15N chemical shift changes (CCSCs) on a previously determined X-ray crystallographic model (15) (Fig. 1, A and B). The largest chemical shift changes were not localized to any single site on the surface of N30β but did cluster in the C domain at the conserved Cα– or pTα-Cβ interface (19, 20). Additional changes were seen in the C domain AB and EF helices, which are located proximal to the C terminus of the TCR. There were some scattered changes within the V domain, but these were not clustered as tightly as those identified in the C domain. This result suggested that any V domain interaction was weaker than the nonspecific interactions within the C domain, which is not known to interact with pMHC in the TCRαβ, making it unlikely that we were observing physiologically relevant interactions. To understand whether the C domain–mediated nonspecific interactions were a general phenomenon, we next studied the interaction of pMHC with N15β. Because N15β is known to self-associate (15) via Vβ, we used cross-saturation transfer (CST) NMR experiments, which highlight only regions of heteromolecular association, in contrast to chemical shift perturbation analyses, which also highlight changes in surfaces due to self-association or allosteric changes. In CST experiments, the observed protein is 15N-labeled and perdeuterated, which implies that it will have no resonances in the aliphatic region of the 1H NMR spectrum. The binding partner, in this case pMHC, will be protonated and will harbor resonances in the aliphatic region, but because this protein is not 15N-labeled it will be invisible in a 15N HSQC spectrum. In CST experiments, we selectively magnetize the unlabeled protein and monitor transfer of magnetization to the labeled protein, which will reveal the direct interaction interface. In the case of N15β, we found a significant CST effect in the Vβ as well as Cβ (Fig. 1, C and D). Because the pre-TCR contains a pTα subunit that lacks a Vα-like subunit (Fig. 2A), we posit that the changes seen in the V domain were more physiologically relevant than those seen in the C domain (20).

Structure-aided design of β as a pre-TCR model

Fig. 2A shows the overall topology of the pre-TCR pTα-β heterodimer (20), illustrating that the interaction surfaces highlighted in the N15β-VSV8/Kb interaction are occluded by pTα in the C domain but are not occluded in the V domain (Fig. 1, C and D). When examining the crystallographic model of the pre-TCR, we noted a hydrophobic cluster, including residues Phe-131, Val-147, and Leu-149 that are occluded in the pre-TCR (Fig. 2B). We hypothesized that mutating the equivalent residues in N15β or N30β would abrogate the interaction within this non-physiologically relevant binding surface. It was thus clear that to accurately judge the binding capacity of TCRβ, measuring only interaction potential of elements exposed on cell surfaces, we needed to remove the non-physiologic inter-
action within the Cβ interface normally contacting Cα or pTα (19, 20). Due to the placement of the hydrophobic residues Phe-131, Val-147, and Leu-149 in β-sheet secondary structures, we chose replacements with known β-strand–forming propensities (21), F131R, V147Q, and L149Q, each of which places a charged or hydrophilic residue at this interface.

**Mutagenesis of Cβ**

The 15N HSQC spectrum of the N30β mutant F129R/V145Q/L147Q, denoted N30β-c1, is quite similar to that of the WT (Fig. 2C) as is the case with N15β F128R/V144Q/L146Q (N15β-c1) compared with WT (Fig. 2D), indicating an overall similar fold of the molecule. When examining residues in the Cβ domain of N30β-c1, these resonances are shifted compared with those in N30β, whereas those in the V domain remain unchanged (Fig. 2E), indicating that any changes due to the mutation are local and do not affect V domain–binding regions. We conclude that the replacement of the Phe/Val/Leu hydrophobic cluster resulted in a chain with a fold similar to that of the WT (9, 18).

**Rotational correlation time** (τc) **measurements show diminished self-association in β-c1**

Using NMR, one can measure relaxation rates from which molecular correlation times such as τc can be extracted. The τc may report on the oligomeric states of the observed protein, in solution, at the given concentration and in equilibrium where the interacting partners are not distinguished. Utilizing transverse relaxation correlation time (TRACT) experiments (22), we were able to measure τc values, which are positively correlated to the apparent size of a molecule, inclusive of oligomeric state, for N30β and N15β (Fig. 2F). τc increased with increasing
Figure 2. Mutagenesis targets within the pre-TCR β subunit. A, pre-TCR crystal structure model (Protein Data Bank code 3OF6) in schematic representation in the same orientation as Fig. 1B, expanded view of boxed area of A showing only TCRβ. The view is rotated −15° about x and y to highlight the contact surface. Side chains of residues contacting (i.e. within 4 Å of) pTa are shown in stick representation. Three hydrophobic residues targeted for mutagenesis are colored yellow. Numbering is according to that found in the crystal structure 3OF6 and is offset from those of N15 and N30β by three or two residues, respectively. C and D, overlay of 1H-15N TROSY-HSQC of N30β (18) and N30β-c1 (9) (C) or N15β (18) and N15β-c1 (9) (D) to illustrate similarity of spectra. E, region of overlaid spectra shown in C with labeled peaks indicating lack of chemical shift changes in the V domain (Thr-6, Thr-32, Asp-25, Leu-111, and Leu-117), whereas those in the C domain (Val-125, Tyr-184, Arg-189, Gln-209, and Phe-210) show significant changes. F, rotational correlation times of WT and mutant TCRβs suggest abrogation of self-association within Cβ. The plot shows τc versus concentration for N15 (blue diamonds) and N30β (red triangles). Lines show exponential fits of points to guide the eye. WT is shown with filled symbols and solid lines, and c1 mutants are shown with open symbols and dotted lines.
pre-TCR–pMHC interaction



**Figure 3.** Three-dimensional correlations for backbone assignment of VSV8/Kb. A, selected regions of the HNCA experiment illustrating the quality of the spectrum for a given section within the α1 helix. B, selected regions of the HNACAB experiment for the same section as in A.

concentration for each protein, ranging from 18 to 23 ns for N30β at 50–500 μM, whereas N15β varied from 23 to 32 ns over similar concentrations. At the lowest concentrations measured here, the τc was 18.6 and 22.7 ns. These values were greater than the predicted value of 16.3 ns for these 27-kDa proteins as calculated using the software HYDRONMR (23). Although we cannot rule out self-association via Vβ, we posit that at least part of this self-association is the result of intermolecular interactions of Cβ (15). N15β self-associates more than N30β (p = 0.028), supporting the idea that N15β utilizes the V domain more in self-association. τc for N30β-c1 was found to be 15.2 and 15.8 ns for 100 and 200 μM samples, respectively (Fig. 2F). This is consistent with essentially monomeric TCRβ as determined from hydrodynamic radius calculations using HYDRONMR (23) wherein a value of 16.3 ns is predicted for the 27-kDa monomer. This supports the idea that N30β-c1 exists as a monomer at these concentrations and self-associates significantly less than WT (p = 0.035). In contrast, N15β-c1 exhibits an increase in τc at concentrations from 100 to 500 μM (Fig. 2F) with values comparable with those seen for the WT N30β but significantly less self-association than WT N15β (p = 0.032). It is expected that N15β-c1 continues to self-associate more than N30β-c1 (p = 0.042) because X-ray crystallographic analyses have revealed Vβ-Vβ association in N15β-c1, whereas N30β-c1 contained contacts mainly utilizing Cβ (15). This analysis also agrees with observations that N15β will utilize the Vβ hydrophobic patch more readily in the context of pre-TCR recognition of pMHC (9).

**Backbone resonance assignments of full-length VSV8/Kb**

In our previous study, we were able to demonstrate an interaction between N15β-c1 and labeled Kb within the VSV8/Kb complex (9). To characterize the binding site on pMHC, we required the backbone resonance assignment of VSV8/Kb. Utilizing non-uniform sampling (24), we collected high resolution TROSY versions of HNCA/HNCOCA/HNCO/HNCACO/HNCACB spectra on perdeuterated 2H/13C/15N-labeled heavy chain with perdeuterated but otherwise unlabeled β2m, the smaller protein subunit of the MHC heterotrimer, and unlabeled VSV8. As shown in Fig. 3A, signals of cross-peaks were quite strong in the HNCA experiment with signals for both the HN correlated and its preceding (i and i−1, respectively) residue in each spin system. Conversely, for the HNCACB (Fig. 3B), most spin systems lack cross-peaks for Cβ−1, whereas ~30% lack cross-peaks for Cα. Similarly, the HNCO experiment provided robust Co−1 cross-peaks, whereas the HNACO lacked cross-peaks for most Co resonances. Because of the sparse coverage in sequential connectivities, we pursued orthogonal means of backbone assignment using the program RASP (25) to iteratively utilize chemical shift predictions to find potential resonance assignments. These assignments were cross-checked manually against a high-resolution 15N dispersed TROSY-HSQC-NOESY experiment for confirmatory cross-peaks indicative of through-space contacts. This procedure was critical in assigning the β-sheet that forms the base of the peptide-binding groove and those that comprise the α3 domain and allowed the completion of 73% of the non-proline backbone amide resonance assignments (Fig. 4A). Depicted in Fig. 4, B and C, is the completeness of assignment overlaid on the X-ray structure of VSV8/Kb (26). Notably, the α2 helix, proximal to the peptide-binding loop, with a likely role in pre-TCR binding, remained largely unassigned. Whether this is related to the specific peptide in the MHC groove or a more common feature of Kb per se requires further study but implies that mobility in this region might have important consequences for dynamics of αβ TCR–pMHC as well as pre-TCR–pMHC interaction.

**Large size of VSV8/Kb–N15β-c1 complex impacts measurement of binding interface**

The backbone assignments were then used to interpret the previously reported 1H-15N TROSY-HSQC measurement of
unlabeled N15β addition to VSV8/Kb wherein the heavy chain of Kb was 15N-labeled (9). As shown in Fig. 5A, chemical shift perturbation data analysis suggests a binding interaction involving the canonical peptide-binding groove previously reported for all TCRαβ–pMHC interactions. However, the data are not unequivocal, with a number of residues distal to the canonical surface and within the α3 subunit revealing changes similar to those seen at the putative interfacial region. The regions associated with the largest peak intensity losses (Fig. 5B) conform to the region surrounding the peptide, clustering more closely to the canonical binding interface. Whether due to the large size of the complex or the weak nature of the binding event, a more precise interaction map could not be obtained. This, combined with the incomplete assignment of the region surrounding the peptide-binding pocket, led us to attempt to engineer improvements in the biophysical behavior of the pMHC complex.

**Engineering a truncated pMHC model**

The inability to obtain the complete backbone resonance assignments of the receptor-binding portion of the MHC may have been due to the large size of the pMHC complex or the conformational heterogeneity within the complex. TRACT measurements of VSV8/Kb at 100 μM indicate a τc of 32 ns, corresponding to a predicted molecular mass of 85 kDa for a spherical molecule, nearly double that of the 45-kDa monomeric pMHC. This long correlation time may be caused by the oblong shape of the molecule but regardless explains quite well the poor quality of the triple-resonance backbone experiments. To obtain more complete backbone assignments while improving the quality of spectra when measuring the β-pMHC complex, we designed a truncated Kb molecule (termed Kb-t) lacking the α3 domain and β2m. TRACT measurements at 67 μM indicate that VSV8/Kb-t is monomeric with a τc of 15 ns, corresponding to an expected molecular mass of 28 kDa, which is in better agreement with its molecular mass of 22 kDa.

**Backbone resonance assignment of VSV8/Kb-t**

We proceeded to assign backbone resonances for 2H/13C/15N-labeled VSV8/Kb-t, acquiring triple-resonance experiments as for the full-length VSV8/Kb. The HNCA shows a complete cross-peak complement for each HN-correlated spin system (Fig. 6A) as does the HNACAB (Fig. 6B). This allowed a more facile assignment of the majority of HN resonances present within the TROSY-HSQC spectrum, assigning 139 of 178 non-proline residues (78%; Fig. 7A). Surprisingly, similar to the full-length VSV8/Kb (Fig. 4), the α2 helix flanking the peptide-binding groove remained recalcitrant to assignment with 19 residues within the α2 helix as well as nine more directly contacting this region remaining unassigned. We suspect, given the high quality of the spectra as well as the known mobility of the α2 helix (27), that this inability to assign this region of the protein reflects an intermediate (−ms−s)-timescale motion (28) on the part of the peptide and α2 helix, broadening these resonances. That it is unassigned in both full-length and truncated constructs argues that this is an intrinsic property of the molecule and not the result of experimental manipulation.
As the truncated K\textsuperscript{b} construct appeared folded in a manner consistent with the fold of the full-length heterotrimeric molecule, we tested pre-TCR recognition of VSV8/K\textsuperscript{b}-t by incubation with \textsuperscript{15}N-labeled N15\textbeta-c1. Spectral changes (Fig. 8A) are consistent with the binding event reported previously (9) with
chemical shift and intensity changes consistent with binding kinetics in the fast-to-intermediate exchange regime (28). The changes are almost exclusively located in Vβ (Fig. 8, B and C) with significant overlap between regions with chemical shift perturbation and intensity loss. The largest chemical shift and intensity changes are located in the CDR1 and -3 loops as well as Vβ/H9252 patch region. Note that the CDR2 loop is not assigned likely due to conformational exchange. A region with smaller chemical shift changes is found distal to the interaction surface but still within Vβ/H9252, clustered within a region encompassing residues 15–25 (the AB loop and B strand) and residues 74–85 (E strand and EF loop). These changes may occur as a result of secondary rearrangements within the binding interface and are consistent with a dynamic view of pre-TCR or TCR interactions with pMHC (4, 6, 9, 18, 29). The magnitude of chemical shift changes are similar when 200 μM N15β-c1 is added to full-length VSV8/Kb compared with VSV8/Kb-t with the 95th percentile of CCSC equal to 0.064 (9) versus 0.060, respectively. Peak intensity losses, as expected, were lower overall due to the smaller size of the β-pMHC complex when truncated pMHC was used with median intensity loss (thus measuring residues not directly interacting with TCRβ) of 47% (9) when the full-length MHC is utilized. Less overall peak broadening allowed more precision in measuring peak positions for chemical shift perturbation analysis and allowed a greater dynamic range for distinguishing exchange broadening leading to peak intensity losses (Fig. 8, B and C).

**Pre-TCR recognizes a surface similar to TCRαβ on pMHC**

Because we were able to assign a significant portion of the putative interface, we proceeded to measure 1H-15N TROSY-HSQC spectra of 1H/15N-labeled Kb-t bound to unlabeled VSV8. With titration of unlabeled N15β-c1, we measured chemical shift changes and intensity losses within Kb-t (Fig. 9, A and B). We observed significant chemical shift changes localized to the α1 helix, which comprises residues 57–85, and assigned portions of the α2 helix, which comprises residues 138–176, as well as residues forming the “floor” of the peptide-binding groove (Fig. 9, C and D). Despite the lack of assignments within the α2 helix and the peptide VSV8 itself, these chemical shift perturbations define a contiguous binding interface. The chemical shift perturbation map compares favorably with that measured using the full-length Kb molecule (Fig. 5), demonstrating the validity of improvements made by shortening the construct. Indeed, if one overlays the chemical shift perturbation map obtained with the truncated pMHC onto the structure of VSV8/Kb bound to N15β(30), the extent of the pre-TCR–pMHC interaction matches with the TCRαβ–pMHC interface determined by X-ray crystallography (Fig. 10). This strongly suggests that the pre-TCR recognizes a similar surface on pMHC as does TCRαβ. To highlight the distinctness...
of geometries that the respective interfaces place upon the molecular complex, we utilized HADDOCK2.2 web server (31) to generate a preliminary model of the pre-TCR–pMHC interaction interface (Fig. 10D). Given the lack of orientational restraints, it is not surprising that the low-energy conformations clustered into two distinct docking orientations offset by ~180° as typified by the three lowest-energy conformations shown here. Each of the models utilizes the CDR2 and -3 loops as well as the surface that normally interacts with Vα within the αβ heterodimer, i.e. the Vβ patch and surrounding regions. It is
likely that interaction of CDR3 and the \( \gamma/\beta \) patch also requires neighboring CDR2 to participate in the modeled complex despite the absence of NMR-generated constraints. The models also provide conformations that do not result in obvious clashes with known pre-TCR (15, 20, 32) or pMHC glycosylation sites (33) (Fig. S1). The apposition of these elements results in a more

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**Figure 9. NMR detection of interaction between N15\( \gamma \)-c1 and VSV8/K\( \kappa \)-t.** A, selected regions of \(^{1}H^{15}N\) TROSY-HSQC spectra highlighting changes with addition of unlabeled N15\( \gamma \)-c1 to \(^{15}N\)-labeled VSV8/K\( \kappa \)-t. Spectra are overlaid for VSV8/K\( \kappa \)-t alone (red) and with addition of 200 (pink), 300 (lavender), or 500 (blue) \( \mu \)M N15\( \gamma \)-c1. B, CCSCs for spectra outlined in A for 200 (dark blue), 300 (blue), or 500 (cyan) \( \mu \)M N15\( \gamma \)-c1. Residues that differ at the \( p < 0.01 \) (**) or \( p < 0.05 \) (*) level between the CCSC for that residue and the median CCSC for each experiment are indicated. C and D, schematic representation of VSV8/K\( \kappa \)-t with assigned residues colored blue, unassigned residues in white, prolines in cyan, and VSV8 in purple. Significantly perturbed residues by CCSC at \( p < 0.01 \) are highlighted as red spheres, and those that differ at \( p < 0.05 \) are shown as orange spheres. Residue identities are indicated. D, view rotated by 90° about the \( x \) and \( y \) axes relative to C with the \( \alpha2 \) helix proximal.

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**Figure 10. Modeling pre-TCR–pMHC complex.** A–C, location of residues participating in interaction with N15\( \beta \) in the N15\( \beta \)-VSV8/K\( \kappa \) X-ray structure (30). A, interacting residues indicated by red spheres were defined by CCSC differing at the \( p < 0.01 \) level between the CCSC for that residue and the median CCSC for each experiment. The VSV8/K\( \kappa \)-t structure is depicted with assigned residues colored blue, unassigned residues as well as \( \alpha3 \) and \( \beta2m \) subunits not present in this construct colored white, and VSV8 in purple. B and C are zoomed in to highlight the interaction site with \( \alpha2 \) proximal in B and rotated 180° about the \( y \) axis and \( \alpha1 \) proximal in C. D, three representative conformers of N15\( \gamma \)-c1 interacting with VSV8/K\( \kappa \) as determined by molecular modeling using CST data for the N15\( \gamma \)-c1 interaction surface (9) and VSV8/K\( \kappa \)-t CCSC data (Fig. 9) as ambiguous restraints input for HADDOCK2.2 (31). The conformers are representative of the three lowest-energy clusters of binding conformations and are color-coded with one shown as van der Waals surface representation and two shown in ribbon. The C\( \beta \) domain membrane-proximal end for each conformer is indicated with a color-coded "C." The models are aligned by overlay of the pMHC, shown in gray schematic with \( \alpha1 \) and \( \alpha2 \) labeled. E, side view with only conformer 1 of \( \beta \) shown and differing by a rotation of 90° about the \( x \) axis relative to D. VSV8/K\( \kappa \)-t is in the same approximate orientation as VSV8/K\( \kappa \) shown in A with \( \alpha2 \) facing front.
Pre-TCR–pMHC interaction

horizontal orientation of the β subunit relative to the MHC peptide-binding groove, suggesting an acute angle of contact between pre-TCR and pMHC. This is readily observed by comparing the long axis of the β subunit shown in Fig. 10A with that in Fig. 10E.

Discussion

This study highlights the rational design of recombinant prokaryotically expressed proteins, which in the wild-type form are not ideal for NMR studies. Specifically, domains that self-associate or interact within non-physiologically relevant regions (Figs. 1 and 2) in comparison with their eukaryotically expressed transmembrane protein counterparts were rendered less reactive, whereas proteins that are simply too large for NMR to measure with sufficient precision (Figs. 3–5) have been pared down to their essential components. Structure-based design considerations allowed these hurdles to be overcome, thereby acquiring more accurate measurement of their binding capacity, thus leading to more biologically important observations (Fig. 10 and Refs. 6 and 9). TRACT measurements clearly report the predicted decrease in τc and thus apparent molecular size at any given concentration with mutation of Cβ residues. Additionally, the increase of τc with increasing concentration is lesser for the c1 mutant, making readily apparent the difference in molecular behavior conferred by these mutations. Similarly, building on prior design of pMHC for molecular studies (16), we were able to improve an existing model system for NMR study through modification of pMHC (Figs. 6–9).

The chemical shift perturbation analysis of N15β binding to VSV8/Kb-t implies a binding mode in which the pre-TCR utilizes a similar binding interface on pMHC as seen for TCRαβ. As shown by the overlay of chemical shift perturbation data for the pre-TCR model with crystallographic data for the TCR system (Fig. 10, A–C), the interaction surface is similar between the systems. Once high-resolution structures are available, it will be valuable to compare the determinants of binding for the pre-TCR system versus the TCR, particularly focusing on the complementarity of the interacting surfaces as the weaker binding as measured by biomembrane force probe (9) or optical tweezers (6) for the N15β-c1 for VSV8/Kb as compared with that of N15αβ (9) implies suboptimal contacts in the former despite apparently similar contact area with pMHC ligands. The fact that the only other pre-TCR tested, N30β, exhibits even more attenuated binding than N15β (9) suggests that weaker interactions are the norm for pre-TCR. In contrast, we have noted that the pre-TCR−N15β exhibits a force-bond lifetime profile similar to that of the OT1 αβTCR-Ova/Kb system (9). It should be noted, additionally, that there are likely an array of self-pMHCs that are the physiologic ligands for any given pre-TCR (6, 9) and that some may possess as high or even higher affinity than that found for the cognate ligand of the respective TCRαβ. Because of the convoluted and transient procession from thymic progenitor to T cell (34), it may be difficult to define the “true” ligand for any pre-TCR. Although there is good evidence for self-pMHC promotion of pre-TCR-mediated proliferation and progression (6, 9), the existence of non-MHC ligands for pre-TCR (35, 36) has not yet been ruled out.

Meanwhile, as implied by single-molecule studies, the Vβ patch region actively participates in binding by replacing Vα (6). In support of this hypothesis, the molecular modeling using ambiguous NMR restraints (Fig. 1D) shows conformations that uniformly predict a more horizontal apposition of pre-TCR to pMHC. In the pre-TCR, the long axis of the β subunit is roughly parallel to the MHC groove, whereas in the TCR it is perpendicular. If this is the case, then this implies an initial contact orientation prior to the application of mechanical force that is rather tilted relative to the more upright TCR when bound to pMHC. This tilted interface would affect the load-sustaining ability and transmission of force through the pre-TCR with implications for the mechanisms of catch-bond formation and mechanical triggering. The sensitivity of pre-TCR-baring thymocytes to triggering may be potentiated by such a mechanism. It has been seen that the pTαβ is more compliant than the TCRαβ, undergoing a catch-bond requisite reversible transition more readily (6). The unpaired Vβ domain structure in the case of the pre-TCR relative to the paired VαVβ module in the TCRαβ can explain this compliance difference. As we have previously suggested (15), the location of glycans on murine pTα rules out some homodimerization models (20) of pre-TCR function. Although glycan sites do not appear to directly conflict with ligand binding measured here, the proximity of the β chain (Figs. 10, D and E, and S1) or the pTα subunit (Fig. S1) in these models to the conserved glycan sites on pMHC means that some guiding role for glycan modulation of early thymocyte progression directly through the impact of these interactions or indirectly via recruitment of other cell surface proteins (37) cannot be excluded. The initial pre-TCR structural binding model presented here in conjunction with single-molecule biophysical analysis performed previously (6) suggests that the biology of the pre-TCR and αβTCR will manifest important distinctions built around a common β chain module.

Experimental procedures

Protein production

Unlabeled and isotopically labeled N15β, N15β-c1, N30β, and N30β-c1 were produced as detailed (18). Briefly, β chains were expressed using Escherichia coli as inclusion body (ib) preparations. These ib preparations were washed thoroughly in 50 mM Tris-Cl, pH 8, 150 mM NaCl (TBS); dissolved in 6 M guanidine hydrochloride; refolded by dilution in 5.4 M guanidine hydrochloride, pH 8, 1 M Arg, 1 mM reduced glutathione, 0.1 mM oxidized glutathione; and subsequent dialysis in TBS. VSV8/Kb was prepared as detailed (9, 38). VSV8 was chemically synthesized (United Biosystems, Inc.), and Kb and βm were produced via E. coli as ib preparations. The ib preparations were washed in TBS and dissolved in 8 mM urea. VSV8, Kb, and βm were mixed in 20 mM Tris-Cl, pH 8, 8 mM urea buffer and dialyzed serially against 2, 1, 0.5, and 0 mM urea in 20 mM Tris-Cl, pH 8, for 2 h or overnight with a final dialysis against 0 mM urea, 20 mM Tris-Cl, pH 8, overnight. VSV8/Kb- was prepared identically to VSV8/Kb with the exception that 1 mM reduced glutathione and 0.1 mM oxidized glutathione were included prior
to dialysis-mediated refolding. Proteins were purified by successive rounds of size-exclusion chromatography.

**NMR**

Standard pulse sequences were utilized on Bruker 750-MHz and Varian 600-MHz spectrometers equipped with cryogenically cooled probes or a Bruker 500-MHz spectrometer with a room temperature probe. Concentrations of proteins were in the range of 50–500 µM, and NMR experiments were conducted at 298 K in PBS (50 mM NaPO₄, 150 mM NaCl, pH 7.0) unless otherwise indicated. CCSC was calculated using the formula $CCSC = \frac{5 \times \Delta \delta^{(1)}(H)}{\Delta \delta^{(13)}(N)}$. All spectra were processed with NMRPipe (39) and visualized using CARA (40).

**Cross-saturation transfer experiments**

Unlabeled 175 µM VSV8/Kb was added to uniformly $^{2}H/^{13}C/^{15}N$-labeled 350 µM N15β in PBS (20 mM NaPO₄, 50 mM NaCl, pH 7.0) in 80% ($\psi/\psi$) $^{2}H_{2}O/H_{2}O$. The protonated VSV8/Kb was saturated with an adiabatic WURST (wideband, uniform rate, and smooth truncation) pulse (25 ms) with an excitation bandwidth of 750 Hz. CST experiments were performed in an interleaved fashion with the on-resonance saturation pulse centered at 1 ppm (sat) and the off-resonance saturation pulse centered at −19 ppm (ref). The CST effect was calculated as a ratio of resonance peak intensities (sat/ref). Optimal saturation was found to occur at 3-s saturation time. The top 10th or 25th percentile was taken as the affected region.

**Backbone assignment**

TROSY versions of HNCA, HNCOCA, HNCO, HNCACO, and HNCACB (41) were used for backbone assignment of Kb and Kb-t. Assignments were completed using CARA (40) in conjunction with RASP (25), utilizing chemical shift predictions generated by SPARTEA + (42).

**Statistical analysis**

Statistical significance ($p$) was calculated by linear regression analysis using the R software package (43).

**Author contributions**—R. J. M. conducted experiments, analyzed results, and drafted the paper. R. J. M., H. A., E. L. R., M. J. L., and G. W. conceived the research, planned experiments, analyzed results, and wrote the paper.

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