Peptide Centric Vβ Specific Germline Contacts Shape a Specialist T Cell Response

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Certain CD8 T cell responses are particularly effective at controlling infection, as exemplified by elite control of HIV in individuals harboring HLA-B57. To understand the structural features that contribute to CD8 T cell elite control, we focused on a strongly protective CD8 T cell response directed against a parasite-derived peptide (HF10) presented by an atypical MHC-I molecule, H-2Ld. This response exhibits a focused TCR repertoire dominated by Vβ2, and a representative TCR (TG6) in complex with Ld-HF10 reveals an unusual structure in which both MHC and TCR contribute extensively to peptide specificity, along with a parallel footprint of TCR on its pMHC ligand. The parallel footprint is a common feature of Vβ2-containing TCRs and correlates with an unusual Vα-Vβ interface, CDR loop conformations, and Vβ2-specific germline contacts with peptides. Vβ2 and Ld may represent “specialist” components for antigen recognition that allows for particularly strong and focused T cell responses.

Keywords: MHC, TCR, elite controller, structure, germline contacts

INTRODUCTION

A key strategy for adaptive immune recognition in mammals is to generate enormous diversity by somatic rearrangements of antigen receptor genes during T and B cell development. While B cells can realize the full potential of this diversity, allowing them to recognize virtually any molecular structure, T cell recognition is highly constrained by the requirement for the presentation of antigenic peptides by Major Histocompatibility Complex (MHC) proteins, and the need for the T cell antigen receptor (TCR) to recognize both the antigenic peptide and polymorphic self-MHC proteins. How T cells achieve both broad coverage and high specificity, given the constraints imposed by MHC restriction, is a central question in T cell biology.

Part of the answer to this question may come from the binding orientation of the TCR on its peptide MHC (pMHC) ligand (1). In the vast majority of known TCR-pMHC structures, the TCR docks in an approximately diagonal orientation (2–4), such that the highly variable
complementarity determining (CDR) 3 loops, which are encoded by somatic rearrangement joints, are positioned primarily over the peptide, the most variable part of the pMHC ligand. In contrast, CDRs 1 and 2, which are germline-encoded within individual variable (V) gene segments, primarily contact the MHC α-helices that make up the sides of the peptide-binding groove with the TCR α chain positioned over the top of the MHCI α2 helix or MHCIÍ β chain near the amino-terminal end of the peptide, and the TCR β chain over the MHCI α1 helix or MHCIÍ α chain carboxy-terminal region of the peptide (2, 4–8). There is evidence that tyrosine residues within the TCR CDR1 and 2 loops help to impose this characteristic docking angle (9), although this remains controversial, and random selection models have also been proposed (2, 4, 5). Due to the large number of allelic forms of MHC, the impact of CDR3, and the flexible geometry of TCR-pMHC interactions, identifying conserved germline contacts is not straightforward, and requires comparing the same V segment TCRs complexed with multiple pMHC structures. Indeed, most of our current understanding of conserved germline contacts comes from mouse Vβ8 containing TCRs, and related Vβs in humans, which contain tyrosine residues within their CDR1 and 2s, and are the most represented TCR-pMHC structures in the Protein Data Bank (4, 9, 10). It remains unclear whether TCRs that use divergent Vβs segments have a similar docking orientation and germline contacts with pMHC.

In addition to the TCR docking orientation, the peptide binding characteristics of MHC also contribute to the broadness and specificity of antigen recognition. Peptides generally bind in an extended conformation within a deep groove of MHC, and the ability to bind to distinct allelic forms of MHC is largely determined by two or three peptide “anchor” residues. This arrangement ensures that peptide binding is sufficiently broad that a handful of MHC molecules in an individual can present peptides from virtually any pathogen, with the fine specificity for peptide determined largely by the TCR. On the other hand, there are indications that certain MHC-I molecules have more restricted peptide binding, and may contribute to strong CD8 T cell responses to particular pathogens (11). For example, the ability of certain individuals to control HIV infection without anti-retroviral therapy, termed “elite control”, is associated with HLA-B alleles (e.g., B27 and B57) with limited ability to bind peptides (12, 13). Restricted peptide binding by mouse H2-Ld is correlated with resistance to CD8 T cell exhaustion during chronic infection (14), and is controlled by amino acid polymorphisms that also correlate strongly with HIV control (12, 15). The paradoxical association between restricted peptide binding and elite control may be due to the combination of weak binding to self-peptides coupled with strong binding to particular antigenic peptides (13, 14). However, precisely why certain MHC-I molecules favor the development of particularly potent CD8 T cell responses remain a mystery.

The potent T cell response to the intracellular protozoan parasite, *Toxoplasma gondii* in resistant (H2d) mouse strains is dominated by CD8 T cells specific for a single peptide, HF10, (derived from the parasite protein GRA6) presented by Ld (16). The Ld-HF10 specific T cell response exhibits a number of similarities to CD8 T cell responses in HIV elite controller patients, including the lack of T cell exhaustion in the face of persistent infection and continuous production of armed effector T cells via a proliferative intermediate T cell population (17–19). This unusually potent T response may serve as a model for understanding CD8 T cell responses that underlie strong resistance to viral infection in certain individuals.

Here we show that the Ld-HF10-specific T cell response displays a focused TCR repertoire dominated by Vβ2. Crystallographic studies of a representative TCR (TG6) in complex with Ld-HF10 reveal an unusual parallel footprint on the pMHC complex, a feature that is also observed in other Vβ2 containing TCRs, and which promotes germline-encoded TCR contacts with bound peptide. In addition, the HF10 peptide binds tightly and with high complementary to Ld, in a conformation that optimizes peptide side chain interactions with both the MHC and TCR. Thus, this example of T cell elite control uses a strategy in which both TCR and MHC contribute substantially to peptide specificity and expands the prevailing view of T cell germline recognition. We discuss these results in terms of a model in which both Vβ2 and MHC-I Ld represent “specialist” recognition components that sacrifice broad coverage in order to provide unusually strong and focused responses to particular pathogens (20).

**MATERIALS AND METHODS**

**Animals**

B6 (C57BL/6) and B6.C (B6.C-H2d/bByJ) were obtained from The Jackson Laboratory (Bar Harbor, ME, USA). In order to monitor multiple *T. gondii* epitopes, F1 mice (B6xB6.C) expressing both the H-2b and H-2d MHC class I molecules were used for all experiments. For *in vitro* T cell activation assays, we used mice expressing a rearranged TCR transgene derived from a CD8 T cell clone specific for the HF10 peptide (HPGSVNEFD; corresponding to the C-terminus of the parasite protein Gra6) presented by mouse MHC class I molecule Ld (called TG6). The generation of TG6 mice was previously described (17). Six- to 10-week-old mice were used in all experiments. All mice were bred in the UC Berkeley animal facility and were used within the approval of the Animal Care and Use Committee of the University of California.

**Infection**

Mice were orally fed 70-80 cysts or injected intraperitoneally (i.p.) with 1x10^5 live tachyzoites from the type II Prugniaud-tomato-OVA strain (Pru) (21). This parasite strain harbors immunogenic T cell epitopes derived from the parasite proteins, GRA6 (HF10 peptide) (16), GRA4 (22), and ROP5 (23), and is engineered to express a red fluorescent protein (RFP).

**Flow Cytometry**

All antibodies were from eBioscience (San Diego, CA, USA), Biolegend (San Diego, CA, USA), or Tonbo (San Diego, CA, USA). All tetramers were obtained from the NIH tetramer facility.
(Atlanta, GA, USA). The tetramers were made by conjugating the biotin-labeled monomers with PE-labeled streptavidin (Prozyme, Hayward, CA, USA) according to protocols from the NIH tetramer facility. All flow cytometry data were acquired by BD LSR Fortessa analyzers (BD Biosciences) and were analyzed with FlowJo software (Tree Star, Ashland, OR, USA). Fluorescent AccuCheck counting beads (Invitrogen) were used to calculate the total numbers of live lymphocytes.

**Single Cell TCR Sequencing**
Mice were infected i.p. with the Pru strain of *T. gondii*. Splenocytes were harvested at 3 weeks post infection and Ld-HF10 tetramer positive CD8 T cells were single-cell sorted into 96-well plates. TCRα and TCRβ sequences were obtained by reverse transcription and nested PCRs as described (24).

**MHC I stabilization Assay**
RMA-S.Ld cells were obtained from N. Shastri (UC Berkeley). This assay was performed as previously described (25). In brief, RMA-S.Ld cells were incubated at 37°C, 5% CO2 for 8 h to saturate the culture medium with CO2, and then at room temperature overnight. The next day, cells were washed with PBS and plated at 3x10⁵ cells/well in a 96-W plate. Peptides of interest were added to the cells in serial dilutions. The plate was incubated for 1 h at RT and 3 h at 37°C. Cells were stained with the 30-5-7 antibody (specific for conformed, peptide-bound Ld) and a goat anti-mouse IgG phycoerythrin (PE)-conjugated secondary antibody and analyzed by flow cytometry.

**Analysis of T Cell Activation**
For analysis of the potency of HF10 peptide variants on TG6 T cell activation, RBC lysed splenocytes from TG6 TCR transgenic mice containing 10⁵ TG6 T cells were cultured in triplicate wells at 37°C in 5% CO2. HF10 peptide variants were added to the cells in serial dilutions. Samples were harvested 48 h later, stained for surface CD8, Ld-HF10 tetramer, CD25, and CD44, and then analyzed by flow cytometry.

**Vβ Usage of *T. gondii* Epitope-Specific CD8 T Cells**
F1 (H₂Kbx) mice were infected with Pru strain *T. gondii* parasites. Three weeks post infection, RBC lysed splenocytes were stained for surface CD8, peptide-MHC tetramer, CD44, and individual Vβs, and then analyzed by flow cytometry.

**Immunization With Peptide-Loaded Dendritic Cells**
Bone marrow-derived dendritic cells from male mice were incubated with 1μM of peptide for 3 hours at 37°C. Cells were washed and 5x10⁶ peptide-loaded dendritic cells were injected subcutaneously into naive F1 (H₂Kbx) female mice. Mice were sacrificed 7 days post immunization. Peptide sequences: HF10: HPGSVNEFD; ROP5: YAVANYFFL; GRA4: SPMNNGYYM.

**Protein Expression and Purification**
We used two systems to generate Ld-HF10. For biophysical and crystallographic studies of pMHC, we produced soluble Ld-HF10 by baculovirus-infected insect cell expression (26). The DNA encoding Ld (α1-α3) and HF10 (or alanine substituted variants) fused via a linker to β2m were cloned into pbac plasmid under polyhedrin and p10 promoters to produce secreted soluble Ld-HF10 from Hi5 insect cells. In this construct, Ld Tyr 84 and Gly at peptide p12 position from the linker that attaches β2m to pHF10 were mutated to cysteine to form a disulfide bond (27). Secreted Ld-HF10 in insect cell medium was captured with immunoaffinity chromatography and further purified by GE Healthcare FPLC Superdex 200 10/300 GL size exclusion column. For the crystallography of the TCR/peptide/MHC ternary complex, we produced the wild type Ld variable region (α1-α2). Ld (α1-α2) was expressed in E. coli BL21 as an inclusion body, solubilized in 8M urea, and refolded with synthetic HF10 peptide (HPGSVNEFD, ordered from Peptide 2.0 Inc.) (28, 29). Refolded Ld-HF10 was further purified with a HiLoad Superdex 200 26/600 size chromatography column.

TG6 α and β TCR chains were also produced by both baculovirus insect cell and bacterial expression systems. Acid-base leucine zipper stabilized, soluble TG6 molecules were produced in baculovirus-infected Hi5 insect cells and enzymatically biotinylated for SPR study. For the structural study, Vα and Vβ TG6 TCR sequences were fused to the pET30 vector with human αα chain as previously described (10). The TG6 α and β TCR vectors were transformed separately into E.coli BL21. TG6 α and β proteins inclusion bodies were solubilized, mixed, and refolded by dialysis. The refolded TCR was further purified with a HiLoad Superdex 200 26/600 size chromatography column, followed by a Mono Q ion exchange chromatography.

**Surface Plasmon Resonance Measurements**
Approximately 2000 RU of biotinylated TG6 TCR was captured in the flow cells of a BIACore streptavidin (SA) BIAsensor chip. Various concentrations of insect cell-produced Ld-HF10 and its mutated variant peptides were injected into the sensor chip and the association and dissociation kinetics were recorded and then corrected for the fluid phase SPR signal using the data from the biotinylated mouse BDC2.5 TCR. Kinetics was analyzed with BIACore BIAEval 4 software. Different SPR fitting models showed that the two-state reaction (conformational change) gave the best fit and better chi2 as shown in Supplementary Figure S8. We have repeated The BIACore experiments of TG6 TCR and WT Ld-HF10 twice, and the affinity calculation and curve fitting were very similar.

**Protein Crystallization**
All crystals for data collection were produced by the hanging-drop vapor-diffusion method. Crystals of Ld-HF10 were obtained at room temperature at a concentration of 7 mg/ml. The crystallization condition was 25% PEG 3350, 0.1M citrate pH5.5. TG6 TCR alone was crystallized at a concentration of 10 mg/ml at 4°C in 20% w/v PEG10K, 0.1M Sodium citrate tribasic dihydrate pH5.5 and 1M Lithium sulfate monohydrate. For the Td6/Ld-HF10 complex crystallization, refolded Ld-HF10 protein and TG6 TCR were mixed at a 1:1 molar ratio at a final concentration of 10 mg/ml in 11% w/v PEG 8K, 0.1 M MES 6.0, 0.24 M Ammonium sulfate.
Data Collection, Data Processing, and Structural Analysis

All diffraction data sets were collected at synchrotron beamline ID-24C at the Advanced Photon Source, Argonne National Laboratory using the Pilatus detector. Initial models were solved by molecular replacement. Data collection and refinement statistics are shown in Supplementary Table S2. The X-ray diffraction data were collected under liquid-nitrogen cryo conditions at 100°K. The protein crystals were flash-cooled in liquid nitrogen after a short soak in a cryo-protection solution consisting of the crystallization solution with 25% glycerol added. The data were indexed, integrated, scaled, and merged using the HKL2000 program (30), the structures were solved by molecular replacement method using Phaser (31) software and further refined by refmac5 (32), and rebuilding of the structure was performed by Coot (33). NCONT in CCP4 was used to analyze the atom-to-atom contacts between the TCRs and pMHC (34). Buried surface area (BSA) (Å²) is calculated using the PISA program from the CCP4 package (35). Graphical software and further refining were done using PyMol (Schrodinger, LLC). The atomic coordinates and structure factors have been deposited in the Research Collaboratory for Structural Bioinformatics Protein Data Bank, https://www.rcsb.org (PDB ID codes are shown in Supplementary Table S2).

TRAngle Determination and Comparison

TRAngle values of the TCR structures in the PDB were obtained from STCRDab on April 26, 2019. New structures presented in this manuscript are annotated using the same pipeline as all structures in STCRDab. The TRAngle was calculated using the protocol described in previous studies (36, 37). In brief, the algorithm uses a defined set of the most structurally conserved positions in both the Vα and Vβ domains. It then fits reference frames through interface positions and computes the deviation from the pivot axis, C. The length of C is the dc distance. BA describes a torsion angle between the Vα and Vβ domains. BC1 and AC1 are angles that describe the tilt, while BC2 and AC2 capture the twist, between the two domains (Figure 4C).

Docking Angle Calculation

The conventional method to characterize the docking angle of a TCR to pMHC, as described before (8, 38), fails to model the importance of the CDR loops in TCR-pMHC interactions. Furthermore, the conventional method does not allow for independent scrutinization of the TCR alpha (TRA) and beta (TRB) chains. To account for these shortcomings, we have developed an extension of the conventional docking angle to further characterize TCR-pMHC interaction geometry.

To model the TRA chain, the coordinates for the atoms within the CDR1/2α loops were selected and fit with linear regression. This line along with the center of mass for the CDR3 variable and joining regions of the TRA chain were used to define the equation of a plane where the TRA chain sits in space. The directionality of the TRA plane was defined as that the normal vector of the TRA plane faces the TRB chain. The TRB chain was modeled in the same manner as TRA, using the CDRb loops and center of mass of the TRB variable, diversity, and joining regions. For the TRB chain, directionality is defined so that the normal vector of the TRB plane faces the TRA chain. The TCR was modeled as a whole using a plane consisting of a linear regression through all atoms of all CDR loops and the center of mass of the variable region of the TCR. The direction of the TCR plane was defined by the direction of the normal vector, where the cross-product is taken from the CDRa’s to the CDRb’s.

The binding groove was modeled as a line using a linear regression of the coordinates of the alpha carbon atoms in the MHC helices that form the binding groove. The helix residues of the binding groove were defined by an alignment of IMGT MHC reference sequences to the structure. A1 helix residues for class I structures selected for the regression correspond to residues 50-85 of the IMGT nomenclature (38); for class II structures, residues 50-84 were selected. A2 helix residues correspond to residues 50-90. B1 helix residues correspond to residues 50-85. The directionality of the binding groove is defined by N-terminus to C-terminus.

The angle between the normal vector of a plane and a line is defined as follows:

\[ \cos \theta = \frac{m \cdot n}{|m||n|} \]

where \( \theta \) is the angle between the normal vector of the plane (m) and the line (n).

The angle between a plane and a line is the complement of the angle between the normal vector of the plane and the line. Therefore, when we let the angle between a plane and a line be represented by \( \varphi \):

\[ 90° - \theta = \varphi \]

\[ \cos \theta = \sin(90° - \theta) \]

\[ \cos \theta = \sin \varphi \]

\[ \cos \theta = \frac{m \cdot n}{|m||n|} = \sin \varphi \]

\[ \varphi = \sin^{-1} \frac{m \cdot n}{|m||n|} \]

The conventional docking angle and incident angle, as described by Rudolph, Stanfield, and Wilson (8), were obtained from the TCR3d database (https://tcr3db.ibbr.umd.edu/) for murine Vβ2 (PDB IDs: 1FO0, 1KJ2, 1NAM, 2OL3, 6X31, and 6DFS) and murine Vβ8 containing TCRs (PDB IDs: 4N5E, 3RDT, 3C6L, 3CSZ, 3RGV, and 6DFW). Additionally, the newly modeled TRA docking angle, TRB docking angle, and TCR docking angle were calculated as described for the sets of murine Vβ2 and Vβ8 containing TCRs with publicly available structures in PDB. To test for differences in angles between the Vβ2 and Vβ8 TCRs, Mann-Whitney U tests were performed with Bonferroni multiple testing corrections.
RESULT

The L\textsuperscript{d}-HF10 Specific T Cell Response Is Characterized by Early Activation, Strong Expansion, and a Focused TCR Repertoire

T. gondii infection of genetically resistant (H-2\textsuperscript{d}) mice elicits a potent CD8 T cell response directed against a single parasite-derived peptide, associated with the continuous production of armed effector T cells during chronic infection (16, 17). To examine the priming of this T cell response in more detail, we performed pMHC tetramer staining of splenic T cells during the 2 weeks following i.p. infection of F1(B6xB6.C) mice. Consistent with previous results, T cells specific for the GRA6 derived 10mer peptide HF10 presented by MHC-I L\textsuperscript{d} expanded \(>10^4\) fold, compared to a <100x expansion by the subdominant responses (Figure 1A). The strong expansion of L\textsuperscript{d}-HF10 specific T cells corresponds to a greater upregulation of activation and effector markers at day 5 post infection compared to subdominant T cell responses (Figure 1B). A similar expansion and immunodominance hierarchy of T. gondii epitopes was observed following oral infection (Supplementary Figure S1).

Previous studies have shown that the secretion pattern of the antigenic precursor protein and the C-terminal location of the HF10 epitope contribute to, but do not fully account for, the strong L\textsuperscript{d}-HF10 response (23, 25). To determine the impact of TCR-pMHC interactions, we examined T cell responses following immunization of naïve mice with peptide-loaded dendritic cells (DC) (Figure 1C). T cells specific for L\textsuperscript{d}-HF10 expanded 4000x upon peptide-DC immunization, whereas T cells specific for the other epitopes showed substantially lower expansion (Figure 1C). These data indicate that the interactions between TCR, peptide, and MHC, as well as parasite biology and antigen presentation, contribute to the potency of the L\textsuperscript{d}-HF10 T cell response.

We previously demonstrated that L\textsuperscript{d}-HF10 specific T cells from infected mice show preferential usage of V\textbeta2 (16). This preference for V\textbeta2 is also observed upon immunization of mice with DC loaded with the HF10 peptide, whereas T cell responses to other peptides displayed a V\textbeta profile that closely matched that of bulk CD8 T cells (data not shown, and Supplementary Figure S1). L\textsuperscript{d}-HF10 specific T cells from naïve mice also showed a V\textbeta2 frequency significantly above that of bulk CD8 T cells (Figure 1D) suggesting that a preference for V\textbeta2 is already evident after thymic selection and increases after T cell priming with the HF10 antigenic peptide.

To further investigate the TCR repertoire of the L\textsuperscript{d}-HF10 response, we sequenced paired TCR \alpha and \beta genes from 80 individual T cells from two different chronically infected mice, yielding 32 unique paired \alpha/\beta TCR sequences. We analyzed the unique sequences using the TCRdist algorithm (40). The majority of T cells used the TRBV1 segment (which encodes V\textbeta2) together with TRBJ2-1 and displayed a strong selection for a GRG motif in the TCR\beta CDR3 (Figure 1E). We also noted a preference for TCR\beta CDR3 length of 13 amino acids, a trend that was particularly prominent for V\textbeta2 containing T cells (Supplementary Figure S1). Finally, principal component analysis based on TCR distances showed a predominant cluster of similar TCRs which included the previously identified TG6 TCR (Figure 1F) (17). Indeed, the TG6 TCR\beta coding sequence from the original L\textsuperscript{d}-HF10 specific T cell hybridoma was found independently in the two additional mice examined, each time paired with a closely related TCR\alpha. Likewise, the TCR\alpha gene of the TG6 TCR was found independently in one additional mouse, paired with a closely related TCR\beta gene (Supplementary Table S1). These data indicate that the response to L\textsuperscript{d}-HF10 displays a highly focused TCR repertoire dominated by V\textbeta2. Moreover, TG6 is a V\textbeta2-containing TCR that provides a good representative of this response.

Unusual Peptide Conformation and Non-Armor Contacts Characterize HF10 Binding to L\textsuperscript{d}

Previous studies revealed unusual features of L\textsuperscript{d}, including a constrained peptide binding site and the requirement for non-anchor residues for optimal peptide binding (15, 41, 42). To characterize the binding of the HF10 peptide to L\textsuperscript{d}, we expressed recombinant soluble L\textsuperscript{d} molecules containing a covalently linked HF10 peptide and solved the crystal structure at 1.8 Å resolution. As described for other L\textsuperscript{d}-bound peptides, HF10 does not lie flat but instead bends within the peptide-binding groove to accommodate an obstruction formed by aromatic stacking interactions between elite-control associated residues including 97W in L\textsuperscript{d} (Figure 2A, Movie S1) (13, 15). However, while previously described L\textsuperscript{d}-bound peptides (all 9mers) have a bend at either p5 or p6 of the peptide (3, 15, 28, 42), HF10 displays pronounced bends at both p5 and p7 (Figure 2B, Supplementary Figure S2). Thus, the extra length of the HF10 peptide is accommodated by small bends in the peptide and a close fit with MHC, without the pronounced bulge outside of the MHC that is often observed with longer than optimal peptides.

In line with the published structures of L\textsuperscript{d}, p2P and p10F of HF10 are buried in the B and F pockets, respectively, and serve as anchor residues. The side chain of p6D interacts with the base of the groove, occupying the C pocket. In addition to these buried contacts, residues 4, 5, 7, 8, and 9 project to the sides of the groove (Figure 2B), with p5V and p9D making extensive contacts with the L\textsuperscript{d} \alpha1 helix, and p4S and p8F contacting the L\textsuperscript{d} \alpha2 helix. Because these peptide side chains project to the sides of the groove, portions of the residues are exposed to solvent, providing potential TCR contacts. Only a single amino acid side chain, that of p7E, is facing away from H2-L\textsuperscript{d}, and fully available to engage a TCR (Movie S1). The portion of the single-chain peptide-MHC corresponding to linker sequences was either disordered or located outside of the Ld peptide-binding groove, away from the canonical TCR–pMHC interaction surface as observed before (27).

To confirm the interactions between HF10 and L\textsuperscript{d}, we performed alanine scan mutagenesis of the peptide (Figure 2C). We measured binding based on the ability of peptides to stabilize surface L\textsuperscript{d} expression on the TAP-deficient RMA-S cell line (25). As expected, alanine substitution of the two anchor positions, p2P or p10F, abolished peptide binding, as did substitution at position 8. In addition, substitution at positions 1,
**FIGURE 1** | Characteristics of the L^4^HF10 specific T cell response. (A) T. gondii-specific CD8 T cells were quantified by pMHC tetramer staining and flow cytometry of splenocytes at different time points after intraperitoneal infection of F1 (B6xB6.C) mice. Fold change between naive and expanded T cells was calculated using the average number of tetramer+ cells in each population in naive mice (GRA6 = 74.4, ROP5: 654.9, GRA4: 1330). (B) Flow cytometric analysis of size (FSC or forward scatter) or expression of activation and effector markers (CD71, KLRG1, and CD25) on gated tetramer+ splenic CD8 T cells at day 5 post infection. (C) Mice were immunized with bone marrow-derived dendritic cells loaded with the indicated peptides. Expanded tetramer+ CD8 T cells were quantified by tetramer staining of splenocytes 7 days post immunization (open circles). Numbers of tetramer+ CD8 T cells in the spleen were quantified by tetramer enrichment of naïve mice and were used to calculate the fold expansion of each antigen-specific T cell population (closed circles). Statistical significance of differences in fold change between the three groups was calculated using Mann-Whitney tests. GRA6 vs. GRA4: p < 0.0001, GRA6 vs. ROP5: p < 0.0001, GRA4 vs. ROP5: p=0.0002. Statistical significance between tetramer+ T cell populations was calculated by two-way ANOVA. The interaction p-values are as follows: GRA6 vs. GRA4: p < 0.0001, GRA6 vs. ROP5: p < 0.0001, GRA4 vs. ROP5: p=0.99. (D) Top-scoring CDR3^b^ motif. Results of a CDR3 motif discovery algorithm are shown using a TCR logo that summarizes V and J usage, CDR3 amino acid enrichment, and inferred rearrangement structures. The bottom panel shows the motif enriched by calculating against a background dataset of non-epitope specific TCR sequences. (E, F) Principal components analysis (PCA) projection of the TCRdist landscape colored by V_a (left panel) and V_b (right panel) gene usage. The groups of TCRs that correspond to the top scoring CDR3^b^ motif are indicated with a dashed circle, and the TG6 TCR is indicated with an arrow.
5, 6, 9 substantially reduced binding. These data are consistent with the 3D crystal structure and confirm the close complementarity between peptide and MHC molecules, including non-anchor residue contacts.

An Unusual TCR Footprint and a Dominant Peptide Contact Characterize TG6 TCR Interaction With the Ld-HF10 Complex

To investigate the structural features underlying the potent T cell response to Ld-HF10, we solved the crystal structure of the TG6 TCR bound to the Ld-HF10 complex. We prepared soluble TG6 TCR by expressing and refolding the extracellular domains as described (10). We used a refolded version of Ld-HF10 consisting of the a1 and a2 domains with five mutations to improve its stability, and with the original tryptophan at position 97 to preserve HF10 peptide binding (3) (data not shown). We co-crystallized the refolded Ld-HF10 and TG6 at a 1:1 ratio and solved the complex structure at a resolution of 2.5Å. The electron densities of HF10 peptide are well-defined, and the structures of covalent and the non-covalent Ld-HF10 complexes are almost identical (Supplementary Figure S9).

In the majority of reported TCR-pMHC structures, the TCR has a characteristic diagonal docking orientation, in which CDR3 of the TCR α and β chains are positioned over the peptide, CDR1 and 2 of TCRβ are positioned over the MHC α1 helix, and CDR1 and 2 of TCRα are positioned over the α2 helix of MHC-I (or the β1 helix of MHC-II) (1, 4, 43). In contrast to this consensus, TG6 displays an unusual footprint, in which all the TCR β CDR loops shift toward the Ld a2 helix. As a result, there are 121 TCR contacts to Ld a1, but only 23 contacts to Ld α2 (Table 1, Figure 3A). The buried surface between TG6 and Ld-HF10 is 1131.8 Å²; relatively small compared to the known TCR-pMHC-I complexes (1).

The TCR interaction with the HF10 peptide is centered around TCRβ chain CDR3 contacts with p7E: the only peptide side chain that points away from the MHC molecule (Figure 2). This interaction involves a complex hydrogen bond network with the CDR3 backbone (Figure 3B). The close wrapping of the CDR3 loop around p7E is consistent with the conserved length of 13 amino acids, and two strongly selected glycine residues in the consensus TCRβ CDR3 determined by TCR sequencing (Figure 1E, Supplementary Figure S1, Supplementary Movie S2). Additional TCR contacts are formed with peptide residues that project toward the sides of the MHC groove and are sandwiched between the Ld α1 and α2 helices and the TCR.

FIGURE 2 | Features of the antigenic HF10 peptide bound to H2-Ld. Soluble H2-Ld containing a covalently linked HF10 antigen peptide was crystallized and the structure was solved at 1.8 Å (Experimental Procedures and Supplementary Table S2). (A) The electrostatic surface charge of the Ld molecule (with bound HF10 peptide) is shown colored by the relative charge of the surface atoms (red - negative and blue - positive). A stick representation of the HF10 peptide is colored as: carbon, green; oxygen, red; nitrogen blue. The location of W97 is indicated by an arrow. (B) Conformation of the bound HF10 peptide (green with blue side chains) in comparison to other Ld-bound peptides. Peptides with a bend at p5 (HF10 and 2O19 in pink) are on the left, and peptides with a bend at p6 (p7 for HF10 (HF10 and STJH in white) are on the right. See Movie S1. Additional Ld-bound peptides are shown in Supplementary Figure S2. (C) H2-Ld binding to HF10 peptide alanine substitution variants. Flow cytometry surface expression (MFI) of Ld on TAP-deficient RMA-S-Ld cells incubated with increasing concentrations of the indicated HF10 or peptide variants. Data are representative of three independent assays.
TABLE 1 | Contacts between TG6 TCR and Ld-pHF10.

| V Domain | CDR Loop | Amino to | Ldx1 | HF10 | Ldx2 | Ligand | Amino to | Vα | Vβ | TG6 | Ld-pHF10 |
|----------|----------|----------|------|------|------|--------|----------|-----|-----|------|---------|
|          |          | Atom     |      |      |      |        | Atom     |     |     |      |         |
| Vα       | CDR1     | 29Y      | –    | –    | 50   | Ldx1   | 62R      | 9   | –   | –    | α29Y    | 166E    |
|          | CDR2     | 50S      | –    | –    | 3    | –      | 66V      | 7   | –   | –    | α51R    | 161E    |
|          |          | 51R      | –    | –    | 27   | –      | 73W      | –   | 7   | –    | α65T    | 62R     |
|          |          | 54E      | –    | 5    | 1    | –      | 79R      | –   | 1   | –    | α66G    | p5V     |
|          | CDR3     | 95T      | 10   | 17   | –    | –      | HF10     | p4S | 11  | –    | α98N    | p5V     |
|          |          | 96G      | 5    | 17   | –    | –      | p6V      | 20  | 2   | 2    | β28Q    | 149Q    |
|          |          | 97A      | 1    | 7    | –    | –      | p6N      | 2   | 6   | 6    | β50R    | p9D     |
|          |          | 98N      | 1    | 17   | –    | –      | p7E      | 8   | 67  | 67   | β96R    | p6N     |
| Vβ       | CDR1     | 28Q      | –    | –    | 13   | –      | P8F      | –   | 15  | –    | β96R    | p7E     |
|          |          | 31W      | –    | 3    | –    | –      | p9D      | –   | 14  | –    | β96R    | p9D     |
|          | CDR2     | 50R      | –    | –    | 1    | –      | Ldx2     | 149Q| –   | 13   | β98G    | p7E     |
|          |          | 51S      | –    | –    | –    | –      | 151G     | –   | 3   | –    | β99Y    | p7E     |
|          | CDR3     | 94A      | –    | 6    | –    | –      | 154E     | 1   | 7   | –    | –       |         |
|          |          | 95G      | –    | 12   | –    | –      | 155Y     | –   | 17  | –    | –       |         |
|          |          | 96R      | 6    | 44   | –    | –      | 157R     | 3   | –   | –    | –       |         |
|          |          | 97G      | –    | 9    | –    | –      | 158A     | 12  | –   | –    | –       |         |
|          |          | 98G      | –    | 9    | –    | –      | 161E     | 12  | –   | –    | –       |         |
|          |          | 99Y      | –    | 14   | 27   | –      | 162G     | 3   | –   | –    | –       |         |
|          |          | 100A     | –    | 8    | –    | –      | 163E     | 32  | –   | –    | –       |         |
|          |          |          |      |      |      |        |          |     | 166E| 167W| 170R   | 1       |
| Total    |          |          | 23   | 146  | 121  | 138    | 152    |     |     |     |         |         |

FIGURE 3 | Features of TG6 TCR bound to Ld-HF10 complex. (A) TCR footprint on the solvent-accessible surfaces of the Ld-HF10 complexes (α, cyan; Ld₂, magenta; peptide, white). The TCR CDR loops are colored as TCR footprint. Areas of TCR contact with pMHC (≤ 4.5Å) are colored as: CDR1α, red; CDR2α, blue; CDR3α, yellow; CDR1β, gray; CDR2β, orange; CDR3β, green. (B-E) Interactions between TG6 CDRs and Ld-HF10. HF10 residues are shown in white carbon stick; residues on TG6 are shown in pale yellow carbon stick; residues of Ld are shown in magenta (α1) and cyan (α2) carbon stick; H-bonds and salt bridges are indicated by green lines. (B) Extensive contacts between TCR CDR3β and p7E of the HF10 peptide. (C) Residues of both TCR CDR3α and Ld contact HF10 p5V. (D) Hydrophobic stacking of TCR CDR3β and Ld with HF10 p8F. (E) Both TG6 CDR2β and CDR3β form a salt bridge to p9D. Ld also provides contacts. See Movie S2.
residues (Figures 3C–E). These include p5V: which forms hydrogen bond interactions with TG6 CDRx3 via its main chain N and O atoms, as well as van der Waals contacts with its aliphatic side chain (Figure 3C), p8F: which makes van der Waals contacts with CDR3β (Figure 3D), and p9D: which is contacted by 96R of CDR3β and 50R from CDR2β (Figure 3E, Movie S2). Interestingly, 96R is a prominent part of the enriched motif (GRG) in CDR3 of TCRβ (Figure 1D), implying that Lα-HF10 specific TCRs are highly selected to preserve this interaction.

Measurements of TCR binding to pMHC by surface plasmon resonance are in good agreement with the ternary complex structure. The affinity for the wild-type peptide is at the high end of the range reported for TCRs (0.4 μM) (Supplementary Figure S3) (44). In addition, alanine substitution of p6N, p7E, p8D, and p10F all greatly reduce both TCR binding and T cell activation (Supplementary Figure S3). Thus, the C-terminal peptide residues (p6-10) are all crucial for TCR recognition, with p7E contributing exclusively to TCR contacts, and the remaining residues affecting both MHC and TCR interactions (Figure 2C, Supplementary Figure S3).

The overlay of the Lα-HF10 structure before and after engagement of the TG6 TCR shows that, while most of the Lα-HF10 surface changes little upon TCR binding, there is a shift in the side-chain rotamers of Lα 155Y and 62R. The side chain of 155Y rotates to point toward the HF10 peptide in the binding groove (Supplementary Figure S3), allowing the 99Y from TG6 Vβ CDR3 to contact the Lα α2 domain via hydrogen bond and van der Waals interactions (Table 1). On the other hand, Lα 62R rotates toward the TCR Vα CDR1, allowing the 27S from TG6 to form a hydrogen bond with the Lα α2 domain. The large rotamer changes from the Lα 155Y and 62R upon TCR binding is consistent with the two-state binding observed in SPR of Lα-HF10 binding to TG6 (Supplementary Figure S3), indicating an initial low affinity binding step, followed by a conformational change leading to a more stable complex. Similar second-order kinetics are also observed in HF10 peptide variants (Supplementary Table S3).

**Vβ2-Specific Germline Contacts Generate Strong Protection**

To understand the basis of the unusual Vβ2 footprint of the TG6 TCR on Lα-HF10, we compared the structure of TG6 TCR to Yae62 TCR (PDB: 3RGV), a Vβ8-containing TCR that binds to Kβ/pMW with a classic docking orientation and footprint (10). TG6 and Yae62 TCRs use the same Vα4 segments, facilitating a comparison of the impact of the different germline-encoded Vβ segments. As expected, the 29Y of TG6 Vα4 shared the conserved germline contacts with MHCs (Supplementary Figure S6). We noted a shift in the positions of the TCRβ CDR loops of TG6 relative to Yae62, leading to the repositioning of the loops away from the MHC α1 helix, and toward the peptide and MHC α2 helix (Figure 4A). To explore the basis for the shift in CDR loops, we superimposed the TG6 and Yae62 structures (Figure 4B, Movie S3). We noted a clear difference in the conformation of the TCRβ CDR1 and 2 loops, with kinks in the Vβ2 CDR1 and 2 loops due to proline residues at positions 30 and 52. While CDRs are normally flexible to accommodate the antigens, residues 30P and 52P of Vβ2 likely limit the flexibility of CDR loops due to the confined phi angle of proline (45) and contribute to the re-positioning of the loops toward the peptide and MHC α2 helix. Similar CDR1 and 2 conformations are observed in three other Vβ2 containing TCRs from six different structures (Supplementary Figure S4). Consistent with conformational constraints imposed by proline residues, the average root mean square deviation (RMSD) of the Vβ2 CDR1 and are only 0.421 and 0.277, respectively, compared to a RMSD of around 1 for the CDR1/2 loops of Vβ8 (46).

Another striking difference between TG6 and Yae62 TCRs is the shift in the position of the Vβ domain relative to Vα (Figure 4B). To quantify the difference in the Vα-Vβ domain interface, we used a method called TRangle, which defines variations in the geometry of Vα-Vβ interface based on one distance and five angle measurements (36). Interestingly, TG6, as well as the other Vβ2 containing TCRs in the Protein Data Bank (https://www.rcsb.org/), have an unusually low DC1 distance, and an unusually high BC1 angle compared to other published mouse TCR structures, most of which use Vβ8 (Figure 4C). Moreover, these parameters do not show any obvious correlation with the TCRα usage of these same TCRs (Supplementary Figure S4). Thus, both an altered Vα-Vβ interface and the conformation of the CDR1 and 2 loops contribute to the shift in the TG6 footprint toward the MHC α2 helix.

Given the striking difference between the CDR1/2 loop conformations and Vα-Vβ interface in Vβ2- compared to Vβ8-containing TCRs, we considered that the unusual footprint of TG6 on pMHC might be a common feature with other Vβ2-containing TCRs. Superimposing the TCR footprint of three additional Vβ2 TCRs from five different structures onto pMHC revealed a similar shift in the TCRβ contacts toward the peptide and MHC α2 helix compared to Vβ8 TCRs (Figure 5). Typically, TCR docking angles on pMHC are calculated using the positions of conserved V domain cysteines to determine the TCRαβ axis. Since this approach would not capture changes in the Vβ2 footprint due to the unusual CDR2 and 3 conformations, we defined a new parameter that we call the “footprint angle”. First, we selected the CDR residues involved in binding pMHC (using a contact radius of 4.5 angstroms, Table 1). We then defined a TCR vector between the two mass centers of the Vα and Vβ CDR contact regions and calculated the angle between the TCR vector and the peptide vector (defined by the position of the α carbons from residues p1H and p10F). The TCR vectors of all Vβ2 TCRs were relatively parallel to their peptide vectors (footprint angle 14.5 - 27.7 degrees), compared to relatively diagonal TCR vectors for a set of Vβ8 containing TCRs (footprint angle 37 - 53 degrees) (Figure 5). Thus, the shift in the TCRβ footprint toward a parallel binding orientation on pMHC appears to be a conserved feature of Vβ2 containing TCRs.

In addition to the footprint angle measurement, we also defined a new set of docking parameters that allows us to separately measure and quantify the impact of germline encoded CDR1/2 and the TCRα and β chains on the angle of TCR docking on pMHC (Supplemental Figure S5 and Methods). In agreement with our
footprint angle calculation, the conventional docking angle (TCR3d_DA) as well as the germline orientation of the TCR and TRB chain to the binding groove (TRB_germ, TCR_germ) were determined to be different between Vb2 and Vb8 bearing TCRs. Based solely on the conventional docking angle, this parameter would suggest that Vb2 takes on a less steep angle to the pMHC surface than Vb8. Interestingly, a different story is revealed when TRB is examined in isolation to better characterize its orientation. Vb2 demonstrates a steeper angle to the pMHC surface than Vb8; this difference can be attributed specifically to the orientation of germline components, CDR1b and CDR2b. This information also suggests that differences in TCR orientation are largely influenced by the TRB germline angles, as neither component of the TRA chain angles is different between the two groups of TCRs. Parameters determined to be statistically insignificant were the conventional incident angle (TCR3d_IA), the angle between the TRA chain and the binding groove (TRA), the germline orientation of the TRA chain to the binding groove (TRA_germ), the angle between the TRB chain and the binding groove (TRB), and the angle between the TCR and the binding groove (TCR) (Supplementary Figure S5).

It has been proposed that the footprint of TCR on pMHC is influenced by germline-encoded contacts, which may differ between particular Vβ segments (4). We compared the interactions of CDR1 and 2 with pMHC in TG6, and three other unique Vβ2 containing TCRs (Figure 6). In all four structures, germline-encoded residue 28Q from the Vβ2 CDR1 contacts the α2 helix of MHC-I or the equivalent β1 helix of MHC-II (Figure 6A, Table 1). In addition, 50R from the Vβ2 CDR2 contacts both the α1 helix and the peptide in each of the structures. Interestingly, while the aliphatic portion of 50R contacts α76V of MHC-I or α67A of MHC-II, the amino group forms a salt bridge with an acidic residue of the bound peptide in three out of four of the structures (Figure 6B, Table 1). The previously described Vα germline contact between tyrosine at position 29 with the α2 helix of MHC-I is preserved in TG6 (Supplementary Figure S6), consistent with the similar TCRα footprint for Vβ2- and Vβ8- containing TCRs on pMHC (Figure 4A). Thus, Vβ2 specific germline contacts are associated with a shift in the TCRβ footprint that leads to a parallel footprint angle and conserved germline contacts with peptide.

The footprint of TCR on pMHC is

FIGURE 4 | Structural differences between Vβ2 and Vβ8 containing TCRs lead to an altered footprint for pMHC binding. (A) A comparison of the positions of CDR loops from TG6 and YAe62 TCRs over their pMHC ligands. CDR loops from TG6 are shown in blue and Yaee62 are in orange. (B) Ribbon diagrams of TG6 TCR (Vβ2) and YAe62 TCR (Vβ8) overlaid with their TCRα chains aligned (both Vα4 encoded by TRAV6). Note that there is a shift in the juxtaposition of TCRα and TCRβ domains that contributes to the shift in the position in the CDR loops of TCRα relative to TCRβ. In addition, proline residues in the CDR1 and CDR2 loops of Vβ2 lead to a further shift in the CDR1 and 2 loops away from the α1 helix of MHC and toward the peptide and α2 helix of MHC-I (or β1 helix of MHC-II). See Movie S3. (C) TRangle parameters dc distance and BC1 angle for the TG6 TCR (indicated by arrow) compared to non-redundant TCR structures in the PDB. Vβ2 TCRs are shown in blue and Vβ8 TCRs are in orange. Right panel shows TRangle parameters used to define the Vα-Vβ interface geometry superimposed over a ribbon diagram of TCR.
FIGURE 5 | Footprint for binding of Vβ2 and Vβ8 containing TCRs to pMHC. The TCR footprints for four different Vβ2 TCR/pMHC complexes (A) and six different Vβ8 TCR/pMHC complexes (B) are shown, along with footprint angles, calculated based on a vector for the peptide (grey line) and a vector for the TCR-pMHC contact regions (black line) as described in the text. The center of the TCRα and TCRβ footprints are indicated by dots. Structures are: TG6 TCR binding to Ld-HF10 (PDB: 6X31); BM3.3 TCR binding to Kβ-pBM1 (PDB: 1FO0); KB5-C20 TCR binding to Kβ-pKB1 (PDB: 1JK2); TCR I29 binding to IAg7-insulin B:9-23 (PDB: 5JZ4); TCR Yae62 binding to Kb-pWM (PDB: 3RGV); TCR 42F3 to Ld-pCPA12 (PDB: 4N5E); TCR 2w20 to IAb-3k (PDB: 3C6L); TCR Yae62 to IAb-3k (PDB: 3C60); TCR ANI2.3 to DR52c-pHIR (PDB: 4H1L); TCR D10 to IAα-pCA (PDB: 1D9K). (C) Summary of all calculated Vβ2 and Vβ8 TCR footprint angles. Statistical significance was determined by a t-test (***p < 0.001).
DISCUSSION

It has been proposed that conserved germline contacts between TCR CDR1/2 residues and MHC help to impose the characteristic diagonal footprint of TCR on pMHC, although this remains controversial (2, 4, 5, 47). While investigating the structural basis of a potent CD8 T cell response to the parasite antigen HF10, we noted that the L^4-HF10 specific TCR TG6, as well as other Vß2-containing TCRs, adopt a parallel footprint on pMHC due to an unusual Vß-Vß domain interface and TCRß CDR1 and 2 loop conformations. This parallel footprint corresponds to a distinct set of conserved Vß2-specific germline contacts, including one between CDR2 and the peptide. Thus, Vß2 represents “the exception that proves the rule” and solidifies the concept that conserved germline-encoded contacts help to define the binding orientation of a TCR on its pMHC ligand.

Why has it taken more than a decade since the discovery of MHC-centric germline-encoded interactions (9, 10) to uncover the peptide-centric germline-encoded interaction pattern reported here? Identifying conserved TCR germline contacts requires comparing multiple structures involving different MHC molecules. Thus far, examples of conserved germline-encoded pMHC contacts come from analyses of Vß8-containing TCRs and structurally related Vßs in humans (Vß13, 6, 7, 8 encoded by TRBV6, 7, 4, and 12 subfamilies respectively) (4, 9, 10). Vß8 is expressed on ~50% of T cells in mice and represents >70% of published TCR-pMHC structures. This is in spite of the fact that Vß8 is encoded by only three gene segments (TRBV13.1, 2, 3), out of more than 30 functional TCRß gene segments in the mouse genome (Supplementary Figure S4). With the addition of the TG6/HF10/ L^4 structure, crystal structures of four Vß2 containing TCR bound to pMHC have now been determined (48–52), making Vß2 the next most well-represented Vß segment amongst non-redundant mouse TCR-pMHC structure in the Protein Data Bank (https://www.rcsb.org/). Only by comparing all four unique Vß2 containing TCR-pMHC structures, using different Vα chains (TRAV6D-7, 16, 14-1, 10) and interacting with different MHC molecules (L^4, K^b, I-A^κ), was it possible to identify the conserved features of germline-encoded pMHC interactions. While we were unable to identify an obvious counterpart to mouse Vß2 amongst human TCR structures (data not shown), it seems likely that additional Vß-specific pMHC docking patterns will emerge once more pMHC ligated structures using divergent Vß family members are determined. It is also important to note that the germline-encoded view of TCR recognition of pMHC invokes generally similar, but not identical, contacts between TCR CDR1/2 and MHC (7, 10, 53). As a result, the Vß8 TCR may also occasionally adopt an unorthodox angle on its pMHC ligand, such as the parallel docking angle of Vß8.2 TCR 2C on H-2Ld (3). Interestingly, this TCR was nonstimulatory, suggesting that there are docking geometry limits for Vß8 TCRs (Adams, 2011 #72). In spite of individual variations, the docking angles for Vß8 in the structure database are generally diagonal (Figure 5, Supplemental Figure S5).

For the Vß8-specific interaction pattern, the most prominent germline contacts involve tyrosine residues (e.g., Vß8 Tyr 48, Vß4 Tyr 29, and Vß3 Tyr 29/50), which form extensive van der Waals contacts with the MHC ß helices that make up the sides of the peptide-binding groove (7, 9, 10). Tyrosine residues are also highly represented in antibody CDRs and it has been suggested that the hydrophobicity and geometric flexibility of these contacts provides for relatively broad specificity, allowing for interactions with many allelic forms of MHC (4, 54–56). In
contrast, an arginine residue in CDR2 of Vß2 forms a prominent germline contact with an acidic residue in the bound peptide in three out of four of the existing structures. Thus, the positioning CDR1 and 2 over the peptide due to the unusual Vα-Vß domain interface and CDR1/2 loop conformations, together with conserved germline-encoded ionic interactions with peptides, result in a greater contribution to the fine specificity of antigen recognition by Vß2 compared to Vß8. It is tempting to speculate that Vß8 and Vß2 containing TCRs may fill different niches in the immune repertoire, with Vß8 representing “generalists”, with relatively broad specificity, and Vß2 representing “specialists”, optimized for binding particular pMHC complexes (Supplementary Figure S7). While TCR Vß generalists would provide adequate responses and reliable coverage for many different pathogens, Vß specialists could allow for “jackpot” responses that provide strong protection for particular pathogens. This is in line with the motif-driven, focused repertoire of Ld-HF10 specific T cells reported here.

The notion of generalist versus specialist may also be applicable to the MHC-I molecule Ld, which presents the HF10 peptide (Supplementary Figure S7). In contrast to the broad peptide binding exhibited by most MHC-I molecules, Ld forms highly specific interactions with the antigenic HF10 peptide with bends in the peptide stabilized by multiple non-anchor residue contacts. Moreover, six of the side chains are contacted by both the MHC and TCR, such that the specificity for peptide is shared between the MHC and the TCR. The highly specific binding between HF10 and Ld is consistent with earlier studies indicating that Ld has a constrained peptide binding site, binds poorly to self-peptides, and requires particular antigenic peptides to stabilize its cell surface expression (15, 41, 42). Interestingly, human HLA-B alleles associated with elite control of HIV share polymorphisms with Ld that contribute to constrained peptide binding (12, 15) and are also predicted to bind poorly to self-peptides (13). Moreover, a recent study showed that human MHC-I alleles, particularly HLA-B alleles, vary substantially in the proportion of pathogen-derived peptides that they can bind, and that this appears to correlate with pathogen specialization (18). Thus Ld, as well as certain human MHC alleles associated with HIV control, may represent specialist MHC molecules which sacrifice broad coverage for the potential to generate highly protective T cell responses to particular pathogens (20).

The ability of MHC molecules to bind broadly to self-peptides has potential implications for how germline-encoded TCR-MHC reactivity is utilized in the mature TCR repertoire. It has been proposed that unfavorable interactions between CDR3 and self-peptides may counteract germline-encoded reactivity to MHC in order to avoid negative selection (4, 10). In support of this idea, a TCR that was selected by a single peptide MHC complex, and therefore not subject to negative selection by diverse self-peptides, displayed exaggerated germline reactivity for MHC, leading to a high degree of cross-reactivity (10). TCRs that are selected in the thymus by specialist MHC molecules may largely avoid the impact of negative selection due to the lack of binding of self-peptides (13). This may set the stage for jackpot T cell responses since TCRs that recognize rare peptides that are able to bind to and stabilize the specialist MHC can take full advantage of both germline reactivity and peptide specificity to generate high-affinity responses. This strategy may be particularly potent for Vß2- containing TCRs, since specialist MHC molecules would be less likely to present self-peptides with acidic residues near the C-terminus, and thus would avoid strong self-reactivity leading to negative selection of thymocytes bearing these TCRs. Thus, while specialization in MHC and TCR may independently contribute to pathogen control, they may also synergize to generate particularly robust responses.

Previous efforts to understand T cell antigen recognition have largely focused on the most prevalent “generalist” strategies, which provide broad and adequate coverage for most infections. In contrast, the current study highlights alternative “specialist” strategies for generating rare, but highly effective responses. Less commonly used Vß gene segments and MHC alleles may provide a reservoir of recognition components that have the potential to provide highly focused and effective responses, and which could confer a selective advantage when populations are faced with particularly challenging pathogens. A better understanding of the recognition strategies used by specialist MHC alleles and Vß segments should aid in the rational design of TCRs and improve our ability to target T cell responses in individual patients.

DATA AVAILABILITY STATEMENT

The datasets presented in this study can be found in online repositories. The coordinate and structure factors files for the five structures presented here have been deposited in the RCSB Protein Data Bank. Accession numbers are 8D5N, 8D5P, and 8D5Q.

AUTHOR CONTRIBUTIONS

YW, AT, WG, HC, ER, and SDe designed the research. YW, AT, WG, HC, YZ, and WL conducted experiments. YW, AT, ER, and SDe analyzed data. YW, AT, ER, and SDe wrote the manuscript. JS provided essential reagents. DN assisted in the synchrotron X-ray data collection. WW and CD performed the TRangle data collection. WG, HC, YZ, and WL conducted experiments. YW, AT, ER, and SDe supervised the work. All authors contributed to the article and approved the submitted version.

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fimmu.2022.847092/full#supplementary-material

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