Increased peptide contacts govern high affinity binding of a modified TCR whilst maintaining a native pMHC docking mode

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INTRODUCTION

CD8+ αβ T cells recognize mainly intracellularly expressed antigens through an interaction mediated by the cell surface expressed T cell receptor (TCR). Intracellular proteins, processed by the proteasome into short peptides (generally 8–13 amino acids in length), are presented to CD8+ T cells on the surface of almost all nucleated human cells by class I major histocompatibility complex proteins (pMHC). TCR recognition of pMHC initiates CD8+ T cell activation and the adaptive immune response. The ability of CD8+ T cells to scrutinize the intracellular environment provides an important mechanism to target aberrant disease epitopes that would be otherwise hidden from the immune system. Thus, CD8+ T cells play an important role during viral infections (Miles et al., 2010), which typically elicit strong CD8+ T cell responses, many of which have been well-characterized, including those to HTLV-1 (Bieganowska et al., 1999; Vine et al., 2004), although some viruses can escape CD8+ T cell mediated clearance (Kleinerman and Zinkernagel, 1998; Overbaugh and Bangham, 2001). Cancers too, as a result of their malignant transformation, have altered protein expression causing the presentation of tumor-associated peptide antigens (TAPAs) (Renkvist et al., 2001; Cole et al., 2009). However, although these TAPAs can elicit a host CD8+ T cell response, this is often insufficient to cause tumor rejection (Blohm et al., 2002; Parkhurst et al., 2004). CD8+ T cell responses are also integral to the initiation and progression of many autoimmune diseases, possibly through the unwanted recognition of self-peptide antigens (Bulek et al., 2012). The TCR/pMHC interaction, which governs CD8+ T cell responses, is therefore an attractive therapeutic target in many varied diseases, particularly...
in cases where a disease-associated peptide antigen “target” has been unambiguously established. However, unlike antibodies that can undergo somatic hypermutation and bind with high affinity ($K_D = nM–pM$) and long half-lives (typically hours), TCRs are selected in the thymus to bind with weak affinity ($K_D = 100 nM–270 \mu M$) and short half-lives (typically seconds) (Cole et al., 2007; Bridgeman et al., 2012). Why TCRs are selected to bind within this weak affinity range is not fully understood, but may represent a balance between self-tolerance and a requirement for T cell cross-reactivity (Mason, 1998; Sewell, 2012; Wooldridge et al., 2012). However, the weak affinity and short half-lives of natural TCR/pMHC interactions imposes limitations on the use of TCRs for targeting cell surface expressed pMHCs, primarily because the short half-life is not adequate for the delivery of therapeutic interventions to target antigens. In order to overcome this limitation, we have recently implemented phage display to generate TCRs with an enhanced affinity for cognate antigen (Li et al., 2005; Dunn et al., 2006; Sami et al., 2007; Varela-Rohena et al., 2008; Liddy et al., 2012) that can be used to target cell surface expressed MHC molecules displaying any disease epitope of interest. Using this technique, we have generated several high affinity TCRs derived from the αβ TCR A6 (A6wt) that are specific for the HTLV-I1-19 peptide, presented by HLA-A*0201 (A2-Tax) (Li et al., 2005). High affinity TCRs generated using this method can be used for two distinct type of therapies. The first involves genetically reprogramming host T cells so that they express a modified TCR (adoptive therapy) (Morgan et al., 2006; Varela-Rohena et al., 2008). The second involves using a soluble high affinity TCR to deliver a therapeutic payload intravasitionally (soluble therapy) (Liddy et al., 2012). The optimal TCR affinity for these two types of therapy, in terms for retaining specificity and reactivity, will probably be distinct and will likely be lower for adoptive therapy because of the polyvalent nature of T cell antigen recognition at the cell surface versus a soluble therapy in which the TCR reagent will likely require a longer half-life to effectively target intended disease markers. Understanding how these reagents function at the molecular level is key to determining these parameters and optimizing these types of T cell directed therapies.

Previous structural investigations of TCR/pMHC interactions have shown that TCRs bind with a relatively conserved diagonal orientation (Garboczi et al., 1996; Rudolph et al., 2006), with the α-chain focused toward the N-terminus of the peptide and the β-chain toward the C-terminus. Although exceptions occur (Burrows et al., 2010), this orientation enables the TCR complementarity determining region (CDR)2 loops to be positioned over mainly the MHC surface, the CDR3 loops to be positioned primarily over the peptide and the CDR1 loops positioned in between. This binding mode, and the low native TCR binding affinity, is presumably important for maintaining T cell specificity and antigen sensitivity, and is possibly important in T cell signaling (Adams et al., 2011). However, we have previously shown that just a small number of mutations in the TCR CDR loops can improve the low natural TCR/pMHC binding affinity dramatically (Li et al., 2005; Dunn et al., 2006; Haswe et al., 2012). Thus, it is likely that high affinity TCRs are generated in the thymus, but they are not selected for release into the periphery. In order to better understand the consequences of high affinity TCR interactions, and to provide further insight into: (1) how high affinity TCR binding is mediated and (2) what effects this binding is likely to have on TCR specificity, we solved the atomic structures of a high affinity TCR, A6c134, free, and in complex with A2-Tax. By comparing this structure with the previously published structures for the A6wt TCR (Garboczi et al., 1996; Scott et al., 2011), we provide a molecular explanation for the improved binding of this high affinity TCR.

**RESULTS**

**DECONSTRUCTION OF HIGH AFFINITY A6 TCR VARIANTS SPECIFIC FOR A2-TAX**

In order to generate a high affinity version of the A6wt TCR, we implemented phage display as previously described (Li et al., 2005). This process generated a number of high affinity TCRs, including the mutant A6c134 that varied from the A6wt TCR parental sequence at only four codons, all located within the CDR3β loop (Table 1). It has been previously determined that the A6wt TCR binds to A2-Tax with an affinity of ~1–3 nM and an off-rate ($t_{1/2}$) of ~7–10 s (Davis-Harrison et al., 2005; Armstrong and Baker, 2007; Cole et al., 2007). In contrast, the engineered high affinity A6c134 TCR bound to A2-Tax with an affinity of 4 nM (nearly 1000 times greater than the A6wt TCR, or $\Delta\Delta G^\circ = -3.96$ kcal/mol) and an off-rate ($t_{1/2}$) of ~3900 s (~400 times longer than the A6wt TCR), as determined by surface plasmon resonance (SPR) (Table 1). A6c134 did not bind to other HLA-A2 restricted peptides that were used as negative controls in different SPR experiments, including as A2-ILAKFLHWL, A2-ELAGIGITUV, and A2-YLEPGPVTA demonstrating that A6c134 retained peptide specificity (data not shown). In order to investigate the molecular basis for how this high affinity was generated, we used reverse engineering to construct a range of A6wt-based TCRs containing different combinations of amino acids from the A6c134 CDR3β sequence and, conversely, a range of A6c134-based TCRs containing different combinations of amino acids from the A6wt CDR3β sequence (Table 1). All of these TCRs exhibited very similar on-rates, but showed marked differences in their off-rates (Table 1). In general, the mutations appeared to act cooperatively to enhance affinity. The mutation of the A6wt TCR from R102 to Q102 had only a small effect upon binding affinity and may have been selected because of the lower toxicity of the TCR to the TGI phage host (Li et al., 2005).

**A6c134 TCR BOUND TO A2-TAX USING A SIMILAR CONFORMATION TO THE A6wt TCR**

In order to determine the structural basis of the high affinity binding for the A6c134 TCR, we solved the A6c134/A2-Tax complex structure to 2.74 Å. Molecular replacement was successful only in space group C121, consistent with the presence of one molecule of the complex per asymmetric unit, and the resolution was sufficiently high to show that the interface between the two molecules was well ordered and contained well defined electron density. The crystallographic R/RFactors were 22 and 26%, within the accepted limits shown in the theoretically expected distribution (Tickle et al., 2000) (Table S1 in Supplementary Material). The overall buried surface area (BSA) of
Table 1 | Biophysical analysis of different combinations of “knock-in” and “knock-out” mutations (in bold type and underlined) of A6wt and A6c134 TCRs binding to A2-Tax.

| Mutant  | CDR3β | K_D   | k_on (s⁻¹ M⁻¹) | k_off (s⁻¹) | τ₁/₂ (s) | ΔG° (kcal/mol) | ΔΔG° (kcal/mol) |
|---------|-------|-------|---------------|-------------|----------|----------------|-----------------|
| A6wt    | AGGR  | 3.2 µM | 2.3 × 10⁴     | 7.4 × 10⁻²  | 9.3      | −7.49         | n/a             |
| A6c134M | MGGR  | 1.9 µM | 1.8 × 10⁴     | 3.5 × 10⁻²  | 20       | −7.8          | −0.31           |
| A6c134S | AGGR  | 1.8 µM | 2.3 × 10⁴     | 4.1 × 10⁻²  | 17       | −7.83         | −0.34           |
| A6c134E | ASAE  | 9.4 nM | 2.3 × 10⁴     | 5.2 × 10⁻⁴  | 1320     | −10.94        | −3.45           |
| A6c134E | MSAE  | 4.4 nM | 5.5 × 10⁴     | 2.2 × 10⁻⁴  | 3120     | −11.39        | −3.90           |
| A6c134R | MSAAR | 8 nM   | 1.9 × 10⁴     | 1.5 × 10⁻⁴  | 4500     | −11.04        | −3.55           |
| A6c134  | MSAQ  | 4 nM   | 4.5 × 10⁴     | 1.8 × 10⁻⁴  | 3900     | −11.45        | −3.96           |

Off-rates were determined by least squares fitting of single-component exponential decay equation to the decay curve following TCR binding to A2-Tax. Half-lives were calculated using the equation t½ = ln2/k_off.

ΔΔG° = ΔG° of high affinity A6c134 variant binding to A2-Tax − ΔG° of A6wt binding to A2-Tax.

2445.4 Å (TCR/pMHC) was within the range observed for previously characterized TCR/pMHC interactions (Rudolph et al., 2006). The high affinity A6c134 TCR bound with a diagonal docking geometry to A2-Tax and showed one to one stoichiometry as previously reported of other TCR/pMHC complexes (Rudolph et al., 2006). We observed a high level of similarity between the A6wt/A2-Tax and A6c134/A2-Tax complexes, suggesting that overall conformation was unaffected by the mutations in A6c134. Importantly, the Tax peptide conformation was virtually identical in both complexes, discounting the possibility that structural changes in the peptide contributed to the high affinity observed (Figures 1A,B). Similarly, the architecture of the CDR loops was unaffected by the mutations in the A6c134 TCR (Figure 1C), and the crossing angle of both TCRs was identical at 34° (Figure 1D) and fell within the previously observed range (Rudolph et al., 2006). Thus, differences in binding affinity between the A6wt and A6c134 TCRs could not be explained by a large conformational change in geometry, consistent with observations in similar studies with other systems (Dunn et al., 2006; Sami et al., 2007; Madura et al., 2013).

THE A6c134 TCR CDR LOOPS UNDERGO LARGE CONFORMATIONAL ADJUSTMENTS DURING A2-TAX ENGAGEMENT

We next solved the structure of the A6c134 TCR at 2 Å (Table S1 in Supplementary Material; Figure 2). The crystallographic R/Rfree factors were 23.4 and 29.5%, consistent with the expected ratio range (Tickle et al., 2000). In many cases, although not all (Borbulevych et al., 2011b; Holland et al., 2012), TCR CDR loops have been shown to undergo numerous, and sometimes large, conformational changes upon pMHC binding (Armstrong et al., 2008b). Superposition of the free and the bound A6c134 TCR showed that, although the overall conformation of the TCR was virtually identical (Figure 2A), the CDR loops underwent substantial movements (Figure 2B). The CDR3 loops of both chains of the A6c134 TCR were poorly ordered and could not be fully resolved, indicating a large amount flexibility in this region of the TCR. Although the apex of the CDR3 loops could not be accurately located, the portions of the CDR3α and CDR3β loops that were visible underwent large hinge movements of ~5.9 and ~4.6 Å, respectively (Figures 2C,D). These changes were substantial compared to other structural studies in which the largest loop movement observed for a human MHC-I restricted TCR was 5.6 Å (Kjer-Nielsen et al., 2002; Stewart-Jones et al., 2003). Interestingly, a nearly identical observation was made for A6wt TCR (Scott et al., 2011), for which the dynamics of the CDR3 loops were shown to have a large influence on the specificity and cross-reactivity of the TCR. This occurrence with A6c134 leads to the somewhat counterintuitive conclusion that substitution of the sequence AGGR with MSAQ in CDR3β does not greatly impact the overall dynamics of the loop, and leads us to suggest that the enhanced affinity of A6c134 was not attributable to “preorganization” of CDR3β, i.e., the MSAQ mutation did not alter the non-bound form of the A6c134 TCR to a conformation closer to that of the bound form. This conclusion is consistent with the observation that the binding
of A6c134 to A2-Tax is characterized by a much more favorable enthalpy change along with a less favorable entropy change (as pre-organization would have resulted in a more favorable binding entropy) (Armstrong and Baker, 2007; Piepenbrink et al., 2009).

The CDR1 and CDR2 loops underwent a smaller rigid body shift and hinge movements of 2.5–4 Å (Figures 2E–H). On average, the A6c134 TCR CDR loops moved by 4.1 Å. Altogether, this degree of conformational plasticity is high compared to other TCRs (Armstrong et al., 2008b), and demonstrated that the A6c134 TCR undergoes a high degree of conformational melding during binding, as does the A6wt TCR (Borbulevych et al., 2011a). These movements enable direct contacts between the A6c134 TCR and A2-Tax and resolved steric clashes that would have occurred between the unbound TCR and the MHC surface, as observed for the A6wt TCR (Scott et al., 2011).

The high affinity binding of the A6c134 TCR was governed by increased peptide contacts

As the overall free and bound conformations of A6wt and A6c134 were nearly identical, we decided to investigate differences in atomic interactions at the A6wt and A6c134 interfaces. The binding footprints of the A6wt and A6c134 TCRs on A2-Tax were similar, but not identical, resulting in the involvement of different peptide and MHC residues at the interface (Figures 3A,B). Although the A6c134 TCR α-chain contained no mutations, it utilized a different combination of TCR residues for binding A2-Tax compared with A6wt. As a result, the A6c134 TCR α-chain made a number of new and different contacts with A2-Tax compared with the A6wt TCR (Tables S2 and S3 in Supplementary Material; Table 2; Figures 3C,D). Thus, the mutations in the A6c134 TCR β-chain mediated a knock-on, or indirect effect resulting in a modified binding mode for the TCR α-chain.

In contrast with the A6c134 TCR α-chain, the A6c134 β-chain containing the mutated MSAQ motif formed a virtually identical footprint on A2-Tax compared to the A6wt TCR (Figure 3D). The A6c134 β-chain made a similar number of contacts with the MHC compared to the corresponding residues in the A6wt TCR (AGGR) (Figures 4A,B). This was reflected by the observation that the A6c134 TCR made only 11 more contacts with the MHC surface compared to the A6wt/A2-Tax complex. However, the A6c134 TCR made 26 extra contacts with the Tax peptide compared to the A6wt TCR, suggesting a TCR-peptide mediated mechanism for the enhanced affinity observed. This observation was also supported by the increase in shape complementarity index (Lawrence and Colman, 1993; Reinherz et al., 1999) (SC = 0.74) for the A6c134/A2-Tax complex compared to the A6wt/A2-Tax complex (SC = 0.63), and is consistent with the
Table 2 | Direct contacts made by the A6wt TCR, or A6c134 TCR CDR3 loops.

| TCR    | vDW (3.2–4 Å) | H-bonds (≤ 3.4 Å) |
|--------|---------------|-------------------|
| A6wt TCR | Thr93         | 0                 | 1                  |
| CDR3α  | Thr98         | 2                 | 1                  |
|        | Asp99         | 15                | 3                  |
|        | Ser100        | 7                 | 2                  |
|        | Trp101        | 5                 | 0                  |
|        | Gly102        | 0                 | 0                  |
| A6c134 TCR | Thr93      | 1                 | 0                  |
| CDR3α  | Thr98         | 2                 | 1                  |
|        | Asp99         | 15                | 2                  |
|        | Ser100        | 12                | 2                  |
|        | Trp101        | 9                 | 0                  |
|        | Gly102        | 3                 | 0                  |
| A6wt TCR | Arg95         | 0                 | 1                  |
| CDR3β  | Leu98         | 14                | 1                  |
|        | Ala99         | 0                 | 0                  |
|        | Gly100        | 1                 | 0                  |
|        | Gly101        | 6                 | 2                  |
|        | Arg102        | 14                | 1                  |
|        | Pro103        | 5                 | 0                  |
| A6c134 TCR | Arg95      | 2                 | 1                  |
| CDR3β  | Leu98         | 17                | 0                  |
|        | Met99         | 4                 | 0                  |
|        | Ser100        | 6                 | 0                  |
|        | Ala101        | 17                | 1                  |
|        | Gln102        | 7                 | 0                  |
|        | Pro103        | 11                | 0                  |

Mutated residues are shown in red.

Table 2 observation that the enhanced affinity of A6c134 was enthalpically driven as noted above (Armstrong and Baker, 2007; Pieltenbrink et al., 2009).

Overall, the mutated MSAQ motif directly accounted for 11 of the 37 new contacts with the surface of A2-Tax (Figures 4C,D; Table 2). For instance; the A99–M99 mutation generated 4 additional van der Waals contacts (Figure 5A), G100–S100 generated an additional 5 additional van der Waals contacts (Figure 5B) and G101–A101 generated 10 additional contacts (Figure 5C), with A2-Tax. Interestingly, the R102–Q102 mutation resulted in the loss of seven van der Waals contacts and one hydrogen bond (Figure 5D). However, the overall affinity was stronger for A6c134 (MSAQ) compared to A6c134R (MSAR) suggesting that the R102–Q102 mutation contributed indirectly to binding. Thus, 26 new contacts were generated through indirect interactions with non-mutated residues in the c134 TCR. The majority of these (21 new contacts) were made between non-mutated residues in the c134 TCR CDR3α and CDR3β loops demonstrating that the proximity of residues to the mutated MSAQ motif in the CDR3β loop was probably important for enabling the formation of these new interactions (Table 2).

DISCUSSION

Thymic selection generates T cells that express TCRs with a weak binding affinity ($K_D = 100$ nM–$270$ µM) for cognate antigen (Cole et al., 2007; Bridgeman et al., 2012). Presumably, this affinity range is important to ensure host protection against foreign invaders, whilst maintaining tolerance to self-antigens from a T cell repertoire of around 25 million (Arstila et al., 1999). In order to perform this function, growing evidence suggests that TCRs must be highly cross-reactive within the confines of MHC-restriction (Mason, 1998; Sewell, 2012; Wooldridge et al., 2012). We have previously shown that it is possible to enhance TCR binding affinity by phage display (Li et al., 2005). These engineered high affinity TCRs represent a potentially useful tool to target specific disease molecules, such as cancer (Liddy et al., 2012) or HIV (Varela-Rohena et al., 2008). However, how enhanced affinity affects TCR binding and specificity is not fully understood, and there are likely multiple mechanistic routes through which affinity can be enhanced. A greater understanding of such mechanisms is particularly important when developing T cell therapies that involve genetically modifying T cells with enhanced affinity TCRs or when using soluble high affinity TCR therapies, to limit potentially dangerous self-reactivity.
A6c134 TCR demonstrated that the TCR CDR3 loops underwent a face was greater for A6c134/A2-Tax than for A6wt/A2-Tax. For instance, in the A6c134/A2-Tax complex, there were a total of 154 contacts, including 72 to peptide and 82 to the MHC. In the A6wt/A2-Tax complex, there were total of 117 contacts, including 46 to peptide and 71 to the MHC. Both the MHC and peptide were involved in generating these new contacts, and they were mediated by both mutated and non-mutated TCR residues. However, the majority of the new contacts arose from interactions between the peptide and the c134 TCR CDR3 loops. Thus, we concluded that higher affinity was mediated predominantly by new TCR-peptide interactions. This conclusion, that new contacts mediated the stronger binding affinity, is consistent with previous thermodynamic analyses, which showed that the A6c134 TCR bound to A2-Tax with a substantially more favorable enthalpy compared to the A6wt TCR \( (\Delta H = -10 \text{ kcal/mol}) \) (Armstrong et al., 2008a; Piepenbrink et al., 2009).

Our generation of a high affinity TCR that contained mutations at only four residues in the CDR3β loop compared to the wt TCR sequence raises the question of why high affinity TCRs are not naturally selected in the thymus (Holler et al., 2003). Clearly, the structural framework of the TCR allows for high affinity binding, and the mutations we have identified fall within the rearranged gene segment of the TCR rather than the predefined germline encoded segments. Furthermore, the mutations in the TCR CDR3β chain generated an increase in peptide contacts within the boundaries of the native A6wt TCR binding mode and would thus be unlikely to alter MHC-restriction. Therefore, it seems very likely that high affinity TCR variants could be generated during the process of TCR rearrangement in the thymus. Yet such high affinity TCRs have not been observed during the peripheral immune response, implying that they are negatively selected (Holler et al., 2003). Presumably, this process is designed to limit self-reactivity. However, weaker affinity TCRs may also be selected to ensure a level of T cell cross-reactivity capable of fully protecting the host against all possible disease epitopes (Mason, 1998; Sewell, 2012; Wooldridge et al., 2012). In support of this notion, our data indicate that A6c134 and other high affinity TCRs can retain extremely high levels of specificity, and may be more specific than their wild-type parents (Laugel et al., 2005; Dunn et al., 2006; Madura et al., 2013). Generally, therapeutic TCRs do not need the capacity to be cross-reactive as they are designed to target a single disease epitope. This difference in desired function (immune response versus specific therapy) may represent an opportunity to improve the affinity of natural TCRs in a safe manner. Thus, our structural investigation of A6c134/A2-Tax, showing that the high affinity interaction was mediated by a native binding mechanism that was peptide-focused, may represent an important approach for developing antigen specific high affinity TCR reagents for use in T cell based therapies.

**MATERIALS AND METHODS**

**PHAGE DISPLAY**

Selection of high affinity A6wt TCR variants was performed as previously described (Li et al., 2005).

**PROTEIN PURIFICATION**

A2-Tax peptide-MHC complexes was prepared as previously described (Garboczi et al., 1992), by expressing HLA-A*0201 heavy-chain truncated at residue Pro-276 and β2 microglobulin separately in E. coli in the form of inclusion bodies, followed by in vitro refolding with synthetic peptide. pMHC for binding analysis was prepared similarly, but with the MHC fused to a biotinylation tag (Cull and Schatz, 2000) which was biotinylated in vitro by the BirA enzyme (O’callaghan et al., 1999). Disulfide-linked A6c134 TCR was prepared as previously described (Boulter et al., 2003; Li et al., 2005).

**BINDING ANALYSIS BY SURFACE PLASMON RESONANCE (BIACORE™)**

Binding analysis was performed on a Biacore™ 3000 machine using a CM-5 (research grade) chip as previously described (Cole et al., 2008; Miles et al., 2011). Streptavidin was immobilized on all flow cells using amine coupling to a level of >1000 RU (response units). Biotin tagged peptide-MHC was flowed over...
the streptavidin coated surface at a concentration of approximately 10 μg/ml until ~150 RU pHLa was bound. Control surfaces were coated with non-cognate pMHCs (A2-ILAKFLHWL, A2-ELAGIGILTV, and A2-YLEPGPVTA) or were left coated with streptavidin. Kinetic binding data were generated using the KINJECT program to inject 10 nM TCR over the flow cells. Data were analyzed using BLAevaluation™ software by kinetic fitting to calculate $k_{on}$ and $k_{off}$ rates. Binding affinities were calculated using the following equation: $K_D = k_{off}/k_{on}$.

CRYSTALLIZATION AND X-RAY DATA COLLECTION
A6c134/A2-Tax crystals were grown in MES 25 mM pH 6.5, 24% PEG 3350 and 10 mM NaCl; A6c134 free crystals were grown in MES 25 mM pH 6.5, 24% PEG 3350 and 10 mM NaCl. All crystals were soaked in 30% ethylene glycol before cryo-cooling. Data were collected at 100 K at the Advanced Photon Source at Argonne National Laboratory, USA. Reflection intensities were estimated with the XIA2 package (Winter, 2010) and the data were scaled, reduced, and analyzed with SCALA and the CCP4 package (Collaborative Computational Project, Number 4, 1994). Structures were solved with molecular replacement using Phaser (McCoy et al., 2007). Sequences were adjusted with COOT (Emsley and Cowtan, 2004) and the models refined with REFMAC5. Graphical representations were prepared with PMOL (Delano, 2002). The reflection data and final model coordinates were deposited with the PDB database (A6c134/A2-Tax, PDB: 4FTV; A6c134 free, PDB: 4GRM).

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SUPPLEMENTARY MATERIAL
The Supplementary Material for this article can be found online at http://www.frontiersin.org/T_Cell_Biology/10.3389/fimmu.2013.00168/abstract

Table S1 | Data collection and refinement statistics (molecular replacement).

Table S2 | Structural analysis of A6wt/A2-Tax contacts.

Table S3 | Structural analysis for A6c134/A2-Tax contacts (mutant residues in red).

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