

Thermodynamics of T-cell receptor–peptide/MHC interactions: progress and opportunities

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$\alpha\beta$ T-cell receptors (TCRs) recognize peptide antigens presented by class I or class II major histocompatibility complex molecules (pMHC). Here we review the use of thermodynamic measurements in the study of TCR–pMHC interactions, with attention to the diversity in binding thermodynamics and how this is related to the variation in TCR–pMHC interfaces. We show that there is no enthalpic or entropic signature for TCR binding; rather, enthalpy and entropy changes vary in a compensatory manner that reflects a narrow free energy window for the interactions that have been characterized. Binding enthalpy and entropy changes do not correlate with structural features such as buried surface area or the number of hydrogen bonds within TCR–pMHC interfaces, possibly reflecting the myriad of contributors to binding thermodynamics, but likely also reflecting a reliance on van't Hoff over calorimetric measurements and the unaccounted influence of equilibria linked to binding. TCR–pMHC binding heat capacity changes likewise vary considerably. In some cases, the heat capacity changes are consistent with conformational differences between bound and free receptors, but there is little data indicating these conformational differences represent the need to organize disordered CDR loops. In this regard, we discuss how thermodynamics may provide additional insight into conformational changes occurring upon TCR binding. Finally, we highlight opportunities for the further use of thermodynamic measurements in the study of TCR–pMHC interactions, not only for understanding TCR binding in general, but also for understanding specifics of individual interactions and the engineering of TCRs with desired molecular recognition properties. Copyright © 2008 John Wiley & Sons, Ltd.

Keywords: T-cell receptors; peptide/MHC complexes; thermodynamics; enthalpy/entropy compensation; conformational selection

INTRODUCTION

Recognition of an antigenic peptide presented by a class I or class II major histocompatibility complex (MHC) protein is required for the initiation and propagation of a cellular immune response as well as generation and maintenance of the T-cell repertoire. Peptide/MHC complexes (pMHCs) are recognized by $\alpha\beta$ T-cell receptors (TCRs) expressed on the surface of CD4⁺ or CD8⁺ T-cells (Figure 1). In some respects, TCRs are similar to antibodies; important here is that their antigen-binding sites are composed of multiple complementarity-determining region (CDR) loops generated via genetic recombination processes similar to those used in antibody generation. However, one of the many key differences between antibodies and TCRs is the nature of the ligand recognized. Whereas antibodies recognize linear or non-linear epitopes of seemingly unlimited chemical and structural diversity, TCRs recognize a composite surface consisting of elements of the antigenic peptide as well as the α -helices of the MHC peptide-binding groove (recently reviewed in Rudolph *et al.*, 2006). Thus, unlike antibodies, the ligand for the TCR consists of both the self (the MHC) and non-self (the peptide).

A second key difference between antibodies and TCRs is that TCRs are cross-reactive, capable of recognizing multiple peptides bound to one or more MHC molecules (recently reviewed in Wucherpfennig *et al.* (2007), where use of the term “polyspecific” was suggested for describing TCR recognition of multiple ligands). TCR cross-reactivity (or polyspecificity) is necessary for the development and continued maintenance of the T-cell arm of the immune system and is crucial given the fixed size of the T-cell repertoire relative to the vast universe of potential peptide

antigens (Mason, 1998). These two features, cross-reactivity and dual recognition of self/non-self, present special challenges and opportunities in the study of TCR molecular recognition. Here, we review the use of thermodynamic measurements in the study of TCR–pMHC interactions, with particular attention to how thermodynamic measurements have instructed our understanding of the determinants of TCR binding and specificity. We discuss opportunities for the further use of thermodynamic measurements, not only for understanding TCR binding in general, but also for understanding specifics of individual interactions and the engineering of TCRs with desired molecular recognition properties. We also discuss the use of binding kinetics as a complementary tool to investigate molecular recognition properties, although we do not address the ongoing debate regarding the relative importance of TCR-binding kinetics versus affinity in T-cell signaling and activation (e.g., Rosette *et al.*, 2001; Tian *et al.*, 2007).

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Abbreviations: TCRs, T-cell receptor; MHC, major histocompatibility complex; pMHC, peptide/MHC complex; CDR, complementarity-determining region.

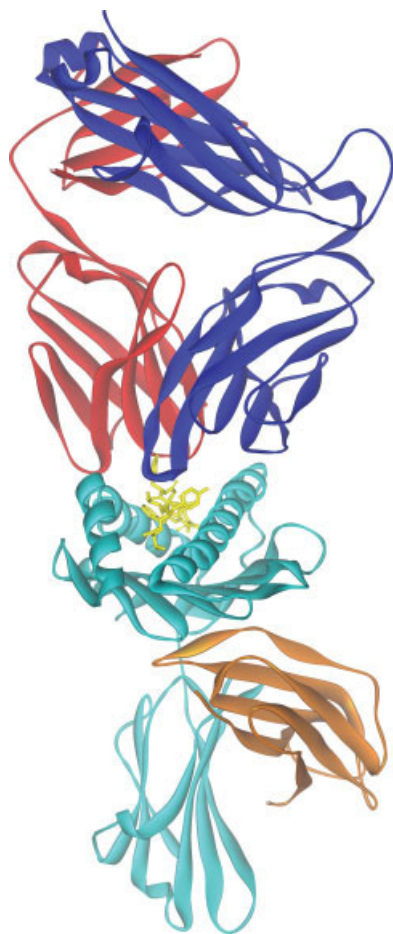


Figure 1. Ribbon diagram of a complex between an $\alpha\beta$ T-cell receptor and a peptide/MHC complex. The α chain of the TCR is red and the β chain is blue. The peptide in the peptide/MHC complex is in yellow stick format, the heavy chain cyan, and β_2m is light brown. The dual recognition of peptide (non-self) and MHC (self) is evident from the figure. The figure is of the B7 TCR recognizing the Tax peptide presented by HLA-A2, PDB file 1BD2 (Ding *et al.*, 1998).

GENERAL FEATURES OF TCR-BINDING THERMODYNAMICS: IS THERE A THERMODYNAMIC SIGNATURE FOR TCR-pMHC BINDING?

Surface plasmon resonance studies of TCR-pMHC interactions performed in the mid-to-late 1990s with soluble ectodomains indicated that TCRs bind ligand weakly with slow-to-moderate association rates (usually $\leq 10^5 \text{ M}^{-1} \text{ s}^{-1}$) (Davis *et al.*, 1998), slower than those expected for a diffusion limited, geometrically constrained protein-protein interaction (Janin, 1997; Vijayakumar *et al.*, 1998). As slow kinetics can result from conformational adjustments required for binding and because little or no conformational differences were seen between bound and free pMHC (Garboczi *et al.*, 1996; Garcia *et al.*, 1996, 1998), some authors suggested that TCR CDR loops must undergo conformational adjustments upon recognition of ligand (Matsui *et al.*, 1994). This suggestion was supported by crystallographic studies with the $\alpha\beta$ TCR 2C, for which multiple CDR loops were shown to populate different conformations in the free and bound states (Garcia *et al.*, 1996, 1998). Around the time of these observations,

the inherent cross-reactivity of TCRs was becoming increasingly appreciated (e.g., Bhardwaj *et al.*, 1993; Evavold *et al.*, 1995; Wucherpfennig and Strominger, 1995), highlighted by Mason's estimation that any given T-cell is capable of reacting "productively with approximately 10^6 different MHC-associated minimal peptide epitopes" (Mason, 1998). TCR cross-reactivity, sometimes discussed as binding degeneracy, fits well with the notion that receptor binding occurs with conformational shifts in one or more CDR loops, as the availability of multiple conformations for unbound TCRs could broaden the reactivity of any given receptor.

The first measurements of TCR-binding thermodynamics closely followed, with van der Merwe and colleagues showing in 1999 that the human TCR JM22 and the murine TCR F5 bound pMHC with favorable enthalpy changes and unfavorable entropy changes (Willcox *et al.*, 1999). Although these measurements were performed via van't Hoff analysis not accounting for the influence of a heat capacity change, the measurement with JM22 was corroborated by a direct measurement of the enthalpy change using titration calorimetry (Willcox *et al.*, 1999). Considered with the kinetic and structural data, the unfavorable entropy changes for the binding of JM22 and F5 suggested that rather than simply adjusting conformation upon binding, TCR CDR loops are flexible in the unbound state. Such flexibility was proposed to be a key feature of TCRs and their biology, allowing the receptor-binding site to adapt to different ligands.

The measurements of van der Merwe were closely followed by a report from the Davis lab, who showed that the 2B4 TCR bound the mouse class II MHC I-E^k presenting the MCC peptide with thermodynamics similar to those for the JM22 and F5 TCRs (a favorable enthalpy change and an unfavorable entropy change, reported at 25°C) (Boniface *et al.*, 1999). Van't Hoff analysis was used again, although the data analysis incorporated a heat capacity change, which at -660 cal/mol/K was reasonably large for a protein-protein interaction that buried about 2000 \AA^2 (Stites, 1997). The availability of measurements of both ΔS° and ΔC_p allowed Davis and co-workers to perform a structure-based thermodynamic dissection of the binding thermodynamics, albeit using modeled structures. The approach used, developed by Spolar and Record in their analysis of protein-DNA interactions (Spolar and Record, 1994), extracts the contribution of changes in solvation and the loss in translational and rotational degrees of freedom from overall binding entropy changes, with the residual attributed to changes in protein backbone or side chain conformational entropy. Application of this method to the 2B4 TCR-binding data suggested a significant conformational entropy penalty must be overcome for binding, estimated to result from the need for approximately 30 amino acids to "fold" in order for the 2B4 TCR to bind. Again, considering prior kinetic and structural observations, the overall interpretation of the data was that flexibility in the TCR CDR loops permitted a TCR repertoire of limited size to interact with a much larger array of pMHC ligands, that is, the cross-reactivity inherent in the TCR was attributable at least in part to the capacity for one or more CDR loops to adopt multiple conformations.

Additional reports of unfavorable entropy changes for TCR-pMHC interactions followed (Garcia *et al.*, 2001; Anikeeva *et al.*, 2003; Krosggaard *et al.*, 2003; Lee *et al.*, 2004), and for a brief period the notion that unfavorable binding entropy changes reflect a loss of TCR conformational flexibility was generally well accepted, having been bolstered by structural studies indicating conformational differences between free and bound receptors

(or conformational differences in the same receptor bound to different ligands) (Garcia *et al.*, 1996, 1998; Ding *et al.*, 1999; Reiser *et al.*, 2002, 2003). TCRs were presumed to bind ligand with a “thermodynamic signature” consisting of a favorable enthalpy change and an unfavorable entropy change, with strong implications for T-cell immunobiology.

However, this thermodynamic signature was soon questioned with the description of entropically favorable TCR–pMHC interactions. The first example was the binding of the A6 TCR to the Tax peptide presented by HLA-A2 (Davis-Harrison *et al.*, 2005). Notably, the CDR loops of the A6 TCR had previously been shown to change conformation upon recognition of different ligands (Ding *et al.*, 1999), an observation used to support the idea of structural plasticity within TCR-binding sites. The second observation of an entropically favorable TCR–pMHC interaction was the binding of the LC13 TCR to the FLR peptide presented by HLA-B8 (Ely *et al.*, 2006). Since these two observations, numerous other TCR-binding reactions have been shown to be entropically favorable, conclusively demonstrating that unfavorable binding entropies are not a signature of TCR recognition of pMHC (Colf *et al.*, 2007; Gakamsky *et al.*, 2007; Mazza *et al.*, 2007; Miller *et al.*, 2007), irrespective of whether conformational changes occur upon binding.

Yet should these results have been surprising? The overall binding thermodynamics associated with molecular recognition in any system are influenced not only by what needs to occur for binding to proceed (e.g., conformational changes or reductions in flexibility), but also the specific details of the resulting interface (e.g., hydrophobic/hydrophilic surface buried, hydrogen bonds and salt bridges formed, water or ion incorporation, pK_a shifts, etc.). In that respect, due to the extensive variation in the makeup of the CDR loops, the different peptides and MHC molecules, and the variations in TCR-binding modes, the chemistry in every TCR–pMHC interface is different. Loop conformational changes or reductions in flexibility could indeed occur in any given interaction, but the thermodynamic consequences may be masked by other contributors, the costs of which are difficult to predict—even the deceptively simple entropic cost for forming a complex out of two proteins is not well established given differing capacities for residual motion at the binding site (Brady and Sharp, 1997; Lu and Wong, 2005) and observations that protein dynamics can redistribute or even increase upon ligand binding (e.g., Zidek *et al.*, 1999; Fayos *et al.*, 2003; Grunberg *et al.*, 2006).

Thus, the notion that TCR–pMHC interactions should have an enthalpic/entropic “signature” is too much of a generalization. A good example is provided by the A6 and B7 TCRs. Both receptors recognize the Tax/HLA-A2 ligand as a strong agonist with similar structural topologies and using the same $V\beta$ segments, yet the binding thermodynamics for the two receptors are markedly different: whereas the binding of A6 is entropically favored, the binding of B7 is entropically opposed, and the binding enthalpy changes differ by more than 10 kcal/mol at 25°C.

THE VARIATION IN TCR–pMHC-BINDING THERMODYNAMICS: ENERGETICALLY, BIOLOGY DOES NOT CARE HOW YOU FORM THE COMPLEX, JUST THAT YOU DO

If there is no enthalpic or entropic signature for TCR–pMHC interactions, are there trends in the thermodynamic data likely to yield insight into T-cell immunobiology? All of the TCR–pMHC

interactions whose underlying binding thermodynamics have been characterized to-date are tabulated in Table 1, and plotted graphically as binding ΔH° versus binding ΔS° in Figure 2 (all data are shown at a common reference temperature of 25°C). Figure 2A shows data for only “wild-type” TCRs, peptides, or MHC molecules; Figure 2B adds thermodynamic data for all interactions involving peptide variants, MHC mutants, and altered TCRs (again, only for those interactions for which ΔH° and ΔS° are available). A number of details are clear from this analysis. Examining only the “wild-type” interactions (Figure 2A), the enthalpy/entropy changes vary over a wide range (from -30 to -2 kcal/mol for ΔH° and from -80 to 15 cal/mol/K for ΔS°), yet the binding affinities (reported by the ΔG° values) are all very similar. Binding enthalpy changes are all favorable, consistent with the majority of protein–protein interactions (Stites, 1997), although there are several with enthalpy changes very close to zero. A close comparison of the data in Figure 2A with Table reveals there are no commonalities dictated by either TCR or MHC subtype (discussed further below).

Is there any distinction in the data between recognition of class I or class II MHC? The majority (17/20) of the interactions in Figure 2A are for recognition of peptides presented by class I MHC molecules. The three interactions involving class II MHC are indeed characterized by unfavorable entropy changes offset by very favorable enthalpy changes, but these values do not by themselves distinguish recognition of class II from class I MHC molecules, as the values for the three class II interactions are bracketed by the values for the class I interactions. It remains possible that more thermodynamic data for class II will reveal more clustered enthalpy and entropy values, amounting to an MHC class II-specific thermodynamic signature; however, considering that the chemical variation in class II interfaces is no less than the chemical variation in class I interfaces, this would be unexpected. Overall, the data seem clear in that as far as enthalpy and entropy are concerned, it matters not *how* you form the TCR–pMHC complex, just that you *do*.

The trends in the TCR–pMHC thermodynamic data remain unchanged if thermodynamic data involving peptide variants, MHC mutants, and altered TCRs are included (Figure 2B). While the class II data are still clustered in the favorable enthalpy/unfavorable entropy regime, this arises because of the large number of altered peptides examined with the 2B4 TCR (Krogsgaard *et al.*, 2003), most of which maintain the general overall thermodynamic trend of the wild-type interaction when compared at the common reference temperature of 25°C.

The linearity of the plots in Figure 2 reveals a striking degree of enthalpy/entropy compensation in the TCR–pMHC thermodynamic binding data. Enthalpy/entropy compensation is often described with regard to compensating thermodynamics in molecular events such as flexibility versus specificity, hydrogen bonding, etc. However, in the case of the TCR–pMHC data, the compensation arises solely because the range of binding free energies for the various interactions studied is small relative to the range of binding enthalpies: for all the data available, ΔH° spans 40 kcal/mol, whereas ΔG° spans only 3 kcal/mol. Thus, compared to the range in ΔH° , ΔG° is essentially constant. A plot of ΔH° as a function of ΔS° as in Figure 2 consequently takes the form of $\Delta H^\circ = T\Delta S^\circ + \langle \Delta G^\circ \rangle$, or the equation of a line with slope equal to the reference temperature of the data and a y -intercept equal to the average ΔG° . In the case of Figure 2, the slope and intercepts for Figures 2A,B are 291 and 295 K and -7.3 and -7.1 kcal/mol, respectively, with an actual reference temperature

Table 1. Published TCR–pMHC-binding thermodynamics at 25°C

TCR–pMHC interaction	ΔS° (cal/mol/K)	ΔH° (kcal/mol)	ΔG° (kcal/mol)	ΔC_p (cal/mol/K)	MHC class	Notes	Reference
2C–dEV8/H-2K ^b	–54.33	–22.7	–6.5	–1100	I		Degano <i>et al.</i> (2000), Krogsgaard <i>et al.</i> (2003)
2C–p2Ca/H-2K ^b	–77	–29	–6.0	–1500	I		Degano <i>et al.</i> (2000), Krogsgaard <i>et al.</i> (2003)
2C–QL9/H-2L ^d	11.58	–4.19	–7.6	—	I	a	Colf <i>et al.</i> (2007)
2C–SIYR/H-2K ^b	–4.1	–8.4	–7.2	–1100	I		Degano <i>et al.</i> (2000), Krogsgaard <i>et al.</i> (2003)
A6–Tax/HLA-A2	11.6 ± 3.5	–4.2 ± 1.1	–7.7	–560 ± 120	I		Davis-Harrison <i>et al.</i> (2005)
A6–Tax/HLA-A2	25 ± 1	–1.8 ± 0.3	–9.2 ± 0.1	–520 ± 50	I	b	Armstrong and Baker (2007)
AHIII–p1049/HLA-A2	14.76	–3.9	–8.3	—	I	c	Miller <i>et al.</i> (2007)
AHIII–p1049/HLA-A2 K66A	21.13	–0.6	–6.9	—	I	c	Miller <i>et al.</i> (2007)
B7–Tax/HLA-A2	–22 ± 3.1	–14.6 ± 0.9	–8.0	–750 ± 90	I		Davis-Harrison <i>et al.</i> (2005)
BM3.3–pBM1/H-2K ^b	–19.8 ± 1	–13.4 ± 0.4	–7.5 ± 0.1	–535 ± 76	I		Mazza <i>et al.</i> (2007)
BM3.3–pBM8/H-2K ^{bm8}	8 ± 1	–3.1 ± 0.4	–5.5 ± 0.03	–141 ± 7	I		Mazza <i>et al.</i> (2007)
BM3.3–pBM8(Y4D)/H-2K ^{bm8}	4 ± 1	–5.4 ± 0.1	–6.5 ± 0.1	–108 ± 3.3	I		Mazza <i>et al.</i> (2007)
CMV–pp65/HLA-A2	13.89 ± 2.34	–2.97 ± 0.5	–7.1	–190 ± 300	I		Gakamsky <i>et al.</i> (2007)
D3–SL9/HLA-A2	–10 ± 2	–10.4 ± 0.6	–7.5	–363 ± 80	I		Anikeeva <i>et al.</i> (2003)
F5–NP/H-2D ^b	–41	–19	–6.8	—	I	d	Willcox <i>et al.</i> (1999)
G10–SLF/HLA-A2	–10.7 ± 1.7	–10.4 ± 0.4	–7.2	–620 ± 50	I		Lee <i>et al.</i> (2004)
G10–SLY/HLA-A2	–9.39 ± 1.3	–10.5 ± 0.4	–7.7	–630 ± 50	I		Lee <i>et al.</i> (2004)
LC13–FLR/HLA-B8	14 ± 8.7	–2.4 ± 2.8	–6.7 ± 0.05	–620 ± 270	I		Ely <i>et al.</i> (2006)
2C m6–QL9/H-2L ^d	–16.25	–13.8	–9.04	—	I	a	Colf <i>et al.</i> (2007)
P14–gp33/H-2D ^b	–8	–10.8	–8.4	–600	I		Boulter <i>et al.</i> (2007)
JM22–Flu/HLA-A2	–50 ± 2	–23 ± 0.6	–7.1 ± 0.2	–640 ± 100	I		Willcox <i>et al.</i> (1999), Ishizuka <i>et al.</i> (2008)
JM22(I53βV)–Flu/HLA-A2	–50 ± 2	–22 ± 0.6	–7.0	–630 ± 100	I		Ishizuka <i>et al.</i> (2008)
JM22(N55βA)–Flu/HLA-A2	–40 ± 3	–18 ± 0.9	–6.1	–330 ± 110	I		Ishizuka <i>et al.</i> (2008)
JM22(D56βA)–Flu/HLA-A2	–57 ± 5	–24 ± 1.6	–7.1	–710 ± 150	I		Ishizuka <i>et al.</i> (2008)
JM22(Q58βA)–Flu/HLA-A2	–37 ± 3	–18 ± 0.8	–6.7	–320 ± 70	I		Ishizuka <i>et al.</i> (2008)
JM22(Q58βE)–Flu/HLA-A2	–44 ± 1	–20 ± 0.4	–7.2	–340 ± 50	I		Ishizuka <i>et al.</i> (2008)
JM22(S99βA)–Flu/HLA-A2	–34 ± 2	–18 ± 0.5	–7.2	–420 ± 60	I		Ishizuka <i>et al.</i> (2008)
JM22(Y101βA)–Flu/HLA-A2	–74 ± 3	–29 ± 0.8	–7.0	–1100 ± 110	I		Ishizuka <i>et al.</i> (2008)
JM22(Y101βF)–Flu/HLA-A2	–44 ± 1	–20 ± 0.3	–6.7	–410 ± 40	I		Ishizuka <i>et al.</i> (2008)
JM22(S32αA)–Flu/HLA-A2	–60 ± 3	–24 ± 0.8	–6.2	–480 ± 70	I		Ishizuka <i>et al.</i> (2008)
JM22(Q34αA)–Flu/HLA-A2	–57 ± 4	–23 ± 1.1	–6.2	–1100 ± 130	I		Ishizuka <i>et al.</i> (2008)
172.10–MBP/I-A ^u	–48	–21.2	–6.9	–159	II	e	Garcia <i>et al.</i> (2001)
1934.4–MBP/I-A ^u	–32	–15.7	–6.1	–1248	II	e	Garcia <i>et al.</i> (2001)
2B4–102S/I-E ^k	–23.0 ± 10.2	–12.6 ± 1.8	–5.7 ± 3.5	–300 ± 300	II	f	Krogsgaard <i>et al.</i> (2003)
2B4–K2/I-E ^k	–22.2 ± 5.2	–13.6 ± 1.3	–7.0 ± 2.0	–2100 ± 200	II	f	Krogsgaard <i>et al.</i> (2003)
2B4–K3/I-E ^k	–42.1 ± 16.8	–18.5 ± 2.6	–6.0 ± 5.7	–4000 ± 500	II	f	Krogsgaard <i>et al.</i> (2003)
2B4–K5/I-E ^k	–54.9 ± 5.2	–23.6 ± 1.6	–7.2 ± 2.2	–1200 ± 100	II	f	Krogsgaard <i>et al.</i> (2003)
2B4–MCC ₈₈₋₁₀₃ /I-E ^k	–18.7 ± 0.6	–12.7 ± 0.2	–7.1	–663 ± 53	II		Boniface <i>et al.</i> (1999)
2B4–MCC ₉₅₋₁₀₃ /I-E ^k	–16.0 ± 5.7	–11.8 ± 1.8	–7.0 ± 2.5	–600 ± 800	II	f	Krogsgaard <i>et al.</i> (2003)
2B4–PCC/I-E ^k	59.6 ± 41.0	11.5 ± 3.5	–6.3 ± 12.7	–1800 ± 200	II	f	Krogsgaard <i>et al.</i> (2003)
2B4–PCC(A103K)/I-E ^k	–25.1 ± 3.5	–14.4 ± 1.1	–6.9 ± 1.5	–1000 ± 100	II	f	Krogsgaard <i>et al.</i> (2003)

In the “Notes” column, the letters indicate:

a, values reported at 20°C, no heat capacity value to scale ΔH° and ΔS° to 25°C;

b, calorimetric measurement; values are the “intrinsic” binding thermodynamics corrected for the influence of linked protonation;

c, calorimetric measurement; errors in ΔH° reported to be <5%;

d, van’t Hoff analysis without ΔC_p term; F5 ΔH° reproduced via titration calorimetry;

e, reference temperature not indicated, assumed to be 25°C;

f, values reported at temperatures ranging from 12 to 28°C, presented here at 25°C using reported ΔC_p values.

of 298 K and average ΔG° values of –7.2 and –7.1. Results such as this are commonly seen in discussions of entropy/enthalpy compensation and are one of the main contributors to a vigorous debate regarding the phenomenon (Jen-Jacobson *et al.*, 2000; Cooper *et al.*, 2001; Sharp, 2001). The appearance of enthalpy/entropy compensation within the entire TCR–pMHC thermodynamic database therefore indicates only that the measured

interactions are all of very similar affinities and the underlying thermodynamics for the various TCR–pMHC interactions vary widely. Two questions that arise then are (a) why are the affinities for the various interactions in Table 1 and Figure 2 so similar (i.e., why is there a narrow free energy window), and (b) can the variation in enthalpy and entropy be related to properties of the individual interactions?

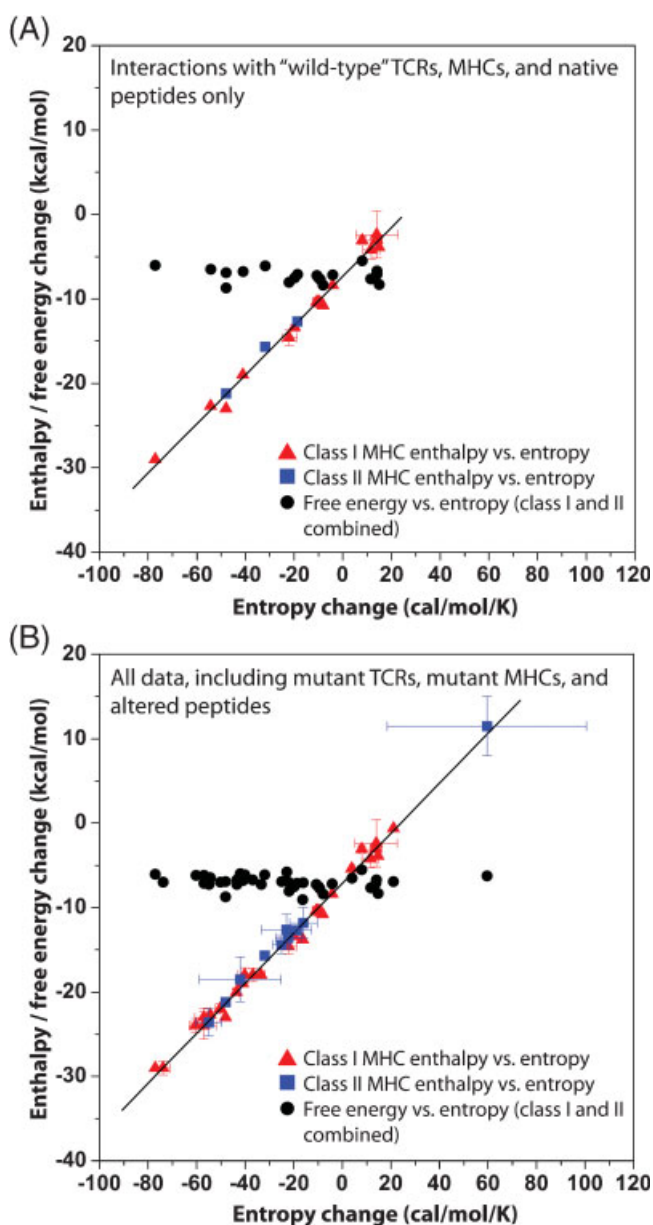


Figure 2. Diversity in TCR-pMHC-binding enthalpy and entropy changes revealed by plotting ΔH° and ΔG° versus ΔS° at 25°C. (A) Only data for interactions involving wild-type TCRs, wild-type MHCs, and native peptides. (B) All data including mutant proteins and modified peptides. For both (A) and (B), data involving recognition of class I pMHC are shown as red triangles, data involving recognition of class II MHC are shown as blue squares, and ΔG° values for recognition of both class I and class II pMHC are shown as black circles. Errors were included when available. The lines represent linear fits to the ΔH° versus ΔS° data. The clear linearity reflects the presence of enthalpy/entropy compensation within the data, but this only arises because ΔG° is relatively invariant with respect to ΔH° as described in the text. For (A), the slope and intercept are 291 K and -7.3 kcal/mol and $R^2 = 0.99$. For (B), the slope and intercept are 295 K and -7.1 kcal/mol and $R^2 = 0.99$. All data are tabulated in Table 1, and unless otherwise indicated reported at a reference temperature of 298 K (25°C).

With regard to the first question, most of the interactions shown represent TCR recognition of efficiently activating ligands, the exceptions being recognition of p2Ca/H-2K^b by the 2C TCR and p1049/HLA-A2 by the K66A mutant of the AHIII TCR. Do the similar affinities reflect a biological requirement, perhaps

reflecting the need to ensure efficient recognition while biasing against autoimmunity (Holler *et al.*, 2003)? Is a tight window of binding free energy necessary for efficient TCR signaling, perhaps reflecting optimal kinetic parameters (McKeithan, 1995)? Or are the interactions in Figure 2 simply TCR-pMHC interactions which can easily be investigated biophysically, that is, the "low-hanging fruit" of TCR interaction thermodynamics? The observation that none of the measured affinities are substantially stronger than $1 \mu\text{M}$ would seem to argue for a biological limit on how strong an affinity a positively selected TCR can have toward a fully activating ligand (although T-cells expressing receptors engineered for very high affinity can retain specificity and function; see Weber *et al.*, 2005). At the other end of the scale, however, some very weak interactions can activate T-cells (e.g., Gagnon *et al.*, 2006). Further biophysical investigations, including those involving weak or partial agonists and pre-selected TCRs are needed to fully address this question. Emerging technologies for accurately measuring low affinities and calorimeters with enhanced sensitivity and substantially reduced sample requirements will be helpful in this regard.

With regard to the second question, so far there are no clear correlations between structural or dynamical features and TCR-binding enthalpy or entropy changes. Table 2 summarizes features of the TCR-pMHC interactions whose thermodynamics have been characterized and for which structures of the ternary complexes are available. Binding enthalpy changes are not correlated with the number of hydrogen bonds or salt-bridges in the interface, nor are they well predicted by empirical prediction methods that account for buried polar and apolar surface area (Murphy and Freire, 1992; Baker and Murphy, 1998). As discussed below, there may be multiple reasons for this, including not only conformational differences between free and bound TCRs, but also inaccuracies in van't Hoff measurements and the potentially significant influence of linked equilibria. Importantly though, structural data are not available for all the TCR-pMHC interactions whose thermodynamics have been measured, and, excluding a single NMR study that examined dynamics on the picosecond timescale (Hare *et al.*, 1999), there have been no direct assays of CDR loop dynamics with which to compare binding entropy changes.

As per phenomenological correlations with TCR-binding thermodynamics, again, there are no clear correlations, but equivalent specificity data are not available for all the interactions that have been thermodynamically characterized. Interestingly, for the A6 and B7 TCRs, relaxed specificity is not correlated with a less favorable binding ΔS° . Binding of the A6 TCR is entropically favored, whereas binding of the B7 TCR is entropically opposed (Davis-Harrison *et al.*, 2005). Yet a detailed investigation of the fine specificity of the A6 and B7 TCRs indicated that A6 is considerably less specific than B7 (Hausmann *et al.*, 1999). Thus, the A6 and B7 data are contrary to the general expectation that finer specificity arises from a "tightening" of conformational mobility, insofar as reported on by binding entropy changes.

DATA QUALITY AND LINKED EQUILIBRIA

An important issue when comparing binding thermodynamics for different interactions concerns the accuracy and precision of the data. All but two of the entries in Table 1 were collected via van't Hoff analysis (i.e., analysis of measurements of affinities as a function of temperature). Although in all but one case a heat capacity term was included, van't Hoff analysis requires a level of precision and accuracy in free energy measurements which can

Table 2. Structural features and predicted thermodynamics for TCR-pMHC interactions characterized both structurally and thermodynamically

TCR-pMHC Interaction	Buried apolar SASA ^a (Å ²)	Buried polar SASA ^a (Å ²)	Hydrogen bonds ^b	Salt bridges	ΔH° experimental ^c (kcal/mol)	ΔH° predicted ^d (kcal/mol)	ΔC_p experimental ^c (cal/mol/K)	ΔC_p predicted ^e (cal/mol/K)	ΔC_p predicted ^{2f} (cal/mol/K)	PDB ID
2C-dEV8/H-2K ^b	1139	841	4	1	-22.7	-6.8	-1100	-270	-250	2CKB
2C-QL9/H-2L ^d	1057	659	7	0	-4.19	-1.7	—	-280	-250	2O19
2C-SYR/H-2K ^b	1074	773	5	0	-8.4	-5.6	-1100	-260	-240	1G6R
A6-Tax/HLA-A2	1259	817	11	4	-4.2 ± 1.1 (-1.8 ± 0.3) ^g	-3.2	-560 ± 120 (-520 ± 50) ^g	-330	-290	1A07
AHIII-p1049/HLA-A2	1299	681	5	1	-3.9	+2.9	—	-380	-320	1LP9
AHIII-p1049/HLA-A2 (K66A)	1296	708	5 (3)	4 (2)	-0.6	+1.8	—	-370	-320	2J8U
B7-Tax/HLA-A2	1248	737	4	3	-14.6 ± 0.9	-0.3	-750 ± 90	-350	-300	1BD2
BM3.3-pBM1/H-2K ^b	677	698	8	0	-13.4 ± 0.4	-11.6	-535 ± 76	-110	-120	1F00
BM3.3-pBM8/H-2K ^{cm8}	880	657	6	1	-3.1 ± 0.4	-5.5	-141 ± 7	-210	-190	2OL3
2C m6-QL9/H-2L ^d	1103	765	7	0	-13.8	-4.7	—	-280	-250	2E7L
JM22-Flu/HLA-A2 ^h	932/897/ 934	633/610/ 648	8	0	-23 ± 0.6	-3.2/-2.8/ -3.7	-640 ± 100	-250	-210/-210/ -200	1OGA/ 2VLJ/2VLK
JM22(S99βA)-Flu/HLA-A2	929	662	7	0	-18 ± 0.5	-4.1	-420 ± 60	-250	-200	2VLR
LC13-FLR/HLA-B8	1196	1040	8	1	-2.4 ± 2.8	-13.0	-620 ± 270	-270	-240	1MI5
172.10-MBP/I-A ^u	1080	824	5	0	-21.2	-7.2	-159	-270	-230	1U3H
1934.4-MBP/I-A ^u	1055	938	8	2	-15.7	-12.4	-1248	-230	-200	2PXY

^aBuried apolar and polar solvent accessible surface area (SASA) calculated assuming a rigid body interaction, using a 1.4 Å probe radius and a 0.05 Å slice width using the Naccess program (Hubbard and Thornton, 1993).

^bHydrogen bonds calculated using HBPLUS (McDonald and Thornton, 1994), integrating the data tabulated in Rudolph *et al.* (2006).

^cValues from Table 1.

^dBinding enthalpies predicted from buried apolar and polar solvent accessible surface area as described in Baker and Murphy (1996). Scaled to 25°C using predicted ΔC_p values.

^eBinding heat capacity changes predicted from buried apolar and polar solvent accessible surface area as described in Baker and Murphy (1997).

^fBinding heat capacity changes predicted from buried apolar and polar solvent accessible surface area as described in Spolar and Record (1994).

^gValues in parenthesis are the "intrinsic" binding thermodynamics removed from the influence of protonation (see Armstrong and Baker, 2007).

^hThree structures are available for the JM22-Flu/HLA-A2 complex (see Stewart-Jones *et al.*, 2003; Ishizuka *et al.*, 2008).

be difficult to achieve with weak or moderate affinity interactions, particularly when protein concentrations are limiting (Zhukov and Karlsson, 2007). This point has been raised in discussions of differences between calorimetric and van't Hoff enthalpies (Naghbi *et al.*, 1995), where the discrepancy was eventually shown to be largely attributable to issues of accuracy, precision, error propagation, and data analysis (Chaires, 1997; Horn *et al.*, 2001, 2002; Mizoue and Tellinghuisen, 2004). Although some of the TCR-binding data in Figure 2 were corroborated by calorimetric measurements, and in general the data in Figure 2 should be expected to be of high quality, it is worth appreciating that obtaining accurate thermodynamics via van't Hoff analysis places high demands on data acquisition and analysis. On this note, although calorimetry may be preferable to van't Hoff analyses, calorimetric measurements can likewise be inaccurate in the absence of a well-defined sigmoidal titration curve (Wiseman *et al.*, 1989; Tellinghuisen, 2008), a problem which has limited the application of titration calorimetry to the studies of TCR–ligand interactions due to the need for very high protein concentrations. New instruments with higher sensitivity and reduced sample requirements should help obviate this concern (on a related note, due to concerns about the accuracies of both van't Hoff and calorimetric data, the trend to publish thermodynamic values without showing relevant binding data should be strongly discouraged).

However, even if binding ΔH° and ΔS° values are highly accurate, equilibria linked to binding can cause the measured values to differ significantly from the true thermodynamics of binding. A frequently encountered form of linkage is proton linkage, which occurs when binding induces a shift in the pK_a of an ionizable group (Baker and Murphy, 1996). The contribution to the binding thermodynamics results from the proton release or uptake associated with the pK_a shift: protons that are released due to a pK_a decrease will be bound by the buffer, and protons taken up due to a pK_a increase will be released by the buffer. As most biological buffers have very large enthalpies of ionization (e.g., 5 kcal/mol for HEPES at 25°C), the contribution to the observed binding enthalpy change can be substantial. Further, because buffer ionization influences the binding ΔH° but not ΔG° (Baker and Murphy, 1996), there will be a compensatory shift in the binding ΔS° . Thus, proton linkage can dramatically influence both the enthalpy and entropies of binding. In extreme cases (or in cases where the intrinsic binding enthalpy is close to zero), the sign of ΔH° or ΔS° can vary with buffer choice (Baker and Murphy, 1997). Although proton linkage is most commonly associated with calorimetric measurements, a pK_a shift occurring upon binding will result in proton release/uptake regardless of which technique is used to monitor the process, and van't Hoff measurements are just as susceptible to the influence of proton linkage as calorimetric measurements (Horn *et al.*, 2002; Armstrong and Baker, 2007).

It remains to be seen how much influence proton linkage will have on the thermodynamics of TCR–pMHC interactions. However, because of its potentially large influence, as TCR–pMHC interactions are probed in more detail, measurements of binding thermodynamics in buffers with different ionization enthalpies will be an ever more important control. In the TCR–pMHC data summarized in Table 1, the influence of proton linkage has only been examined in two cases: recognition of p1049/HLA-A2 by the AHIII TCR and recognition of Tax/HLA-A2 by the A6 TCR. In the case of AHIII recognition, no evidence for proton linkage was found (Miller *et al.*, 2007). For A6 recognition of Tax/HLA-A2,

proton linkage resulting from a pK_a shift from 7.5 to 6.9 was found (Armstrong and Baker, 2007), resulting in a fourfold variation in binding enthalpy and a twofold variation in binding entropy depending upon buffer choice. Although the overall thermodynamic profile of the reaction was not changed upon correcting for the influence of proton linkage (i.e., correcting for proton linkage still yielded a TCR–pMHC interaction with favorable enthalpy and entropy changes), the availability of the “intrinsic” thermodynamics removed from the influence of protonation allowed for a much more detailed interpretation of the binding thermodynamics in terms of interface structure and protein conformational shifts.

HEAT CAPACITY CHANGES AND MOLECULAR FLEXIBILITY OR CONFORMATIONAL CHANGES IN TCR RECOGNITION

As noted above, interpretation of binding entropy changes in terms of conformational changes or reductions in molecular flexibility can be difficult. A clearer parameter for gauging the presence of changes in conformations is the heat capacity change, first applied to TCR–pMHC binding by Davis and colleagues in examining the binding of the 2B4 TCR to MCC/HLA-E^k (Boniface *et al.*, 1999). Heat capacity changes in protein binding reactions are strongly influenced by changes in solvation (Prabhu and Sharp, 2005), with the burial of hydrophobic surface contributing negatively to ΔC_p and the burial of hydrophilic surface contributing positively to ΔC_p (Murphy and Gill, 1991; Spolar *et al.*, 1992). Often the ΔC_p for a rigid body interaction can be reliably estimated from the change in solvent exposed surface area upon binding, and it has become routine to conclude that binding reactions which have ΔC_p values more negative than those estimated from the structure of the complex proceed with conformational changes that alter the solvent exposed surface area (although such conclusions are not uniformly accepted; see, e.g., Henriques *et al.*, 2000).

The reported heat capacity changes for TCR–pMHC interactions are generally large and negative, clustering between –400 and –800 cal/K/mol, although both larger (Garcia *et al.*, 2001; Krosggaard *et al.*, 2003; Ely *et al.*, 2006; Ishizuka *et al.*, 2008) and smaller (Garcia *et al.*, 2001; Krosggaard *et al.*, 2003; Gakamsky *et al.*, 2007; Mazza *et al.*, 2007; Ishizuka *et al.*, 2008) values have been reported (Table 1). As shown in Table 2, many ΔC_p values are substantially more negative than those calculated from the structures of the TCR–pMHC complexes. Thus, given the caveats that have been noted (particularly regarding the accuracy of van't Hoff determined heat capacity changes), conformational differences between bound and free TCRs may be common, if not ubiquitous (see Chen *et al.*, 2005 for a receptor that is nearly identical in the bound and free states).

But what are these conformational differences? Are they attributable to defined conformational changes or partial folding of CDR loops? As noted above, Davis and colleagues used heat capacity to dissect the binding entropy change for the recognition MCC/HLA-E^k by the 2B4 TCR, concluding that ~30 amino acids need to “fold” for binding to proceed (Boniface *et al.*, 1999). But as of yet, there is little evidence for highly flexible CDR loops occurring with regularity—the electron density in most crystallographic structures of unligated TCRs has shown the various CDR loops in clearly defined positions, the two exceptions

being the ELS4 and the 1.D9.B2 TCRs, for which missing electron density was observed in the crystal structures for CDR3 α (ELS4) or CDR3 β (1.D9.B2) (Tynan *et al.*, 2007; McBeth *et al.*, 2008). Until detailed spectroscopic assays of CDR loop dynamics are available, we may be best advised by Garcia's suggestion that TCR CDR loops are unlikely to be "easily accommodating limp noodles" (Garcia and Adams, 2005), instead adjusting their positions in more rigid conformational shifts. For the A6 TCR, analysis of the thermodynamic data suggest exactly this—after correcting for the influence of proton linkage, a thermodynamic dissection of the intrinsic binding thermodynamics showed that any TCR conformational rearrangements occurring in the A6 TCR upon binding Tax/HLA-A2 were likely to be *entropically driven* and *enthalpically opposed*, arguing against the ordering of highly flexible CDR loops upon binding (Armstrong and Baker, 2007).

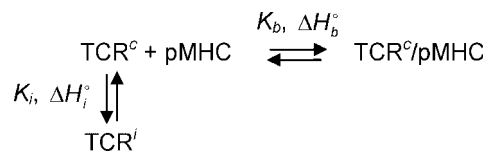
Unusually large negative heat capacity changes (greater than -1 kcal/mol/K) have been measured for a few TCR–pMHC interactions (Garcia *et al.*, 2001; Krosggaard *et al.*, 2003; Ely *et al.*, 2006; Ishizuka *et al.*, 2008), and these values have likewise been interpreted to result from conformational differences between bound and free receptor. However, using the two most widely used relationships between surface area and heat capacity (Spolar and Record, 1994; Baker and Murphy, 1998), a ΔC_p of -1 kcal/mol would require burial of between 2000 and 3000 \AA^2 of hydrophobic surface area. Even more buried surface would be needed if we consider the concomitant need to bury at least some polar surface along with apolar surface. To put these values in perspective, a ΔC_p of approximately -1 kcal/mol/K is on the order of that observed for the folding of small proteins. For example, cytochrome *c*, a 103 amino acid protein with a folding ΔC_p of -1.3 kcal/mol, buries 5039 \AA^2 of apolar surface and 3726 \AA^2 of polar surface upon folding (Murphy and Freire, 1992). While the very large TCR–pMHC binding heat capacity changes in Table 1 are provocative, it seems unlikely these can be attributed to changes in solvation resulting from small-scale conformational shifts in the antigen-binding site or changes in domain orientation. It may be useful to consider other sources for large TCR–pMHC-binding heat capacity changes such as linked equilibria, molecular strain, association/aggregation (Sturtevant, 1977; Guinto and Di Cera, 1996; Jen-Jacobson *et al.*, 2000; Prabhu and Sharp, 2005), or as discussed below, the difficulties of measuring heat capacity changes for weak-to-moderate affinity interactions by van't Hoff analysis, a point recently emphasized by Zhukov and Karlsson (2007).

Indeed, when considering TCR–pMHC-binding heat capacity changes, in all but one case the values have been determined by van't Hoff analysis. As ΔC_p is determined from the second derivative of ΔG° (or K) with respect to temperature, van't Hoff heat capacity changes can vary with the exclusion of only a single data point or even a change in the error associated with a data point (Zhukov and Karlsson, 2007). Given recent concerns regarding van't Hoff *enthalpies* and the eventual attribution of discrepancies between van't Hoff and calorimetric values to issues of data quality, statistics, and error analysis (Naghibi *et al.*, 1995; Chaires, 1997; Horn *et al.*, 2001, 2002; Mizoue and Tellinghuisen, 2004), van't Hoff-determined heat capacity changes should be interpreted even more cautiously. At this time, calorimetrically measured heat capacity changes are available only for the A6-Tax/HLA-A2 interaction (Armstrong and Baker, 2007). Notably, the calorimetric value of -330 cal/mol/K (uncorrected for linked protonation) is almost one-half of the van't Hoff value of -560 cal/mol/K (Davis-Harrison *et al.*, 2005)

(although the agreement is closer when the influence of proton linkage is accounted for, this agreement is fortuitous as discussed in Armstrong and Baker, 2007).

Although direct assays of dynamics are needed to clearly gauge the flexibility of TCR CDR loops, thermodynamics may still provide further insight into conformational differences between free and bound TCRs, particularly if the conformational differences arise due to a pre-existing conformational equilibrium in the unbound receptor as hypothesized by Holler and Kranz and recently observed by James and Tawfik in the SPE7 antibody (James and Tawfik, 2003; James *et al.*, 2003; Holler and Kranz, 2004). As noted below, as a recognition mechanism distinct from induced-fit (Koshland, 1958), conformational selection from pre-existing equilibria has received much attention in recent years and can be traced back to the Monod, Wyman, and Changeux model of allosteric regulation (Monod *et al.*, 1965).

Formally, we can describe a conformational selection binding reaction via the following scheme, where the horizontal reaction describes binding of pMHC to a binding competent state of a TCR (TCR^c) and the vertical reaction describes a conformational equilibrium in the TCR between binding the competent state and at least one other binding incompetent state (termed TCR^i). The terms K_b and ΔH_b° are the equilibrium constant and enthalpy change for pMHC binding, and the terms K_i and ΔH_i° are the equilibrium constant and enthalpy change for the TCR conformational shift (defined in the direction of TCR^c to TCR^i).



Taking into account the conformational equilibrium between TCR^c and TCR^i , the measured thermodynamics for pMHC binding the unligated TCR will be:

$$K_{\text{obs}} = \frac{K_b}{1 + K_i} \quad (1)$$

$$\Delta H_{\text{obs}}^\circ = -\Delta H_i^\circ \frac{K_i}{1 + K_i} + \Delta H_b^\circ \quad (2)$$

Equations 1 and 2 indicate that the observed affinity and binding enthalpy change are functions of not only the TCR–pMHC-binding reaction, but also the TCR^i to TCR^c transition. If the temperature dependencies of these two processes differ, the result would be an observed binding enthalpy change that varies nonlinearly with temperature; that is, a temperature dependent apparent heat capacity change.¹ The extent of this temperature dependence will depend upon the magnitude of the K_i equilibrium constant (i.e., how much of the TCR is incompetent vs. competent for binding pMHC) and the enthalpy and heat capacity changes associated with the conformational transition. If the conversion of TCR^c to TCR^i involves a significant conformational rearrangement, we would expect K_i to be greater than 1 (i.e., the binding competent state will

¹ This can be demonstrated formally by examining the temperature derivative of Equation 2, where the bracketed term is identical to the equation describing the excess heat capacity for two-state protein unfolding as would be measured by differential scanning calorimetry (Freire, 1995):

$$\Delta C_{p,\text{obs}} = - \left[\frac{\Delta H_i^{\circ 2} K_i}{RT^2(1+K_i)^2} + \Delta C_{p,i} \frac{K_i}{1+K_i} \right] + \Delta C_{p,b}$$

not be overly dominant in the TCR conformational ensemble) and large values for ΔH° . Provided sufficient enthalpy measurements are performed over a wide temperature range, the resulting nonlinearity in measurements of ΔH° versus temperature may be detectable in calorimetric measurements. A complete analysis of the data according to the equations above could allow for a full characterization of the binding reaction, including the equilibrium constant and enthalpy/entropy changes of the transition between the binding incompetent and binding competent states of the TCR.

The scheme described above does not account for motions occurring upon binding after initial association. In its simplest form, this mechanism can be represented as:



where the bracketed term represents an initial encounter complex that isomerizes to a higher affinity, fully bound state. This is a classical "induced-fit" mechanism as hypothesized by Davis and colleagues in the "two-step" binding model for TCR recognition (Wu *et al.*, 2002). Recently, Gakamsky and colleagues provided kinetic evidence for the operation of an induced-fit mechanism in TCR recognition of the pp65 peptide presented by HLA-A2 (Gakamsky *et al.*, 2007). Rapid kinetic experiments such as those employed by Gakamsky *et al.* can best identify the existence of such mechanisms in protein-binding reactions, as thermodynamic measurements report only on the differences between the free and most stable bound states. Thus, the temperature-dependent thermodynamic studies described above could not rule out the presence of induced-fit binding. This highlights the importance of using complementary approaches to investigate binding, particularly in the study of conformational dynamics and its role in recognition. Importantly, *a priori*, there is no reason why conformational selection from a pre-existing equilibrium and induced-fit could not both be operating in any given TCR-pMHC interaction, particularly if conformational dynamics and structural changes are not limited to the TCR as noted below. Notably, James and Tawfik recently demonstrated that the SPE7 antibody can recognize some ligands through a combination of both mechanisms (James and Tawfik, 2005), and recent computational studies suggest such hybrid recognition mechanisms may be common in protein-protein interactions (Grunberg *et al.*, 2004). Several recent studies and reviews address pre-existing conformational equilibria and induced-fit in protein-protein interactions in more depth (Tsai *et al.*, 1999; Kumar *et al.*, 2000; James and Tawfik, 2003; Goh *et al.*, 2004; Tobi and Bahar, 2005; Gunasekaran and Nussinov, 2007; Keskin, 2007).

Importantly, structural changes occurring upon TCR recognition of pMHC may not be confined to the TCR. At least three studies have shown conformational differences in the peptide between the free and bound complex. While two of these are small conformational shifts (Garboczi *et al.*, 1996; Lee *et al.*, 2004), recognition of an extensively bulged Epstein-Barr virus peptide presented by HLA-B35 by the ELS4 TCR results in a dramatic "flattening" of the peptide (Tynan *et al.*, 2007). Other structural studies have found peptides that adopt multiple conformations in MHC peptide-binding groove or disordered peptide side chains or backbones that in some cases become ordered upon TCR binding (Dessen *et al.*, 1997; Kuhns *et al.*, 1999; Hillig *et al.*, 2001; Sharma *et al.*, 2001; Speir *et al.*, 2001; Fremont *et al.*, 2002; Hulsmeyer *et al.*, 2004; Wucherpfennig, 2004; Gagnon *et al.*, 2006). Pohlmann *et al.* (2004) directly observed significant

conformational dynamics in a peptide presented by HLA-B*2709 that presumably are frozen out upon receptor binding. While it may be tempting to dismiss many of these peptide motions as inconsequential, Borbulevych *et al.* (2007) observed via both experiment and simulation that dynamics in a modified MART-1 peptide directly influence T-cell recognition. Importantly, many of the peptide conformational changes seen in peptide/MHC structures are of similar magnitude to those observed for TCR CDR loops. Thus, conclusions that complex binding data (whether thermodynamic or kinetic) result from conformational changes occurring solely in the TCR should be made cautiously in the absence of supporting structural information.

THERMODYNAMIC COMPARISONS AMONG MORE CLOSELY RELATED TCR-pMHC INTERACTIONS

The wide range of binding enthalpy, entropy, and heat capacity changes seen in Table 1 and Figure 2 reflects the diversity in TCR-pMHC interactions, and indicates that broad comparisons between very different interactions provides little information beyond highlighting the differences. Comparisons between more closely related interactions, on the other hand, can be more revealing. Two studies have compared the recognition of the same pMHC ligand by different TCRs: recognition of the MBP1-11 peptide presented by the class II MHC I-A^d by the 172.10 and 1934.4 TCRs (Garcia *et al.*, 2001) and recognition of the Tax peptide presented by the class I MHC HLA-A2 by the A6 and B7 TCRs (Davis-Harrison *et al.*, 2005). Both studies found considerable differences in the binding thermodynamics, revealing that a given pMHC molecule does not dictate any one TCR recognition mechanism. Colf *et al.* found similar results in comparing recognition of different pMHC by the same TCR: recognition of the QL9 peptide presented by H-2L^d and the dEV8 peptide presented by H-2K^b by the 2C TCR proceeded with opposing thermodynamics (Krogsgaard *et al.*, 2003; Colf *et al.*, 2007), indicating that a given TCR can recognize different ligands via distinct thermodynamic mechanisms. Malissen and colleagues observed the same when comparing recognition of pBM1/H-2K^b and pBM8/H-2K^b by the BM3.3 TCR (Mazza *et al.*, 2007).

More detailed results are available from studies comparing recognition by the same TCR of related or modified peptides presented by the same MHC. The first such study by Krogsgaard *et al.* revealed that subtle modifications to the MCC peptide could have unpredictable thermodynamic consequences for recognition by the 2B4 TCR (Krogsgaard *et al.*, 2003). The most dramatic reported was an unprecedented -3 kcal/mol/K shift in ΔC_p for a proline to phenylalanine substitution in a peptide variant (surprisingly, the changes in ΔS° and ΔH° were much more modest). These results were related to peptide immunological potency and the possibility of TCR flexibility and conformational change. In a second study, Lee *et al.* (2004) identified a modified HIV gag epitope incorporating a Phe \rightarrow Tyr substitution that, although being recognized by the G10 TCR with binding thermodynamics almost identical to the native peptide, adopted a different structure in the unligated MHC molecule, HLA-A2. These results imply compensatory changes in the activation energetics for TCR association and dissociation, which were related to the timing of the peptide conformational change along the receptor-binding pathway.

Miller *et al.* (2007) recently measured the thermodynamic consequences of the loss of a positive charge on the $\alpha 1$ helix in the interface between the AHIII TCR and the p1049 peptide presented by HLA-A2. Although complicated by the presence of a conformational shift in the CDR3 β loop, the calorimetrically measured loss in binding enthalpy was consistent with the loss of hydrogen bonds within the TCR–pMHC interface, providing insight into the distribution of energy within the AHIII-p1049/HLA-A2 interface. In addition to providing insight into individual interfaces, the approach of measuring the thermodynamic consequences of small changes can also be used to guide the engineering of TCR–pMHC interfaces, perhaps through the introduction of favorable interactions, elimination of unfavorable interactions, or the biasing of conformational equilibria.

By van't Hoff analysis, Ishizuka *et al.* (2008) recently examined the thermodynamic effects of 10 separate substitutions in the interface between the JM22 TCR and the influenza MP₅₈₋₆₆ peptide presented by HLA-A2. Although the thermodynamic measurements were not discussed in detail, all the mutations retain the overall thermodynamics of the wild-type interaction, perhaps not surprisingly as all the mutants characterized affected the binding free energy by ≤ 1 kcal/mol.

Finally, there is considerable interest in using thermodynamics to optimize affinity and specificity in drug design, perhaps the best published examples being inhibitors of HIV protease (Ohtaka and Freire, 2005; Lafont *et al.*, 2007). Similar approaches may be used in the optimization of affinity or specificity in TCR–pMHC interactions, perhaps for generating altered peptides designed to activate certain T-cell subsets or modifying TCRs to either direct immune responses against particular antigens or recognize antigens with high affinity for imaging or drug delivery purposes. Colf *et al.* (2007) have provided some initial data toward these goals, demonstrating that the m6 variant of the 2C TCR, generated via directed evolution (Holler *et al.*, 2000), achieves an approximately 100-fold enhancement in binding affinity though entirely enthalpic gains, offset by a shift from a favorable binding entropy change to an unfavorable binding entropy change. This result is in contrast with recent data indicating that the affinity maturation process can enhance antibody binding affinity by reducing the entropic cost for binding (Zimmermann *et al.*, 2006; Thorpe and Brooks, 2007), presumably by pre-optimizing the antigen combining site toward the structure of the antigen. Thus thermodynamics may provide a means to guide the design of TCRs with enhanced affinity in a fashion that mimics that used naturally by the humoral immune system.

CONCLUSIONS

From the first measurements in 1999 (Willcox *et al.*, 1999), the database of binding thermodynamics for TCR–pMHC interactions has grown to include 41 interactions as of this writing. Although early measurements suggested a thermodynamic “signature” for

TCR recognition consisting of a favorable enthalpy change and an unfavorable entropy change, the collective data clearly indicate that this is not the case. Rather, the binding thermodynamics vary considerably, reflecting the diversity in TCRs, pMHCs, and their interfaces. There is insufficient data to distinguish recognition of class I from class II pMHC, although if the class I data are a guide, we might expect recognition of class II pMHC to vary as broadly as class I. Although early thermodynamic data were interpreted as indicating a need for TCR CDR loops to organize or “fold” upon recognition of pMHC, the accumulation of more thermodynamic and structural data have led to a more refined view, where although CDR loop adjustments may occur, and these most certainly contribute to TCR cross-reactivity, the adjustments are not always of similar magnitude and seem more likely to consist of more defined structural shifts rather than the ordering of highly flexible backbones.

The variation in binding thermodynamics highlights one of the most remarkable aspects of TCR–pMHC interactions: despite the diversity in the molecules and interfaces, the interactions proceed with similar structural topologies, and, for the majority of activating ligands that have been investigated thus far, with similar affinities and kinetics. Whether this is due to selection mechanisms operating on a pre-selection repertoire (Buslepp *et al.*, 2003; Huseby *et al.*, 2006), a deeper self-recognition code not yet discerned from structural, physical, or genetic data (Feng *et al.*, 2007), or some combination of both is still unclear. As individual interactions are probed in more detail, thermodynamic measurements will provide further insights into the molecular driving forces behind the remarkable molecular recognition properties of TCRs. As the field evolves, calorimetric experiments should be strongly encouraged over van't Hoff measurements; new calorimeters with higher sensitivity and reduced sample requirements should facilitate this. As more probing experiments are performed, investigating and controlling for the influence of linked equilibria will be crucial. Application of other techniques such as rapid kinetics (Gakamsky *et al.*, 2007), computational chemistry (Michielin and Karplus, 2002; Gagnon *et al.*, 2005; Zoete and Michielin, 2007), and spectroscopic assays of loop dynamics in conjunction with thermodynamics and structural biology will be further illuminating.

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