

The molecular basis of TCR germline bias for MHC is surprisingly simple

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The elusive etiology of germline bias of the T cell receptor (TCR) for major histocompatibility complex (MHC) has been clarified by recent ‘proof-of-concept’ structural results demonstrating the conservation of specific TCR-MHC interfacial contacts in complexes bearing common variable segments and MHC allotypes. We suggest that each TCR variable-region gene product engages each type of MHC through a ‘menu’ of structurally coded recognition motifs that have arisen through coevolution. The requirement for MHC-restricted T cell recognition during thymic selection and peripheral surveillance has necessitated the existence of such a coded recognition system. Given these findings, a reconsideration of the TCR–peptide-MHC structural database shows that not only have the answers been there all along but also they were predictable by the first principles of physical chemistry.

By focusing the molecular ‘lens’ through which immunologists look at experimental questions, structural biology has had a considerable effect on immunology. There is a relatively high degree of structural sophistication among immunologists, as molecular recognition lies at the heart of immunology. However, although many complexes of T cell antigen receptor (TCR) and peptide–major histocompatibility complex (pMHC) have been published in the past 10 years¹, there is far from uniform agreement among structural biologists and immunologists about how to interpret this structural information^{2–12}. We believe the cause of most of this confusion is rooted in misunderstandings about very basic physicochemical aspects of molecular interactions between proteins. The intention of this perspective is to clarify some of these principles and to illustrate that both recent and previous structural data are actually very much in accord with functional biological observations. The main issue addressed here is often referred to as ‘TCR-MHC bias’ or ‘germline-encoded recognition’ of MHC. These terms are used to describe the original idea that TCR and MHC molecules coevolved a ‘preference’ to interact with one another¹³. Although this hypothesis has been supported by several cellular, biochemical and functional studies^{14–17}, the molecular basis of this phenomenon, and even its existence, remains

controversial^{3,10,12,18,19}. Here we attempt to provide a clear structural explanation of both the questions and the answers.

Unexplained features of TCR-pMHC interactions

Ten years after the first complex structures were reported^{20,21}, TCR-pMHC interactions still present one of the most interesting and enduring structural puzzles in biology, mainly because the binding interface is functionally and structurally segregated into four distinct components that collectively form composite surfaces²² (Fig. 1a,b). The pMHC-binding site of the TCR is segregated into invariant germline variable (V) gene–encoded components (complementarity-determining region 1 (CDR1) and CDR2) and components encoded by junctions between somatically rearranged variable-(diversity)-joining regions (CDR3). The MHC-binding site ‘visible’ to the TCR is also segregated into a mostly conserved helical scaffold, which presents within it a diverse array of peptides to the TCR. The TCR-pMHC recognition event ‘pairs’ invariant and variant structural components of the TCR and pMHC, in that the most variable regions of the TCR (CDR3) are positioned in the center of the binding interface where they contact the peptide, whereas the more conserved elements of the TCR (CDR1 and CDR2) and the tops of the MHC helices engage in contacts that surround the central CDR3-peptide region like a gasket. Thus, the structures of the individual molecules, together with their mode of interaction (the TCR ‘footprint’ on MHC), beautifully mirror the genetic makeup of the proteins involved, as originally predicted²². The idea that most of the binding interface (75–80%) involves contact between the germline-encoded CDR1 and CDR2 TCR regions and the MHC helices (Fig. 1b) seems logical, given that the TCR V-gene repertoire is specific for MHC molecules regardless of the peptide presented¹⁶. This ‘focus’ on the MHC may facilitate the rapid, TCR-mediated ‘scanning’ of MHC molecules^{15,23} to identify those presented peptides that stabilize the half-life of the TCR-pMHC complex sufficiently for signaling to occur.

Intriguing to biophysicists is the use of the term ‘bias’ to describe the apparent ‘preference’ of the $\alpha\beta$ TCR repertoire for engaging MHC molecules. In contrast, antibodies show no bias for a particular antigen and can be polyspecific²⁴. The ‘jury is still out’ for the $\gamma\delta$ TCR repertoire^{25,26}. Also fascinating is the idea that an entire repertoire of V genes encodes proteins that interact, at least in the context of an intact immune system, only with a particular structural class of ligands (MHC), which are highly diverse themselves. There are no other analogous protein-protein interaction systems in the mammalian genome that present such intriguing structural questions. What could be the structural solution to such a conundrum? Bias is a ‘fuzzy’ term for a biophysicist and is an idea about which immunologists tend to have highly polarized opinions.

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Published online 16 January 2008; doi:10.1038/ni.f.219

This need not be the case. 'Bias' is simply a euphemism for 'specificity', and specificity in protein-protein interactions is something that is well understood from the standpoint of structure and chemistry, although understanding of the relationship between structure and binding energetics is still a long way off. Thus, 'bias' can more correctly be referred to as TCR 'specificity' for MHC. That is, the natural TCR repertoire seems to physically react with MHC proteins. Given that the TCR is 'specific' for the MHC, then the most logical prediction, from first principles of physical chemistry, is that TCR V-domain CDR1 and CDR2 loops have evolved specific sets of contacts with residues found exclusively on MHC helices. Thus, the basis of MHC restriction is simply the coevolution of specific, conserved constellations of contacts between TCR V-gene products and MHC gene products.

However, it is not quite as simple as that. The TCR is faced with a considerable challenge, as it is specific for the MHC but is also 'cross-reactive' with many MHC molecules¹⁶. That is, each $\alpha\beta$ TCR can in principle recognize any MHC allele; there does not seem to be class (I or II) or allotype specificity by particular TCR germline segments as there is, for example, in natural killer cell receptors²⁷. This cross-reactivity is essential in that it enables the TCR to briefly dock and 'scan' the peptide contents of many different MHC molecules. Although there do seem to be TCR V_α and V_β TCR chain usage 'preferences' in certain immune responses²⁸, in general, the $\alpha\beta$ chains pair randomly, and practically any $\alpha\beta$ combination can be used to recognize most MHC molecules during an immune response. Given the combinatorial diversity of the $\alpha\beta$ chain repertoire (approximately 2,000–3,500 possible pairs), how is the TCR specific for only MHC and many MHC molecules at the same time? This question has captured the interest of structural biologists and immunologists alike.

Until recently, the database of crystal structures was apparently not very informative about an answer to this problem^{1,12,29} (Fig. 1c). Although they vary considerably in the details, the known TCR-pMHC structures share a roughly diagonal docking mode ($\pm 75^\circ$) with a uniformly stereotyped binding polarity, in which the V_α domain lies mainly over the amino-terminal end of the peptide and the $\alpha 2$ helix (MHC class I) or $\beta 1$ helix (MHC class II), and the V_β domain lies mainly over the carboxy-terminal region of the peptide and the respective $\alpha 1$ MHC helices. Notably, there have not seemed to be obviously conserved contacts between TCR V regions and MHC helices among the TCR-pMHC complexes; such a broadly conserved structural determinant might have constituted the 'smoking gun' of MHC 'bias'. Such a lack of conservation supports the idea that the prethymic TCR repertoire has unbiased, random specificity¹⁹ and that 'extrinsic influences', such as the antigenic peptides, constraints encountered during thymic selection³⁰, or coreceptors^{2,18} focus the docking orientations of the V-domain interactions on the MHC helices. This calls into question the idea that there is indeed a specific germline-derived structural origin of the TCR bias for MHC. The implication is that the V-domain contacts with MHC helices are an energetically continuous or flat landscape that forms as a secondary consequence of CDR3-peptide interactions and/or coreceptor-imposed steric influences.

Nevertheless, although the existing collection of TCR-pMHC structures has not shown conserved contacts, the conservation of the docking polarity (that is, V_α always lies over the $\alpha 2$ - $\beta 1$ helix, and V_β always lies over the $\alpha 1$ helix) is a clear indication, dating back to the first structures of complexes, of germline specificity between the V_α and V_β domains and particular helices of the MHC (Fig. 1c). Given such specific docking polarity, the answer to this problem, consequently, seems predictable on the basis of basic structural principles: as interatomic contacts in protein interfaces are specific and persistent, the contacts of CDR1 and CDR2 with the MHC helices represent germline-encoded energetic minima

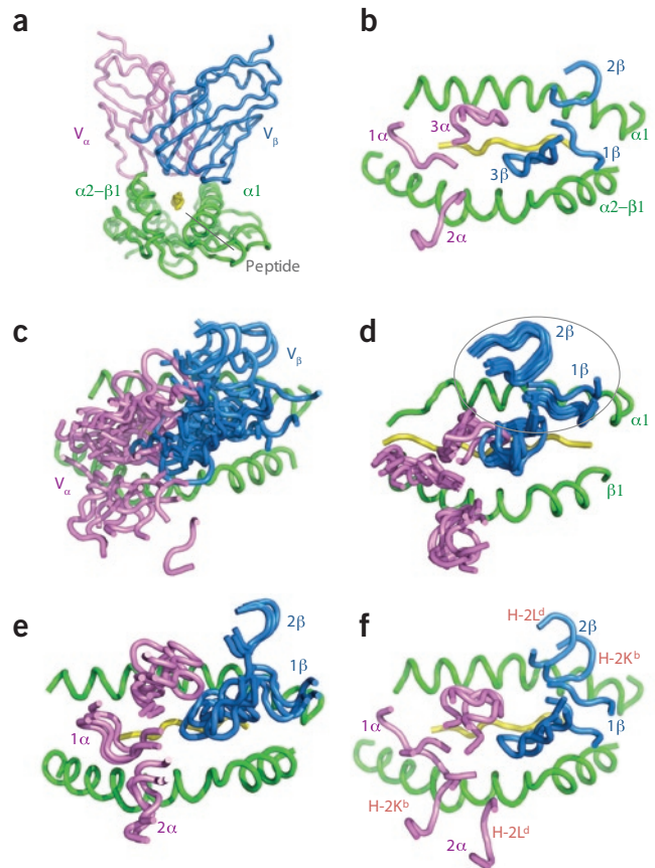


Figure 1 Divergence and convergence of TCR footprints on MHC molecules. (a) Interaction of $V_\alpha V_\beta$ with peptide-MHC, viewed down the MHC groove (Protein Data Bank accession number, 2CKB). (b) 'Footprint' view of **a** showing the stereotyped polarity of the V_α and V_β CDR loops on pMHC. (c) Convergent footprint polarity but diverse CDR loop positions in nine different TCR-pMHC complexes (Protein Data Bank accession numbers, 1A07, 1F00, 1J8H, 1KJ2, 1ZG1, 2NX5, 1MI5, 10GA and 1U3H). (d) Close superpositions (in circle) of the contacts of $V_\beta 8$ CDR1 and CDR2 with the I-A MHC $\alpha 1$ helix in six different TCR-pMHC complexes (Protein Data Bank accession numbers, 1U3H, 2Z31, 2PX1, 1D9K, 3C60 and 3C61). (e) Retention of similar germline-mediated contacts by the BM3.3 TCR with H-2K^b in three different peptide complexes (Protein Data Bank accession numbers, 1F00, 2OL3 and 1NAM). (f) Use of alternative 'codons' for interaction of the 2C TCR V_α and V_β with H-2K^b versus H-2L^d (Protein Data Bank accession numbers, 2CKB and 2O19). H-2K^b and H-2L^d (in red) adjacent to the respective loops indicate the positions of CDR2 α and CDR2 β in the structures.

('clicks', if thought of as analogous to a socket wrench) rather than slippery surfaces devoid of innate specificity. Thus, despite the 'blurred' structural data, from first principles, a TCR-MHC recognition 'code' seems likely to orchestrate germline bias⁹. For a better understanding of this, clarification of some basic ideas is in order.

Protein-protein interactions are governed mainly by van der Waals interactions, hydrogen bonds and charges (salt bridges)^{31,32}. Interactions between TCRs and MHC molecules are no different from any other protein-protein interactions in that their engagement is governed by the same first principles of molecular recognition. Protein-protein interactions generally bury several thousand square angstroms of surface area in the interfaces. As with all protein-protein interactions, most contacts in TCR-pMHC interfaces are mediated by van der Waals interactions^{1,7}, which are individually weak but, when present in large numbers across

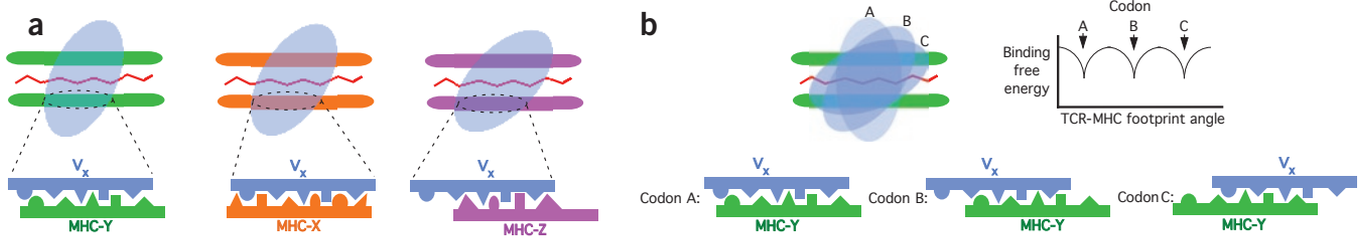


Figure 2 The 'codon hypothesis' for germline TCR-MHC interactions. (a) Each V-gene product (where 'V_x' is V_α or V_β) interacts with diverse MHC surface residues on the tops of the helices of different MHC molecules (Y, X and Z) by distinct yet specific mechanisms. That is, each CDR1 and/or CDR2 engages different MHC surfaces in diverse ways: the pairwise interactions that form each codon need not be shared by different MHC molecules. The CDR-MHC interface is presented as 'teeth' on the respective interacting surfaces. The same 'teeth' in a particular V_α or V_β CDR1-CDR2 engage different MHC surface structures (opposing 'teeth') in each complex in unique, highly specific ways. (b) An individual TCR germline V segment can engage one MHC molecule using several distinct codons (A, B and C) that are influenced by the interactions of CDR3 with the MHC-bound peptide. This is presented here as different docking geometries ('footprints') on the MHC that are mediated by the interaction of common residues on the TCR CDR1-CDR2 with different 'registers' of the MHC helix in each peptide complex. Inset, free energy diagram indicating that each footprint represents a low-energy binding solution ('click') rather than an energetic continuum.

broad interfaces, 'sum' to substantial binding energies. Hydrogen bonds are important in specificity because of their exquisite energetic dependence on the stereochemistry and geometry of the bond, as is also true for salt bridges³³. Proteins associate mainly because water is expelled from the interacting surfaces (the hydrophobic effect), which generates a gain in entropy³⁴. These 'desolvated', interacting surfaces then engage in many van der Waals contacts, whose specificity is enhanced by hydrogen bonds and salt bridges. Protein interfaces with good 'shape complementarity' (such as 'knob-in-hole' interfaces) maximize the number of van der Waals contacts and generally have higher affinity³⁵. Although TCR-pMHC interactions do bury large amounts of surface area, they seem to have relatively less optimal shape complementarity (dissociation constants in the micromolar range)³⁶ than do cytokine-receptor interactions (dissociation constants in the nanomolar range); this low shape complementarity no doubt contributes to the low average affinity of TCR-pMHC interactions. Of importance to the argument about germline bias is the fact that interatomic contacts in low-affinity protein-protein interactions are no less specific than those in high-affinity interactions³². Thus, the collection of interatomic contacts mediating the several thousand square angstroms of contact between TCR germline-derived regions and the MHC helices are probably not formed passively or secondarily as a result of 'steering' or 'steric exclusion' by a coreceptor or MHC-bound peptide, respectively.

While this may explain TCR specificity for one MHC molecule, how can the TCR cross-reactivity for so many different MHC molecules be explained? Is this not evidence of a lack of germline MHC specificity, given how diverse MHC alleles are? Again, no matter how cross-reactive, protein complexes bind each other only with highly specific, evolutionarily refined docking chemistries. Cases of true cross-reactivity are more correctly called 'polyspecificity'³⁷. There are many examples in the immune system of specific recognition by one protein of a spectrum of diverse ligands^{24,38-40}. The fact that the TCR has specificity for many MHC molecules does not indicate that this polyreactivity is due to a flat energetic landscape mediated by 'promiscuous' interactions between the V segments and the MHC helices. Structurally, the idea of 'promiscuity' is a complete misnomer and runs counter to the basic first principles of physical chemistry: there is no such thing as a 'promiscuous interface' from a structural chemistry standpoint. If there is a measurable binding event between two macromolecules, even very weak binding, it is mediated by specific interatomic contacts. In addition to mutational data that have identified focused energetic interaction 'hot spots' on MHC helices and CDR1 and CDR2 domains^{6,9,15,17,41}, evidence for the specificity of the interatomic contacts is that low-affinity protein complexes can be crystallized and structures can be determined to high

resolution. Crystal structure determination requires that hundreds of millions of different complexes in the crystal lattice be in the exact same structural arrangement to within fractions of an angstrom. Nonspecific complex interfaces simply cannot be crystallized because of the requirement that each complex in the lattice be an exact duplicate of every other molecule. Thus, were germline-encoded interactions between TCR and MHC nonspecific and merely a consequence of a loosely constrained shape complementarity, degenerate main-chain contacts or 'extrinsic factors', TCR-pMHC complexes simply could not be crystallized.

Clarification provided by recent observations

The fact that the database of approximately 20 TCR-pMHC crystal structures consists of complexes containing many different $\alpha\beta$ chains, peptides and MHC molecules has blurred and obscured any structural conservation (Fig. 1c). Recently, however, some clarity has emerged for this problem (Fig. 1d). One study focusing on the interaction of one particular germline V_β segment (V_β8.2) with a particular MHC molecule (I-A^A) has determined crystal structures of three V_β8.2 TCR-I-A^A MHC complexes⁹. Comparisons among these structures has revealed a notably coincident pair-wise interaction motif, which was termed a 'codon', between the V_β8.2 CDR1 and CDR2 loops, involving mainly Glu54 β , Tyr48 β and Asn31 β , contacting residues on the top of the I-A^A α 1 helix that are conserved across I-A alleles: Lys39, Gln57 and Gln61, respectively. This constellation of amino acids engages in hydrogen bonds, van der Waals interactions and charged contacts; notably, these interactions occur between amino acid side chains and are therefore specific contacts. Another study determined the structures of three additional V_β8 TCR-I-A^B complexes (two with V_β8.2 and one with V_β8.1) that, notably, were derived from mice with impaired negative selection⁴². Thus, these TCRs might represent 'preselection' TCRs and could demonstrate whether thymic selection 'filters' an unbiased TCR repertoire into one that binds MHC exclusively with the typical geometry. These complexes have shown binding geometries similar to 'conventional' TCRs, which suggests that these interfacial contacts are indeed 'engrafted' in the TCR germline before selection⁴². Even more notably, the complexes crystallized in these studies, together with a published D10 TCR (V_β8.2)-I-A^K complex, constitute a set of six different V_β8.2 TCRs and one V_β8.1 TCR in complex with I-A (Fig. 1d). Superposition of these complexes shows close convergence of the CDR1 β and CDR2 β contacts with MHC, which presents a very different picture than that of previous superpositions (Fig. 1c). Although there is nearly exact correspondence of the V_β contacts in four of the complexes, in two of them, the CDR1 and CDR2 interactions with MHC show some 'wobble' in the positions of side chains, as would be expected given the pairing of different peptides

and V_{β} with different V_{α} chains (as well as differing resolutions of the structures). Nevertheless, all of the complexes show a similar ‘knob-in-hole’ complementarity in which Tyr48 of CDR2 β occupies a depression on the α 1 helix of I-A; this interaction is then surrounded by a ‘halo’ of side chain-specific interactions. The coincidence is striking. The appearance of this V_{β} 8-interaction motif in the different structures with distinct MHC alleles (I-A^u, I-A^k and I-A^b), V_{α} gene segments (V_{α} 4.2, V_{α} 4.1, V_{α} 3.1 and V_{α} 2.3) and peptides (MBP1-11, ConA and 3K) is the strongest evidence to date that this interaction is germline ‘encoded’ at the level of pairwise amino acid contacts.

Further evidence supporting the idea of germline-encoded specificity is provided by crystallized complexes of TCR molecules bound to allogeneic MHC class I molecules. The 2C TCR (also V_{β} 8.2), together with several high-affinity variants of 2C generated by randomization of the CDR3 α , have been crystallized in complex with H-2L^d-QL9 (refs. 8,43). These studies have shown identical germline-encoded contacts between the CDR1 and CDR2 domains and the helices of H-2L^d in 2C and its variants. The importance of this experiment is that the selection of the high-affinity variants is done by yeast display in a cell-free system, in which the docking orientation of the TCR on H-2L^d is not influenced by coreceptors or any other extrinsic influences⁴⁴. After selection for high-affinity binding to H-2L^d-QL9 by yeast display, each of the CDR3 α loops of the variants has an entirely different sequence. In the complex structures, these CDR3 α conformations and QL9 peptide contacts have been substantially remodeled, yet the surrounding ‘gasket’ of germline contacts with MHC remains unperturbed. A similar CDR3-randomization experiment with a series of human MHC class I-specific TCRs has produced the same end result⁴⁵. Finally, a series of studies of the BM3.3 TCR bound to H-2K^b molecules presenting very different peptides has shown that the CDR1 and CDR2 docking positions with H-2K^b helices are mostly preserved, whereas the CDR3 contacts differ to accommodate the different peptides⁴⁶ (Fig. 1e). Notably, the germline contacts are not exactly similar but instead seem to adjust slightly to the different peptides. As with the V_{β} 8-I-A MHC class II examples, these results collectively indicate that germline-encoded interactions represent energetic minima (‘clicks’) that are not superceded or dictated by divergent peptide-CDR3 contacts.

Returning to the issue of germline-encoded cross-reactivity for different MHC molecules, it is also clear from the structural database that each V segment will have distinct interaction codons with each MHC (Fig. 2). That is, there does not seem to be a universally shared or ‘cryptic’ structural epitope on all MHC molecules that orchestrates MHC recognition by the TCR. Structural ‘mimicry’ by different MHC molecules is not needed to explain their shared ability to engage a single TCR V segment⁴⁷. Instead, a given MHC can use entirely structurally distinct codons to engage different V segments (Fig. 2b), as one V segment can engage different MHC molecules through entirely unrelated chemistries⁹. For example, V_{β} 8.2 has completely different binding modes on H-2L^d and on H-2K^b despite the high degree of amino acid conservation on the tops of the H-2K^b and H-2L^d helices⁸ (Fig. 1f). Furthermore, the amino acid pairwise interactions of the V_{β} 8.2-containing 2C TCR with H-2K^b and H-2L^d do not even remotely resemble the interactions of V_{β} 8.2 with I-A molecules^{9,42}. The structural database also shows that each V-segment type can use more than one codon with a given MHC haplotype (Fig. 2b). For example, two V_{β} 2-containing TCRs with H-2K^b presenting different peptides have been crystallized, showing that the V_{β} 2 docking footprints on the α 1 helix differ considerably^{48,49}; therefore, we interpret this as showing that the V_{β} 2 uses alternative codons for H-2K^b recognition. The peptide is probably involved in ‘editing’ germline-encoded TCR docking codons (‘clicks’) by selecting the most energetically and/or functionally optimal for that particular

peptide. In this way, the peptide is certainly involved in the final docking geometry in that it selects from a limited ‘menu’ of defined codon options (Fig. 2b).

The extension of the codon idea is that in principle, a table of interaction codons between each type of V_{α} or V_{β} segment and each MHC molecule could be constructed that would allow prediction of a docking footprint for any TCR-pMHC complex for which the V-gene segment usage and MHC type is known. The difficulty in this will no doubt be in understanding the nature and extent of the peptide-mediated editing of codon ‘choice’.

The coded nature of TCR-MHC interactions is a unique feature of T cell recognition not found in any other receptor-ligand system in the mammalian genome. In contrast, antibody-antigen interactions need not have coded interactions because the antibody is not restricted to binding antigen presented in a composite structural scaffold that is partly conserved and partly variable. Thus, the purpose of the coding in the TCR repertoire is to ensure that every TCR has some residual affinity for most MHC molecules for the purposes of positive selection and peripheral scanning. In addition, by ‘hard-wiring’ the TCR germline to recognize MHC helices and straddle the peptide in the MHC groove, TCR CDR3 domains are ideally positioned to ‘read out’ the centrally located peptide contents.

If the idea that the TCR is structurally specific for MHC is accepted, what can be made of results showing that T cells from mice lacking MHC class I, MHC class II, CD4 and CD8 (quadruple-knockout mice) can be activated by non-MHC ligands, apparently in the absence of coreceptors¹⁸? Does this prove that TCRs are not biased for MHC recognition? The suggestion of that paper¹⁸ is that the monomorphic coreceptors, which associate their cytoplasmic domains with the kinase Lck on T cells and physically engage their extracellular domains with MHC in the TCR signaling complex, lead to a misleading appearance of TCR bias for MHC. However, when coreceptors are removed, they can no longer sequester Lck away from the TCR and the TCR can now signal through engagement of non-MHC ligands. Thus, that study concludes that TCRs are being focused on MHC molecules merely by coreceptors (that is, an ‘extrinsic factor’)¹⁸. Surely a chief function of coreceptors is to focus the TCR on MHC and to limit TCR ligand recognition ‘options.’ However, from a purely structural standpoint, one would never say that the TCR can bind only MHC among the ‘universe’ of potential protein ligands in an artificial system. When taken out of its normal biological context (for example, through the removal of coreceptors and MHC ligands), the TCR simply becomes a binding protein, like an antibody, with a diverse ‘library’ of central CDR3 sequences. It is very likely that if this combinatorial ‘library’ of CDR3 domains is ‘panned’ against a broad spectrum of proteins on the surface of a mammalian cell, there will be reactivity. For example, a TCR reactive with the non-MHC ligand fluorescein has been reported⁵⁰. When an antigen-specific Fab fragment is displayed on phage with its CDR3 domains randomized, Fab fragments with binding specificities for practically any antigen can be recovered. Thus, the quadruple-knockout experiment is essentially the surface display of a variable antigen receptor ‘panned’ against naive antigens on mammalian cell surfaces. No detection of reactivity would have been a more unexpected result. The finding of interactions between TCRs and non-MHC molecules in this system does not indicate that the TCR repertoire is not biased in the germline toward MHC specificity.

Outstanding questions

While we now have ‘glimpses’ of codons of MHC restriction, many questions remain in order to refine this concept. For example, there are several cases of TCR-pMHC complexes that have minimal germline interactions yet still retain the stereotypical orientation and

polarity⁵¹; one particularly intriguing example is the 'superbulged' peptide complex that forms a 'triad' of MHC contacts⁵. Is this minimal triad sufficient to 'encode' germline bias for MHC? Will the codon idea apply to TCR interactions with nonclassical MHC molecules and/or CD1 (refs. 52,53)? These are all open questions. The structural basis of TCR-MHC recognition, like the structural foundations of most complicated biological processes, is a continuum of more or less 'canonical' binding modes, and surprises and exceptions to the 'rules' will no doubt appear. This perspective has highlighted our views of the most parsimonious explanations for how the system works based on fundamental principles of physical chemistry, now buttressed by structural results that have allowed us to reexamine the structural database with fresh eyes.

ACKNOWLEDGMENTS

We thank J. Kuriyan and D. Kranz for discussions. Supported by the National Health and Medical Research Council (CJ Martin Fellowship to L.K.E.), the Canadian Institute of Health Research (J.A.), N.I.H. (D.F., K.C.G.) and the Howard Hughes Medical Institute (K.C.G.).

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