The Molecular Basis for Recognition of CD1d/α-Galactosylceramide by a Human Non-Vα24 T Cell Receptor

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Abstract

CD1d-mediated presentation of glycolipid antigens to T cells is capable of initiating powerful immune responses that can have a beneficial impact on many diseases. Molecular analyses have recently detailed the lipid antigen recognition strategies utilized by the invariant Vα24-Jα18 TCR rearrangements of iNKT cells, which comprise a subset of the human CD1d-restricted T cell population. In contrast, little is known about how lipid antigens are recognized by functionally distinct CD1d-restricted T cells bearing different TCRs chain rearrangements. Here we present crystallographic and biophysical analyses of α-galactosylceramide (α-GalCer) recognition by a human CD1d-restricted TCR that utilizes a Vα3.1-Jα18 rearrangement and displays a more restricted specificity for α-linked glycolipids than that of iNKT TCRs. Despite having sequence divergence in the CDR1α and CDR2α loops, this TCR employs a convergent recognition strategy to engage CD1d/αGalCer, with a binding affinity (~2 μM) almost identical to that of an iNKT TCR used in this study. The CDR3α loop, similar in sequence to iNKT-TCRs, engages CD1d/αGalCer in a similar position as that seen with iNKT-TCRs, however fewer actual contacts are made. Instead, the CDR1α loop contributes important contacts to CD1d/αGalCer, with an emphasis on the 4′OH of the galactose headgroup. This is consistent with the inability of Vα24+ T cells to respond to α-glucosylceramide, which differs from αGalCer in the position of the 4′OH. These data illustrate how fine specificity for a lipid containing α-linked galactose is achieved by a TCR structurally distinct from that of iNKT cells.

Introduction

Natural killer T (NKT) cells are a highly conserved lineage of T lymphocytes found in both human and mice that are involved in the modulation of the immune response in autoimmunity, infection, and tumor development [1]. Unlike conventional CD4+ and CD8+ αβ T cells that recognize peptides presented by MHC molecules, NKT cells are reactive to a broad range of self and foreign lipids displayed by the MHC class I–like molecule CD1d [2,3]. This reactivity is initiated by the recognition of the CD1d-lipid complex via the NKT T cell receptor (NKT-TCR) followed by Th1 and/or Th2 biased cytokine secretion that can regulate the activity of other immune cells such as conventional αβ T cells, B cells, and Natural Killer (NK) cells [4].

The most extensively studied NKT cells in humans and mice are invariant (iNKT) or type 1 NKT cells that express TCRs composed of a highly conserved α chain encoded by a Vα24-Jα18 rearranged gene segment in humans and Vα14-Jα18 in mice. This invariant α chain is covalently paired with a β chain in which the variable region is encoded in humans by the Vβ11 gene and can be Vβ8, Vβ7, or Vβ2 in mice [1]. NKT cells expressing these TCRs have a pre-activated phenotype that is due to the expression of the transcription factor pro-myelocytic leukemia zinc finger (PLZF) [5,6] and are also characterized by high reactivity towards the potent stimulatory lipid antigen α-galactosylceramide (αGalCer) [7]. In both humans and mice there are additional classes of T cells that respond to CD1d, one that expresses diverse TCRs but do not respond to αGalCer; these are generally called Type II or non-invariant NKT cells [8]. These NKT cells are typically reactive to lipid antigens such as sulfatide and use an entirely different molecular strategy for recognizing the CD1d/lipid complex [9,10]. A third group of T cells exist that do respond to CD1d presenting αGalCer and also express TCRs different from that of the iNKT-TCR. In mice these NKT cells express a TCR comprised of a Vα10-Jα50/Vβ8 pair [11]. These cells are called Vα10 NKT cells and show a preference for α-glucosylceramide (αGlcCer) over αGalCer; indeed, Vα10 NKT cells can produce a several magnitudes greater cytokine response relative to iNKT cells when stimulated by the related α-glucuronosyldiacylglycerol (α-GlcA-DAG) [11].
Author Summary

Certain lineages of T cells can recognize lipids as stimulatory antigens when presented in the context of CD1 molecules. We know how most Natural Killer T (NKT) cells react with this unusual ligand because they use a single invariant T cell receptor (TCR) alpha chain to do the job. NKT cells place particular emphasis on their CDR3α and CDR2β loops in recognition of antigen—these complementarity determining regions (CDRs) are the hypervariable parts of the TCR that “complement” an antigen’s shape. How do these other T cells recognize closely related yet distinct lipid antigens? Here we show that human CD1d-restricted T cells, typically called Vα24−TCRs, use similar molecular strategies to respond to lipid antigens presented by CD1d. To this end we present a 2.5 Å complex structure of a Vα24−TCR complexed with CD1d presenting the prototypical lipid, α-galactosylceramide (αGalCer). The TCR examined in this study notably shifts its binding slightly, placing more emphasis on the interaction with the CDR1α loop as revealed through alanine scanning mutagenesis. This shift explains the inability of these T cells to respond to lipids that vary at this site of contact (the 4’OH), like the related α-linked glucosylceramide. These results provide a molecular basis for the fine-specificity of different CD1d-restricted T cell lineages.

In humans this third group of CD1d reactive T cells express TCRs with many different Vα domains joined with Jα18, paired with the VB11 domain [12,13]. In contrast to both Type I and Type II NKT cells, these T cells do not typically express CD161, a Natural Killer cell marker found on NKT cells [13]. They have been called Vα24−NKT cells or CD1d-restricted, Vα24−T cells due to their use of diverse Vα domains in their TCRs, use similar molecular strategies to respond to lipid antigens presented by CD1d. To this end we present a 2.5 Å complex structure of a Vα24−TCR complexed with CD1d presenting the prototypical lipid, α-galactosylceramide (αGalCer). The TCR examined in this study notably shifts its binding slightly, placing more emphasis on the interaction with the CDR1α loop as revealed through alanine scanning mutagenesis. This shift explains the inability of these T cells to respond to lipids that vary at this site of contact (the 4’OH), like the related α-linked glucosylceramide. These results provide a molecular basis for the fine-specificity of different CD1d-restricted T cell lineages.

Results

Structure of a Vα24−TCR in Complex with CD1d/αGalCer

In order to understand the molecular basis of Vα24−TCR recognition of CD1d, we expressed a soluble, heterodimeric version of the extracellular domains of the Jα24/N22 TCR [12], which uses the Vα3.1 (TRAV17) gene segment rearranged with Jα18 complexed with VB11, in insect cells. The purified TCR was co-crystallized with recombinant, soluble CD1d loaded with αGalCer; X-ray data were collected to 2.5 Å, and the structure was solved via molecular replacement. Data collection and refinement statistics are listed in Table 1. One TCR/CD1d/αGalCer ternary complex was identified in the asymmetric unit. All components of this complex were well resolved in the electron-density, enabling unambiguous assignment of TCR-CD1d/lipid antigen contacts.

Sequence Variability between Vα24+ and Vα24−TCRs

Table 2 presents a comparison between the amino acid sequences of the α and β CDR loops of the Vα24−(Vα3.1+) TCR studied here and an iNKT Vα24+ TCR studied previously [25]. Vα3.1 and Vα24 share 46% amino acid identity overall, with only 33% (2/6) identity at the CDR1α and 15% (1/7) at the CDR2α loop. However, the shared usage between these TCRs of the Jα18 segment and the canonical DRGSTLGR motif that it encodes gives high sequence identity to the CDR3α loops of these TCRs with different residues encoded only at the Vα24−junction, with ATY and VVS motifs in the Vα24−and Vα24+TCRs,
respectively. The Vβ11 domain is also shared between these TCRs; therefore, the CDR1β and CDR3β sequences are identical. However, the rearranged CDR3β loops differ due to differences introduced during the rearrangement process.

**Recognition of CD1d/αGalCer by the Vα24—TCR**

Overall, the Vα24—TCR recognizes CD1d/αGalCer with the α and β chains oriented on CD1d in a parallel fashion unlike the typical diagonal mode of MHC-I peptide-TCR complexes and similar to that of iNKT-TCR and Vα10 NKT-TCR in complex with CD1d/αGalCer (Figure 1A and 1B) [11,16,19]. However, the binding angle of the Vα24—TCR in relation to the CD1d/αGalCer surface is more acute than the almost perpendicular orientation observed with the Vα24+ iNKT TCR-CD1d/αGalCer structure (Figure 1A) [16,19]. The CDR2 loops adopt a similar yet slightly shifted footprint for the α-chain, yet the β-chain CDR loop positioning is counter-clockwise rotated compared with the Vα24+ TCR complexed with αGalCer [16,19], which is even more extreme than rotations observed in structures of human NKT-TCRs complexed with CD1d presenting LPC or βGalCer (Figure 1B) [23,25]. The TCR-CD1d-lipid contacts mostly fall in the F′ pocket area of the CD1d molecule (Figure 1C), where there are slight differences in TCR contact surface between the Vα24— and Vα24+. The total buried surface area (BSA) between the Vα24—TCR and the CD1d-αGalCer complex was 747 Å², which is slightly smaller than the previously reported interface area for the Vα24+ TCR, ~910 Å². This difference is more pronounced in the β-chain loops with ~37% less contribution in the Vα24+ complex (205.7 Å² versus 325.3 Å² for the Vα24— and Vα24+, respectively).

**αGalCer Positioning in the Complex with the Vα24—TCR**

The conformation and positioning of αGalCer presented by CD1d is almost identical in both complexes with the Vα24+ and the Vα24—TCRs. The sphinosine base and acyl chain of αGalCer fall in the F′ and A′ pockets, respectively (Figure 1D). The αGalCer headgroup also adopts a very similar conformation, with solvent exposed with the sugar oxygens displayed for recognition by the TCR. The conformation of the α helical side chains of CD1d were also highly conserved between the Vα24+ and Vα24— complex structures, with only a few exceptions that are noted later in the text.

**Convergent Recognition Strategy of a Vα24—TCR**

In all three human iNKT TCR-CD1d/lipid complexes that have been resolved to date, the CDR1α loop makes important contacts with the lipid headgroup [16,19,23,25]. In recognition of αGalCer and βGalCer the O′ of Ser30 and the mainchain carbonyl oxygen of Phe29 make hydrogen-bonds (some water-mediated) with the 3′OH of αGalCer and βGalCer, and in the case of LPC, the O′ Ser27 and the mainchain carbonyl oxygen of Phe29 establish hydrogen bonds with the phosphate oxygens of the phosphorylcholine headgroup. Pro28 establishes van der Waals (VDW) contacts with the galactose headgroup; mutagenesis of this residue has a marked effect on recognition but is likely due to global structural changes in the conformation of the TCR as this mutation also disrupted binding of a conformational-specific antibody [17]. In our structure the Vα24— CDR1α loop is slightly shifted from the Vα24+ CDR1α loop (Figure 1B); therefore, the equivalent structural positions to the Vα24+ S27P28F29S30 motif are T26S27I28N29 in Vα24—. Despite the

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**Table 1.** Data collection and refinement statistics (molecular replacement).

| Data Collection | Vα24— TCR-CD1d-αGalCer |
|-----------------|------------------------|
| Space group     | P 1 2 1                |
| Cell dimensions | a, b, c (Å) 57.11, 72.57, 113.75 |
| β (°)           | 103.3                  |
| Resolution (Å)  | 50–52.55 (2.59–2.55)    |
| Rsym (Å)        | 0.065 (0.495)          |
| I/m (Å)         | 24.0 (2.4)             |
| Completeness (%)| 100 (100)              |
| Redundancy (%)  | 3.8 (3.7)              |

| Refinement       |                        |
|------------------|------------------------|
| Resolution (Å)   | 2.55                   |
| Total number of reflections | 109,937               |
| Number of unique reflections | 28,919               |
| Rwork/Rfree     | 0.208/0.266            |
| Number of atoms  | 6,132                  |
| Protein         | 6,132                  |
| Ligand/ion      | 148                    |
| Water           | 126                    |
| β-factors       | 60.7                   |
| Protein         | 60.7                   |
| Ligand/ion      | 71.9                   |
| Water           | 53.9                   |
| R.m.s. deviations |                       |
| Bond lengths (Å)| 0.002                  |
| Bond angles (°) | 0.570                  |

Values in parentheses are for highest resolution shell. doi:10.1371/journal.pbio.1001412.t001

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**Table 2.** Alignment of the Vα24— and Vα24+ NKT TCR CDR loop sequences.

| TCR       | CDR1α | CDR2α | CDR3β |
|-----------|-------|-------|-------|
| J24N22    | TSIINN| IRSNERE| ATRY DRGSTMGLYFGRTGQLTWP |
| iNKT Vα24+| VSPFSN| MTFSENTE| VVS DRGSTMGLYFGRTGQLTWP |
| TCR       | CDR1β | CDR2β | CDR3β |
| J24N22    | MGHDK | YSYGVNST | CASSE NSGTRGI YEQYFGPGLTLYT |
| iNKT Vβ11 | MGHDK | YSYGVNST | CASSE GLRDGRGL YEQYFGPGLTLYT |

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| TCR       | CDR1β | CDR2β | CDR3β |
| J24N22    | MGHDK | YSYGVNST | CASSE NSGTRGI YEQYFGPGLTLYT |
| iNKT Vβ11 | MGHDK | YSYGVNST | CASSE GLRDGRGL YEQYFGPGLTLYT |

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chemical and structural differences of the CDR1α loops between these TCRs, specific side-chain-mediated hydrogen bonds are still formed in the Vα24+ CDR1α loop, both with the galactose headgroup of αGalCer and through VDW contacts with CD1d’s Val72 (Figure 2A and Table 3). The shifted position of Ser27 in this complex enables a hydrogen bond between its Oγ with the 6′OH of αGalCer, whereas the Nδ2 of Asn29 hydrogen bonds with the 3′OH and 4′OH of αGalCer and Asn29 also forms VDW contacts with the galactose headgroup. Therefore, alternative residues in the CDR1α loop are effectively used in recognition of αGalCer with a focus on the 4′OH of the galactose ring, with a novel contact with CD1d also noted.

We have also noted residues in the CDR2α loop that make water-mediated contacts with the αGalCer galactose headgroup: Ser50 and Asn51 both establish water-mediated hydrogen bonds with the 4′OH of αGalCer (Figure 2B). In the other human complexes, Phe51 of the Vα24+ CDR2α loop makes VDW contacts with both βGalCer and LPC, however hydrogen bonds have not been noted for the CDR2α loop of Vα24+ TCRs. In contrast to the sequence and contact differences at the CDR1α and CDR2α loops, the residues of the CDR3α loop in the Vα24+ TCRs adopt a similar conformation to that of the Vα24+ iNKT TCRs (Figure 2C). Yet despite the similarity in footprint, the Vα24+ CDR3α loop establishes fewer contacts with CD1d and αGalCer than does the CDR3α loop of the iNKT TCR (Table 3) (25 instead of 32, respectively, for CD1d and eight instead of 19, respectively, for αGalCer). There are fewer hydrogen bonds (two versus eight with CD1d and one versus four with αGalCer) and, in the case of αGalCer, fewer than half (seven versus 15) VDW contacts of those observed in the Vα24+ complex. The residues of
the Vα24+ CDR3α were previously shown to be energetically critical for CD1d/αGalCer recognition [17], a finding recapitulated in our data (discussed further below) despite the lower contact number.

A Shifted Vα24—TCR β Chain Maintains Conserved Contacts through the CDR2β

While the CDR3α loop serves to anchor human iNKT TCRs on the CD1d/lipid platforms with highly similar conformations [16,19,23,25], the remaining loops have demonstrated rotational flexibility in how they are positioned over the CD1d/lipid surface, in particular at the CDR2β, which establishes energetically critical contacts with CD1d [17]. A similar rotation is seen in the Vα24—TCR docking on the CD1d/αGalCer platform in the complex structure presented here (Figure 1B and Figure 3A). As in the Vα24+ complexes, the involvement of the CDR2β loop in CD1d binding is predominantly mediated by Tyr48 and Tyr50. Despite an average shift of 4.6 Å between the Vα24— and Vα24+ CDR2β CA backbones, the rotationally flexible tyrosine side chains maintain highly similar contacts between the two complexes (Figure 3A). Glu83 on CD1d takes a central role in contact with the CDR2β in both complexes, establishing a hydrogen-bonded network with both Tyr48 and Tyr50 hydroxyls. Met87 also contributes VDW contacts with Tyr50 in both complexes. However, in contrast to the Vα24+ complex, where Glu36 of the CDR2β establishes a robust salt-bridge with Lys86 of CD1d (3.7 Å distance), in the Vα24—complex Lys86 has shifted such that it is 4.6 Å from Glu36 (Figure 3A). Thus, the critical contacts of the CDR2β loop are maintained in the Vα24—complex despite large main chain shifts of the CDR2β backbone.

The highly variable CDR3β loop has been demonstrated to confer reactivity to specific lipids presented by CD1d by both human [26] and mouse [27] iNKT cells. In the Vα24—complex, the CDR3β loop is well resolved in the electron density and establishes only one weak hydrogen bond and a VDW contact with Gln150 on CD1d’s β2-helix via Ser97 (Figure 3B). Thus, unlike the murine Vα24+ NKT TCRs, which have CDR3β sequence specificity and use this loop in CD1d binding, this Vα24—TCR does not appear to rely heavily on its CDR3β loop for binding.

Conformational Flexibility of Vα24—CDR3α Loops

The availability of a Vα24—TCR also expressing a Vβ3.1 domain (named 5B) [28] in the unliganded state allows a direct comparison between the loop structures between the TCR examined here (bound to CD1d) and a Vα24—, Vβ3.1+, TCR in its unbound state. Due to the use of different β gene segments that results in global domain orientation shifts, the TCRs are not perfectly superimposable (Figure 4A) and there are two amino acid differences in the CDR3α sequences of these TCRs due to junctional diversity (Figure 4B). Alignment of the two Vβ3.1 domains shows the CDR1 and CDR2 loops are essentially identical structurally (Figure 4B), yet examination of the CDR3α loops (Figure 4B) shows significant structural differences. While the unliganded structure of J24-N22 is not known, modeling of the 5B TCR onto our complex structure suggests a large shift in loop conformation would need to occur in the CDR3α loop for it to dock onto CD1d/αGalCer in a similar fashion. Because of the similarities between these TCRs in all other loops save the CDR3β, it is very likely that the 5B TCR would dock in a similar fashion as seen here. Thus in contrast to the Vα24+ NKT TCRs’ recognition of CD1d/αGalCer, where loop conformation was highly conserved in the liganded and unliganded state, we suggest that the CDR3α loop can be flexible in Vβ3.1+, Vα24—TCRs,
Table 3. Atomic contact comparison of iNKT-TCRs, CD1d, and lipid ligands.

| CDR1α | αGalCer | Bond Type |
|-------|---------|-----------|
| Ser27 | O5A     | VDW       |
| Ser27O1 | O5A | HB |
| Asn29N12 | O3A, O4A | HB |
| Asn29 | O4A, C3A | VDW |

| CDR3α | αGalCer | Bond Type |
|-------|---------|-----------|
| Asp92 | C1A     | VDW       |
| Arg93 | C2A, O2A, O3 | VDW |
| Gly94 | C2A | VDW |
| Gly94 | O2A, O3A, C2A | VDW |

| CDR1α | CD1d | Bond Type |
|-------|------|-----------|
| Ser27 | Val72 | VDW |

| CDR2α | CD1d | Bond Type |
|-------|------|-----------|
| Ser50O | O4A | Water-mediated HB |
| Asn51O1 | O4A | Water-mediated HB |

| CDR3β | CD1d | Bond Type |
|-------|------|-----------|
| Ser97O | Gln150N12 | HB (3.67) |
| Ser97 | Gln150 | VDW |

| CDR2β | CD1d | Bond Type |
|-------|------|-----------|
| Tyr48 | Glu83, Lys86 | VDW |
| Tyr48O1 | Glu83O1, Glu83O2, Lys86N1 | HB |
| Tyr50 | Glu83, Met87 | VDW |
| Tyr50O1 | Glu83O1 | HB |
| Glu56 | Lys86 | VDW |
| Glu56 | Lys86 | VDW |

| CDR3α | CD1d | Bond Type |
|-------|------|-----------|
| Asp92 | Arg79 | VDW |
| Asp92O1 | Arg79 N1, Arg79 N12 | SB |
| Asp92O2 | Arg79 N1, Arg79 N12 | SB |
| Arg93 | Ser76, Arg79, Asp80 | VDW |
| Arg93N13 | Asp80 O1, Asp80 O2 | SB |
| Gly94 | Gln150, Asp151 | VDW |
| Ser95 | Val147, Gln150 | VDW |
| Thr96N1 | Gln150 O1 | HB |
| Thr96 | Val147, Gln150 | VDW |
| Leu97O | Arg79 N1 | HB |
| Leu97 | Asp80, Glu83, Phe84, Met87, Val147 | VDW |
| Arg99 | Arg79 | VDW |
| Tyr101 | Arg79 | VDW |

All contacts are shown with bond typeHB (3.67) for hydrogen bonds.

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Figure 3. The role of the TCR β chain in Vα24—TCR engagement of CD1d/αGalCer. (A) Contacts made by the CDR2β loops of the Vα24— and Vα24+ NKT TCRs with CD1d. CD1d is shown as grey ribbons and the Vα24— and Vα24+ NKT TCR CDR2β loops in yellow-orange and pale-green color, respectively. Hydrogen bonds (<3.3 Å) and salt bridges are shown as yellow dotted lines for the Vα24— and Vα24+ NKT TCRs. (B) Electron density (Fo-Fc omit map, contoured at 3σ) for the CDR3β loop of the Vα24— TCR is shown as blue mesh together with the CDR3β in stick representation in yellow-orange; CD1d is shown in grey ribbons and αGalCer in yellow sticks. Potential H-bond is displayed as dotted, yellow line. doi:10.1371/journal.pbio.1001412.g003

similar to what was previously seen in the iNKT TCR recognition of CD1d/LPC [25].

Residues Contributing to Vα24—TCR Binding of CD1d/αGalCer

To evaluate the kinetics involved in binding of our Vα24—TCR with CD1d/αGalCer, we used surface plasmon resonance to measure the association (k_{on}) and dissociation rates (k_{off}) of this interaction and determine the dissociation constant (K_D) (Figure 5A). We also used this to calculate K_D by equilibrium analysis (Figure 5A, insets). We included an iNKT (Vα24+) TCR in our kinetic measurements such that we could compare these values to a representative of the iNKT population. The affinity of the Vα24— TCR used in this study for CD1d/αGalCer (2.1 μM kinetic, 2.5 μM equilibrium) was similar to the affinity we measured for the iNKT TCR (2.1 μM kinetic, 1.9 μM equilibrium) as well as affinities from previous measurements with Vα24—TCRs (using Vα3.1 and Vα10.3 domains) [28]. Stronger affinities (0.5 μM) have been noted for other human iNKT TCRs [17].

We sought to further evaluate the residues contributing most to Vα24—TCR binding to CD1d/αGalCer. We chose key TCR residues identified as interacting with CD1d/αGalCer in our complex and evaluated their contribution to binding via alanine-scanning mutagenesis and SPR. We first evaluated the CDR1α loop residues Ser27 and Asn29, as these appeared to mediate the side-chain-specific contacts that differed most from the Vα24+ TCRs. While mutation of Ser27 to Ala (S27A) did not drastically change Vα24—TCR binding kinetics, mutating Asn29 to Ala (N29A) resulted in a significant disruption to binding with changes in both the association and dissociation rates and an increase in the K_D by an order of magnitude (Figure 5B). Thus the CDR1α loop provides a clear contribution to Vα24—TCR binding to CD1d/αGalCer. Previous mutational analysis of the CDR1α loop of a Vα24+ TCR [17] of Pro28 to Alanine disrupted binding, however this was assumed to be due to changes in the TCR architecture as conformational-specific antibodies failed to bind this mutant.

Mutation of the CDR2β side chains Ser50 and Asn51 had subtle effects on k_{on} and k_{off} (Figure 5B) yet did not appear to have a substantial effect on the overall affinity of CD1d/αGalCer binding, similar to what we observed with mutation of Ser97 in the CDR3β loop sequence. Because of the similarities in CDR2β loop contacts between Vα24— and Vα24+ TCRs, we included a mutation of Arg95 of the CDR3β as a positive control; this side chain has been shown to be central to iNKT TCR binding to CD1d/αGalCer [17]. We also observed that mutation of this side chain to Ala (R95A) abrogated binding of the Vα24—TCR and thus supports the importance of the CDR3β loop to Vα24—TCR docking.

Discussion

Our complex structure of a Vα24—TCR with CD1d/αGalCer provides a model by which to understand how this diverse population of CD1d-restricted human T cells recognize antigen. These cells differ from iNKT cells in their specificity, effector function, and the markers expressed on their cell surface; these factors combined argue that these cells provide another arm of T-cell-mediated lipid recognition in humans. Here we provide a structural and biophysical foundation upon which to understand the molecular basis of differential reactivity observed at the cellular level in this NKT cell population.

Despite the divergent amino acid sequences encoded by the Vα3.1 domain for the CDR1α and CDR2β loops, the Vα24—
TCRs reveals CDR3 loop structural differences. This docking orientation is primarily dictated by the conserved docking of the CDR3α loop, containing the highly similar sequence encoded by the Jα28 segment of iNKT TCRs. The contacts mediated by the other loops, while not identical to those of iNKT TCRs, were very similar, suggesting that despite sequence differences in the Vα loops they could establish contacts with similar regions of the CD1d/αGalCer surface. The αGalCer headgroup position was almost identical to that observed in the iNKT complex structures [16,19]. This docking mode, also shared with that of the murine Vα10 NKT TCR [11], is strikingly different from that of the recently resolved type II NKT TCR structures [9,10], where the TCRs dock on an entirely different surface of CD1d (the Aβ pocket) and use all six of the TCR’s CDR loops in recognition (similar to what is observed in conventional ββ TCR recognition of MHC/peptide). These structures demonstrate that CD1d-restricted T cells can use at least two divergent ways to recognize their antigens [29].

Our complex structure provides a useful model to compare other Vα24− TCRs’ structures, notably the structure of a highly related unliganded TCR called 5B [28]. If we assume the 5B TCR would dock similarly to the Vα24− TCR examined in our study, a significant conformational change would have to occur in 5B’s CDR3α loop. This conformational flexibility was a feature we also observed in human iNKT TCR binding to CD1d/LPC [25]. In contrast to what was observed with the iNKT TCR complex structure with CD1d/αGalCer [16,19], this suggests that not all CD1d-TCR interactions are “lock and key” and that changes to CDR3α loop conformation may contribute to differences in binding kinetics and thermodynamics. A similar phenomenon of loop movement was observed in the murine Vα10 NKT TCR upon binding [11].

The CDR3α loop footprint on CD1d/αGalCer is conserved in all the iNKT-TCR/CD1d structures noted to date as it is here. However, the number of contacts in this complex structure were less than that observed in the iNKT-TCR CD1d/αGalCer complex structure, yet the binding affinities measured for the Vα24+ and Vα24− TCRs in this study did not differ substantially (<2 μM for both TCRs). The alanine-scanning mutagenesis revealed important contributions from the CDR1α loop (in particular, residue N29) in the Vα24− TCR binding that were not noted in Vα24+ TCR binding (mutation of the equivalent position, S30 in the Vα24+ TCR, showed little effect [17]). This shift of importance toward the CDR1α likely compensates for fewer CDR3α loop contacts and would explain the altered reactivity patterns of Vα24− TCRs for lipids that are recognized similarly by Vα24+ TCRs (such as αGlcCer and αGalCer, discussed more below). We cannot rule out that contributions from other loops, such as the CDR2α and CDR3β, contribute as well; while individual mutagenesis of these residues had small effects upon TCR binding, in combination they may have a cumulative effect in binding CD1d/lipid, evident only when they are mutated in concert.

Extensive studies in the mouse iNKT cell system have revealed how lipid ligands are structurally modified during recognition by the iNKT TCR. Even though extensive structural variability exists in the glycolipid headgroups, each carbohydrate structure adopts a similar orientation when bound by the TCR [20–24]. Therefore, contributions of the CDR1β in recognition of alternative lipids, both α- and β-linked glycolipids, could be an important factor in Vα24− T cell reactivity towards different lipids. Directly relevant to this point is the clear distinction between Vα24− T cells and Vα24+ iNKT cells in their differential reactivity to the α-linked glycolipids αGlcCer and αGalCer. Vα24+ iNKT cells respond well to both lipids, whereas Vα24− T cells do not respond to αGlcCer. The only difference present between these two lipids is the orientation of the 4′OH group on the sugar ring (glucose versus galactose). Our structural and biophysical data provide an explanation for this difference in reactivity. Asn29, a residue in the Vα24− CDR1α, establishes both VDW and hydrogen bonds with

Figure 4. Superposition of unliganded and liganded Vα24− TCRs reveals CDR3 loop structural differences. (A) Superposition of the unliganded Vα24− (Vα24−[1]) TCR 5B (grey ribbon) [28] (PDB ID: 2CDG) and the Vα24− TCR of this study (α chain in light blue ribbon and β chain in yellow orange ribbon). (B) Close-up view of the α and β CDR loops of the unliganded and liganded Vα24− TCRC. The CDR loops are colored according to their TCR chain coloring in (A). The CDR3α loop sequences of the 5B and J24.N22 TCRs are shown at bottom.

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the 3'OH and 4'OH. Mutation of this residue to alanine results in an order of magnitude decrease in binding of the Vα24− TCR, presumably due to disruption of these contacts. Furthermore, the CDR2ζ loop residues Ser50 and Asn51 establish water-mediated hydrogen bonds with the 4'OH that may help to stabilize the interaction despite lacking clear energetic contributions (as

Figure 5. SPR analysis of the binding of CD1d-αGalCer to the Vα24− and Vα24+ NKT TCRs. (A) SPR binding curves of CD1d-αGalCer. Shown are the curves and fits used for kinetic analysis to surface-immobilized Vα24− (clone J24N.22) (top) and Vα24+ (clone J24L.17) (bottom) NKT TCRs. For the Vα24− TCR: $k_{on} = 1.62 \times 10^5 \pm 0.11 \times 10^5$ (M s)$^{-1}$, $k_{off} = 0.342 \pm 0.007$ s$^{-1}$, and $K_D = 2.1$ μM; for Vα24+ TCR: $k_{on} = 1.94 \times 10^5 \pm 0.16 \times 10^5$ (M s)$^{-1}$, $k_{off} = 0.414 \pm 0.010$ s$^{-1}$, and $K_D = 2.1$ μM. Grey traces represent experimental data and black lines fittings to a Langmuir 1:1 kinetic model. Curves represent the following concentrations of analyte: 0, .037, .111, .333, 1, and 3 μM. In addition, equilibrium analysis was performed on these curves and those for 9 μM and 27 μM; the fits and calculated $K_D$s for this analysis are shown as inserts. (B) Alanine scanning mutants of key residues in the Vα24− TCR interface were screened by SPR, and their $k_{on}$ and $k_{off}$ values are shown plotted on a $k_{on}$ versus $k_{off}$ plot with $K_D$ isotherms shown along with the values for the wild-type Vα24− TCR and the Vα24+ iNKT TCR (J24.L17). doi:10.1371/journal.pbio.1001412.g005
assessed in our alanine-mutagenesis studies). We therefore propose that modification to the 4′-OH between the galactose (αGalCer) and glucose (αGlcCer) structure is the primary molecular factor mediating the differences in reactivity of the Vα24+ population of CD1d-restricted T cells. The alternative contacts with the carbohydrate headgroup in the iNKT TCR/CD1d/αGalCer structure may explain why iNKT cells can respond to both lipids; the main contacts with the 4′-OH are mediated by Ser30, which when mutated to alanine only had a minimal effect on binding [17]. The greater number of contacts and BSA of the Vα24+ TCR CDR3α loop on CD1d/αGalCer may make these T cells relatively insensitive to variation in the glycolipid headgroup at other positions. The difference in 4′-OH recognition may translate to alternative reactivity to other glycolipid and non-glycolipid lipid structures both in development of these T cells in the thymus and their effector functions in the periphery. Despite their shared use of Jα18 and Vβ11, the Vα24+ T cells are differentiated from iNKT cells in their development and activation state; presumably altered TCR recognition of a selecting antigen during thymic development plays a role in these differences. Our structure provides a model by which to understand the molecular basis of this altered reactivity. Our results, which focus much of the differences in reactivity to αGlcCer on the CDR1α loop and its interaction with the 4′-OH, contrast with the murine Vα10 NKT cell preferred reactivity to αGlcCer [11], where preference in binding appears due to many factors. The highly convergent recognition of αGlcCer by these TCRs distributes the binding contacts over much of the CDR loop surfaces [11]. While mutagenesis data for these residues are not available, it is clear there are differences in the nature of the contacts between the Vα10 and iNKT TCR with CD1d (VDW versus hydrogen bonds), that many new contacts are established with CD1d, and therefore modification to the sugar ring may have more of a distributed effect over the Vα10 NKT interaction than what we observe in our Vα24+ TCR complex structure. Both structures, however, provide molecular models for the observed differences in lipid reactivity and demonstrate how divergent NKT TCR structures can convergently recognize similar CD1d/lipid antigen structures. The molecular basis of the differences in reactivity we have described here are the first clues into understanding why Vα24+ cells are developmentally and functionally distinct from the iNKT population.

**Materials and Methods**

### Human Wild-Type CD1d—β2m Expression and Purification

The ectodomain region of human CD1d and human β2m microglobulin (β2m) were co-expressed in insect cells and purified as described [25].

### Vα24+ and Vα24− TCR Expression and Purification

The cDNAs corresponding to the α and β chains of the Vα24+ NKT TCR clone J24L.17 and the α and β chains of Vα24− TCR clone J24N.22 were separately cloned into different versions of the pAG67A vector each containing a 3C protease site followed by either acidic or basic zippers and a 6xHis tag. Both chains were co-expressed in Hi5 cells via baculovirus transduction. The heterodimeric TCRs was captured with Nickel NTA Agarose (Qiagen) and further purified by anion exchange and size-exclusion chromatography.

### Generation of Vα24− TCR Mutants

Mutants of the Vα24− TCR (S27A, N29A, S50A, N51A, R95A for the alpha chain, and S97A for the beta chain) were generated through overlapping PCR with specific primers containing the desired mutation. Mutant heterodimeric TCR was expressed in insect cells as described above.

### CD1d Loading with αGalCer

Purified human CD1d was used for loading with αGalCer at room temperature with a three molar excess of lipid for 16 h. The excess of lipid was then removed with a Superdex 200 (10/30) column (GE Healthcare).

### Surface Plasmon Resonance Measurements

A human CD1d construct bearing a 3C protease site + 6X-Histidine tag at the C-terminus was expressed in Hi5 cells and purified as described [25]. All interaction experiments were performed in a BLAcore 3000 Instrument (GE Healthcare). Three hundred RU of wild-type Vα24− NKT TCR or a mutant version of it were captured in a flow channel of an Ni-NTA sensor chip (GE Healthcare) previously treated with NiCl₂. Insect-cell-derived recombinant IgFc was used to block unbound sensor chip surface to minimize nonspecific binding events. Increasing concentrations (0, 0.037, 0.111, 0.333, 1, 3, 9, and 27 μM) of CD1d/αGalCer were injected at a flow rate of 30 μl/min in 10 mM Hepes pH 7.4, 150 mM NaCl, and 0.005% Tween-20. Both kinetic and equilibrium parameters were calculated off of these curves using BIAevaluation software 3.2RCl (GE Healthcare) and GraphPad Prism.

### Ternary Complex Formation and Crystallization

Nickel agarose-purified Vα24− TCR was digested with 3C protease for 16 h at 4°C to remove the zippers and His tags and purified by anion exchange chromatography in a MonoQ column (GE Healthcare). Endoglycosidase F3 (EndoF3) was used next at a 1:10 enzyme-to-protein ratio for 2 h at 37°C in order to minimize the sugar content present in the protein. The digested protein was purified by a new round of anion exchange followed by size-exclusion chromatography. Both αGalCer-loaded CD1d and EndoF3-treated Vα24− TCR protein samples were mixed in HBS at 1:1 molar ratio and concentrated in Nanosep Centrifugal Devices ( Pall Life Sciences) to 10 mg/ml. Initial hits were found in 0.1 M sodium acetate, 20% PEG 4000, and were optimized to birefringent crystals that grew in 0.1 M sodium acetate, 20% PEG 4000, and 0.1 M ammonium acetate.

### Crystallographic Data Collection, Structure Determination, and Refinement

Crystals were cryo-cooled in mother liquor supplemented with 20% glycerol prior to data collection. All data sets were collected on a MarMosaic 300 CCD at the LS-CAT Beamline 21-ID-G at the Advanced Photon Source (APS) at Argonne National Laboratory and processed with HKL2000 [30].

The structure of the ternary complex was solved by molecular replacement with the program Phaser [31] using the human CD1d-β2m (Protein Data Bank (PDB) accession number 1Z74) and an iNKT Vα24+ TCR (2EYS) as search models. Refinement with Phenix software suite [32] was initiated through rigid body and followed with XYZ coordinates and individual B-factor refinement. These first steps of refinement yielded clear unbiased and continuous density for αGalCer. Next, extensive cycles of manual building in Coot [33] and refinement were carried out and ligands such as αGalCer or covalently bound sugars were introduced guided by Fo–Fc positive electron density. Ligands structures and chemical parameters were defined with C.C.P.P.’s Sketcher [34] and included in subsequent refinement and manual
building steps. Translation/libration/screw (TLS) partitions were calculated and incorporated at later refinement stages. All the refinement procedures were performed taking a random 5% of reflections and excluding them for statistical validation purposes (Rfree).

Structure Analysis

Intermolecular contacts and distances were calculated using the program Contacts from the CCP4 software package [34], interface surface areas were calculated using the PISA server [http://www.ebi.ac.uk/msd-srv/prot_int/pistart.html], and all structural figures were generated using the program Pymol (Schrodinger, LLC).

Accession Numbers

Coordinates and structure factors for the J24.N22 Vα24—TCR/CD1d/αGalCer complex have been deposited in the Protein Data Bank under the accession code 4EN3.

References

1. Bendelac A, Savage PB, Teyton L (2007) The biology of NKT cells. Annu Rev Immunol 25: 297–336.
2. Bril M, Brenner MB (2006) CD1: antigen presentation and T cell function. Annu Rev Immunol 24: 817–890.
3. Godfrey DI, Rosjohann J (2011) New ways to turn on NKT cells. J Exp Med 208: 1121–1125.
4. Joyce S, Girardi E, Zajonc DM (2011) NKT cell ligand recognition logic: molecular basis for a synaptic duet and transmission of inflammatory effectors. J Immunol 187: 1081–1089.
5. Kovalovsky D, Uche OU, Eladad S, Hobbs RM, Yi W, et al. (2008) The BTB-zinc finger transcriptional regulator PLZF controls the development of invariant natural killer T cell effector functions. Nat Immunol 9: 1035–1044.
6. Savage AK, Constantinides MG, Smyth MJ, Van Kaer L (2004) TCR/CD1d/αGalCer complex have been deposited in the Protein Data Bank under the accession code 4EN3.

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Author Contributions

The author(s) have made the following declarations about their contributions: Conceived and designed the experiments: JLS JG EJA. Performed the experiments: JLS EJA. Analyzed the data: JLS EJA. Contributed reagents/materials/analysis tools: PBS JG. Wrote the paper: JLS EJA.