Molecular design of the γδT cell receptor ectodomain encodes biologically fit ligand recognition in the absence of mechanosensing


High-acuity αβT cell receptor (TCR) recognition of peptides bound to major histocompatibility complex molecules (pMHCs) requires mechanosensing, a process whereby piconewton (pN) bioforces exert physical load on αβTCR-pMHC bonds to dynamically alter their lifetimes and foster digital sensitivity cellular signaling. While mechanotransduction is operative for both αβTCRs and pre-TCRs within the αβ T cell lineage, its role in γδT cells is unknown. Here, we show that the human DP10.7 γδTCR specific for the sulfoglycolipid sulfatide bound to CD1d only sustains a significant load and undergoes force-induced structural transitions when the binding interface-distal γδ constant domain (C) module is replaced with that of αβ. The chimeric γδ-αβTCR also signals more robustly than does the wild-type (WT) γδTCR, as revealed by RNA-sequencing (RNA-seq) analysis of TCR-transduced Rag2−/− thymocytes, consistent with structural, single-molecule, and molecular dynamics studies reflective of γδTCRs as mediating recognition via a more canonical immunoglobulin-like receptor interaction. Absence of robust, force-related catch bonds, as well as γδTCR structural transitions, implies that γδT cells do not use mechanosensing for ligand recognition. This distinction is consonant with the fact that their innate-type ligands, including markers of cellular stress, are expressed at a high copy number relative to the sparse pMHC ligands of αβT cells. We posit that mechanosensing emerged over ~200 million years of vertebrate evolution to fulfill indispensable adaptive immune recognition requirements for pMHC in the αβ T cell lineage that are unnecessary for the γδT cell lineage mechanism of non-pMHC ligand detection.

Significance

TCR mechanosensing is thought necessary for digital sensitivity of αβT cell response to scant pMHC antigens. We use bioinformatic analysis, molecular dynamics, single-molecule optical tweezers techniques, cellular activation, and RNA-seq analysis to explore this paradigm in the γδT cell lineage. We find that, in keeping with its role in recognizing abundant cell-surface ligands, the γδTCR lacks force-dependent hallmarks of mechanosensing in αβT cells.

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T cell receptor | mechanosensor | optical tweezers | T cell activation | γδT cells

Within jawed vertebrates (Gnathostoma), αβ and γδT cells utilize somatic genomic rearrangement to generate a receptor repertoire large enough to recognize the enormous diversity of viral or other pathogen-derived antigens and then mount a protective immune response (1–4). αβT cells are prominent in blood and lymph nodes while γδT cells are more abundant in barrier tissues including skin, intestinal and other epithelia, suggesting distinct, nonredundant roles for each T lineage subset (5, 6). Whereas the vast majority of γδT lymphocytes recognize sparse “foreign” peptides within a vast array of normal self-peptides processed and presented by classical major histocompatibility complex (MHC) molecules on the surface of pathogen-infected or otherwise damaged cells, this is almost without exception not the case for γδT cells (7–9). Instead, γδT cells recognize ligands, including self-derived and plentiful stress-induced nonpeptide ligands, diverse in structure and distinct from classical MHC molecules (6, 8, 10–17). Furthermore, γδT cells are generally in an activation state capable of mediating rapid (i.e., minutes- to hours-long time frames) innate-like cytolysis and cytokine release, in contrast to αβT cells, which exist in distinct naïve, effector, and memory states (6, 18). Naïve αβT cells become cytotoxic or mediate cytokine/chemokine-based inflammation only following differentiation that requires exposure over several days to antigen and costimulatory molecules on professional antigen-presenting cells (APCs).

Over the last decade, it has become increasingly clear that contrary to conventional receptor-ligand interactions exemplified by antigen-antibody binding, bioforces are essential for nonthermodynamic equilibrium, mechanosensor-based αβT cell activation (19–26). αβT cell motility and the local cytoskeletal machinery
place physical load on individual αβTCR–pMHC bonds, which tunes both the sensitivity and specificity of αβTCR recognition (19). In fact, chemical thresholds in the absence of external load require a 1,000- to 10,000-fold higher number of pMHC molecules than observed physiologically to trigger cellular αβTCR cell activation (20). In contrast, under force the ligand-mediated induction of the αβTCR cell biological response can be essentially digital. Mechanistically, physical load fosters stability and interfacial matching as well as a temporally correlated structural transition in the αβTCR heterodimer ectodomain that strengthens bond lifetime (i.e., a “catch bond”), energizes the αβTCR and induces αβTCR complex quaternary change, conformationally altering the transmembrane (TM) segments and ligands surrounding the TCR and thereby exposing the immunoreceptor tyrosine-based activation motifs (ITAMs) in the cytoplasmic tails of the associated signaling CD3 for phosphorylation (19, 27–29 and refs. therein). As the structural transition of the TCRαβ heterodimer is itself reversible in the context of relevant bioforces, a ligated TCR can be repetitively energized by the same pMHC in the absence of its release, thereby perpetuating the activation geometry of the αβTCR and surrounding membrane to optimize cognate antigen-dependent signaling performance, accounting for high acuity recognition (21, 30).

The role of bioforces in γδTCR recognition has not been addressed. Here, we examine the character of physical load in γδTCR signaling comparing αβTCR signaling, using a combination of single-molecule (SM) optical tweezers-based technology, molecular dynamics (MD) simulation, transcriptomics, and functional analyses. For this purpose, we selected the DP10.7 γδTCR, since it is structurally well characterized and binds the MHCib molecule, CD1d, in complex with a defined ligand, sulfatide. This TCR–ligand pair is similar in overall three-dimensional topology and size to complexes of peptides bound to MHC Ib Kβ or Dα molecules used in related work in αβTCR systems (31). Our findings reveal that the DP10.7 γδTCR, unlike αβTCRs, does not bear the force-sensitive hallmarks of a functional mechanosensor. Instead, although DP10.7 binds ligand with submicromolar affinity under zero force, it readily dissociates from its ligand under force loading. By creating a chimeric Vγδ–αβTCR in which Cαβ replaces Cγδ, we define a gain of function that supports mechanotransduction comparable to the level of a wild-type (WT) αβTCR. Moreover, RNA-sequencing (RNA-seq) analysis at the double negative (DN) thymocyte stage following retroviral transduction of TCRs into Rag2−/− thymocyte progenitors reveals that such a chimeric TCR complex augments signaling compared to thymocytes expressing the wild-type γδTCR. Collectively, these findings inform that mechanotransduction associated with structural transitions and dynamic bond formation are linked to the constant domains in αβTCRs rather than a property of their variable domains per se. We reconcile our data with those of prior studies suggesting that γδTCR signaling is stronger than that of αβTCR during tyrosine kinase. Lastly, we discuss how structural differences between these two lineages of TCRs are well aligned with their distinct ligand specificities and attendant biology.

Results

The CγδFG Loop Structurally Distinguishes αβ from γδTCRs. We have previously characterized the mechanotransducing properties of the αβTCR at the SM and single-molecule–single-cell (SMSC) level and have noted signaling correlates with several structural features (19, 21, 27, 28, 30). Tied to preservation of the force-mediated catch bond, a requisite feature in pMHC ligand discrimination is the CγδFG loop. We observe that the γδTCR, while sharing many features with the αβTCR, including CD3 components and general ectodomain organization, lacks the large FG loop in its homologous Cγ domain as compared to the CγδFG loop within the αβTCR (Fig. L1). The CγδFG loop but not that of Cγ appears to buttress the region joining the V and C domains (Fig. L4). Multiple sequence alignment of mammalian C domains, comparing Cβ and Cγ with C regions of human or murine immunoglobulin heavy (IgH) subtypes clearly shows that the extensive FG loop (containing a 12- to 13-amino-acid insertion relative to the heavy chain constant region domain 1 [CH1]) is a conserved feature only within the αβTCR (Fig. L1B). This contrasts with otherwise high structural conservation throughout the domains. Lack of the distinct FG loop feature within Cγ could plausibly indicate either that equivalent mechanotransduction is not a property of the γδTCR, or alternatively that Cδ may provide the avenue of mechanosensing, particularly given the predominance of the Vδ in several characterized γδTCR–ligand interactions (10, 31, 32).

The CγδFG Loop Stabilizes the V–C Interface in αβ and Chimeric γδ–αβTCRs Relative to γδTCR. As we (19, 33) and others (34) have noted, the CγδFG loop provides significant structural contact between Vβ and Cβ that is absent in the antibody V–C interface (~350 Å2 versus ~150 Å2 buried surface areas, respectively). Our previous MD simulation on TCRαβ showed that the CγδFG loop influences the motion of the variable domains relative to the constant domains, as well as imposes an orientational restraint of the former while facing pMHC, which enables a load- and ligand-dependent control of the bond lifetime (29). Comparative MD simulations of the TCR structures used in the present study show that the Vγ–Cδ interface has significantly fewer high-occupancy contacts as compared to Vβ–Cβ (Fig. 2A, V–C(γδ); also see SI Appendix, Fig. S1), and similar to the α-chain of TCRαβ, the δ-chain has few V–C contacts (Fig. 2A, V–C(αβδ) group). Thus, the TCRγδ constant domains are unlikely to have any strong influence on the conformational motion of the variable domains. Another property important for mechanosensing in TCRαβ is the compliance of the Va–Vb interface, leading the interface with pMHC to be responsive to and controlled by an applied load. In our previous simulation, TCRαβ lacking the CγδFG loop had an increased number of Va–Vb contacts (29). The higher number of nonpolar Vγ–Vδ contacts (Fig. 2A, V–Vγδ group) also suggests a reduced compliance, which further makes TCRγδ unsuitable for mechanosensing. In contrast, when Cγδ was substituted for Cγδ the number of Vγ–Vγδ contacts increased, comparable to that of the full TCRαβ [blue in Fig. 2A, V–C(γβ); SI Appendix, Fig. S1], perhaps explaining why chimeric Vγδ–Cαβ ectodomain constructs have utilized the CγδFG loop as a substitute for the CγδFG loop in its homologous Cγ domain as well, forming nearly half of its high-occupancy contacts (Fig. 2B). Our simulations thus suggest that not only will the γδTCR lack the requisite allosteric connections for mechanotransduction, but also that these connections should form in chimeric Vγδ–Cαβ constructs mainly via nonpolar contacts (SI Appendix, Fig. S1). By this analysis it would also appear that the Vδ–Cδ interface is unlikely to compensate for the relative paucity of contacts in the Vγ–Cδ interface and shift the mechanosensing potential to the δ-subunit.

The γδTCR Lacks Strong Mechanotransduction Elements, Which Are Rescued by Replacement of γδ C Domains with Those of αβ. To experimentally determine whether or not the TCRγδ has the potential for mechanosensing by direct biophysical analyses, we utilized the DP10.7 γδTCR (DP10.7γδ) in SM experiments to test the hypothesis that the γδTCR is a mechanosensor (Fig. 3A). Sulfatide–CD1d has been shown to engage DP10.7 with only weak interactions with CD1d in the absence of sulfatide lipid (31). To probe the DP10.7 TCRγδ–ligand interaction, CD1d ± sulfatide was bound to a functionalized surface in a tethered bead configuration (19) (Fig. 3A). To this end, DP10.7γδ was cloned and produced as a leucine zipper-paired heterodimer (LZ) with N15δ–VS8/Kβ used as comparison (Fig. 3B) (19, 31). CD1d or VS8/Kβ ligands were bound through streptavidin to a PEG-pacified surface containing sparse PEG–biotin. The TCRs fused to LZ at the C terminus were tethered via a long DNA molecule
**Fig. 1.** Structural features of γδ and αβ TCRs. (A) Structural comparison of TCRαβ (Top) with TCRγδ (Bottom). The Left two panels in each row show surface representations with individual subunits and domains delineated. The FG loop is shown in light blue (TCRαβ) or yellow (TCRγδ) and highlighted in the boxed region. The Right two panels of each row present a zoomed view of the boxed region at approximately the same magnification for each TCR to illustrate the relative size and structure of the respective FG loops. (B) Multiple sequence alignment of mammalian TCR Cγ, Cβ domains, and equivalent Ig CH1 domains of selected isotypes. Secondary structure as assigned in the murine N15αβ X-ray structure 1NFD are indicated. The FG-loop region is boxed. Invariant cysteines are highlighted in red-brown and conserved residues denoted in black and gray.

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**Table 1:**

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**Mallis et al.**

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such that force could be applied to the optically trapped bead (Materials and Methods and ref. 19). Lifetime measurements were performed by translating each sample relative to the fixed trap using the piezo stage, then holding at a fixed position/force until bond rupture. Bond rupture is identified as an abrupt snap back of the bead to the trap center (distance = 0 nm) while conformational extensions are observed as smaller displacements of the bead toward the trap center.

Testing the WT DP10.7 against CD1d alone in the absence of sulfatide we found lifetimes of less than 0.5 s down to 0.3 s at 10 pN (Fig. 3C and SI Appendix, Fig. S2 A and B) while sulfatide-complexed CD1d resulted in similar bond lifetimes with the exception of a narrow force window in which lifetimes around 1 s were measured (Fig. 3D and SI Appendix, Fig. S2C). In the absence of sulfatide loading, bond formation was of lower frequency, with 1 to 2 tethers per field of view found in comparison to ~25 when presented with sulfatide–CD1d. Whereas the CD1d without exogenous ligand resulted in a slip-bond profile in force-bond lifetime plots of the collected binding data, a narrow enhancement of bond lifetime was measured for sulfatide–CD1d with forces under 10 pN, suggesting force may minimally organize the interface (29). There was no continued enhancement above 10 pN; instead, the system transitioned to a slip bond. This force threshold corresponds to loads generated by cells even under static culture conditions, and so this does not appear to be a physiologically relevant catch bond as observed in other systems (19, 26, 35–37) (Fig. 3E).

In contrast, when the chimeric γδ–αβTCR is tested versus sulfatide–CD1d, longer bond lifetimes were measured (Fig. 3F and SI Appendix, Fig. S2D) with organization/bond strengthening continuing on to form a typical catch bond of 5 s lifetime at 15-pN force, comparable to N15αβ interacting with its cognate ligand VSVK-Kβ (Fig. 3E). Incorporating the Cβ domain in lieu of Cγδ thus leads to force-responsive properties in the context of the same Vγδ domain module and ligand. Of note, γδ–αβTCR does not form a catch bond with CD1d lacking sulfatide (Fig. 3 E and G and SI Appendix, Fig. S2E) in keeping with previous observations of ligand gating specificity by TCRαβ and, more specifically, requisite allosteric control within Cβ, another hallmark of mechanosensing (19, 26, 29, 30). Indeed, the SM sensitivity index (the ratio of a TCR–ligand bond lifetime for specific to nonspecific MHC-bound interaction) (19) shows discrimination of the γδ–αβTCR to be much greater than that of the γδTCR, and more similar to the b7–α2 (SI Appendix, Fig. S3). The appearance of sustained bond lifetime under force was consistently accompanied by a structural transition for the γδ–α2TCR (Fig. 3F), analogous to that seen previously for the αβTCR and pre-TCR in response to specific ligands (19, 30) but essentially unobserved in traces for the γδTCR (Fig. 3 C and D and SI Appendix, Fig. S2C).

**Catch Bond and Structural Transition Rescue by αβ C Domains on the T Cell Surface.** To confirm that the biomechanical observations above translate to the fully assembled and membrane-embedded γδTCRs on T cell surfaces, TCR γδ or γδ–αβ heterodimers (Fig. 3H) were retrovirally transduced and resultant TCRs were interrogated with bead-bound sulfatide–CD1d in the SMSC format as previously described (19). To this end, BW5147 T cells lacking endogenous TCRs were transduced using a 2A peptide-containing construct for simultaneous expression of paired TCR subunits (38). Murine C domains and TM regions were substituted for the human counterparts to attain optimal surface expression in this mouse-derived thymocyte lymphoma line as previously described (39, 40). As shown in Fig. 3I, the γδTCR exhibits only slip bond, whereas in contrast, the chimeric γδ–αβTCR manifests a significant catch bond up to 15 pN and slip bond thereafter, in parallel to the SM behavior, demonstrating the same force sensitivity. The significant bond lifetime over a range of forces for both TCRs is likely due to the presence of CD3 molecules in the TCR complex (ref. 28 and refs. therein) as well as an increase in compliance and stress relaxation of the cell membranes and connecting linkages compared to the biotinylated surfaces in the SM assay (Fig. 3I).

The SM is more similar to a force clamp, while the SMSC behaves more like a stress relaxation test. The cell mechanical linkage pathway may also participate in modulating bond lifetime, even actively through actin–myosin-based coupling and feedback (20). We note that SM and SMSC assays are executed differently, yielding a slower effective “instrument response time” for SMSC. In the absence of sulfatide we observed lower frequency of bond formation, with fewer than 10% of beads forming effective tethers for either γδ (1 in 25) or γδ–αβ (2 in 25), as compared to ~50% when presented with CD1d–sulfatide. Moreover, the structural transition previously observed in αβTCRs and pre-TCRs (19, 30) is present here in over 50% (23/44) of the individual traces and is similar in length to that of the N15αβ TCR (19) (Fig. 3J). All transitions occurred at 10 pN or higher force for the chimeric γδ–αβTCR, consistent with behavior of the N15αβ TCR (19). In contrast, only 1 in 45 traces had evidence of a transition for the γδTCR. These data strongly suggest that the γδTCR lacks specific adaptations for mechanosensing, a result consistent with absent force-dependent ligand signaling threshold sensitivity enhancement (SI Appendix, Fig. S4).

**Enhanced Thymic Signaling through TCR Is Impacted by αβ C Domains.** To assess the biological consequences of TCR mechanosensing function, we exploited an in vitro stromal cell-lymphoid progenitor coculture experimental system. Fetal liver–derived thymic progenitors isolated from Ragg2−/− B6 mice were transduced with either full-length γδTCR, γδ–αβTCR, or αβTCR heterodimers as detailed in Fig. 4A. Stromal cells used in this assay were OP9–DL4 applied previously for both γδT cell and αβT cell development (35,
In order to test the effect of ligand binding, an OP9–DL4 cell line, which expresses single-chain human CD1d/β2m at levels comparable to those of murine CD1d on these same cells, was developed (SI Appendix, Fig. S5 A and B). DP10.7 tetramer binding analysis (31) confirmed robust binding to this OP9–DL4–CD1d cell line when exogenous sulfatide was added (SI Appendix, Fig. S5). Some tetramer binding was detected for the γδ TCR-transduced DN3 and DN4 subsets used in SM experiments. (7) SMSC optical trap assay are indicated. Note that the TCR γδ TCR cultures on stroma with sulfatide (pink curve, n = 92) with CD1d, and N15γδ interaction with its cognate ligand VSV8/H-2Kb (30) (green curve, n = 192). Error bars indicate SEM. (F) SM traces at indicated forces for DP10.7 γδ TCR chimera interaction with CD1d-sulfatide. Initial binding dwell is shown in green and posttransition dwell in blue. Transition points are indicated in each trajectory with blue arrows. Black and red sections are as in C. (G) SM traces at indicated forces for DP10.7 γδ TCR chimera interaction with CD1d with color coding as in F. A transition was identified in the 25-pN trace (blue arrow). (H) Constructs used in SMSC optical trap assay are indicated. Note that the TCR γδ TCR is depicted with a bend analogous to that of TCR (28), although there is currently no data to confirm either a bipartite or single helix structure. (7) SMSC optical trap assay force vs. lifetime plot for the DP10.7 TCR γδ TCR interaction with sulfatide (purple curve, n = 43) or DP10.7 γδ TCR chimera (red curve, n = 44) interaction with sulfatide-CD1d. Error bars indicate SEM. (J) Transition distances for DP10.7 γδ TCR chimera (red curve, 23 of 44) interaction with sulfatide–CD1d from traces acquired by SMSC as compared to N15γδ (19) (green curve, n = 15). Error bars indicate SEM. For reference only a single, shorter transition was found in 45 traces for DP10.7 TCR γδ TCR interaction with sulfatide–CD1d (purple triangles, see key).

41). In order to test the effect of ligand binding, an OP9–DL4 cell line, which expresses single-chain human CD1d/β2m at levels comparable to those of murine CD1d on these same cells, was developed (SI Appendix, Fig. S5 A and B). DP10.7 tetramer binding analysis (31) confirmed robust binding to this OP9–DL4–CD1d cell line when exogenous sulfatide was added (SI Appendix, Fig. S5). Some tetramer binding was detected for the OP9–DL4–CD1d cell line in the absence of sulfatide addition. WT OP9–DL4 + sulfatide showed a slight increase over untreated WT OP9–DL4, while the OP9–DL4–MHC knockout (KO) cell line, which lacks class I MHC expression (30) including CD1d, showed no detectable effect of sulfatide treatment (SI Appendix, Fig. S5).

Following an 8-d coculture of γδ or γδ–αβ TCR-transduced DN3 Rag2−/− thymocytes with either WT OP9–DL4 or OP9–DL4–CD1d, we observed transition of thymocytes from the CD4−CD8−CD25+CD44+ (DN3) population through CD4−CD8−CD25−CD44− (DN4) with a subset also progressing beyond DN4 to CD4+CD8+ (DP) (SI Appendix, Fig. S6). When sulfatide was added to each culture, a differential loss in surface CD3 staining was measured in association with human CD1d-expressing stromal cells (Fig. 4B). This effect was significantly more pronounced for γδ–αβ TCR transduced thymocytes versus those expressing WT γδ TCR and is present both in DN and DP thymocytes (Fig. 4 B and C). Surface CD3 loss is most likely due to an enhanced responsiveness of these thymocytes imprecated by the αβ TCR C regions leading to down-modulation of the TCR complex with activation (42). The down-modulation occurs in both DN3 and DN4 subsets for γδ–αβ TCR, while the effect is not observed for γδ TCR in DN3 (Fig. 4D). Of note, post-DN4 cell numbers following culture were significantly reduced for the γδ–αβ TCR cultures on stroma with sulfatide present (SI Appendix, Fig. S6 B and C). This was not the case for γδ TCR thymocytes. In addition, the OP9–DL4 stroma expressing endogenous murine CD1d-only also showed this reduction, consistent with the ability of 10.7 TCR tetramers to bind weakly but nevertheless clearly to OP9–DL4 (SI Appendix, Fig. S5). Loss of DP thymocytes in γδ–αβ TCR- but not γδ TCR-transfected thymocytes on both OP9–DL4 stroma implies negative selection linked to the CoCβ module (SI Appendix, Fig. S6). The strong down-regulation of CD3 on γδ–αβ TCR transduced thymocytes following sulfatide exposure implied that active TCR signaling was occurring for γδ–αβ TCR thymocytes. Given that higher TCRγδ–ligand signaling strength was required to induce IFNγ as opposed to IL-17 T cell differentiation (43–45), we assessed cytokine production within hours of calcium ionophore plus phorbol myristate acetate (PMA) stimulation following 8 d of γδ TCR and γδ–αβ TCR thymocyte–stromal cultures with or without exogenous sulfatide addition (SI Appendix, Fig. S7). While no enhanced IFNγ production was observed when comparing γδ–αβ TCR to γδ TCR, an absence of cellular elements in epithelial cultures present in thymus and required to recapitulate the cytokine

Mallis et al. Molecular design of the γδ T cell receptor ectodomain encodes biologically fit ligand recognition in the absence of mechanosensing.

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PNAS | 5 of 12

Molecular design of the γδ T cell receptor ectodomain encodes biologically fit ligand recognition in the absence of mechanosensing.
phenotype could not be excluded. As a consequence, we selected a more global and unbiased approach to interrogate signaling differences.

**Chimeric γδαβTCR Signaling Generates Stronger Transcriptome Changes than Those Through WT γδTCR.** To this end, we analyzed transcriptome signatures of DN3 and DN4 thymocytes from lymphoid progenitors previously transduced with γδTCR or γδαβTCR and then cultured on OP9–DL4–CD1d with or without sulfatide addition for 8 d. Three independent experiments analyzing DN3 and DN4 cells ± sulfatide for the γδTCR- and γδαβTCR-transduced thymocytes yielded 24 cDNA libraries that underwent next generation sequencing (Dataset S1 and SI Appendix, Tables S1–S7). Global principal component analysis (PCA) separated clearly between the DN3 and DN4 populations (Fig. 5 A, Upper) where the drivers of this transition (e.g., Samhd1, Lgals2, Stsulf6, Gpr15, and Slk064; Dataset S1 and SI Appendix, Table S4) dominated the PCA, masking underlying transitions induced by the response of the wild-type and chimeric TCR to CD1d–sulfatide. Nevertheless, at the DN3 stage, only 20 genes were significantly fold-change regulated in γδTCR thymocytes following sulfatide stimulation (Dataset S1). In contrast, in DN3 γδαβTCR-expressing thymocytes responding to sulfatide, similar changes were observed not only for the 20 altered in the γδTCR cells but for >4,000 other genes (Dataset S1 and SI Appendix, Table S5). PCA alone, however, could not separate the DN3 γδTCR and γδαβTCR sulfatide-stimulated populations as the gene expression patterns were almost identical (Fig. 5 A, Upper and Dataset S1); the differences resided in the extent of the fold change. Additionally, even in the absence of exogenous sulfatides, the Carβ̂B̂ domains in the γδαβTCR-expressing thymocytes influenced the signaling background. For example, using Cδ69 up-regulation as a proxy for TCR stimulation (46), the γδTCR background for Cδ69 at the DN3 stage was 153.6 ± 25.4 expression reads while, in γδαβTCR, levels were 218.3 ± 26.6 with both rising to ~270 on sulfatide stimulation with a similar representation at the DN4 stage (Dataset S1 and SI Appendix, Table S1).

Analysis at the DN4 stage, however, removing the influence of the DN3 to DN4 drivers, permitted discrimination of significant differences in expression programs for γδαβTCR and γδTCR (Fig. 5 A, Lower). For the unstimulated condition, PCA revealed a slight shift on the PC1 axis for γδαβTCR, likely representing the background elevation of discrete gene transcripts, as discussed further below. Upon addition of sulfatide, the γδαβTCR DN4 program shifts strongly on the PC2 axis as well as on the PC1 axis. In contrast, the centroids for the γδTCR-unstimulated and γδTCR-stimulated populations appear similar. Immune gene expression analysis (47), a system trained on subsets of mature single-positive CD4 and CD8 cells both resting and responding to a variety of stimuli, as well as on phenotypically defined and thymic region-localized thymocyte populations, identifies the DN4 γδαβTCR sulfatide-stimulated population as highly and significantly similar to subcapsular cortical thymocytes (Fig. 5B).

Comparison of the gene expression profiles for γδTCR and γδαβTCR thymocytes responding to sulfatide at the DN4 stage reveals 117 significantly regulated gene transcripts for which 20 appear to be preferentially regulated in the γδαβTCR condition (group I), 27 are shared (group II), and 70 are preferentially regulated in the γδTCR condition (group III) (Fig. 6A and Dataset S1). Analysis of the γδαβTCR group III profile, with significantly regulated transcripts grouped according to functional activity, shows that many of the gene transcript levels are directionally regulated similarly in γδTCR-expressing thymocytes but that the degree of regulation is much greater in γδαβTCR-expressing thymocytes (Fig. 6B). Several of the genes represented in group I (specifically Cδ69, Egr1, Egr2, Nr4a1 [Nat77], Cδ200 variants and Kld1 [Cδ94], and Kln5 [Ly9e]) are key markers for TCR stimulation (43, 46, 48–52). Given that TCR signaling, measured as reduced cell surface expression of CD3, was greater in the γδαβTCR condition than in the γδTCR condition, the inclusion of such TCR signaling-associated transcripts in group I was unexpected. As depicted in Fig. 6C, however, genes expected to be down-regulated or up-regulated upon TCR signaling had already moved in these respective directions in the γδαβTCR cells in comparison with γδTCR cells prior to sulfatide stimulation. Since
the maximal stimulation changes for this signaling-associated gene group were similar for both cell types, the apparent fold change upon sulfatide stimulation is greater for the γδTCR cells than for the γδ–αβTCR cells. In the absence of exogenous sulfatides, the background binding of DP10.7 TCRγδ tetramer is greater to OP9–DL4–CD1d than parental OP9–DL4 (SI Appendix, Fig. S5). Hence, endogenous sulfatide presentation may be sufficient to stimulate those genes noted in Fig. 6C through the chimeric γδ–αβTCR but not the γδTCR before signaling induction by exogenously added sulfatides. Overall, the transcriptomics results support the notion of enhanced signaling sensitivity and function of the γδ–αβTCR.

Discussion

Our results demonstrate that γδ T lineage cells exhibit differential signaling sensitivity to αβ T lineage cells by virtue of their respective C modules. The relevance for γδ T cell biology generally, and analog signaling in particular, is highlighted below. Pointedly, γδT cells cannot exploit TCR mechanosensing as used by αβT cells to harness physical load and extend TCR–ligand bond lifetime under nonequilibrium conditions. Load in vivo results from a range of bioforces that lymphocytes experience during both cellular (cell movement) or internal (cytoskeletal rearrangement) processes (reviewed in ref. 21). This differential behavior is striking, given shared use of CD3 dimeric signaling components by both TCRs (ref. 27 and refs. therein), although some distinctions in CD3 composition, may further tune signaling (53–56). Replacement of CyC6 in WTDPI0.7 γδTCR with CyCβ in the γδ–αβTCR construct confers αβTCR mechanosensing properties to the chimeric receptor whose VγV6 module (i.e., ligand interaction surface) remains the same. MD simulations show that the Vβ–Cβ interface is stabilized by the CjFG loop unique to mammalian pre-TCRs and αβTCRs (29) and thus likely implicates the β-subunit in this gain of function. In this regard, our earlier studies using optical tweezers on isolated TCR αβ heterodimers as well as αβTCR complexes on T lymphocytes documented how the VαVβ module was allosterically controlled by the CjFG loop (19, 30) to dictate TCR–pMHC bond lifetime as well as peptide discrimination. Concordantly, deletion of the CjFG loop created αβTCRs, whose recognition and signaling function were attenuated in vitro and in vivo (57, 58). αβTCR mechanosensing affords virtually digital responsiveness in signaling; only one or a handful of TCR–pMHC interactions is required for cellular activation, while chemical thresholds in the absence of load require pMHC numbers higher by orders of magnitude to trigger a cellular response (20). The pre-TCR, consisting of a pTα–β heterodimer, also manifests mechanosensing function that is CjFG loop dependent (30, 35).

Within Gnathostomata there was coevolution of the elongated CjFG loop and molecular speciation of CD3γ and CD3δ genes from a single CD3 precursor (58, 59). Thus, mammals, but likely not birds, amphibians, reptiles, or bony fish, are capable of mechanosensing in their respective αβTCR lineages. Given that the evolutionary distance between human and mouse from a common mammalian ancestor is 75 million years and that between human and birds from a common vertebrate ancestor is 300 million years, roughly 200 million years of vertebrate evolution was required for mechanosensing to emerge as the solution within the αβT lineage system to high acuity adaptive immune recognition.

Catch bonds are observed over a wide range of receptor–ligand systems (60). In these systems, the receptor is assumed to take two alternate conformations, one with a low affinity and the other with a high affinity to the ligand. Without load, the low-affinity state is prevalent. An applied load causes a conformational change to the high-affinity state, thereby initiating the catch-bond pathway (61). An essential aspect in this mechanism is allostery, where load-induced conformational change of the receptor alters the ligand-binding domain. While alternate conformations responsible for the two states have been observed in other systems, our earlier studies using optical tweezers on isolated TCR αβ heterodimers demonstrated how the VαVβ module was allosterically controlled by the CjFG loop (19, 30) to dictate TCR–pMHC bond lifetime as well as peptide discrimination. Concordantly, deletion of the CjFG loop created αβTCRs, whose recognition and signaling function were attenuated in vitro and in vivo (57, 58). αβTCR mechanosensing affords virtually digital responsiveness in signaling; only one or a handful of TCR–pMHC interactions is required for cellular activation, while chemical thresholds in the absence of load require pMHC numbers higher by orders of magnitude to trigger a cellular response (20). The pre-TCR, consisting of a pTα–β heterodimer, also manifests mechanosensing function that is CjFG loop dependent (30, 35).

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(60), the structural origin for the catch bond behavior in TCRγδ has remained enigmatic, as no clear conformational states have been observed in X-ray structures of TCRγδ that might affect its interaction with the pMHC molecule. Two recent studies propose a catch-bond mechanism based on behaviors of a few hydrogen bonds between TCRγδ and pMHC that formed when the complex was rapidly pulled apart in MD simulation (36, 62). Aside from the use of very large forces within short simulation times where conformational relaxation cannot occur, analyses based only on a handful of transiently formed contacts cannot address the question of allostery, which involves conformational motion of the whole protein.

More recently, our own MD simulation study has illuminated a possible mechanism of catch-bond formation within the Vαβ–pMHC interface that is potentiated by V–C interactions, particularly those at the Vβ–Cβ interface, including the CβFG loop (29). An essential aspect of this mechanism is that the capacity to activate the catch bond is endowed by the conformational properties of the entire TCRγδ complex rather than only by residues that immediately contact pMHC. More specifically, the four-domain organization leads to relative motion between Vα and Vβ, which can be stabilized by the applied load and in the presence of the cognate antigenic peptide. The CβFG loop is a structural element that is crucial for the allosteric control. By forming additional contacts with Vβ, the CβFG loop not only influences the Vα–Vβ motion, but it also supports its orientation amenable to form an interface with pMHC in loaded conditions.

Based on the above, the MD simulations in the present study indicate that the TCRγδ chassis is not well suited for mechanosensing. In particular, the Vγδ and Cγδ domains do not include sufficient contacts to establish allostery responding to load (Fig. 2 and SI Appendix, Fig. S1). Further, the greater number of contacts between Vγδ and Vβδ suggest a lower compliance. In contrast, the Vγδ–Cγδ chimera has the number of contacts between Vγδ and Cβ, as well as between Vγδ and Vδ comparable to those for the corresponding interfaces of TCRγδ (Fig. 2), which is fully consistent with our experimental results, demonstrating the chimera responding to load similar to TCRγδ. The reduction in the number of Vγδ–Vδ contacts is due to the orientational constraint imposed by the CβFG loop, as observed between the wild-type TCRγδ and a mutant lacking the CβFG loop (29). Since a majority of the contacts between Vγδ and Cβ are nonpolar, a steric constraint imposed by the CβFG loop further elucidates the steric nature of the contact, we built a model of Vγδ–Cγδ chimera and performed MD simulation (29). As expected, the CβFG loop is more important than forming specific contacts (Fig. 2). To further elucidate the steric nature of the contact, we built a model of Vγδ–Cγδ chimera and performed MD simulation (SI Appendix, Fig. S1E). There were little contacts at the Vα–C6 interface, and a small number of contacts formed at the Vβ–Cγ interface. Since extensive contacts form at the Vβ–Cγ interface of TCRγδ, some of the residues in Vβ are amenable to form nonpolar contacts with Cγ in the chimera. However, the contacts are not extensive and we do not expect the Vαβ–Cγδ chimera to exhibit a catch bond as strongly as the Vγδ–Cγδ chimera. The increase in the Vαβ–Vβ contacts in this case is also consistent with the behavior of the FG-loop deletion mutant where the Vα–Vβ motion is suppressed (29). While additional insight will be gained from future simulations of TCRγδ complexed with sulfatide–CD1d, the present simulations elucidate conformational properties of TCRs that facilitate understanding of the current experiments.
The conserved γδT cell lineage chassis implies that mechanosensing is not a feature of γδTCRs, but detailed assessment of other TCRs using biophysical methods performed here is warranted. That said, TCR ligation-based exposure of the CD3ε proline-rich cytoplasmic region in αβTCRs but not γδTCRs mapping to their respective constant regions reported previously (63) is consistent with the generality of differential mechanotransduction revealed here.

Anticipating that mechanosensing would augment TCR signaling upon sulfate exposure, we observed that the chimeric γ-αβTCR-transfected thymocytes showed greater activation than the WT γδTCR-transfected thymocytes in CD1d–OP9 DL4 epithelial cultures at DN stages and beyond. This was initially revealed as a reduction in cell surface CD3 expression (Fig. 4). Subsequently, a global, unbiased assessment of signaling differences was determined by RNA-seq transcriptome analysis of γδTCR and γ-αβTCR thymocytes in CD1d–OP9DL4 epithelial cultures with and without sulfate addition. These data clearly showed that stimulation of γ-αβTCR, relative to that of γδTCR, induced greater regulated expression of a multiplicity of genes in the DN4 compartment, many of which are specifically associated with T cell stimulation, adhesion control, chemotaxis, signaling, and cellular metabolism (Fig. 6).

Increasing evidence supports a model of gene network-driven lymphocyte lineage diversification preceding antigen receptor expression (64, 65). Although TCR signaling might contribute to γδT lineage fate, gene network drivers per se are a dominant component. Along these lines, we find by PCA that DP10.7 γδTCR manifest small differences in gene expression in the presence or absence of sulfate in stromal cultures, relative to the γ-αβTCR chimera. It is important to view the cellular results presented herein strictly as an indicator of the signaling capacity of the given receptors, and not as a study of developmental pathways of γδT cells per se, since aside from hints of preferential chimeric receptor deletion with sulfate addition, evidence for ligand-directed progression was not unequivocal within the OP9–DL4 stromal system for this γδTCR.

γδT lineage cells are the first to exit the thymus, having already acquired effector function and been programmed to populate different anatomical epithelial locales linked to their Vγ usage (reviewed in ref. 66). These cells demonstrate ligand recognition straddling both innate and adaptive immune spaces. A majority of γδTCR ligands are self-derived and stress-induced in lymphoid and nonlymphoid cells, including epithelial cells. For example, CD1 molecules as well as T10/T22 have prominent display in human and mouse thymus, respectively, with CD1d shared between species (67). The recognition by TCRs, even αβTCRs, of ligands expressed at high copy numbers does not require mechanosensing (20). Therefore, if γδT cell ligands are densely arrayed constitutively or upon up-regulation by cellular perturbations involving stress responses, including inflammation, then ligand multivalency per se is adequate to stimulate T cell signaling. Reduction in ligand density or attenuation of T cell signaling upon up-regulation by cellular perturbations may be disadvantageous in sensing a gradient or threshold, in the presence of higher ligand concentrations representative of common target antigens for γδT cells. Similar analog αβTCR–pMHC interactions may be involved in positive selection in the thymus, homeostatic T cell proliferation in the periphery, or antiviral responses to high-density ligands on infected cells. Thus, while αβTCR cells may exploit both modalities, γδT cells appear to be designed exclusively for analog signaling. The implications of this distinction and further analysis of their molecular mechanisms may have translational impact in areas involving adoptive cellular therapies as well as vaccine design.
Materials and Methods

Choice of TCR Structures for MD Simulations. For the WT TCRβj1, N15 TCRβj1 Protein Data Bank (PDB), 1NFD (33) and JM22 TCRβj, 1OGA (75), were used. PDB 1NFD corresponds to the N15 TCRβj used in experiments. JM22 TCRβj PDB 1OGA has a bound pMHC and is currently the highest in resolution available (1.40 Å). For the WT TCRγδ, we used SC2, PDB 4HLH (76). For the Vβ5–Cγ1 hybrid, we used DP10.7 TCRδ, PDB 4MNH (31) and replaced the Cβj part with those from PDB 1NFD, to match the construct with the one used in experiments (DP10.7–N15). This was done by aligning TCRs of 1NFD and 4MNH using Modeler (77) and replacing the Cβj part of 4MNH with that of 1NFD. For Vδi, residues up to L119 were kept, after which was Cα of 1NFD. For Vγi, residues up to P119 were kept, followed by Cβ of 1NFD. To ensure that the replacement of constant domains has minimal impact on the interface with the variable domains, we compared interdomain contacts between the original 4MNH structure and the one with Cβj from 1NFD, prior to performing simulation. We found that they have very similar hydrogen bonds and nonpolar contacts, using nearly identical sets of residues, which reflect the sequence homology of the constant domains in 4MNH and 1NFD. The Vβj–Cγ1 chimera was built similarly, using the Vβj domain of PDB 1NFD (up to residue 112 in both α- and β-chains) and the rest of the Cβj of PDB 4HLH, starting from R120 of the δ-chain and K126 of the γ-chain. Merging loops in structures were built using Modeler, and hydrogen atoms were added in CHARMM (78). None of the missing residues are at the interdomain interface, hence they do not affect our interdomain contact analysis. Disulfide bonds were placed as they appear in respective domains.

MD Simulation and Analysis. Simulations were performed using CHARMM (79). Each construct was soluted in a cubic water box of about 98 Å in each dimension, which has boundaries at least 12 Å away from the protein. Sodium and chloride ions were added at about 50 mM concentration to neutralize the system. The simulation system underwent a series of energy minimization procedures (4,000 steps in total), where a set of gradually decreasing harmonic restraints was applied to the protein to remove close contacts and relax the surrounding water molecules and ions. After initial energy minimization, the system was heated from 30 K to 300 K during 100 ps, and equilibrated at 300 K for 200 ps. During heating and equilibration, backbone heavy atoms were harmonically restrained with a spring constant of 5 kcal/mol Å2. Pressure was maintained at 1 atm using the constant volume, temperature ensemble. Finally, a 300-ns production run was performed without any restraint applied (120 ns for the Vβj–Cγ1 chimera). Cutoff distance for nonbonded interaction was 12 Å. The particle-mesh Ewald summation method was used to account for the long-range electrostatic interactions. The SHAKE algorithm was used to fix the length of the covalent bond between hydrogen and heavy atoms. For integration, a 2-fs time step was used, and coordinates were saved every 20 ps. A periodic boundary condition was applied during the simulation. The domain decomposition module was used for efficient parallelization (79).

Contact analysis was performed as described (20, 80). Briefly, a 2.4 Å cutoff distance between a hydrogen atom and hydrogen bond acceptor atoms was used to identify hydrogen bonds. For nonpolar contacts, a 3.0 Å cutoff distance between atoms with the absolute value of partial charge less than 0.3e (e: charge of an electron) was used. Occupancy of a contact was calculated as the fraction of coordinate frames during 100 to 300 ns (50 to 120 ns for the Vβj–Cγ1 chimera; SI Appendix, Fig. S1E). For the contact statistics in Fig. 2A, 10 364-ns windows that overlap 50% (i.e., 100 to 136.4 ns, 118.2 to 154.5 ns, etc.) were used and contact occupancies were calculated in each window. Windows were made to overlap to avoid a contact appearing to have a low occupancy if spread between two nonoverlapping windows.

Single-Molecule Tethering Experiments. Simulations were performed using as previously described (31, 81) except for the insertion of flexible linker and LZ motifs (82) prior to the 3C protease site and 6 (31, 81) except for the insertion of flexible linker and LZ motifs (82) prior to DP10.7 to have a low occupancy if spread between two nonoverlapping windows.

Constructs for Cellular Experiments. DP10.7 γTγR (WT) and γδ–γTγR (chimeric) constructs, for cellular and SMSC experiments were cloned from previously reported constructs (31, 81). Viral 2A-linked system sites were inserted between subunits to create single δ-p2α−γ or α-p2α−γ constructs for WT or chimeric constructs, respectively, as previously published for γδ and γδ TCR (30, 31, 38, 81). Chimeric constructs incorporate the constant domain, connecting peptide region and TM region of the α- or β-subunit as appropriate. Site-directed mutagenesis was used to create appropriate EcORI and NotI restriction sites for insertion into L2RS-IresGFP (addgene) retroviral vectors for use in OP9-DL4 stromal cell cultures (35). DNA and protein sequences are included in Dataset S2.

Generation of BW5147 Cells. BW5147 cells were cotransduced with plasmids pMY1 encoding CD3ζ and pMIG1 (addgene) encoding either DP10.7 γδ or γδ–ζ essentially as described in ref. 28. Additional details are in SI Appendix, Materials and Methods.

SMSC Assay. The SMSC assay was carried out essentially as detailed (19). Additional details are in SI Appendix, Materials and Methods.

OP9–DL4 Stromal Cell Culture. OP9–DL4 and OP9–DL4–CD1d stromal cell cultures were performed as described (35). Additional details are in SI Appendix, Materials and Methods.

Flow Cytometry and Cell Sorting. For cell-surface molecule staining, transfected cells were first treated with anti-mouse CD16/CD32 mAbs (2.4 G2) in staining buffer (2% fetal bovine serum and 0.05% sodium azide in phosphate buffered saline) to block FcR binding and then stained with antibodies. Antibodies used in this study are listed below. Zombie Aqua (BioLegend) was used for staining dead cells. Intracellular IL-17 and IFN-γ staining was performed after stimulation with 50 ng/ml PMA, 500 ng/ml ionomycin, and 5 μM Brefeldin A for 4 h. For intracellular staining, cells were fixed with 4% paraformaldehyde and treated with permeabilization buffer (0.1% saponin in staining buffer) and then incubated with the antibodies against intracellular cytokine. Lymphocytes and thymocytes from a 9-wk-old female C57BL/6 (Taconic) mouse were used for the positive control of cytokine expression. Cells were analyzed on an LSR Fortessa (BD Biosciences) as described below. Data were analyzed with FlowJo (Tree Star). The APC-conjugated DP10.7 TCR tetramer was produced as described (31). For DP10.7 TCR tetramer staining, OP9–DL4, OP9–DL4–MHC KO, and OP9–DL4–CD1d stromal cells were plated at 2.5 × 105 cells followed by overnight incubation. Cells were cultured with or without sulfatide (3 μg/ml) for 2.5 h and then stained by 500 nM tetramer in 10% human serum/ staining buffer.

Transcriptome Analysis. A total of 2,000 γTγR- or γδ–γTγR-transduced GFP+ CD45+ DN3 cells were sorted, plated onto OP9–DL4–CD1d cells, which were plated the day before at 5 × 106 cells in six-well plates and cultured in OP9 medium supplemented with Filt-3, IL-7, and gentamicin with or without sulfatide. After 8 d, 5,000 live DNA (GFP+CD45+CD8+CD44+CD25+) cells and DNA (GFP+CD45+CD8+CD44+CD25+) cells were sorted (SI Appendix, Fig. S4).

For each condition (γTγR or γδ–γTγR, DN3, or DNA, plus or minus sulfatide, yielding eight conditions/experiment), 5,000 cells were deposited into 350 μl of TCL lysis buffer (Qiagen) and stored at −80 °C until RNA isolation. Three independent experiments generated 24 libraries. Total RNA was then
purified from the stabilized lysates using the ARCTURUS PicoPure RNA isolation kit (Thermo Fisher). Following RNA purification, residual DNA was removed by treatment with the Turbo DNA-free reagent kit (Thermo Fisher). Total RNA was quantified using the Qubit RNA assay kit (Life Tech) and RNA quality was determined on a bioanalyzer using the RNA Pico chip (Agilent). The NuGen Ovation Human RNA-Seq Multiplex system (NuGen, part 0341) prep kit, was used to target deletion of unwanted high abundance transcripts and ribosomal RNA. More than 100 ng of total RNA was converted into each DNA library following the manufacturer's protocol with the Gene Watson Library construction. DNA libraries were quantified using the Qubit High Sensitivity DNA kit (Life Tech), and library size was determined using the Bioanalyzer High Sensitivity Chip kit (Agilent). Finally, qPCR was carried out on the libraries using the Universal Library Quantification kit for Illumina (Kapa Biosystems) and run on the 7900 HT Fast qPCR machine (ABI). Libraries passing quality control were diluted to 2 nM in sterile water and sequenced on the NextSeq500 (Illumina) at a final concentration of 12 pM, following all manufacturer protocols.

Antibodies. Antibodies are detailed in SI Appendix, Materials and Methods.

Bioinformatic and Statistical Analysis. The output fastq files were aligned against the Ensembl GRCm38.75 reference genome using STAR aligner (v2.5.1) (86) and the resultant binary alignment map (BAM)-format files were filtered to retain only primary-aligned reads (samtools view -F 0 100). The reads were mapped to the ferret genome using the Turbo DNA-free reagent kit (Thermo Fisher). Total RNA was quantified using the Qubit RNA assay kit (Life Tech) and RNA quality was determined on a bioanalyzer using the RNA Pico chip (Agilent). The NuGen Ovation Human RNA-Seq Multiplex system (NuGen, part 0341) prep kit, was used to target deletion of unwanted high abundance transcripts and ribosomal RNA. More than 100 ng of total RNA was converted into each DNA library following the manufacturer's protocol with the Gene Watson Library construction. DNA libraries were quantified using the Qubit High Sensitivity DNA kit (Life Tech), and library size was determined using the Bioanalyzer High Sensitivity Chip kit (Agilent). Finally, qPCR was carried out on the libraries using the Universal Library Quantification kit for Illumina (Kapa Biosystems) and run on the 7900 HT Fast qPCR machine (ABI). Libraries passing quality control were diluted to 2 nM in sterile water and sequenced on the NextSeq500 (Illumina) at a final concentration of 12 pM, following all manufacturer protocols.

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