A conserved hydrophobic patch on Vβ domains revealed by TCRβ chain crystal structures: implications for pre-TCR dimerization

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INTRODUCTION

The αβ T cell receptor (TCR) is a multimeric complex consisting of the antigen-binding αβ clonotypic heterodimer and six invariant CD3 subunit dimers (Sun et al., 2001 and references therein). Specific interaction between the TCR γδ heterodimer and an antigenic peptide bound to an MHC molecule (pMHC) triggers downstream signaling, resulting in T cell activation (Rudolph et al., 2006; Acuto et al., 2008; Smith-Garvin et al., 2009). TCR signaling is also critical for thymocyte differentiation (Ghendler et al., 1998; Mariathasan et al., 1999; Love et al., 2000; Haks et al., 2002). In early thymic development, the first major checkpoint is referred to as β-selection. During this process TCRβ chain gene rearrangement occurs in the absence of TCRα chain gene rearrangements (Von Boehmer et al., 2003). The expressed TCRβ chain associates with the invariant pre-Tα (pTα), which leads to the formation of pre-TCR on the cell surface. Signaling through the pre-TCR terminates TCRβ locus rearrangement, rescues cells from apoptosis and induces massive proliferation. This process eventually enables CD4−CD8− double-negative (DN) cells to differentiate into CD4+CD8+ double-positive (DP) cells, facilitating TCRα gene rearrangement and thus generating a large αβ TCR repertoire (Von Boehmer et al., 2003). DP thymocytes undergo positive selection and negative selection following self–MHC interaction with their αβ TCR. In the former case, DP cells mature into single positive (SP) thymocytes and may egress into the periphery (Starr et al., 2003; Juang et al., 2010). In the latter case, cell death occurs and those apoptotic thymocytes are purged from the T lineage repertoire (Gallegos and Bevan, 2006; Ashton-Rickardt, 2007; Griesemer et al., 2010).

The development of αβ lineage thymocytes and the antigen recognition of mature T cells are critically dependent on the TCRβ chain. Certain germline-encoded amino acids within the TCR Vβ domain promote “generic” recognition of MHC molecules during the thymic selection process (Scott-Browne et al., 2009). The germline-encoded TCR Vβ–MHC interactions have evolved to pre-determine, or at least strongly influence, the conserved diagonal TCR–MHC binding mode (Feng et al., 2007). The Vβ domain’s unique contribution has also been shown in providing a structural basis for immunodominance (Ishizuka et al., 2008; Wang et al., 2008). Immunodominant CTL responses to EBV infection incorporate a particular rearranged β chain generated during thymic development, favoring the survival, and expansion of T cells bearing it (Argaet et al., 1994; Gras et al., 2009). Another example involves the conservation of one Vβ domain in the context...
of highly variable Vα usage, as found in B6 mice responding to the immunodominant Dp12-39–restricted NP366-374 influenza A peptide following PR8 viral infection (Zhong and Reinherz, 2004; Zhong et al., 2007). The above observations raise the interesting possibility as to whether the Vβ recognition itself during β-selection can mediate a physiologic binding event in a manner analogous to certain VH domains in antibody molecules (Ward et al., 1989).

The alternative view regarding β-selection is that it is an autonomous signaling process driven by the oligomerization of pre-TCR’s extracellular domain (Yamasaki et al., 2006; Yamasaki and Saito, 2007). Very recently, Pang et al. (2010) determined the crystal structure of a pre-TCR and, following crystal packing analysis, proposed a model for autonomous dimerization of pre-TCR. In this model, a pTα domain’s ABED face contacts the β chain’s CB domain to form a pTα/Cβ module, similar to the Ca/Cβ module seen in the αβTCR heterodimer. In addition, the other face of the pTα domain (the CFG face) interacts with a Vβ domain from the neighboring molecule in the crystal lattice, resulting in a dimeric pre-TCR. In this way, the “head-to-tail” pre-TCR dimer referred to by the authors, should it adopt this topology pre-TCR. In this way, the “head-to-tail” pre-TCR dimer referred to by the authors, should it adopt this topology in vivo, would sit almost parallel to the cellular membrane surface such that the Vβ domain becomes virtually unable to access any potential ligand (Pang et al., 2010). This contrasts with the upright model of the mature αβTCR complex, which is arrayed on the T cell surface with six CDR loops pointing upward to facilitate immune recognition (Sun et al., 2004).

We report here crystal structures of two β subunit ectodomains, N15β (Vβ5.2Dβ2Jβ2.6CB2), and N30β (Vβ13Dβ1Jβ1.1CB2), derived from N15 and N30 αβTCRs (Chang et al., 1997; Goyarts et al., 1998). We have compared these structures with the previously published structure of a TCR β subunit (Vβ8.2Jβ2.1CB1; Bentley et al., 1995) and several others in the protein data bank. In the structures surveyed, a few conserved residues were identified on the CFG face of Vβ Ig-like domain. In the biological heterodimeric αβTCR these residues are located at the Vα/Vβ interface. In the Vβ structures studied, the same residues are shielded from exposure by crystal packing in one of several possible ways. Based on these observations, we have proposed an alternative pre-TCR dimer model that is more similar to the heterodimeric αβTCR, and discussed its biological significance.

**MATERIALS AND METHODS**

**CLONING, EXPRESSION AND PURIFICATION OF N15β AND N30β**

Gene encoding the extracellular domain of N15β and N30β subunits was cloned into the expression vector pET-11d (Novagen). The DNA sequence of N15β and N30β constructs were terminated immediately after the cysteine of the respective TCRβ subunit involved in interchain disulfide formation with the cysteine itself mutated to serine. The recombinant plasmid was then transformed into *Escherichia coli* strain BL21 (DE3) followed by the induction of N15β and N30β expression using 1 mM isopropyl-β-D-thiogalactopyranoside. Inclusion bodies were isolated and solubilized in 6 M guanidine–HCl. The refolding of protein was achieved by further dilution of the inclusion bodies in a cold buffer containing 100 mM Tris–HCl (pH 8.0), 5.4 M guanidine–HCl, 1 M L-arginine–HCl, 1 mM reduced glutathione, and 0.1 mM oxidized glutathione, followed by dialysis against 20 mM Tris–HCl (pH 8.0), 10 mM NaCl at 4°C for 12 h. Insoluble particles were removed by centrifugation at high speed. The refolding mixture was concentrated prior to sequential purification of refolded proteins on a Sephacryl S-300 column and a Superdex 75 column (GE Healthcare).

**CRYSTALLIZATION AND STRUCTURE DETERMINATION**

Crystals were grown at room temperature by the hanging-drop vapor diffusion method. Equal volumes of protein solution (15 mg/ml in 20 mM HEPES pH 7.5, 100 mM NaCl) and reservoir solution (100 mM HEPES pH 7.0, 20% PEG3350 for N15β, or 100 mM Tris–HCl pH 8.5, 20% PEG4000 for N30β) were mixed and crystals appeared in 1–2 days. 15% glycerol was added in reservoir buffer as a cryoprotectant solution to soak crystals prior to freezing. Diffraction data were collected at beamline 24-ID-E at the Argonne National Laboratories.

The structures were determined by molecular replacement method. TCRβ structures taken from heterodimeric αβTCR models with PDB accession code 1NFD and 1LP9 were used as the templates for searching N15β and N30β structures, respectively. The solution of N30β structure could only be obtained when Vβ and Cβ domains were used as the templates separately. The models were refined using refmac5 (Vagin et al., 2004) or phenix (Adams et al., 2010), viewed and adjusted by Coot (Emsley and Cowtan, 2004), and validated by MolProbity (Davis et al., 2007). The data collection and refinement statistics are listed in Table 1. The coordinates of the N15β and N30β structures have been deposited in the Protein Data Bank with the code 3Q5Y and 3Q5T, respectively.

**RESULTS**

**STRUCTURES OF N15β AND N30β**

The 240-residue N15 and N30 β subunits were expressed in *E. coli* and crystallized readily. Their structures were determined using molecular replacement, with R-factor of 0.178 and Rfree of 0.218 for N15β to 1.9 Å, and 0.235 and 0.289 for N30β to 2.0 Å, respectively (see Table 1 for statistics). As shown in Figure 1, each of the models encompasses two Ig-like domains, Vβ and Cβ.

The superposition of the isolated N15β protein onto the β subunit of our previously published N15 αβTCR heterodimer (PDB code 1NFD) provides one of the first examples demonstrating that there are no significant conformational changes upon TCRβ pairing with TCRα to form an intact αβTCR. If the superposition is based on the Vβ domain, then the two corresponding Cβ domains only have a few degree rigid-body rotation relative to each other (Figure 2 and vide infra). We previously reported that the quaternary arrangement of Vβ–Cβ varies among different TCR molecules (Wang et al., 1998). Our current results suggest that for each individual TCRβ, the quaternary arrangement of Vβ–Cβ domains might be intrinsic, determining at least in part to which α subunit it can pair. A general survey indicates that upon TCR ligation to pMHC, the major conformational changes are limited to the CDR loops of each subunit (Rudolph et al., 2006). The rigidity of the framework module of TCR molecules seems to be essential for signaling following pMHC bindings (Kim et al., 2009).
Table 1 | Data collection and refinement statistics.

|                      | N15β       | N30β       |
|----------------------|------------|------------|
| **DATA COLLECTION**  |            |            |
| Space group          | P212121    | C2221      |
| Unit cell parameters (Å, °) | a = 78.2, b = 129.6, c = 129.7, α = β = γ = 90 | a = 53.3, b = 69.8, c = 129.4, α = β = γ = 90 |
| Resolution           | 20–1.9 (1.93–1.90) | 20–2.0 (2.05–2.0) |
| Completeness (%)     | 99.4 (97.0) | 99.9 (100) |
| Rsym (%)             | 10.3 (60.5) | 6.0 (61.6) |
| I/σ(I)               | 20.6 (2.6)  | 19.2 (2.5) |
| Redundancy           | 8.3 (3.8)   | 6.5 (6.5)  |
| Unique reflections   | 97,771      | 16,199     |
| **REFINEMENT**       |            |            |
| Rwork (%)            | 178        | 23.5       |
| Rfree (%)            | 21.8       | 28.9       |
| Rmsd bonds (Å)       | 0.008      | 0.007      |
| Rmsd angles (°)      | 1.122      | 1.097      |
| Ramachandran (%)     | 96.9, 3.1, 0 | 92.8, 5.9, 1.3 |

Values in parentheses refer to the highest resolution shell.

a $Rsym = \sum |I| - |<\|\sum |F||$.

b $Rfree = \sum \frac{|F_o| - |<\|\sum |F||}{\sum |F||}$.

c $R_{free}$ is the $R$-factor for a selected subset (5%) of the reflections which are not included in prior refinement calculations.

THE Vβ/Vβ HOMODIMER OF N15β IS SIMILAR TO Vα/Vβ HETERODIMER OF N15 αβ TCR

A particularly intriguing feature of the N15β crystal structure is that two N15β subunits pack to form a Vβ/Vβ homodimer in a fashion very similar to that of the Vα/Vβ module in the αβ TCR heterodimer. In each asymmetric unit of the N15β crystal there are four N15β molecules that pair to form two homodimers, using their outer faces of CFG β sheet for contacts. Figure 2 shows the overlay of a crystallographic homodimer of N15β onto the N15 αβ TCR heterodimer. The superposition is based on the Vβ domains (in red and green color, respectively, for N15β and N15 αβ TCR structures). In this figure, the small movement of the Cβ domain in the TCRβ relative to the αβ TCR can be clearly seen. The respective pairing domains, Vβ (in pink color, from the homodimer) and Vα (in lime color, from the heterodimer) are similarly oriented, differing by only 11° from each other (Figure 2). In both the ββ homodimer and αβ heterodimer, the two pairing V-type Ig-like domains combine in an aligned arrangement with their corresponding β strands' directions at an acute angle to one another.
Figure 3 depicts the key interface residues in the N15 Vβ/Vβ structure (Figure 3A) and those in the Va/Vβ interface in the N15 αβ TCR structure (Figure 3B). Both interfaces involve Tyr35, Gln37, Phe91, and Phe108, as well as Leu43 in Vβ or Pro43 in Va. The three aromatic residues along with Leu43 (or Pro43) form a hydrophobic center at the CFG interface. These residues are conserved in the variable domains of the TCR (Chothia et al., 1988; as shown for selected TCR variable domains in Figure 3C) and antibody molecules (Chothia et al., 1998). Figure 3D is an electrostatic surface representation of Vβ domain’s CFG face and the opposite ABED face for comparison. Whereas the exposed ABED face is fully covered by positive and negative charges, by contrast, a concave hydrophobic patch at the center of CFG outer face is readily discernible. Any exposed hydrophobic patch this size may not be stable, necessitating shielding from the solvent environment. On the other hand, the sole hydrophilic residue on the CFG face within this patch, the Gln37, plays a distinct role. In both the N15β and N15 αβ TCR structures, this Gln37 on the C strand forms two hydrogen-bonds with its dimeric counterpart near the bottom of the interface between two V-domains (Figures 3A,B). This Gln has been reported to be invariant in variable domains of TCRs and antibodies (Chothia et al., 1988, 1998). Presumably the hydrogen-bonds offered by Gln37 provide a correct register for the two variable domains to dock. In Figure 3B the Va domain’s Tyr35 forms an additional direct hydrogen bond with the main chain carbonyl oxygen of Gln106 of the Vβ domain near the top of the Va/Vβ interface. This is also a conserved feature of the αβ TCR Va/Vβ interface in general. In the Vβ/Vβ interface of the N15β structure, the Tyr35’s side chain appears slightly far from Gln106’s carbonyl oxygen (around 4.5 Å) for a hydrogen bond (Figure 3A). It is interesting to note that there is a HEPES [4-(2-hydroxyethyl)-1-piperazinethanesulfonic acid] solvent molecule intercalated into the Vβ/Vβ interface adjacent to Tyr35 for each of two N15β homodimers in the crystal structure, which prevents Tyr35 from forming a hydrogen bond to carbonyl oxygen of Gln106 (Figure A1 in Appendix). Notwithstanding, the N15β Vβ/Vβ homodimer is very similar overall to the N15 αβ TCR Va/Vβ heterodimer.

There are structural differences, however, between the Vβ/Vβ homodimers and Va/Vβ heterodimers. For example, using the buried area option in the CNS program package (Brunger et al., 1998), we calculate that the surface buried area for the N15 Vβ/Vβ homodimer is 1,291 Å², significantly smaller than 1,523 Å² for the N15 Va/Vβ heterodimer. Moreover, the complementary fitness index, or Sc value (Lawrence and Colman, 1993) of the Vβ/Vβ homodimer and Va/Vβ heterodimer is 0.469 and 0.545, respectively, indicating a poorer fit at the homodimer interface relative to that of the heterodimer. This differential result reflects the self-association affinity difference between these dimers. Notably, the asymmetric TCR molecule has a much sharper bending angle between Vβ and Cβ domains compared to that between Va and Ca domains (Wang et al., 1998). Thus, when two N15β molecules’ Vβ domains form a Vβ/Vβ homodimer, the two corresponding Cβ domains do not meet to form a C module like that of Ca/Cβ in the αβ TCR molecule, as evident in Figure 2. These data suggest that it is possible to form a homodimer between two TCR β subunits through their Vβ domains, but that the homodimer so formed is substantially less stable than the αβ TCR heterodimer.

The observed Vβ/Vβ homodimer in the crystal lattice may well be of biological significance. For instance, using RAG-2−/− deficient mice for functional studies, it was found that the introduction of a TCRβ transgene, but not a TCRα transgene restores T cell development, allowing thymocytes to transit from the DN to the DP stage. These DP cells were observed to express monomeric TCRβ chains in association with CD3γδε but not ζζ, as well as small amounts of β homodimers (Shinkai et al., 1993). We shall come back to these points later.

The shielding of the exposed hydrophobic CFG face of the Vβ domain is a common feature in other crystal structures that contain TCRβ

There is only one molecule in the asymmetric unit of the N30β crystals. The symmetry-related molecules pack very differently from those of the N15β crystal, with one molecule’s Cβ domain packing onto the other molecule’s Vβ domain as shown in Figure 4A. Despite the apparent difference, the hydrophobic surface of the N30β Vβ domain (the CFG face) is also shielded from...
Moreover, this protein formed homodimers in solution as well, blinding the Vβ domain's ABED face. Note that F128 of the Cα subunit pokes into the concave hydrophobic patch of Vβ's CFG face, (B) The orthogonal packing between two Vβ domains' CFG faces frequently observed in crystal structures containing TCRβ. In the figure is the first solved TCRβ structure (PDB code 1BEC).

The observations on N15β and N30β structures led us to carry out an extensive survey in the protein data bank to investigate whether, in general, a TCR variable domain's hydrophobic CFG face shown in Figure 3D is shielded from solvent exposure by crystal packing in various individual (i.e., non-heterodimeric) TCR subunit structures. In retrospect, the first glimpse at a TCR structure was the TCR 2C and A6 (Li et al., 1997). Many of these TCR β subunit structures are in complex with ligands such as superantigens. In the structure of 1BEC, the hydrophobic surface of the Vβ domain's CFG face was indeed shielded by crystal packing (Figure 4B). However, in that structure the two contacting Vβ domains' β strands are not aligned in the same manner as seen in the N15β and N15 αβ TCR structures. Rather, in the IBEc structure, molecules are in an orthogonal packing pattern, similar to the CD2/CD58 adhesion molecule pair (Wang et al., 1999). Interestingly, the majority of those TCRβ structures, be they in isolation or in complex with ligands, have an orthogonal packing in crystals to shield their hydrophobic CFG faces from solvent exposure.

It is particularly noteworthy that an aligned homodimer was observed in the first TCR Vα-Vβ structure from the αβ TCR 1934.4 specific for the N-terminal nonapeptide of myelin basic protein bound to the MHC class II molecule I-Au (Fields et al., 1995). The same research group later published a mutant 1934.4Vαmut structure and observed a Vα/Vα homodimer resembling the Vα/Vβ heterodimer of αβ TCR 2C and A6 (Li et al., 1997). Moreover, this protein formed homodimers in solution as well (Fields et al., 1994).

STABLE, SINGLE-SUBUNIT MOLECULES DO NOT HAVE AN EXPOSED HYDROPHOBIC PATCH

We further sought to carry out comparative investigation of the two co-receptors: CD4 and CD8. CD8 exists either as a CD8α homodimer or a CD8αβ heterodimer on the T cell surface, whereas CD4 is a monomeric molecule. The N-terminal domains of these two co-receptors are both V-type Ig-like domain. Figures 5A,B are electrostatic surface representations of the N-terminal V-domain CFG face of CD8β and CD4, respectively. The contrast is obvious. Whereas the single-subunit CD4 does not have a hydrophobic patch at the center of the CFG face of its N-terminal domain, CD8β does, analogous to that observed in the TCRβ subunit. This CD8 feature is linked to the fact that CD8β uses its single Ig-like domain to pair with CD8α in order to form a functional heterodimer. This further provides evidence that any membrane distal V-type Ig-like domain in a transmembrane protein is likely to form a dimer to shield its hydrophobic patch, if it exists, at the center of the CFG face from exposure. Moreover, in the case of the CD8α homodimer the conserved Gln41 plays an identical role as Gln37 for the TCR to form two hydrogen-bonds creating the correct domain–domain interface register (PDB code 1BQH; Kern et al., 1998).

It is interesting that camelds possess a functional class of antibodies devoid of light chain (Desmyter et al., 1996). Figure 5C demonstrates that its VH domain does have the conserved aromatic residues Phe37, Tyr95, and Tyr120 (equivalent positions Tyr35, Phe91, and Phe108 in TCRβ, respectively) at the otherwise-exposed hydrophobic patch. However, this cameld molecule has an extraordinarily long FG loop curving into an extra helix that shields off this hydrophobic patch. This structural feature explains why this single heavy-chain antibody is functionally stable in the absence of a V-domain dimeric structure.

DISCUSSION

The above observation has led to the following key conclusion: for structural stability, a TCR Vβ domain must shield the conserved hydrophobic patch on its CFG face from exposure to solvent. The patterns of shielding can vary, either through Vα/Vβ interaction in
the case of the αβTCR heterodimer, Vβ/Vβ aligned homodimeric interaction in the N15β structure, Vβ/Vβ orthogonal pairing in the structure of 1BEC (Bentley et al., 1995), or anti-parallel Vβ/Vβ interaction in the single α domain structure 2APB (Cho et al., 2005; not shown here) and in the Vβ/pTα interaction observed in the pre-TCR structure (PDB code: 3OF6; Pang et al., 2010), or Vβ/Cβ interaction in the N30β structure, etc. This set of variations may arise in the absence of a receptor’s membrane anchoring force where the domain orientation is under less constraint. Moreover, receptors’ domain/domain interactions are relatively weak (Wang, 2002; van der Merwe and Davis, 2003), and the non-specific hydrophobic interface is therefore not uniquely defined. In this regard, it is worth re-emphasizing the key role that the invariant Gln37 plays, facilitating the orientation register of Vβ with respect to Vα in the αβTCR heterodimer by providing two specific hydrogen-bonds, as shown in Figure 3B.

Based on the above analysis, we propose an alternative dimeric pre-TCR model (Figure 6A), distinct from the “head-to-tail” “flat” model suggested by Pang et al. (2010; Figure 6B). Our assumption is that the N15β homodimer might represent a physiological dimer existing at an early DN stage of thymocyte development. We superimposed the pre-TCR structure’s Vβ and Cβ domains (PDB code: 3OF6) onto the N15β’s Vβ and Cβ domains separately, for each of the protomers within the N15β homodimer. This then places the pTα domain along with the Cβ domain as a module without significant steric collision, creating a distinct dimeric pre-TCR as shown in Figure 6A. Given that the Vβ–Cβ bending angle is actually slightly different between the same β subunit in the mature TCR LC13 (PDB code: 1KGC) and in the pre-TCR (PDB code: 3OF6), it is not surprising that this bending angle differs by a few degrees between N15β and LC13β in the pre-TCR.

The dimeric model we have proposed here has several advantageous features. Structurally, as discussed above, the two hydrogen-bonds provided by the invariant Gln37 remain intact (Figure 3A). In this sense, the dimeric pre-TCR appears more similar to the mature αβTCR heterodimer. These hydrogen-bonds are absent in the “flat” pre-TCR model proposed by Pang et al. (2010). A more interesting structural feature is that the mouse pTα has two potential glycosylation sites with the NXS sequon, one being Asn51-Gly-Ser at the tip of CC’ loop and the other Asn101-Arg-Ser at the middle of G strand (Figure 6A, left panel). In both cases, the potential glycan adduct would point toward the cellular membrane, perhaps helping the receptor to orient properly on the membrane (Casasnovas et al., 1997). In the human pre-TCR structure (Pang et al., 2010), a clear sugar moiety is seen at Asn51, which is exposed in the Pang’s model as shown in Figure 6B. Since the human pre-TCR has Ser101 in lieu of murine Asn101, there is no glycosylation site at the middle of the human G strand. Were the mouse pre-TCR to assume the same “head-to-tail” dimeric structure as the human pre-TCR, then the glycan attached to Asn101 (in the place of human Ser101) would seriously collide with the Vβ domain from its dimeric partner. This can be seen in Figure 6B where the Ser101 (in blue sphere) in human pre-TCR is buried between pTα (in red) and Vβ (in green).

Functionally, our model would allow the CD3 subunits to be incorporated into the pre-TCR dimer complex in a fashion more analogous to that in the mature αβTCR complex (Sun et al., 2004; Kuhns et al., 2010). By contrast, it is difficult to envision where the CD3 subunits can fit in the “flat” dimeric pre-TCR model. Our model also forces the Vβ domains’ CDR loops to project up and out from the membrane in a mode similar to that seen in the mature αβTCR. Subsequently, with further differentiation, the TCRα subunit might easily replace pTα. Finally, our postulated orientation of the β chain would not exclude the possibility that during β selection Vβ recognition in the form of a dimeric pre-TCR could mediate a physiologic binding event with potential ligands in the thymic development process.

In their pre-TCR crystal structure, Pang et al. (2010) actually observed two potential pre-TCR dimers generated by crystal packing. In addition to the flat “head-to-tail” dimer described above, a side-by-side dimer was also discussed and presented in their Figure A2 in Appendix. This dimer is mediated by Cβ/Cβ interaction and appears to stand on the membrane in a vertical orientation. However, this dimeric model was not regarded by the
authors as favorable because it was presumed to sterically block CD3 interaction. By comparison, the dimeric pre-TCR model we have proposed, being more homologous to the mature αβ TCR heterodimer, can permit CD3 complex formation as discussed above. This is readily seen from Figure 6A right panel, in which the space beneath and/or adjacent to the protruding FG loop of Cβ could accommodate juxtaposition of the CD3 subunits (Sun et al., 2004; Kuhns et al., 2010). Moreover, we have also noticed that the charged residues Asp22 and Lys24 (or Arg24 in mice) at the tip of AB loop of pTα, which were reportedly involved in pre-TCR oligomerization (Yamasaki et al., 2006), are located below the FG loop where the CD3 subunits may bind (Figure 6A). Mutations of Asp22 and Lys24 (or Arg24 in mice) may alter pre-TCR signaling and/or internalization through modulating effects of CD3-related activation. That being said, whether the CD3 heterodimers are oriented near the Cβ FG loop (Figure 6A) exactly as in the mature TCR complex, or alternatively, that juxtaposition is modified remains to be determined.

In the report by Pang et al. (2010), the authors have provided some experimental evidences, including site-directed mutagenesis, supporting their pre-TCR dimeric model. They demonstrated that three mutations, namely Y35A, F108A, and W46R, affect pre-TCR cell surface expression. It is particularly intriguing that mutation Y35A in the TCRβ completely abrogate cell surface expression of the pre-TCR, but did not affect LC13 αβ TCR cell surface expression (Pang et al., 2010). As discussed above, Tyr35 may be involved in hydrogen-bonding to main chain atom of Gln106 to facilitate dimer formation for the ββ homodimer in the pre-TCR as well as the αβ TCR heterodimer. Given that in the case of αβ TCR, there are many more hydrophobic interactions between Cα and Cβ domains (Wang et al., 1998), the Y35A mutation may be less detrimental than in the pre-TCR where only Tyr35 and Gln37 determine proper homodimeric orientation. On the other hand, Phe108β is within the critical hydrophobic patch we discussed extensively above. It is not surprising to see the lower level expression for the mutant F108βA on the cell surface. The effect of the W46R mutation is more complicated to interpret. This Trp46 is located at the center of CFG face of pTα domain. In the “head-to-tail” model, this face involves contacts with Vβ domain of the dimeric partner. The W46R mutation would disrupt the dimer. In our model, the pTα’s CFG face is exposed. However, compared with the pTα’s ABED face, the structure of this CFG face is much less regular. The residue W46R’s side chain indole ring lies flat onto the pTα domain, and has extensive hydrophobic interaction with nearby Pro42 on the same C strand and His92 from the F strand (Figure A2 in Appendix). Therefore, this Trp46 may serve to stabilize the pTα domain and a mutation of W46R might be detrimental, impacting surface expression.

Whether a pre-TCR monomer–dimer equilibrium might exist on the DN thymocyte surface with one or both structures competent for ligand-binding requires further analysis. Clearly, there is much to be resolved about the structure and function of the pre-TCR through future experimental endeavors.

CONCLUSION

In this work, we report crystal structures of two TCR β subunits. In conjunction with other similar structures deposited in PDB, we have identified a conserved hydrophobic patch on the CFG face of Vβ domain that is invariably shielded from solvent exposure. In one of our TCR β structures, the N15B, a homodimer is formed between the CFG faces of the two Vβ domains. By incorporating a newly published pre-TCR structure, we propose a pre-TCR dimer model based on this N15B homodimer. We discuss the structural and functional advantages of this model relative to alternative models.

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REFERENCES

Acuto, O., Di Bartolo, V., and Michel, F. (2008). Tailoring T-cell receptor signals by proximal negative feedback mechanisms. Nat. Rev. Immunol. 8, 699–712.

Adams, P. D., Afonine, P. V., Bunkoczi, G., Chen, V. B., Davis, I. W., Echols, N., Headle, J. I., Hung, L. W., Kapral, G. J., Grosse-Kunstleve, R. W., McCoy, A. J., Moriarty, N. W., Oeffner, R., Read, R. J., Richardson, D. C., Richardson, J. S., Terwilliger, T. C., and Zwart, P. H. (2010). PHENIX: a comprehensive python-based system for macromolecular structure solution. Acta Crystallogr. D Biol. Crystallogr. 66, 213–221.

Argaet, V. P., Schmidt, C. W., Burrows, S. R., Silins, S. L., Kurilla, M. G., Doolan, D. L., Suhbier, A., Moss, D. J., Kief, E., Scalley, T. B., and Misko, I. S. (1994). Dominant selection of an invariant T cell antigen receptor in response to persistent infection by Epstein-Barr virus. J. Exp. Med. 180, 2335–2340.

Ashton-Rickardt, P. G. (2007). Studying T-cell repertoire selection using fetal thymus organ culture. Methods Mol. Biol. 380, 171–184.

Bentley, G. A., Boulot, G., Karjalainen, K., and Mariuzza, R. A. (1995). Crystal structure of the β chain of a T cell antigen receptor. Science 267, 1984–1987.

Brunger, A., Adams, P., Clore, G., DeLano, W., Gros, P., Grosse-Kunstleve, R., Jiang, J.-S., Kuszewski, J., Nilges, N., Pannu, N., Read, R. J., Rice, L. M., Simonson, T., and Warren, G. L. (1998). Crystallography and NMR system (CNS): a new software system for macromolecular structure determination. Acta Crystallogr. D Biol. Crystallogr. 54, 905–921.

Casasnovas, J. M., Springer, T. A., Liu, J. H., Harrison, S. C., and Wang, J. H. (1997). Crystal structure of ICAM-2 reveals a distinctive integrin recognition surface. Nature 387, 312–315.

Chang, H. C., Smolyar, A., Spoerl, R., Witte, T., Yao, Y., Goyarts, E. C., Nathenson, S. G., and Reinherz, E. L. (1997). Topology of T cell receptor-peptide:class I MHC interaction defined by charge reversal complementation and functional analysis. J. Mol. Biol. 271, 278–293.

Cho, S., Swaminathan, C. P., Yang, J., Keric, M. C., Guan, K., Kieke, M. C., Kranz, D. M., Mariuzza, R. A., and Sundberg, E. J. (2005). Structural basis of affinity maturation and intramolecular cooperativity in a protein–protein interaction. Structure 13, 1775–1787.

Chothia, C., Boswell, D. R., and Lesk, A. M. (1988). The outline structure of the T cell αβ receptor. EMBO J. 7, 3745–3755.

Chothia, C., Gelfand, I., and Kister, A. (1998). Structural determinants in the sequences of immunoglobulin variable domain. J. Mol. Biol. 278, 457–479.

Davis, I. W., Leaver-Fay, A., Chen, V. B., Block, J. N., Kapral, G. J., Wang, X., Murray, L. W., Arendall, W. B., III, Snievey, J., Richardson, J. S., and Richardson, D. C. (2007). MolProbity: all-atom contacts and structure validation for proteins and nucleic acids. Nucleic Acids Res. 35, W375–W383.

Desmyter, A., Transue, T. R., Ghabrouri, M. A., Thi, M. H., Poortmans, E., Hamers, R., Muyldermans, S., and Wysn, L. (1996). Crystal structure of a camel single-domain VH antibody fragment in complex with lysosome. Nat. Struct. Biol. 3, 803–811.
Emsley, P., and Cowtan, K. (2004). Coot: model-building tools for molecular graphics. Acta Crystallogr. D Biol. Crystallogr. 60, 2126–2132.

Feng, D., Bond, C. J., Ely, L. K., Maynard, J., and Garcia, K. C. (2007). Structural evidence for a germline-encoded T cell receptor-major histocompatibility complex interaction “codon.” Nat. Immunol. 8, 975–983.

Fields, B. A., Ober, B., Malchiodi, E. L., Lebedeva, M. I., Braden, B. C., Ysern, X., Kim, J. K., Siao, X., Ward, E. S., and Mariuzza, R. A. (1995). Crystal structure of the V\(\beta\) domain of a T cell antigen receptor. Science 270, 1821–1824.

Fields, B. A., Ysern, X., Poljak, R. J., Shao, X., Ward, E. S., and Mariuzza, R. A. (1994). Crystallization and preliminary diffraction study of a bacterially produced T cell antigen receptor V\(\beta\) domain. J. Mol. Biol. 239, 339–341.

Galgello, A. M., and Bevan, M. J. (2006). Central tolerance: good but imperfect. Immunol. Rev. 209, 290–296.

Ghendler, Y., Teng, M. K., Liu, J. H., Witte, T., Liu, J., Kim, K. S., Kern, P., Chang, H. C., Wang, J. H., and Reinherz, E. L. (1998). Differential thymic selection outcomes stimulated by focal structural alteration in peptide/major histocompatibility complex ligands. Proc. Natl. Acad. Sci. U.S.A. 95, 10061–10066.

Goyarts, E. C., Witte, T., Liu, J., Kim, K. S., Kern, P., Chang, H. C., Wang, J. H., and Reinherz, E. L. (1998). Functional thymic selection outcomes stimulated by focal structural alteration in peptide/major histocompatibility complex ligands. Proc. Natl. Acad. Sci. U.S.A. 101, 5094–5099.

Lawrence, M. C., and Colman, P. M. (1993). Shape complementarity at protein/protein interfaces. J. Mol. Biol. 234, 946–950.

Li, H., Lebedeva, M. I., Ward, E. S., and Mariuzza, R. A. (1997). Dual conformations of a T cell receptor V\(\beta\) homodimer: implications for variability in V\(\beta\) V\(\beta\) domain association. J. Mol. Biol. 269, 385–394.

Love, P. E., Lee, J., and Shores, E. W. (2000). Critical relationship between TCR signaling potential and TCR affinity for thymocyte selection. J. Immunol. 165, 3080–3087.

Marianthasan, S., Jones, R.G., and Ohashi, P. S. (1999). Signals involved in thymocyte positive and negative selection. Semin. Immunol. 11, 263–272.

Pang, S. S., Berry, R., Chen, Z., Kjer-Nielsen, L., Perugini, M. A., King, G. E., Wang, C., Chen, S. H., LaGrutta, N. L., Williams, N. K., Bedoe, T., Tiganis, T., Cowieson, N. P., Godfrey, D. I., Purcell, A. W., Wilce, M. C., McCluskey, J., and Rossjohn, J. (2009). The shaping of T cell receptor recognition by self-tolerance. Immunity 30, 193–203.

Griesemer, A. D., Sorensen, E. C., and Hardy, M. A. (2010). The role of the thymus in tolerance. Transplantation 90, 465–474.

Haks, M. C., Pepin, E., van den Brakel, J. H. N., Smeele, S. A. A., Belkowski, S. M., Kesels, H. W. H. G., Krimpfort, P., and Kruisbeek, A. M. (2002). Contribution of the T cell receptor-associated CD3\(\gamma\)-TAM to thymocyte selection. J. Exp. Med. 196, 1–13.

Ishizuka, J., Stuart-Jones, G., van der Merwe, A., Bell, J., McMichael, A., and Jones, E. (2008). The structural dynamics and energetics of an immunodominant T-cell receptor are programmed by its V\(\beta\) domain. Immunol. Immunopathol. 28, 171–182.

Jiang, J., Ebert, P. F., Feng, D., Garcia, K. C., Krogsgaard, M., and Davis, M. M. (2010). Peptide-MHC heterodimers show that thymic positive selection requires a more restricted set of self-peptides than negative selection. J. Exp. Med. 207, 1223–1234.

Kern, P. S., Teng, M.-K., Smolyar, A., Liu, J.-H., Liu, J., Hussey, R. E., Spoerl, R., Chang, H.-C., Reinherz, E. L., and Wang, J.-H. (1998). Structural basis of CD8 co-receptor function revealed by crystallographic analysis of a murine CD8\(\alpha\)\(\beta\) ectodomain fragment in complex with H-2K\(\beta\). Immunology 99, 519–530.

Kim, S. T., Takeuchi, K., Sun, Z. Y., Tousma, M., Castro, C. E., Fahmy, A., Lam, J. M., Yang, G., and Reinherz, E. L. (2009). The \(\alpha\)\(\beta\) T cell receptor is an anisotropic mechanosensor. J. Biol. Chem. 284, 31028–31037.

Kuhns, M. S., Girvin, A. T., Klein, L. O., Chen, R., Jensen, K. D., Newell, E. W., Huppa, J. B., Lillemiere, B. E., Fluse, M., Chien, Y. H., Garcia, K. C., and Davis, M. M. (2010). Evidence for a functional sidedness for the \(\alpha\)\(\beta\) TCR. Proc. Natl. Acad. Sci. U.S.A. 107, 5094–5099.

Lawrence, M. C., and Colman, P. M. (1993). Shape complementarity at protein/protein interfaces. J. Mol. Biol. 234, 946–950.

Love, P. E., Lee, J., and Shores, E. W. (2000). Critical relationship between TCR signaling potential and TCR affinity for thymocyte selection. J. Immunol. 165, 3080–3087.

Mariathasan, S., Jones, R.G., and Ohashi, P. S. (1999). Signals involved in thymocyte positive and negative selection. Semin. Immunol. 11, 263–272.

Pang, S. S., Berry, R., Chen, Z., Kjer-Nielsen, L., Perugini, M. A., King, G. E., Wang, C., Chen, S. H., LaGrutta, N. L., Williams, N. K., Bedoe, T., Tiganis, T., Cowieson, N. P., Godfrey, D. I., Purcell, A. W., Wilce, M. C., McCluskey, J., and Rossjohn, J. (2009). The shaping of T cell receptor recognition by self-tolerance. Immunity 30, 193–203.
**APPENDIX**

**FIGURE A1 | Solvent molecule HEPES intercalates in the Vβ/Vβ interface.** Only two Vβ domains are shown. The side chain of Tyr35 and the main chain of Gln106 as well as the HEPES molecule are in stick model. The intercalated HEPES prevents the Tyr35 from forming a hydrogen bond to the main chain of Gln106.

**FIGURE A2 | The CFG face of the pre-α domain.** The view looks down the CFG face. Trp46 is located at the center of the face. Compared with the ABE face, this face is much less regular and with shorter β strands. The side chain indole ring of Trp46 lie flat on the domain, and has hydrophobic contacts with Pro42 and His92, which may help stabilize the domain.