The molecular bases of $\delta/\alpha\beta$ T cell–mediated antigen recognition

Daniel G. Pellicci,1,2* Adam P. Uldrich,1,2* Jérôme Le Nours,3,4 Fiona Ross,1,2 Eric Chabrol,3 Sidonia B.G. Eckle,1 Renate de Boer,3 Ricky T. Lim,1 Kirsty McPherson,1 Gurdyal Besra,6 Amy R. Howell,7 Lorenzo Moretta,8 James McCluskey,1 Mirjam H.M. Heemskerk,3 Stephanie Gras,3,4 Jamie Rossjohn,3,4,9** and Dale I. Godfrey1,2**

1Department of Microbiology and Immunology, Peter Doherty Institute for Infection and Immunity and 2Australian Research Council Centre of Excellence in Advanced Molecular Imaging, University of Melbourne, Parkville, Victoria 3010, Australia
2Department of Biochemistry and Molecular Biology, School of Biomedical Sciences and 3Australian Research Council Centre of Excellence in Advanced Molecular Imaging, University of Melbourne, Parkville, Victoria 3010, Australia
3Department of Hematology, Leiden University Medical Center, 2300 RC Leiden, Netherlands
4School of Biosciences, University of Birmingham, Edgbaston, Birmingham B15 2TT, England, UK
5Department of Chemistry, University of Connecticut, Storrs, CT 06269
6Istituto Giannina Gaslini, 16147 Genova, Italy
7Department of Infection and Immunity, School of Medicine, Cardiff University, Heath Park, Cardiff CF14 4XN, Wales, UK
8Istituto Giannina Gaslini, 16147 Genova, Italy
9Institute of Infection and Immunity, School of Medicine, Cardiff University, Heath Park, Cardiff CF14 4XN, Wales, UK

$\alpha\beta$ and $\gamma\delta$ T cells are disparate T cell lineages that can respond to distinct antigens (Ags) via the use of the $\alpha\beta$ and $\gamma\delta$ T cell Ag receptors (TCRs), respectively. Here we characterize a population of human T cells, which we term $\delta/\alpha\beta$ T cells, expressing TCRs comprised of a TCR-$\delta$ variable gene (V$\delta$1) fused to joining $\alpha$ and constant $\alpha$ domains, paired with an array of TCR-$\beta$ chains. We demonstrate that these cells, which represent $\sim$50% of all V$\delta$1+ human T cells, can recognize peptide- and lipid-based Ags presented by human leukocyte antigen (HLA) and CD1d, respectively. Similar to type I natural killer T (NKT) cells, CD1d–lipid Ag-reactive $\delta/\alpha\beta$ T cells recognized $\alpha$-galactosylceramide ($\alpha$-GalCer); however, their fine specificity for other lipid Ags presented by CD1d, such as $\alpha$-glucosylceramide, was distinct from type I NKT cells. Thus, $\delta/\alpha\beta$TCRs contribute new patterns of Ag specificity to the human immune system. Furthermore, we provide the molecular bases of how $\delta/\alpha\beta$TCRs bind to their targets, with the V$\delta$1-encoded region providing a major contribution to $\delta/\alpha\beta$TCR binding. Our findings highlight how components from $\alpha\beta$ and $\gamma\delta$TCR gene loci can recombine to confer Ag specificity, thus expanding our understanding of T cell biology and TCR diversity.

* J. Rossjohn and D.I. Godfrey contributed equally to this paper.
** D.G. Pellicci and A.P. Uldrich contributed equally to this paper.

Abbreviations used: Ag, antigen; $\alpha$-GalCer, $\alpha$-galactosylceramide; $\alpha$-GlcCer, $\alpha$-glucosylceramide; BSA, buried surface area; CDR, complementarity-determining region; MAIT cell, mucosal-associated invariant T cell; rmsd, root-mean-square deviation; SPR, surface plasmon resonance.
the Ag-binding cleft of molecules encoded by the polymorphic MHC. Here, the αβTCR accommodates a wide range of pMHC landscapes with a polarized and approximately conserved docking mode, whereby the Vα and Vβ domains are positioned over the α2 and α1 helices of MHC-I, respectively (Gras et al., 2012). Alternately, some αβ T cells are activated by lipid-based Ags presented by MHC-I–like molecules belonging to the CD1 family (Brigl and Brenner, 2004). The CD1d system, which presents lipid Ags to type I and type II NKT cells, is the best understood in terms of lipid Ag recognition (Girardi and Zajonc, 2012; Rossjohn et al., 2012). Here, a semi-invariant NKT TCR (Vα24-Jα18 in humans), which typifies type I NKT cells, binds a wide range of chemically distinct ligands in a conserved docking mode, whereby the TCR sits in a parallel manner above the F′ pocket of CD1d (Rossjohn et al., 2012). As such, the NKT TCR has been likened to an innate-like pattern recognition receptor (Scott-Browne et al., 2007). In contrast, type II NKT cells can adopt differing docking strategies in binding to CD1d-restricted lipid-based ligands and exhibit features that more closely resemble that of αβTCR recognition in adaptive immunity (Girardi et al., 2012; Patel et al., 2012; Rossjohn et al., 2012). It has also recently been established that mucosal-associated invariant T cells (MAIT cells), which express a semi-invariant αβTCR, recognize vitamin B–based metabolites presented by the monomorphic MHC-I–related protein (MR1; Kjer-Nielsen et al., 2012; Corbett et al., 2014). Here, the MAIT TCR draws upon features typified by innate and adaptive immunity in recognizing these small molecule metabolites (Patel et al., 2013; Eckle et al., 2014). Accordingly, the αβTCR lineage shows remarkable versatility in recognizing three distinct classes of ligands (Bhati et al., 2014).

The γδ T cell lineage uses γδTCRs that are derived from the γ and δ TCR gene loci (O’Brien et al., 2007; Vantourout and Hayday, 2013). γδ T cells and αβ T cells develop from common intrathymic precursors but branch into separate lineages at the time when they undergo TCR gene rearrangement and differentiation (Xiong and Raulet, 2007; Gifuni and Zúñiga-Pflücker, 2010). γδ T cells rearrange Vγ and Jγ genes that join to the γ constant (Cγ) gene to form the TCR–γ chain, whereas rearrangement of Vδ, Dδ, and Jδ genes join to the δ constant (Cδ) gene to form the TCR–δ chain. Similar to αβTCRs, γδTCRs possess six CDR loops, three from each chain, which mediate Ag recognition (Bhati et al., 2014). The number of Vγ and Vδ genes in humans is relatively low (8 × Vδ and 6 × Vγ genes), and further limitation in repertoire diversity comes from restricted pairing of particular Vδ and Vγ genes. However, the potential to use the three Dδ genes, even in multiple copies, combined with N region modifications, dramatically increases TCR–δ diversity (O’Brien et al., 2007; Born et al., 2013). In contrast to αβ T cells, in which Vα and Vβ TCR chains are generally very diverse, some Vγ and Vδ TCR chains show tissue-specific and functional biases. For example, Vδ2+ γδ T cells tend to produce inflammatory cytokines such as IFN–γ and TNF, predominate in human blood, and migrate to sites of inflammation, whereas Vδ1+ γδ T cells tend to produce regulatory cytokines such as IL-10 and home to noninflamed tissues such as spleen and gut (O’Brien et al., 2007). Compared with αβ T cells, much less is known about what types of Ags are recognized by γδ T cells, although it is generally accepted that γδTCRs confer different specificity and functional characteristics (Vantourout and Hayday, 2013). Some γδTCRs can recognize Ags directly, whereas other studies have demonstrated that γδTCRs can recognize cell surface and soluble protein and peptide Ags and microbial metabolites in the absence of classical Ag-presenting molecules (Born et al., 2013; Vavassori et al., 2013; Sandstrom et al., 2014). Some γδ T cells can respond to Ag-presenting molecules in a ligand-independent manner, such as the MHC-II molecule or the MHC class I–like molecules T10/T22 and endothelial protein C receptor (EPCR), whereas others can recognize lipid-based Ags presented by members of the CD1 family (Born et al., 2013). The molecular bases of γδTCR recognition of CD1d-lipid Ag complexes were recently reported (Luoma et al., 2013; Uldrich et al., 2013).

Thus, αβ T cells and γδ T cells act in concert, using distinct TCRs to survey a wide range of Ags to enable protective immunity. Interestingly, the human Vδ gene locus is embedded within the Vα locus, and some human Vδ genes (Vδ4–Vδ8) encoded by TRDV 4, 5, 6, 7, and 8 are also referred to as Vδ genes (Vα6, 21, 17, 28, and 14.1) encoded by TRAV 14, 23, 29, 36, and 38–2, respectively (Lefranc and Rabbits, 1990), because these are capable of rearranging to either Dδ–Cδ or Jδ–Cα genes. Because these V genes can be used by both γδ and αβ T cell lineages, when paired with Cα, they are termed Vα genes, whereas they are termed Vδ genes when paired with Cδ (Lefranc and Rabbits, 1990). However, the majority of human γδ T cells use the Vδ1, Vδ2, and Vδ3 variable regions, encoded by TRDV 1, 2, and 3 genes (O’Brien et al., 2007; Mangan et al., 2013). Although these do not have alternate TRAV names, these can also rearrange to Jα–Cα genes, and at least Vδ1 and Vδ3 can be expressed as a functional Vα–Jα–Cα TCR chain that can pair with a functional TCR–β chain (Miossec et al., 1990; Peyrat et al., 1995). Here we describe a flow cytometry–based method for identifying Vδ1+ TCR–β+ cells, which we have termed δ/αβ T cells, based on their expression of TCRs comprising a TCR–δ variable gene 1 (Vδ1) joined to a TCR Jα and TCR Cα genes and paired with an array of TCR–β chains. These δ/αβ T cells were readily detectable in most humans and included cells with specificity for both peptide- and lipid-based Ags presented by MHC-I molecules and CD1d, respectively. We have determined the cell surface phenotype, Ag specificity, and functional capacity of a population of these cells. Using x-ray crystallography, we have elucidated the structural architecture of two δ/αβTCRs and show how these TCRs can recognize monomorphic and polymorphic Ag-presenting molecules via distinct mechanisms. Accordingly, we highlight a population of δ/αβ T cells that bind Ag by way of both Vδ and Vβ genes, thus reflecting a greater level of diversity and functional potential within the T cell lineage.
with our previous study (clone TS8.2; Uldrich et al., 2013) revealed that some V$\beta_1$+ cells coexpressed a TCR-$\beta$ chain rather than a TCR-$\alpha$ chain (Fig. 1a).

Single cell TCR sequencing of individual V$\beta_1$+ TCR-$\beta$+ CD1d–GalCer tetramer+ T cells confirmed the expression of the V$\beta_1$ gene recombined to J$\alpha$ gene and C$\alpha$ constant region (Fig. 1b).

**RESULTS**

**Identification of CD1d–$\alpha$-GalCer tetramer–reactive V$\beta_1$+ cells expressing a $\delta$/$\alpha\beta$ TCR.**

We previously identified CD1d–$\alpha$-GalCer cells, expanded in vitro, and analyzed by flow cytometry. CD3+ T cells were analyzed for V$\beta_1$ (clone A13) versus CD1d-endogenous tetramer and CD1d–$\alpha$-GalCer tetramer (left-hand density plots). CD3+ CD1d–$\alpha$-GalCer tetramer– V$\beta_1$+ T cells were analyzed for $\alpha\beta$ TCR and $\gamma\delta$ TCR (middle density plots). CD3+ CD1d–$\alpha$-GalCer tetramer– $\gamma\delta$ TCR– V$\beta_1$+ Type I NKT cells were analyzed for CD4, CD8, and V$\beta_11$ expression (right-hand plots).

Data shown represent five healthy donors. (b) TCRs from CD3+ CD1d–$\alpha$-GalCer tetramer– V$\beta_1$+ $\gamma\delta$ TCR+ cells derived from CD1d–$\alpha$-GalCer tetramer–enriched and in vitro expanded PBMC samples were sequenced. Data shown are unique sequences, derived from five separate donors, performed across three separate experiments.

(c) Percentage of CD1d-restricted $\delta$/$\alpha\beta$ T cells within the in vitro expanded CD1d–$\alpha$-GalCer tetramer– T cell population from 30 healthy donors. Donors in which no clear population of CD1d-restricted $\delta$/$\alpha\beta$ T cells were observed were given an arbitrary value of 0.01%. Horizontal line indicates the mean.

Figure 1. Identification of CD1d–$\alpha$-GalCer tetramer–reactive V$\beta_1$+ cells expressing a $\delta$/$\alpha\beta$ TCR. (a) PBMCs from healthy donors were enriched for CD1d–$\alpha$-GalCer cells, expanded in vitro, and analyzed by flow cytometry. CD3+ T cells were analyzed for V$\beta_1$ (clone A13) versus CD1d-endogenous tetramer and CD1d–$\alpha$-GalCer tetramer (left-hand density plots). CD3+ CD1d–$\alpha$-GalCer tetramer– V$\beta_1$+ T cells were analyzed for $\alpha\beta$ TCR and $\gamma\delta$ TCR (middle density plots). CD3+ CD1d–$\alpha$-GalCer tetramer– $\gamma\delta$ TCR– V$\beta_1$+ Type I NKT cells were analyzed for CD4, CD8, and V$\beta_11$ expression (right-hand plots).
Figure 2. Lipid Ag reactivity of δαβTCR⁺ T cells. (a) PBMCs from healthy donors were treated as in Fig. 1a and analyzed by flow cytometry. Plots show CD1d-α-GalCer tetramer-enriched and in vitro expanded PBMCs, depicting CD1d tetramer versus Vδ1 staining on CD3⁺ γδTCR⁻ cells using a panel of lipid Ag tetramers. Numbers on each plot represent the mean fluorescence intensity within gated regions. (b) The relative binding affinity, based on mean fluorescence intensity, of each lipid Ag is shown for type I NKT cells (CD3⁺ CD1d-α-GalCer⁺ γδTCR⁻ Vδ1⁻, open symbols) and for CD1d-restricted δαβTCR⁺ cells (CD3⁺ CD1d-α-GalCer⁺ γδTCR⁺ Vδ1⁺, closed symbols). Data were normalized against endogenous CD1d tetramer (indicated by the dashed red line). Each donor is represented by a different symbol.
TCR sequence analysis of these cells from five different donors revealed nine distinct hybrid Vβ1–Jα–Ca–TCR-β chains from within the Vβ1+ CD1d–α-GalCer tetramer+ population (Fig. 1 b). Thus, the Vβ1-specific antibody (clone A13) was capable of recognizing TCRs where the Vβ1 gene is recombined to Jα and Ca genes, whereas the other Vβ1-specific antibody (clone TS8.2) did not bind to these hybrid TCRs (not depicted). Accordingly, we termed CD1d–α-GalCer tetramer+ cells that express δ/αβTCRs as CD1d-restricted δ/αβ T cells.

In contrast to many of the CD1d-restricted Vβ1+ γδ T cells identified in our previous study that were only partially dependent on CD1d-bound lipid Ag (Uldrich et al., 2013), CD1d-restricted δ/αβ T cells showed an absolute requirement for CD1d-Ag, as “unloaded” (“CD1d/Endo”) human CD1d tetramers failed to bind these cells (Fig. 1 a). Analysis of in vitro expanded type I NKT cells, CD1d-restricted γδ T cells, and CD1d-restricted δ/αβ T cells revealed that the CD1d-restricted δ/αβ T cells typically expressed low levels of CD161 (not depicted). Despite the ability of Vβ1 to pair with several different TCR-β chains (Fig. 1 b), the TCR-β chain (Vβ11) common to type I NKT cells, was not detected on CD1d-restricted δ/αβ T cells (Fig. 1 a). CD4 and CD8 expression by δ/αβ T cells varied between donors and they were mostly CD4+CD8– or CD4–CD8+ (Fig. 1 a), and this typically differed from CD4 and CD8 expression on type I NKT cells from within the same donors. Similar to CD1d-restricted γδ T cells (Uldrich et al., 2013), human CD1d-restricted δ/αβ T cells did not stain with mouse CD1d–α-GalCer tetramer, unlike human type I NKT cells which showed strong cross-reactivity to mouse CD1d (not depicted; Brossay et al., 1998). We detected a clear population of CD1d-restricted δ/αβ T cells within the expanded CD1d–α-GalCer–reactive T cell population in 13 out of 30 donors (Fig. 1 c). In most cases, CD1d-restricted δ/αβ T cells represented <1% of total CD1d–α-GalCer–reactive cells, although they were higher in some individuals, including one individual in which they were over 50% of CD1d–α-GalCer–reactive cells (Fig. 1 c). Accordingly, CD1d-restricted δ/αβ T cells represent a novel subset of human CD1d–α-GalCer–reactive T cells, which are distinct from both type I NKT cells and CD1d-restricted γδ T cells.

**Ag specificity of δ/αβ CD1d-restricted T cells**

To evaluate the Ag specificity of CD1d-restricted δ/αβ T cells, CD1d–α-GalCer tetramer+ cells were isolated and expanded from six different donor PBMC samples and stained with CD1d tetramers loaded with α-GalCer, α-glucosylceramide (α-GlcCer), sulfatide, 3′-deoxy-α-GalCer, 4′-deoxy-α-GalCer, and OCH, known NKT cell ligands (Wun et al., 2010; Rossjohn et al., 2012). Tetramer staining of CD1d-restricted δ/αβ T cells was compared with tetramer staining of type I NKT cells from the same donors (Fig. 2, a and b). CD1d-restricted δ/αβ T cells bound α-GalCer–loaded CD1d tetramer in an Ag-dependent manner as these cells failed to stain with unloaded CD1d tetramer containing endogenous (“endo”) Ag (Fig. 2, a and b). Most CD1d-restricted δ/αβ T cells, with the exception of these cells from donor 2, did not bind to CD1d–α-GlcCer tetramer, which is in stark contrast to type I NKT cells from the same donors that stained brightly with this tetramer (Fig. 2, a and b). The CD1d-restricted δ/αβ T cells were also unreactive to sulfatide-loaded CD1d tetramer, whereas reactivity against 3′-deoxy- and 4′-deoxy-α-GalCer was highly variable between individual donors (Fig. 2, a and b). We have previously established that human type I NKT cells are very sensitive to the loss of the 3′-OH group on α-GalCer (Wun et al., 2012). In contrast, we show here that in at least some individuals, human CD1d-restricted δ/αβ T cells (from donors 1, 3, and 4) were clearly stained by the CD1d-3′-deoxy-α-GalCer tetramer, whereas most of the type I NKT cells from those same donors were not (Fig. 2, a and b). Conversely, CD1d-restricted δ/αβ T cells from donors 4 and 6 stained poorly with CD1d-4′-deoxy-α-GalCer tetramers, whereas the type I NKT cells from the same donors were brightly labeled (Fig. 2, a and b). Furthermore, CD1d-restricted δ/αβ T cells from donor 5 showed considerably brighter staining with 4′-deoxy-α-GalCer compared with CD1d–α-GalCer tetramer (Fig. 2, a and b). Reactivity of CD1d-restricted δ/αβ T cells against the OCH Ag showed that δ/αβ T cells from four donors (1, 3, 4, and 5) were capable of binding to this Ag (Fig. 2, a and b), whereas, as expected, most type I NKT cells failed to bind OCH (Matulis et al., 2010; Wun et al., 2011). Collectively, these data highlight that CD1d-restricted δ/αβ T cells are capable of recognizing glycolipids presented by CD1d, and the δ/αβTCR composition imbues these cells with a different pattern of glycolipid Ag specificity that distinguishes them from type I NKT cells from the same donors.

**δ/αβ T cells are abundant within human PBMCs**

Having determined that some δ/αβ T cells were present within the CD1d–α-GalCer–reactive T cell population, we next examined δ/αβTCRs within the general population of Vβ1+ T cells in humans by analyzing freshly isolated PBMCs. Using the anti-Vβ1 antibody clone A13, we determined that many Vβ1+ T cells coexpress a TCR-β chain, rather than a TCR-γ chain, and are therefore δ/αβ T cells rather than γδ T cells (Fig. 3 a). The ratio of δ/αβ to γδ T cells within the Vβ1+ population varied widely, from <5% to >80% δ/αβ T cells, with a mean of ~45% (Fig. 3 b). Further investigation of the cell surface phenotype of δ/αβ T cells, in comparison with αβ T cells, revealed that δ/αβ T cells can coexpress CD4 or CD8, although the ratio of CD4 to CD8 was generally different from that observed for αβ T cells, with more CD8+ and less CD4+ δ/αβ T cells than αβ T cells (p < 0.05; Wilcoxon paired ranked test). γδ T cells from the same donors were predominantly CD4+CD8– or CD8+ (Fig. 3 c). TCR-Vβ profiling of these cells indicated that they express a broad range of TCR-β chains, and moreover, the representation of Vβ chains used by δ/αβ T cells did not necessarily parallel that of the αβ T cells from the same donors (Fig. 3 d). This is exemplified by the Vβ8 population, which ranged...
Figure 3. **TCR-V61+ cells consist of γδTCR+ and δαβTCR+ subsets.** (a) CD14-CD19- lymphocytes from a representative healthy blood donor, were analyzed by flow cytometry. Plots show CD3+V61+ and CD3+V61- PBMC subsets, gated as indicated (left plot), and then subsets labeled with anti-αβTCR (identifying δαβ T cells and αβ T cells) versus anti-γδTCR (identifying γδ T cells) are shown (second plots). Expression of CD4 and CD8 on these subsets is also shown (right-hand plots). (b) The percentage of V61+ cells that were αβTCR+ (i.e., δαβ T cells) was calculated for eight donors using the gating strategy shown in a. (c) The percentage of CD4-CD8- (double negative [DN]), CD4+CD8- (CD4+), and CD4-CD8+ (CD8+) cells within V61+ αβTCR+ (δαβ), V61+ αβTCR-αβ (αβ), V61+ γδTCR+ (V61+ γδ), and V61- γδTCR+ (V61- γδ) is shown, with each symbol representing a separate donor (n = 6) and bar graphs depicting the mean value. (d) Healthy human PBMCs were stained with a panel of anti–TCR-β–specific antibodies. Plots show the percentage of V61+ αβTCR+ (δαβ, left) and V61- αβTCR+ (αβ, right) T cells that bound to each Vβ antibody. Graphs depict n = 4 donors with each symbol representing a different donor. (e) δαβ T cells, αβ T cells, V61+ γδ T cells, and V61- γδ T cells (as defined in c) were purified by FACS and stimulated with anti-CD3/CD28 beads for 72 h. Cytokines in culture supernatants were measured by cytometric bead array. Each data point represents the mean of n = 2–4 replicates from one donor, with n = 4 donors and each symbol shape representing a different donor. (f) δαβ, αβ, V61+ γδ, and V61- γδ T cells from a representative donor (as defined in c) were stimulated for 4 h with PMA/ionomycin (top plots) or unstimulated (bottom plots), and IFN-γ and TNF were measured by intracellular cytokine staining. (g) Percentage of IFN-γ- and TNF-producing cells stimulated as in e, with each symbol representing a separate donor (n = 6). Symbols indicating specific donors within panels do not correlate with the same donors between different panels. (b, e, and g) Horizontal lines indicate the mean.
from 3 to 15% of αβ T cells in four donors, whereas δ/αβ T cells from the same four donors were <3% Vβ8+. Thus, these data highlight that V61+ δ/αβ T cells are present in similar frequency to V61+ γδ T cells in human peripheral blood. Moreover, aside from their unusual TCR, δ/αβ T cells are also distinct from both αβ and γδ T cells with regard to their CD4 or CD8 coexpression profiles and TCR-β repertoire.

δ/αβTCRs transmit activation signals

Using flow cytometric sorting, we purified δ/αβ, αβ, and γδ T cells from several human donors and stimulated them in vitro for 3 d in the presence of anti-CD3– and CD28-coated beads to measure their relative ability to produce cytokines in response to TCR cross-linking. These results (Fig. 3 e) demonstrated that δ/αβ T cells are responsive to TCR cross-linking, resulting in abundant cytokine production, including IL-2, IL-4, IL-13, IL-17, IFN-γ, GM-CSF, and TNF, at levels comparable with that of αβ T cells and clearly distinct from that observed for both V61+ and Vδ1+ γδ T cells from matched donors. The response from the γδ T cells was generally lower than that of the δ/αβ T cells for IL-2, IFN-γ, and TNF, whereas the V61+ γδ T cells produced comparable GM-CSF and IL-13 to the δ/αβ T cells after anti-CD3/CD28 stimulation (Fig. 3 e). To examine the cytokine-producing potential of these different cell types further, we compared all four subsets after a brief (4 h) stimulation with PMA and ionomycin and measured IFN-γ and TNF by intracellular cytokine staining (Fig. 3 f and g). These data indicated that both V61+ and Vδ1+ γδ T cells can produce abundant IFN-γ and TNF and thus were functionally similar using this method of stimulation.

We next wanted to determine whether direct ligation of the δ/αβTCR with defined Ag could result in cellular activation. To achieve this, we stably transduced a CD1d-restricted δ/αβTCR (clone 9B4) and an irrelevant pHLA-reactive TCR into the αβTCR-deficient Jurkat-76 cell line and cultured these cells in the presence of α-GalCer or α-GlcCer. As these cell lines express CD1d (not depicted), no Ag-presenting cells were added. CD69 up-regulation was used as an indicator of Ag-mediated cellular activation (Fig. 4, a and b). Type I NKT TCR (NK1T15)–transduced SKW3 cells, which also express CD1d, were included as a positive control in this assay. The control HLA-restricted TCR–transduced cells did not respond to either glycolipid Ag. These experiments demonstrated that the δ/αβTCR was capable of recognizing α-GalCer Ag presented by CD1d and, furthermore, that this recognition event could transmit cellular activation signals (Fig. 4 a). Also, although NK1T15 TCR–transduced cells were capable of recognizing α-GalCer and α-GlcCer equally, the δ/αβTCR–transduced cells were only capable of responding to α-GalCer (Fig. 4 b), consistent with our CD1d tetramer staining of CD1d–restricted δ/αβ T cells from in vitro expanded PBMCs (Fig. 2). Furthermore, the δ/αβTCR–transduced cells responded to three different analogues of α-GalCer with different acyl chains, including C24:1 (Fig. 4 a), C20:2 (Fig. 4 b), and C26 analogues (not depicted). Collectively, these data indicate that δ/αβTCRs are capable of transmitting activating signals in response to specific glycolipid Ags, resulting in cellular activation and diverse cytokine production.

Specificity of 9B4 δ/αβTCR to CD1d-Ag

To study the molecular basis for the binding of a δ/αβTCR to CD1d-Ag, we first determined the specificity of an isolated δ/αβTCR (9B4) for CD1d tetramer–glycolipid, using the same panel of glycolipid Ags as shown in Fig. 2. The pattern was clearly distinct from that of cells transduced with type I NKT TCR (clone NK1T15), and cells transduced with an irrelevant pHLA-specific TCR showed no staining with any of the CD1d-Ag tetramers tested. Namely, 9B4 δ/αβTCR–transduced Jurkat cells recapitulated the pattern of reactivity observed for the δ/αβ T cells from donor 1 (Fig. 2), demonstrating strong staining with CD1d tetramers loaded with α-GalCer and 4’-deoxy-α-GalCer, moderate staining with 3’-deoxy-α-GalCer and OCH, and very little reactivity with α-GlcCer and sulfatide–loaded CD1d tetramer (Fig. 5 a). We next generated a soluble version of this TCR using previously described methods (Kjer-Nielsen et al., 2006), and the soluble 9B4 δ/αβTCR exhibited a molecular mass of 48 kD and reacted with an anti-Vδ1 mAb and an anti-TCR Cα mAb, thereby indicating that the δ/αβTCR had refolded properly (not depicted). Surface plasmon resonance (SPR) was used to investigate the affinity of the interaction between
the 9B4 δ/αβTCR and CD1d loaded with C26:0 α-GalCer (Fig. 5 b). The human NKT15 type I Vα24+ αβTCR (Kjer-Nielsen et al., 2006) was included as a positive control (Fig. 5 b). No notable autoreactivity to CD1d-endogenous was observed for the 9B4 δ/αβTCR, in stark contrast to the autoreactivity previously observed for the CD1d-restricted 9C2 γδTCR (Uldrich et al., 2013). The 9B4 δ/αβTCR bound to CD1d–α-GalCer with very high affinity (Kd = 66 nM), even compared with the type I NKT15 αβTCR (Kd = 190 nM; Fig. 5 b), and markedly higher than the 9C2 V81+ γδTCR (Kd of 15 µM; Uldrich et al., 2013). This higher affinity of the 9B4 δ/αβTCR is partly caused by a slower dissociation rate (half-life of 24 s compared with 3.5 s for the NKT15 αβTCR). Thus, the 9B4 δ/αβTCR binds strongly to CD1d–α-GalCer and is highly sensitive to the type of CD1d-restricted lipid Ags presented.

The δ/αβTCR–CD1d-Ag complex
To understand how the 9B4 δ/αβTCR recognized CD1d–α-GalCer, we determined the crystal structure of the ternary complex (Fig. 6 a and Table S1) as well as the nonliganded 9B4 TCR (Table S1). The 9B4 TCR docked orthogonally (73°) above the A-pocket of CD1d, relative to the CD1d cleft, burying ~1,050 Å², and interacted with residues ranging from 58 to 79 of the α1 helix and from 153 to 171 of the α2 helix of CD1d (Fig. 6, a and b). This mode of recognition differed markedly from type I NKT TCR–CD1d–α-GalCer recognition, whereupon the Vα24-Vδ1 TCR sat parallel over the F-pocket of CD1d (Fig. 6, a and b; Borg et al., 2007). Accordingly, the interatomic contacts and roles of the individual CDR loops at the type I NKT TCR–CD1d–α-GalCer interface were markedly different to those at the 9B4 TCR–CD1d–α-GalCer interface (Fig. 6 c). Indeed, the δ/αβTCR–CD1d–α-GalCer complex was more similar to the 9C2 γδTCR–CD1d–α-GalCer complex (Fig. 6, a–c), which sat orthogonally (83°) over the A-pocket (buried surface area [BSA] of ~950 Å²), although the 9B4 TCR variable regions were shifted more toward to the center of the CD1d cleft (shift of 7 Å for the TCR-β chain and 3 Å for the TCR-Vδ/α chain, relative to the 9C2 TCR-γ and δ chains,
chain, both the CDR1β and CDR3β made contributions to the interface (13% and 11% of the total BSA, respectively; Figs. 6 c and 7 a). Namely, the aliphatic moiety of Arg79 of CD1d was pincered by Gln29δ from the CDR1δ loop and the framework residue Leu77δ (Fig. 7 a). In addition, Arg79 respectively), thereby bringing the δ/αβTCR closer to the α-GalCer headgroup.

The δ/αβTCR-mediated recognition of CD1d was dominated by the Vδ1 chain (66% of the total BSA), relative to the Vβ2 chain (33% of the total BSA; Table S2). For the TCR-β chain, both the CDR1β and CDR3β made contributions to the interface (13% and 11% of the total BSA, respectively; Figs. 6 c and 7 a). Namely, the aliphatic moiety of Arg79 of CD1d was pincered by Gln29β from the CDR1β loop and the framework residue Leu77β (Fig. 7 a). In addition, Arg79...
contacted Thr31β, which, together with the adjacent Thr32β, packed against Val72 from CD1d, with Thr32β and Asn57β also interacting with His68 of CD1d (Fig. 7 a). The CDR3β/α loop (13% of the total BSA) extended toward the α2 helix of CD1d, with its main chain packed against α-GalCer (discussed below) and contacted residues from the α1 and α2 helices (Fig. 7 b). Namely, Trp160 of CD1d was positioned between Val108α and Thr109α, the latter of which also contacted Ile69 and Trp153 of CD1d (Fig. 7 b). Trp153 is a position that differs in mouse CD1d (Gly155; Godfrey et al., 2005), thereby providing a basis of why the δαβTCR does not cross-react onto mouse CD1d–β-GalCer (not depicted).

The Vβ1-mediated contacts with CD1d are dominated by the CDR1β loop (38% of the total BSA), with further contributions from the framework region (15% BSA; Table S2). Although the CDR2β loop did not interact with CD1d, the flanking framework regions did (Fig. 7 c). Here, Arg55β extended down toward the α2 helix to form a salt bridge with Glu156, whereas Asp66β and Lys71β mediated contacts with the glycosylation site attached to Asn163 of CD1d (Fig. 7 c). These framework-mediated interactions with CD1d were analogous to those observed in the 9C2 γδTCR–CD1d–Ag complex (Uldrich et al., 2013).

The CDR1ß loop sat over the distal end of the A’ pocket of CD1d and exclusively contacted CD1d (Fig. 7 d). The interactions were dominated by two Trp residues (Trp29 and Trp30) that were flanked by Ser residues (Ser28β and Ser31β) whose side chains formed vdW interactions with CD1d, while their main chains hydrogen bonded with CD1d residues (Gln62, Gln168, and Thr160; Fig. 7 d). Trp29β was wedged between the α1 and α2 helices of CD1d, packed against the aliphatic side chains of Gln62, Thr65, and Leu66 and the aromatic ring of Trp160 (Fig. 7 d). Trp30β was positioned at the periphery of CD1d, stacked principally against Gln168 (Fig. 7 d).
These CDR1β loop-mediated contacts by the δ/αβTCR were very similar to those observed in the 9C2 γδTCR–CD1d-Ag structure (Fig. 7 e). Thus, the germline-encoded regions of the Vβ1 domain of the δ/αβTCR adopted a very similar mode of CD1d-Ag recognition as the γδTCR. This showed that Vβ1 is not only capable of binding CD1d when rearranged with a Dδ-β-encoded CDR3β motif (in the context of a γδTCR), but also when rearranged with a permissive α segment in the context of a δ/αβTCR.

The structure of 9B4 TCR in the unligated state enabled us to assess the extent of plasticity of the CDR loops upon CD1d–α-GalCer engagement. Upon ligation, the 9B4 TCR variable regions did not significantly change conformation (root-mean-square deviation [rmsd] 0.5 Å for both Vβ1 and Vβ2); however, surprisingly, the positioning of the constant regions was significantly altered by approximately a 17° shift in the elbow angle between the V and C domains in a direction toward the F’ pocket of CD1d (not depicted). In contrast, there was minimal change in the conformation of the CDR loops upon ligation (rmsd ranging from 0.3 to 1 Å), with the biggest alteration observed for Trp30 of the CDR1β loop, which flipped upon ligation to avoid clashing with the α2 helix of CD1d (Fig. 7 f). Thus, analogous to αβTCR and γδTCR recognition of CD1d–α-GalCer, a rigid “lock-and-key” mechanism underpinned δ/αβTCR ligation (Borg et al., 2007; Pellicci et al., 2009; Uldrich et al., 2013).

Lipid Ag recognition
The electron density at the interface and for the α-GalCer Ag in the 9B4 ternary complex was unambiguous (Fig. 8 a), thereby permitting a comparison of how three TCRs arising from distinct lineages, the αβTCR, the γδTCR, and the δ/αβTCR, interacted with the same Ag-presenting molecule bound to the same ligand. Although the 9B4 TCR–Vβ1 chain interaction with CD1d, α-GalCer recognition was mediated entirely by the TCR-β chain, specifically being sequestered by the CDR1β and CDR3β loops via a polar interaction network. This recognition is notably dominated by water-mediated contacts and interactions that involved the main chain of the δ/αβTCR (Fig. 8 b). Namely, all of the hydroxyl moieties of the α-GalCer headgroup, with the exception of the 2′-OH moiety, were involved in hydrogen bonding with the TCR-β chain of the δ/αβTCR. Specifically, Pro107β hydrogen bonded to the 4′-OH, whereas three water-mediated hydrogen bonds were formed between Glu29β, Thr31β, and Gly109β and the 4′-OH, 6′-OH, and 3′-OH moieties of α-GalCer, respectively (Fig. 8 b).

The structural basis for α-GalCer recognition by the δ/αβTCR was markedly distinct to that mediated via the 9C2 γδTCR and the NKT15 αβTCR (Fig. 8, b–d; Borg et al., 2007; Pellicci et al., 2009; Uldrich et al., 2013). Within the γδTCR–CD1d–α-GalCer complex, the CDR3γy loop exclusively contacted the α-GalCer headgroup, with Arg103γy and Tyr111γ hydrogen bonded to the 3′-OH and 4′-OH moieties (Fig. 8 c; Uldrich et al., 2013). In the NKT15 complex, the interactions with α-GalCer were exclusively mediated via the invariant TCR-α chain, where the α-GalCer headgroup sat underneath the CDR1α loop and abutted the CDR3α loop (Fig. 8 d; Borg et al., 2007; Pellicci et al., 2009). Here, the galactose 2′-OH, 3′-OH, and 4′-OH moieties were involved in
polar-mediated contacts with the αβTCR. Accordingly, the αβTCR, γδTCR, and δ/αβTCRs contact the same lipid Ag via markedly distinct binding mechanisms.

Molecular basis of δ/αβTCR recognition of pHLA-I

We determined that there are many other δ/αβTCRs in healthy humans (Fig. 3), although the specificity of these cells is unclear. Nevertheless, previously, we had described a panel of T cell clones that were restricted to HLA-B*35:01–IPS complex (not depicted), which fell within the normal affinity range observed for αβTCR–MHC-I interactions (Gras et al., 2012). Next, we solved the structure of the clone 12 δ/αβTCR in complex with HLA-B*35:01–IPS Ag at 3.0 Å resolution (Fig. 9 and Table S1). The clone 12 TCR docked 56° across the HLA-B*35:01–IPS cleft (Fig. 9 a), within the range of standard αβTCR–pHLA-I complexes (Gras et al., 2012). In addition, the clone 12 TCR ternary complex possessed features typically associated with αβTCR–pHLA-I complexes (Gras et al., 2012). For example, the BSA of the TCR interface was 1,014 Å² (total BSA of the TCR/pHLA interface = 2,040 Å², within the range of αβTCR–pMHC-I interactions).

The clone 12 δ/αβTCR predominantly used its TCR-δ chain (65% BSA) to engage the pHLA complex, where the interaction was dominated by the CDR1β (27%), CDR3β (20%), and Vδ1 framework (18% BSA) residues, whereas the CDR3α loop was the principal contributor (19% BSA) from the β chain (Fig. 9 b). Although the CDR3β loop mainly contacted the peptide Ag (discussed below), Tyr112H bonded to Gln155, an MHC residue which is frequently contacted by αβTCRs (Burrows et al., 2010). The other contacts mediated via the β chain included a salt bridge (Arg30/β to Gln76) and a hydrogen bond with Asn80 and two residues from the CDR3β loop mediating contacts. Thus the germline-encoded TCR-β chain–MHC contacts were very limited (Table S3 and not depicted).

The long CDR3δ/α loop (17 residues) formed an extended hairpin structure that sat above the HLA α1 helix (residues 65–72), while pointing away from the peptide (Fig. 10 a). Here, two residues from the N-region and two residues from the Jα gene segment were involved in hydrophobic interactions with HLA-B*35:01 (Table S3). Additionally, Thr115δ hydrogen bonded to Thr69 of the α1 helix (Fig. 10 a and Table S3). Interestingly, the clone 12 TCR framework residue Arg55δ interacted with the HLA-B*35:01 molecule in a similar fashion to that observed for the 9B4 interaction with the CD1d molecule (Fig. 8 c), whereby the Arg55δ points toward the α2 helix and hydrogen bonded to Ala150 (Fig. 10 b). In addition, Glu67δ formed a salt bridge with Arg151, and Asp66δ made hydrophobic interactions with Glu154, Gln155, and Ala158 (Fig. 10 b).

The CDR1δ loop, with its two large Trp residues (Trp29δ and Trp30δ), contacted the N-terminal side of the Ag-binding cleft of the HLA-I molecule on both helices (Fig. 10 c). Trp29δ packed against Arg52, whereas Trp30δ extended toward the α2 helix, stacking between the short aliphatic chains of Ala158 and Leu163. Furthermore, another aromatic residue from the CDR1δ loop, Tyr33δ, packed against Gln155. Accordingly, the CDR1δ loop played an extensive role in enabling this HLA-I–restricted response. Nevertheless, the placement of the CDR1δ loop atop the HLA-I and CD1d differed

Figure 9. Clone 12 TCR–HLA-B*35:01–IPS structure. (a) Overview of the clone 12 TCR (β/α chain in pale pink, β chain in pale blue) in complex with the IPS peptide (purple sticks) bound to the HLA-B*35:01 molecule (white cartoon) and β-2-microglobulin (β2m, black cartoon). The clone 12 TCR CDR loops are colored in purple, green, and yellow for the CDR1/2/3α and red, blue, and orange for the CDR1/2/3β. (b) Atomic footprint of the clone 12 TCR, colored by CDR loops according to panel a, on the surface of the IPS peptide (gray surface) bound to the HLA-B*35:01 molecule (white surface). The black spheres represent the mass center of the Vδ/α and Vβ domains.
markedly (Fig. 10 d), indicating the versatility of the same germline-encoded region to interact with diverse Ag presentation platforms.

Peptide Ag recognition

Although the TCR-δ chain of clone 12 TCR dominated the interaction with the HLA-I molecule, it made very limited contacts with the IPS peptide. Specifically, the contacts from the δ chain were focused solely onto P4-Ile, which formed hydrophobic contacts with CDR1δ and CDR3δ/α (Fig. 10 e). Notably, however, these Vδ1-driven contacts appeared to play a key role in interacting with P4-Ile, as mutation of the P4-Ile to an Ala dramatically decreased the binding affinity of the δ/αβTCR (Kd > 200 µM; not depicted). The majority of contacts to the IPS epitope were made by the VB chain, principally a framework residue (Arg66β) and the CDR3β loop (Fig. 10 f). Arg66β contacted the P8-His, and interestingly, Arg66β is found in the Vβ5, Vβ1, and Vβ6.8 genes only. The structure of Vβ1+ TCR also showed how this framework TCR–β chain residue directly contacted the Epstein–Barr virus–derived epitope (Gras et al., 2010). The CDR3β loop formed most of the interactions with the IPS peptide, contacting the P4-Ile, P6-Val, P7-His, and P8-His residues, and many of these peptide residues were important for the interaction (Fig. 10 f and Table S3). Here, the Tyr112β inserted its aromatic ring between the main chain of the P4-Ile and P6-Val, whereas Tyr113β sat atop the P7-His and contacted the P6-Val (Fig. 10 f). Furthermore, Glu109β contacted P7-His and hydrogen bonded to the main chain of P8-His (Fig. 10 f). Thus, the TCR–β chain plays a major role in mediating contacts with the peptide, with a key contribution from the CDR1β loop. In summary, we have demonstrated that T cells expressing δ/αβTCRs can recognize both CD1d–glycolipid Ags as well as MHC–peptide Ags and have provided the molecular bases for how these interactions occur.

DISCUSSION

Since the discovery of the TCR genes in the 1980s (Hedrick et al., 1984), considerable insight into αβ T cell–mediated immunity, and to a lesser extent γδ T cell immunity, has been gleaned. αβ T cells are frequently associated with the adaptive immune response, whereas γδ T cells are considered to exhibit more innate-like features (Vantourout and Hayday, 2013). However, such functional division of these T cell subsets has
become blurred. For example, although it was traditionally considered that αβ T cells recognized peptides presented by MHC molecules, innate-like T cells expressing semi-invariant αβTCRs can also recognize distinct nonpeptide-based Ags. Namely, CD1d-restricted type II NKT cells and MR1-restricted MAIT cells, upon activation by lipids and vitamin B metabolites, respectively, rapidly produce an array of cytokines (Rossjohn et al., 2012; Birkinshaw et al., 2014; Gapin, 2014). Furthermore, it is emerging that TCRs can adopt distinct, as well as similar, docking strategies in binding to peptide, lipid, or small molecule metabolites that are bound to their respective Ag-presenting molecules, thereby highlighting the adaptability of the αβTCR scaffold (Bhali et al., 2014).

Whereas most T cells either fall into the αβTCR+ or the γδTCR+ fraction, the existence of TCR gene rearrangements involving unusual combinations of α, β, γ, and δ chain genes have sporadically been reported in the literature (Hochstenbach and Brenner, 1989; Miossec et al., 1990, 1991; Peyrat et al., 1995; Bowen et al., 2014), although the significance of these unusual gene recombination events, the Ags to which they responded to, and the molecular bases for these interactions are unclear. Here, we have identified populations of Vβ1+ δ/αβ T cells, demonstrating that they are present with a similar frequency to Vγ6 T cells. Importantly, δ/αβTCRs appear to be functional, capable of transmitting activation signals to the T cells that express them. To understand the molecular bases for how these TCRs recognize Ag, we have focused on two distinct δ/αβTCRs, a CD1d-restricted α-GalCer-reactive δ/αβTCR and an HLA-restricted viral peptide-reactive δ/αβTCR.

We show that the δ/αβTCR architecture resembles that of αβTCRs and γδTCRs and provide a basis for how the Vβ1 chain can pair with the TCR-β chain. The binding of the δ/αβTCR to CD1d was dependent on the nature of the Ag bound, and the mode of recognition was markedly distinct from that of type I NKT TCR–CD1d–α-GalCer docking, more closely resembling that of the γδTCR–CD1d–α-GalCer complex (Uldrich et al., 2013). Indeed, the CDR1δ loop-mediated contacts between the δ/αβTCR and γδTCR were very similar (Uldrich et al., 2013), suggesting a critical role for this region of the TCR in the CD1d restriction of these cells. In contrast, the interactions of the δ/αβTCR with α-GalCer were markedly different from the γδTCR–CD1d–α-GalCer complex. Thus, although the CDR1δ loop appears to represent a focal point for the CD1d-restricted response, the nature of the docking mode is nevertheless fine-tuned by the pairing of the TCR-β or TCR-γ chain and the hypervariable CDR3 loops, in a manner which is strikingly unique to this hybrid TCR. Furthermore, this is the first example of a CD1d–α-GalCer–reactive TCR that does not bind with a parallel docking mode over the Fpocket of CD1d, a characteristic of all αβTCR+ type I NKT cells (Rossjohn et al., 2012).

We also demonstrate how a δ/αβTCR can engage an HLA-I–peptide complex. Notably, the δ/αβTCR, docking on CD1d-lipid and HLA-I–peptide differed, despite these two δ/αβTCRs sharing the same Vβ1-Jα52+ chain. Nevertheless, in both cases, the Vβ1 framework region, coupled with the Trp-rich CDR1δ loop, played a prominent role in contacting the respective Ag-presenting molecules. These interactions differed in their positioning on the Ag-binding platforms, thereby highlighting how the same germline–encoded region can play key, yet disparate, roles in binding to polymorphic and monomorphic Ag-presenting molecules.

In summary, our study highlights the existence, and molecular bases for Ag recognition, of T cells that express hybrid TCRs comprised of the Vβ1 variable region rearranged to diverse Jα genes and the Cα constant region and paired to a diverse range of TCR–β chains. These hybrid δ/αβTCR+ T cells are surprisingly abundant in humans and have unique characteristics both at the cellular and molecular level, thus adding a level of diversity that extends beyond the αβ T cell and γδ T cell lineages. Collectively, these findings represent a large conceptual advance in our understanding of T cell–mediated immunity.

**MATERIALS AND METHODS**

**Accession codes.** The 9B4 TCR, monomer, 9B4 TCR–CD1d–α-GalCer ternary complex, and clone 12 TCR–HLA-B*3501–IPS ternary complex were deposited in the Protein Data Bank (PDB) under the accession codes 4WNQ, 4WO4, and 4QRR, respectively.

**Human CD1d-restricted δ/αβ T cell identification and isolation.** Healthy human PBMCs were obtained from the Australian Red Cross, ethics approval 13-04VIC-07 (Australian Red Cross) and 1035100.1 (University of Melbourne). PBMCs were isolated using a histopaque-1077 (Sigma-Aldrich) density gradient. CD3+ CD1d–α-GalCer tetramer+ cells were MACS (Miltenyi Biotec) and FACS sort-enriched using a FACSAnA III (BD). Enriched CD3+ CD1d–α-GalCer tetramer+ cells were expanded in vitro, essentially as previously described (Uldrich et al., 2013).

**Flow cytometry.** Enriched CD3+ CD1d–α-GalCer tetramer+ T cells were stained with Vβ1 (clone A13 supernatant, which can bind to Vβ1 when incorporated in hybrid Vβ1-Jα-Cα TCR, chains, was produced in L. Moretta’s laboratory), anti–mouse IgG (clone Poly 4053; BioLegend), 5% normal mouse serum, and then with antibodies specific for αβTCR (clone TV09.1A-31; BD), CD3ε (clone UCHT1; BD), CD8α (SK1; BD), CD4 (RPA-T4; BD), VB11 (C21; Beckman Coulter), CD69 (FN50; BD), γδTCR (11F2; BD), and CD161 (191B8; Miltenyi Biotec). Cells were costained with 7-aminoactinomycin D viability dye (Sigma-Aldrich) and with human CD1d tetramers (produced in-house), as previously described (Uldrich et al., 2013). TCR–Vβ repertoire analysis was performed using a TCR–Vβ repertoire kit (Beckman Coulter). Cells were analyzed using an LSR Fortessa (BD), and data were analyzed using FlowJo software (Tree Star Inc.).

**Lipids.** Lipids were dissolved in 0.5% Tyloxapol (Sigma-Aldrich) or Tween-sucrose-histidine buffer containing 0.5% vol/vol Tween-20, 57 mg/ml sucrose, and 7.5 mg/ml histidine. Lipids included α-GalCer C24:1 (PBS-44; provided by P. Savage, Brigham Young University, Provo, UT); α-GalCer C26 (Alexis); α-GalCer (C20:2); α-GlcCer C20:0, OCH, 3′,4′-dideoxy-α-GalCer (3′-deoxy-α-GalCer; 4′,3′-dideoxy-α-GalCer (4′-deoxy-α-GalCer; the deoxy analogues were provided by S. Keshipeddy and S. Richardson, University of Connecticut, Storrs, CT; Wun et al., 2011); and sulfatide C24:1 (Avanti Polar Lipids). For CD1d tetramer experiments, lipids were loaded into hybrid Vβ1-Jα52+ cells, demonstrating that they can bind to Vβ1 when incorporated in hybrid Vβ1-Jα-Cα TCR, chains, was produced in L. Moretta’s laboratory, anti–mouse IgG (clone Poly 4053; BioLegend), 5% normal mouse serum, and then with antibodies specific for αβTCR (clone TV09.1A-31; BD), CD3ε (clone UCHT1; BD), CD8α (SK1; BD), CD4 (RPA-T4; BD), VB11 (C21; Beckman Coulter), CD69 (FN50; BD), γδTCR (11F2; BD), and CD161 (191B8; Miltenyi Biotec). Cells were costained with 7-aminoactinomycin D viability dye (Sigma-Aldrich) and with human CD1d tetramers (produced in-house), as previously described (Uldrich et al., 2013). TCR–Vβ repertoire analysis was performed using a TCR–Vβ repertoire kit (Beckman Coulter). Cells were analyzed using an LSR Fortessa (BD), and data were analyzed using FlowJo software (Tree Star Inc.).

**Single cell PCR.** CD1d–GalCer tetramer Vβ1+ CD3+ γδTCR+ cells were single cell sorted and cDNA isolated using SuperScriptVII (Invitrogen) as per
the manufacturer’s instructions. Transcripts encoding for Vβ1 were amplified by two rounds of nested PCR, using Vβ1 external primer 5′-CAAGCCCATGTCATCAGTACCC-3′, Vβ1 internal primer 5′-CACTCCTCAGCACAAGAGATTG-3′, Cox external primer 5′-GACACGCTTGACATCAGCAGC-3′, and Cox internal primer 5′-TGTGCTCTTGAAGTCCATAG-3′. Transcripts encoding for the different Vβ genes were identified using multiplex nested PCR as previously described (Wang et al., 2012). PCR fragments were separated using a 1.5% agarose gel and DNA sequenced by Molecular Diagnostics, PCR as previously described (Wang et al., 2012). PCR fragments were separated using a 1.5% agarose gel and DNA sequenced by Molecular Diagnostics.

**Generation of cell lines.** CD1d-restricted β/ββTCR+ Jurkat cells (9B4), CD1d-restricted type I NKT TCR+ Jurkat or SKW3 cells (NKT15), and control HLA peptide–specific TCR+ Jurkat cells were generated essentially as previously described (Uldrich et al., 2013). Parental Jurkat-76 cells and SKW3 cells were provided by L. Kjer-Nielsen (University of Melbourne, Parkville, Victoria, Australia).

**SPR.** SPR experiments were conducted at 25°C on either a BIAcore 3000 (Biacore) or ProteOn XPR36 (Bio–Rad Laboratories) instrument with HBS buffer (10 mM Hepes, pH 7.4, 150 mM NaCl, and 0.005% surfactant P-20). In some instances the HBS buffer was supplemented with 1% bovine serum albumin to prevent nonspecific binding. The human αβTCR–specific mAb (clone 12H8; Borg et al., 2005) or streptavidin was coupled to CMS or GLC chips with standard amine coupling, respectively. Experiments were conducted as previously described (Gras et al., 2009; Pellicci et al., 2009). BIAevaluation version 3.1 (Biacore) or ProteOn Manager version 2.1 (Bio–Rad Laboratories) software was used for data analysis with 1.1 Langmuir binding model. The equilibrium data were analyzed using Prism (GraphPad Software).

**Cloning and expression of TCRs, human CD1d, and HLA.** Soluble TCRs, including CD1d-restricted β/ββTCR (9B4), type I NKT TCR (NKT15), and HLA–B*35–35:01–IPS (clone 12), were generated as previously described (Kjer-Nielsen et al., 2006). In brief, gene fragments encoding the TCR ectodomains were cloned into pET30 expression vector (EMD Millipore) and expressed in BL-21 E. coli cells. Inclusion body protein was prepared and refolded similar to that previously described in Clements et al. (2002) except in the presence of 5 M urea. Human CD1d with and without a Bir-A tag were expressed and purified as previously described (Borg et al., 2007; Uldrich et al., 2013). Soluble pHLA-B*35:01 was prepared as described previously (Gras et al., 2010). The 9B4 TCR–CD1d–α-GalCer and clone 12 TCR–HLA-B*35:01–IPS ternary complexes were purified by gel filtration from mixtures of the respective monomers.

**Structure determination.** Solutions of the 9B4 TCR monomer (at 5 mg/ml in 10 mM Tris-HCl, pH 8.0, and 150 mM NaCl) and 9B4 TCR–CD1d–α-GalCer ternary complex (at 10 mg/ml in 10 mM Tris-HCl, pH 8.7, and 150 mM NaCl) were crystallized at 20°C in 24% PEG 3350, 2% ethylene glycol, 10 mM l-Proline, and 10 mM spermidine. Crystals of 9B4 TCR–CD1d–α-GalCer and clone 12 TCR–HLA-B*35:01–IPS ternary complexes were purified by gel filtration from mixtures of the respective monomers. The versatility of the αβ TCR–HLA-B*35:01–IPS complex was solved as above, using the 9C2 TCR (Uldrich et al., 2013) and HLA-B*3501 without the peptide (pdb code 3LKN [Gras et al., 2010]) as search models. All structures were refined into the experimental maps by iterative rounds of model building using COOT (Emsley et al., 2010) and refinement with BUSTER (Bricogne et al., 2011). The electron density at the Ag interfaces was clear, with unbiased electron density for IPS peptide, the α-GalCer headgroup, and all CDR loops of the 9B4 and clone 12 TCR. Structural models were validated at the Research Collaboratory for Structural Bioinformatics Protein Data Bank Data Validation and Deposition Services website, and molecular illustrations were generated using the PyMOL package (DeLano, 2002). Domain superpositions and rmsd calculations were based on Cα atoms.

**Online supplemental material.** Table S1 shows data collection and refinement statistics. Table S2 shows 9B4 β/ββTCR, contacts with CD1d–α-GalCer. Table S3 shows clone 12 β/ββTCR, contacts with HLA-B*35:01–IPS. Online supplemental material is available at http://www.jem.org/cgi/content/full/jem.20141764/DC1.

We thank John Waddington and Marcin Cula at the University of Melbourne (Parkville, Victoria, Australia) for assistance with protein production, the staff at the Australian Synchrotron for assistance with data collection, and the Macromolecular Crystalization Facility at Monash University for technical assistance. We thank Dr. Richard Birkshaw for assistance with data collection and integration, Professor Paul Savage for kindly providing the PBS-44 α-GalCer analogue and Santosh Keshipeddy and Stewart Richardson for the steoxy-α-GalCer analogues.

The work was supported by the Australian Research Council (ARC: DP110104124, CE140010011, and LE110010106), the National Health and Medical Research Council of Australia (NHMRC; #1013667), the Cancer Council of Victoria (#1042866), and National Institutes of Health RO1 grant (GM 087136). D.G. Pellicci is supported by an NHMRC Biomedical fellowship, A.P. Uldrich by an ARC Future Fellowship (FT140000278), S. Gras by an ARC Future Fellowship (FT120100416), D.I. Godfrey by an NHMRC Senior Principal Research Fellowship (#1020770), and J. Rossjohn by an NHMRC Australia Fellowship (AF50).

The authors declare no competing financial interests.

Author contributions: D.G. Pellicci and A.P. Uldrich identified and performed cellular and molecular characterization of βββ T cells. S.B.G. Eckle, R. de Boer, D.G. Pellicci, A.P. Uldrich, S. Gras, and E. Chabrol produced TCR protein complexes for crystallographic experiments. A.P. Uldrich, J. Le Nours, E. Chabrol, and S. Gras solved the crystal structures. E. Chabrol, M.H.M. Heemskerk, R. de Boer, S.B.G. Eckle, and J. McCluskey undertook or supervised experiments on the MHC-restricted βββTCR. A.P. Uldrich, D.G. Pellicci, and S. Gras undertook SPR investigations, wrote the paper, and generated figures. D.G. Pellicci, A.P. Uldrich, F. Ross, K. McPherson, and R.T. Lim performed cell–based experiments, including glycolipid specificity and functional studies. L. Moretta, A.R. Howell, and G. Besra provided reagents that were crucial to this study. J. Rossjohn and D.I. Godfrey were joint senior authors: they co-led the investigation, devised the project, and wrote the paper.

Submitted: 10 September 2014
Accepted: 5 November 2014

**REFERENCES**

Amir, A.L., D.M. van der Steen, R.S. Hagedoorn, M.G. Kester, C.A. van Bergen, J.W. Drijfhout, A.H. de Ru, J.H. Falkenburg, P.A. van Veelen, and M.H. Heemskerk. 2011. Allo–HLA-reactive T cells inducing graft-versus-host disease are single peptide specific. Blood. 118:6733–6742. http://dx.doi.org/10.1182/blood-2011-05-354787

Bhati, M., D.K. Cole, J. McCluskey, A.K. Sewell, and J. Rossjohn. 2014. The versatility of the αβ T-cell antigen receptor. Protein Sci. 23:260–272. http://dx.doi.org/10.1002/pro.2412

Birkshaw, R.W., L. Kjer-Nielsen, S.B. Eckle, J. McCluskey, and J. Rossjohn. 2014. MAITs, MR1 and vitamin B metabolites. Curr. Opin. Immunol. 26:7–13. http://dx.doi.org/10.1016/j.coi.2013.09.007
Borg, N.A., L. Kjer-Nielsen, W.A. MacDonald, H.H. Reid, C.S. Clements, A.W. Purcell, L. Kjer-Nielsen, J.J. Miles, S.R. Burrows, et al. 2014. The CD1b3 regions of an immunodominant T cell receptor dictate the ‘energetic landscape’ of peptide-MHC recognition. *Nat. Immunol.* 15:617–180. http://dx.doi.org/10.1038/ni1155

Borg, N.A., K.S. Wu, L. Kjer-Nielsen, M.C. Wilce, D.G. Pellicci, R. Koh, G.S. Besra, B. Mahradawil, D.I. Godfrey, J. McCluskey, and J. Rossjohn. 2007. CD1d-lipid-antigen recognition by the semi-invariant NKT T-cell receptor. *Nature.* 448:44–49. http://dx.doi.org/10.1038/nature05907

Bowen, W.K., M. Kemal Aydintug, and R.L. O’Brien. 2013. Diversity of γδ T-cell antigens. *Cell. Mol. Immunol.* 10:13–20. http://dx.doi.org/10.1038/cmi.2012.45

Bowen, S., P. Sun, F. Livak, S. Sharrow, and R.J. Hodes. 2014. A novel T cell subset with trans-rearranged Vγ-ÇB TCRs shows Vß expression is dispensable for lineage choice and MHC restriction. *J. Immunol.* 192:169–177. http://dx.doi.org/10.4049/jimmunol.1302398

Brigl, M., and M.B. Brenner. 2004. CD1: antigen presentation and T cell function. *Annu. Rev. Immunol.* 22:817–890. http://dx.doi.org/10.1146/annurev.immunology.22.012703.104608

Brossay, L., O. Naidenko, N. Burdin, T. Sakai, and M. Kronenberg. 1998. Structural requirements for galactosylceramidase recognition by CD1d-restricted NK T cells. *J. Immunol.* 161:5124–5128.

Burrows, S.R., Z. Chen, J.K. Archbold, F.E. Tynan, T. Beddoe, L. Kjer-Nielsen, J.J. Miles, D.J. Moss, Y.C. Liu, et al. 2010. Hard wiring of T cell receptor specificity for the major histocompatibility complex is underpinned by TCR adaptability. *Proc. Natl. Acad. Sci. USA.* 107:10608–10613. http://dx.doi.org/10.1073/pnas.1004926107

Ciofani, M., and J.C. Zúñiga-Pflücker. 2010. Determining γδ versus αβ T cell development. *Nat. Rev. Immunol.* 10:657–663.

Clements, C.S., L. Kjer-Nielsen, W.A. MacDonald, A.G. Brooks, A.W. Purcell, J. McCluskey, and J. Rossjohn. 2002. The production, purification and crystallization of a soluble heterodimeric form of a highly selected T-cell receptor in its unliganded and ligand bound states. *Acta Crystallogr. D Biol. Crystallogr.* 58:2131–2134. http://dx.doi.org/10.1107/S0907444902015482

Corbett, A.J., S.B. Eckle, R.W. Birkshaw, L. Liu, O. Patel, J. Mahony, Z. Chen, R. Reantragoon, B. Meehan, H. Cao, et al. 2014. T-cell activation by transitory neo-antigens derived from distinct microbial pathways. *Nature.* 509:361–365. http://dx.doi.org/10.1038/nature13160

DeLano, W.L. 2002. The PyMOL Molecular Graphics System. Available at: http://www.pymol.org (accessed June 29, 2014).

Eckle, S.B., R.W. Birkshaw, L. Kostenko, A.J. Corbett, H.E. McWilliam, R. Reantragoon, Z. Chen, N.A. Gherrardin, T. Beddoe, L. Liu, et al. 2014. A molecular basis underpinning the T cell receptor heterogeneity of mucosal-associated invariant T cells. *J. Exp. Med.* 211:1585–1600. http://dx.doi.org/10.1084/jem.20140484

Emsley, P., B. Lohkamp, W.G. Scott, and K. Cowtan. 2010. Features and development of Coot. *Acta Crystallogr. D Biol. Crystallogr.* 66:486–501. http://dx.doi.org/10.1107/S0907444910007493

Gapin, L. 2014. Check MAIT. *J. Immunol.* 192:4475–4480. http://dx.doi.org/10.4049/jimmunol.1400119

Girardi, E., and D.M. Zajonc. 2012. Molecular basis of lipid antigen presentation by CD1d and recognition by natural killer T cells. *Immunity.* 39:1032–1042. http://dx.doi.org/10.1016/j.immuni.2013.11.001

Godfrey, D.L., J.J. McCluskey, and J. Rossjohn. 2005. CD1d antigen presentation: targets for NK T cells. *Nat. Immunol.* 6:754–756. http://dx.doi.org/10.1038/ni0805-754

Godfrey, D.L., J. Rossjohn, and J. McCluskey. 2008. The fidelity, occasional promiscuity, and versatility of T cell receptor recognition. *Immunity.* 28:304–314. http://dx.doi.org/10.1016/j.immuni.2008.02.004
Patel, O., A.G. Pellicci, S. Gras, M.L. Sandoval-Romero, A.P. Ulrich, T. Mallevaey, A.J. Clarke, J. Le Nours, A. Theodossis, S.L. Cardell, et al. 2012. Recognition of CD1d-sulfatide mediated by a type II natural killer T cell antigen receptor. *Nat. Immunol.* 13:857–863. http://dx.doi.org/10.1038/ni.2372

Patel, O., L. Kjer-Nielsen, J. Le Nours, S.B. Eckle, R. Birkinshaw, T. Beddoe, A.J. Corbett, L. Liu, J.J. Miles, B. Meehan, et al. 2013. Recognition of vitamin B metabolites by mucosal-associated invariant T cells. *Nat. Commun.* 4:2142. http://dx.doi.org/10.1038/ncomms3142

Pellicci, D.G., O. Patel, L. Kjer-Nielsen, S.S. Pang, L.C. Sullivan, K. Kyparissoudis, A.G. Brooks, H.H. Reid, S. Gras, I.S. Lucet, et al. 2009. Differential recognition of CD1d-α-galactosyl ceramide by the Vβ8.2 and Vβ7 semi-invariant NKT T cell receptors. *Immunity.* 31:47–59. http://dx.doi.org/10.1016/j.immuni.2009.04.018

Peyrat, M.A., F. Davodeau, I. Houde, F. Romagné, A. Necker, C. Leget, J.P. Cervoni, N. Cerf-Bensussan, H. Vié, M. Bonneville, et al. 1995. Repertoire analysis of human peripheral blood lymphocytes using a human V delta 3 region-specific monoclonal antibody. Characterization of dual T cell receptor (TCR) delta-chain expressors and alpha beta T cells expressing V delta 3j alpha C alpha-encoded TCR chains. *J. Immunol.* 155:3060–3067.

Reantragoon, R., L. Kjer-Nielsen, O. Patel, Z. Chen, P.T. Illing, M. Bhati, L. Kostenko, M. Bharadwaj, B. Meehan, T.H. Hansen, et al. 2012. Structural insight into MR1-mediated recognition of the mucosal associated invariant T cell receptor. *J. Exp. Med.* 209:761–774. http://dx.doi.org/10.1084/jem.20112095

Rosjohn, J., D.G. Pellicci, O. Patel, L. Gapin, and D.I. Godfrey. 2012. Recognition of CD1d-restricted antigens by natural killer T cells. *Nat. Rev. Immunol.* 12:845–857. http://dx.doi.org/10.1038/nri3328

Sandstrom, A., C.M. Peigné, A. Léger, J.E. Crooks, F. Konczak, M.C. Gesnel, R. Breathnach, M. Bonneville, E. Scotet, and E.J. Adams. 2014. The intracellular B30.2 domain of butyrophilin 3A1 binds phospho-antigens to mediate activation of human Vγ9Vδ2 T cells. *Immunity.* 40:490–500. http://dx.doi.org/10.1016/j.immuni.2014.03.003

Scott-Browne, J.P., J.L. Matsuda, T. Mallevaey, J. White, N.A. Borg, J. McChesney, J. Rossjohn, J. Kappler, P. Marrack, and L. Gapin. 2007. Germiline-encoded recognition of diverse glycolipids by natural killer T cells. *Nat. Immunol.* 8:1105–1113. http://dx.doi.org/10.1038/ni1510

Turner, S.J., P.C. Doherty, J. McChesney, and J. Rossjohn. 2006. Structural determinants of T-cell receptor bias in immunity. *Nat. Rev. Immunol.* 6:883–894. http://dx.doi.org/10.1038/nri1977

Uldrich, A.P., J. Le Nours, D.G. Pellicci, N.A. Gherardin, K.G. McPherson, R.T. Lim, O. Patel, T. Beddoe, S. Gras, J. Rossjohn, and D.I. Godfrey. 2013. CD1d-lipid antigen recognition by the γδ TCR. *Nat. Immunol.* 14:1137–1145. http://dx.doi.org/10.1038/ni.2713

Vantourout, P., and A. Hayday. 2013. Six-of-the-best: unique contributions of γδ T cells to immunology. *Nat. Rev. Immunol.* 13:88–100. http://dx.doi.org/10.1038/nri3384

Vavassori, S., A. Kumar, G.S. Wan, G.S. Ramanjaneyulu, M. Cavallari, S. El Daker, T. Beddoe, A. Theodossis, N.K. Williams, E. Gostick, et al. 2013. Butyrophilin 3A1 binds phosphorylated antigens and stimulates human γδ T cells. *Nat. Immunol.* 14:908–916. http://dx.doi.org/10.1038/ni.2665

Wang, G.C., P. Dash, J.A. McCullers, P.C. Doherty, and P.G. Thomas. 2012. T cell receptor αβ diversity inversely correlates with pathogen-specific antibody levels in human cytomegalovirus infection. *Sci. Transl. Med.* 4:128ra42. http://dx.doi.org/10.1126/scitranslmed.3003647

Wun, K.S., G. Cameron, O. Patel, S.S. Pang, D.G. Pellicci, L.C. Sullivan, S. Keshupedy, M.H. Young, M.S. Thakur, et al. 2011. A molecular basis for the exquisite CD1d-restricted antigen specificity and functional responses of natural killer T cells. *Immunity.* 34:327–339. http://dx.doi.org/10.1016/j.immuni.2011.02.001

Wun, K.S., F. Ross, O. Patel, G.S. Besra, S.A. Porcelli, S.K. Richardson, S. Keshupedy, A.R. Howell, D.I. Godfrey, and J. Rossjohn. 2012. Human and mouse type I natural killer T cell antigen receptors exhibit different fine specificities for CD1d-antigen complex. *J. Biol. Chem.* 287:39139–39148. http://dx.doi.org/10.1074/jbc.M112.412320

Xiong, N., and D.H. Raulet. 2007. Development and selection of γδ T cells. *Immunol. Rev.* 215:15–31. http://dx.doi.org/10.1111/j.1600-065X.2006.00478.x