

$\gamma\delta$ 2T cell diversity and the receptor interface with tumor cells

Anna Vyborova,¹ Dennis X. Beringer,¹ Domenico Fasci,¹ Froso Karaiskaki,¹ Eline van Diest,¹ Lovro Kramer,¹ Aram de Haas,¹ Jasper Sanders,¹ Anke Janssen,¹ Trudy Straetemans,¹ Daniel Olive,² Jeanette Leusen,¹ Lola Boutin,³ Steven Nedellec,⁴ Samantha L. Schwartz,^{5,6} Michael J. Wester,⁷ Keith A. Lidke,⁷ Emmanuel Scotet,³ Diane S. Lidke,^{5,6} Albert J.R. Heck,^{8,9} Zsolt Sebestyen,¹ and Jürgen Kuball^{1,10}

¹Center for Translational Immunology, University Medical Center (UMC) Utrecht, Utrecht University, Utrecht, Netherlands. ²Centre de Recherche en Cancérologie Marseille, INSERM, Institut Paoli-Calmettes, Marseille, France. ³Université de Nantes, INSERM, CNRS, CRCINA, LabEx IGO "Immunotherapy, Graft, Oncology," Nantes, France. ⁴Structure Fédérative de Recherche en Santé François Bonamy (SFR-Santé), INSERM, CNRS, CHU Nantes, Nantes, France. ⁵Department of Pathology, ⁶Comprehensive Cancer Center, and ⁷Department of Physics and Astronomy, University of New Mexico (UNM), Albuquerque, New Mexico, USA. ⁸Biomolecular Mass Spectrometry and Proteomics, Bijvoet Center for Biomolecular Research and Utrecht Institute for Pharmaceutical Sciences, Utrecht University, Utrecht, Netherlands. ⁹Netherlands Proteomics Centre, Utrecht, Netherlands. ¹⁰Department of Hematology, UMC Utrecht, Utrecht University, Utrecht, Netherlands.

$\gamma\delta$ 2T cells play a major role in cancer immune surveillance, yet the clinical translation of their in vitro promise remains challenging. To address limitations of previous clinical attempts using expanded $\gamma\delta$ 2T cells, we explored the clonal diversity of $\gamma\delta$ 2T cell repertoires and characterized their target. We demonstrated that only a fraction of expanded $\gamma\delta$ 2T cells was active against cancer cells and that activity of the parental clone, or functional avidity of selected $\gamma\delta$ 2T cell receptors ($\gamma\delta$ 2TCRs), was not associated with clonal frequency. Furthermore, we analyzed the target-receptor interface and provided a 2-receptor, 3-ligand model. We found that activation was initiated by binding of the $\gamma\delta$ 2TCR to BTN2A1 through the regions between CDR2 and CDR3 of the TCR γ chain and modulated by the affinity of the CDR3 region of the TCR δ chain, which was phosphoantigen independent (pAg independent) and did not depend on CD277. CD277 was secondary, serving as a mandatory coactivating ligand. We found that binding of CD277 to its putative ligand did not depend on the presence of $\gamma\delta$ 2TCR, did depend on usage of the intracellular CD277, created pAg-dependent proximity to BTN2A1, enhanced cell-cell conjugate formation, and stabilized the immunological synapse (IS). This process critically depended on the affinity of the $\gamma\delta$ 2TCR and required membrane flexibility of the $\gamma\delta$ 2TCR and CD277, facilitating their polarization and high-density recruitment during IS formation.

Introduction

Cancer immunotherapies with classical checkpoint inhibitors and CAR T cell therapies are limited by their dependency on the mutational load of the tumor and by the lack of targets for solid tumors (1). Within this context, clinical developers currently have a significant, renewed interest in $\gamma\delta$ T cells, because these cells have been reported to be broadly active against multiple solid cancers and hematological malignancies (2, 3). Although very few targets of

$\gamma\delta$ T cells and their receptors have been described to date, the current understanding is that $\gamma\delta$ T cells act on early metabolic changes in cancer cells, which makes $\gamma\delta$ T cells attractive candidates to attack tumors with a low mutational load (for review see ref. 1). However, to date, most clinical trials have shown very limited clinical efficacy, particularly when using expanded $\gamma\delta$ 2T cells (1, 4).

We propose 2 possible causes for the lack of clinical success of adoptively transferred $\gamma\delta$ 2T cells in humans. First, most of the published and ongoing studies use polyclonally derived $\gamma\delta$ 2T cells (1, 4). However, recent work suggests a substantial functional heterogeneity in this cell population (5). It is not clear whether this diversity affects the activity of $\gamma\delta$ 2T cells against cancer cells when used for adoptive immune therapies. A second obstacle might be the poor selection of patients for clinical trials, which is a consequence of the limited understanding of the targets expressed on tumors. Upon encountering the target, $\gamma\delta$ 2T cells are known to sense elevated phosphoantigen (pAg) levels through the small GTPase RhoB (6), which induces a joint spatial and conformational change in surface expression of CD277 (BTN3A) (6–12) via an inside-out signaling mechanism. A direct interaction between the $\gamma\delta$ 2T cells and CD277, involving the isoforms BTN3A1, BTN3A2, and BTN3A3, has been suggested (13). Given the homology of BTNA1, -A2, and -A3, the Abs directed

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against extracellular domains of BTN3A1 usually cross-recognize BTN3A2 and BTN3A3 as well (14); therefore, with CD277, we refer to all 3 isoforms. The rearrangement in CD277 architecture that induces $\gamma\delta$ 2T cell activation is referred to herein as CD277J (14). However, it remains highly controversial whether CD277 interacts directly with the $\gamma\delta$ 2T cell receptor ($\gamma\delta$ 2TCR) and whether additional molecules are involved (7, 8, 13, 14).

In this study, we explored the functional heterogeneity of $\gamma\delta$ 2T cells on a clonal level, as well as the impact of individual $\gamma\delta$ 2TCRs on mediating antitumor responses in relation to their CDR3 usage and the putative ligand CD277. To this end, we isolated and functionally characterized an extensive panel of single $\gamma\delta$ 2T cell clones, and we cloned and expressed individual $\gamma\delta$ 2TCR genes. By engineering $\alpha\beta$ T cells to express a defined $\gamma\delta$ TCR (TEG) as a next generation of chimeric antigen receptor T cells (CAR-T cells), we were able to assess the functional impact of individual $\gamma\delta$ 2TCRs on sensing the target, in the absence of the original functional programming of the parental clone, as well as without other NK and KIR receptors usually expressed on $\gamma\delta$ 2T cells (2). In addition to providing a valuable tool for the molecular characterization of individual $\gamma\delta$ 2TCRs, the TEG concept combines the positive aspects of 2 different cell types: the memory formation and high-proliferation capacity of $\alpha\beta$ T cells and the broad antitumor reactivity of defined $\gamma\delta$ 2TCRs (15, 16). TEGs demonstrated the ability to distinguish between healthy and malignant leukemic stem cells (6) and thereby overcome the obstacle of leukemic stem cells that lack, e.g., NKG2D ligands (17). In addition, we have shown that TEGs eliminate primary multiple myeloma cells in a humanized bone marrow niche (18). As a result of these findings, TEGs are currently being tested in a first-in-human clinical trial (19, 20). Within this context, our current report demonstrates that the failure of previous clinical trials using polyclonal $\gamma\delta$ 2T cells can be partially attributed to a high frequency of $\gamma\delta$ 2T cells with poor tumor reactivity within healthy immune repertoires (14). In addition, after analyzing the molecular interface between cells expressing a $\gamma\delta$ 2TCR and CD277, we propose a paradigm shift in the model of recognition, in which the $\gamma\delta$ 2TCR is only critical for initiating cell conjugation, functioning as an initial signal of T cell activation and maintaining high-density immunological synapse (IS), but is not dependent on pAgs. Conversely, CD277 operates as a mandatory coactivating ligand that functions as a second signal, stabilizes conjugation and synapse formation, and is dependent on pAgs, but not on the presence of the $\gamma\delta$ 2TCR.

Results

Clonal $\gamma\delta$ 2T cell functional diversity. In order to assess the impact of clonal heterogeneity on the antitumor functional activity of the $\gamma\delta$ 2 T cell population in a given individual, we isolated and expanded $\gamma\delta$ 2 T cell clones from the peripheral blood of healthy donors. Next, we determined the *in vitro* antitumor reactivity of individual clones by measuring their response to the Burkitt lymphoma cell line Daudi, which can be considered a model target for studying pAg-mediated and TCR-dependent $\gamma\delta$ 2 T cell responses (reviewed in ref. 21). IFN- γ production was used as a readout, since the capacity to secrete IFN- γ has been shown to be a property of the whole $\gamma\delta$ 2 T cell compartment, regardless of the subset (5). We observed substantial variation in cytokine secretion capacity

among $\gamma\delta$ 2 T cell clones isolated from 3 healthy donors (donors A–C), with approximately 60% of the clones being tumor reactive according to this readout (cutoff 30 pg/mL; Figure 1A). Bisphosphonates such as pamidronate (PAM) are known to boost the $\gamma\delta$ 2 T cell responses by inducing or enhancing CD277J (9). Consistent with these findings, in the presence of 100 μ M PAM, the reactivity of the $\gamma\delta$ 2T cell clones was increased by 10-fold on average (median 8.1, IQR = 15.0–3.4), with more than 95% of the clones showing reactivity above the threshold when PAM was added (Figure 1A and Supplemental Figure 1A; supplemental material available online with this article; <https://doi.org/10.1172/JCI132489DS1>). Other Th1 spectrum cytokines measured in a subset of the investigated clones showed a high correlation with IFN- γ and with each other, indicative of a robust signature (Supplemental Figure 1, B–D). HEK293T, an additional well-characterized $\gamma\delta$ 2 T cell target (6, 9), elicited much lower cytokine secretion in the absence of PAM, however, in the presence of PAM, reactivity of the clones was comparable to their reactivity to Daudi cells (Figure 1, B and C).

Antitumor response of individual $\gamma\delta$ 2T cell clones does not correlate with clonal frequency. As little is known about functional implications of the diversity in the $\gamma\delta$ 2TCR repertoire, it was particularly interesting to explore the antitumor response of the individual $\gamma\delta$ 2T cell clones in the context of their clonotype frequencies. We selected 20 clones for TCR sequencing to track their clonotype within the original bulk population, covering the full range of clonal activities, as depicted in Figure 1A (arrows). Several clones (B2/B5, C4/C14, C7/C11/C15, C6/C9/C13) were found to express the same $\gamma\delta$ 2TCR, yielding a total of 16 unique clonotypes. All δ chains were unique, whereas we observed public (shared between donors) CDR3 γ sequences, as well as pairing of 1 TCR γ 9 with different TCR δ 2 chains in the same donor (Table 1), in line with high-throughput sequencing (HTS) data reported by others (22–24). Next, we performed HTS of the complete TCR δ 2 repertoires of the same donors. The prevalence of a T cell receptor δ locus (TRD) clonotype within the repertoire of a donor did not correlate with reactivity of the respective clone against Daudi or HEK293T cells, as demonstrated by the example of donor C, for whom 2 of the most prevalent TCR δ clonotypes C6/C9/C13 and C17 belonged to 3 of the most reactive $\gamma\delta$ 2 T cell clones isolated from this donor, whereas other prevalent clonotypes (C2/C18, C4/C14, C7/C11/C15) showed low reactivity (Figure 1, A and D). The TCR δ clonotypes of the highly reactive clones from donors A and B (B1, A3–A6) appeared to be rare, as the sequences could not be detected in the HTS data for the respective donor (Figure 1A and Supplemental Figure 1, E and F).

Functional avidity of the parental $\gamma\delta$ 2T cell clone does not correlate with $\gamma\delta$ 2TCR-mediated functional avidity. We aimed to assess whether the antitumor activity of a given $\gamma\delta$ 2T cell clone would associate with its functional avidity mediated by its respective TCR, as suggested by our previous study (15). We defined the binding strength of an individual receptor ligand interaction as affinity, and the combined strength governed by multiple receptor-ligand interactions as avidity. To measure the avidity of a defined $\gamma\delta$ 2TCR, we assessed the functional avidity of TEGs by measuring effector functions, like IFN- γ secretion, against defined targets (25–27) and compared it with the functional avidity of the parental $\gamma\delta$ 2T cell clone, which also harbors other

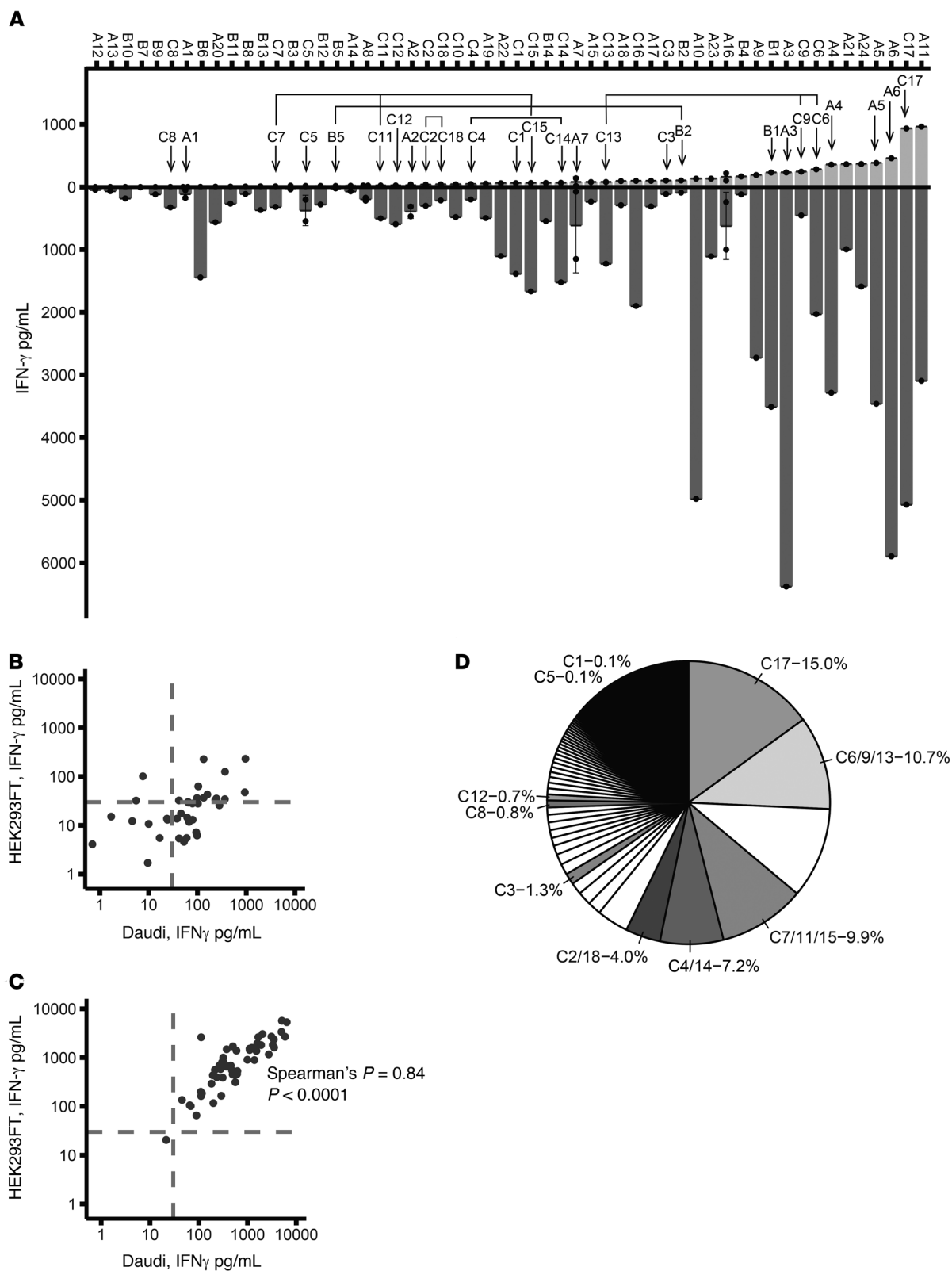


Figure 1. Antitumor reactivity of the $\gamma 9\delta 2$ T cell clones and TRD clonotype frequencies for donor C. (A) Antitumor reactivity of the isolated clones assessed by IFN- γ production ($n = 56$). Bars represent the means of 2 to 4 biological replicates from 1 to 2 independent experiments. Arrows indicate clones with a known TCR sequence. Connecting lines indicate clones expressing the same $\gamma 9\delta 2$ TCR. (B and C) IFN- γ production against Daudi versus HEK293T cell lines without (B) or in the presence of 100 μ M PAM (C). Dashed lines represent the cutoff for reliable ELISA measurements (30 pg/mL). (D) Complete T cell receptor delta variable region (TRDV) repertoire for donor C. Percentages indicate clonotype frequencies. Data were filtered to exclude clonotypes with a frequency of 1 read/clonotype.

Table 1. TCR sequencing of the selected $\gamma 9\delta 2$ T cell clones

Clone ID	TCR γ	CDR3 γ	CDR3 γ length	TCR δ	CDR3 δ	CDR3 δ length
A1	TRGV9*01TRGJP*01	CALWEVKELGKKIKVF	14	TRDV2*03TRDD2*01TRDD3*01TRDJ1*01	CACDTLLLLGDSSDKLIF	16
A2	TRGV9*01TRGJP*01	CALWEV Q ELGKKIKVF	14	TRDV2*03TRDD3*01TRDJ1*01	CACDTAEPGGYKDKLIF	15
A3	TRGV9*01TRGJP*01	CALWEV Q ELGKKIKVF	14	TRDV2*03TRDD3*01TRDJ1*01	CACDAWGHGTDKLIF	12
A4	TRGV9*01TRGJP*01	CALWEVEGLGKKIKVF	14	TRDV2*03TRDD3*01TRDJ1*01	CACDALGDTGSDKLIF	14
A5	TRGV9*01TRGJP*01	CALWEAGELGKKIKVF	14	TRDV2*03TRDD3*01TRDJ1*01	CACDTLGALYTDKLIF	14
A6	TRGV9*01TRGJP*01	CALWEV Q ELGKKIKVF	15	TRDV2*03TRDD3*01TRDJ1*01	CACDQLGDPDKLIF	12
A7	TRGV9*01TRGJP*01	CALWEV RE LGKKIKVF	14	TRDV2*03TRDD3*01TRDJ1*01	CACDSDPWGILNTDKLIF	17
B1	TRGV9*01TRGJP*01	CALWEA Q VELGKKIKVF	15	TRDV2*03TRDD3*01TRDJ1*01	CACDTPWGIAPAAQTDKLIF	18
B2/B5	TRGV9*01TRGJP*02	CALWEVRY Y KKLF	11	TRDV2*03TRDD2*01TRDD3*01TRDJ1*01	CACDPRFLAWGTTDKLIF	16
C1	TRGV9*01TRGJP*01	CALWEV RE LGKKIKVF	14	TRDV2*01TRDD2*01TRDJ1*01	CACDPVPSIHDTKLIF	15
C2	TRGV9*01TRGJP*01	CALWEV RE LGKKIKVF	14	TRDV2*03TRDD3*01TRDJ1*01	CACDQAGGPDKLIF	12
C3	TRGV9*01TRGJP*01	CALWEVGLGKKIKVF	13	TRDV2*01TRDD3*01TRDJ1*01	CACDTPVSGGYQYTDKLIF	16
C4/C14	TRGV9*01TRGJP*01	CALWEVSGELGKKIKVF	15	TRDV2*02TRDD2*01TRDD3*01TRDJ1*01	CACDTLALGDTDKLIF	14
C5	TRGV9*01TRGJP*01	CALWEV Q ELGKKIKVF	14	TRDV2*02TRDD3*01TRDJ1*01	CACDLLAPGDTSFDTKLIF	17
C6/C9	TRGV9*01TRGJP*01	CALWEVKRELGKKIKVF	15	TRDV2*02TRDD3*01TRDJ1*01	CACDTPVVLGGYRDDKLIF	18
C7/C11	TRGV9*01TRGJP*01	CALWEV RE LGKKIKVF	14	TRDV2*01TRDD3*01TRDJ3*01	CACDMGDASSWDTRQMF	16

Paired sequences of the TCR δ and γ CDR3 regions of the selected clones. Sequences were analyzed using the IMGT/V-QUEST tool (http://www.imgt.org/IMGT_vquest/user_guide) in order to determine the identity of the germline segments. Amino acid sequences defining public TCR $\gamma 9$ clonotypes (24, 45) and amino acids at "position 5" of the TCR $\delta 2$ (24) are highlighted in bold. The conserved cysteine of the V region and phenylalanine of the J region were not included when calculating the CDR3 length.

NK-like receptors (Figure 1). First, the individual $\gamma 9\delta 2$ TCRs were expressed in the absence of other receptors present on parental $\gamma \delta$ T cells in the human TCR-deficient T cell line Jurkat-76. The previously reported low functional avidity ("weak") $\gamma 9\delta 2$ TCR CTE-Cl3, high functional avidity ("strong") $\gamma 9\delta 2$ TCR CTE-Cl5, and nonfunctional length mutant 1 (LM1) that were generated by combinatorial TCR chain exchange (CTE) with the chains of the TCR G115 (15, 28) were used as an additional framework. For $\gamma 9\delta 2$ TCRs from all donors, we observed consistent differences in functional avidity, as evidenced by unequal CD69 upregulation and downregulation of the $\gamma \delta$ TCR upon stimulation with the cell line Daudi (Figure 2, A-F, and Supplemental Figure 2, A-F) in cells selected for equivalent $\gamma 9\delta 2$ TCR expression. These cells were able to overcome heterogeneous $\gamma 9\delta 2$ TCR expression from the initial bulk, and later from engineered immune cells (Supplemental Figure 2, G and H). Differences in functional avidity were even more evident in $\gamma 9\delta 2$ TCR-transduced primary $\alpha \beta$ T cells with equivalent $\gamma 9\delta 2$ TCR expression (Supplemental Figure 2, G-J), when we used IFN- γ secretion or tumor cell lysis as a readout (Figure 2, G and H), since primary T cells have a better capacity to produce cytokines and directly kill than do Jurkat cell lines (29). The outcomes of different readouts, such as CD69 upregulation for Jurkat-76, and IFN- γ secretion and tumor cell lysis for TEGs, correlated with each other, indicating that the effect was indeed due to the introduced $\gamma 9\delta 2$ TCRs with defined sequences and was not the property of a defined transduced cell (Supplemental Figure 2I).

Although responses mediated by distinct $\gamma 9\delta 2$ TCRs, such as TCR A3, correlated with the ability of the parental clone to mediate exceptionally high antitumor reactivity upon PAM stimulation (Figure 1A), functional avidity mediated through other TCRs did not correlate with the antitumor response of their parental clone.

For example, the primary clones A1 and A2 showed similarly low reactivity (Figure 1A), whereas TCRs isolated from these clones mediated unequal responses in the TEG format (Figure 2, A, D, G, and H). Conversely, the TCR A5, whose parental clone showed antitumor activity in the higher range, did not differ in activation potential from the TCR derived from the clone A1. These data imply that epigenetic imprints modulate the response of primary $\gamma 9\delta 2$ T cells on a clonal level (2, 5, 30, 31). Although the functional avidity of a given $\gamma 9\delta 2$ TCR CDR3 seems to be of ambiguous relevance for the response of its parental T cell clone, we demonstrate here that $\gamma 9\delta 2$ TCR CDR3 is crucial for the antitumor activity of TEGs. In addition, our data suggest a range of affinities for different $\gamma 9\delta 2$ TCRs that do not, however, exceed a certain threshold. The assumption of these experiments would be that polyclonal $\gamma 9\delta 2$ T cells are inferior in terms of reactivity against tumor targets when compared with TEGs expressing high-affinity $\gamma 9\delta 2$ TCR. To test this hypothesis, we coinoculated $\gamma 9\delta 2$ TCR CTE-Cl5-engineered immune cells and bulk $\gamma \delta$ T cells with a variety of different tumor cell lines and with primary acute myeloid leukemia (AML) blasts. As expected, $\gamma 9\delta 2$ TCR CTE-Cl5-engineered T cells showed superior IFN- γ production in response to various tumor cells and primary AML blasts, thus outperforming bulk $\gamma \delta$ T cells (Supplemental Figure 2, K and L).

Strength of the $\gamma 9\delta 2$ TCR-tumor interaction depends on the tumor type, number of $\gamma 9\delta 2$ TCR molecules, and the CDR3 domain of the TCR δ chain. As TEG antitumor activity was fully dependent on the $\gamma 9\delta 2$ TCR CDR3 region, we next focused on a better characterization of the molecular interface of the $\gamma 9\delta 2$ TCR with its target, by taking advantage of soluble $\gamma 9\delta 2$ TCRs. Because previous reports suggested that $\gamma \delta$ TCR multimers are a suitable tool to study $\gamma \delta$ TCR-ligand interactions (32-35), we first engineered tetram-

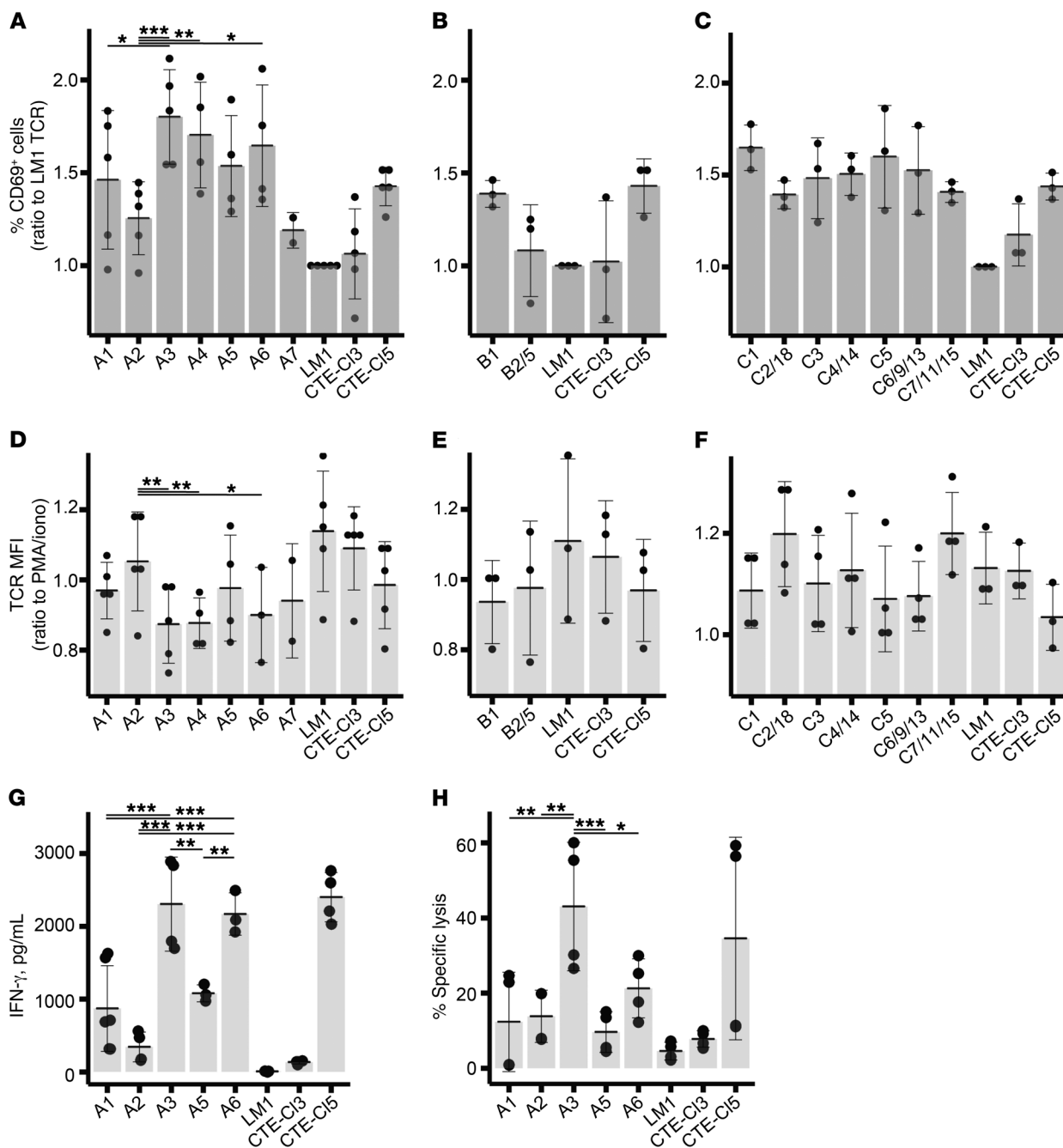


Figure 2. Functional avidity of the isolated TCRs in the TEG format. (A–F) Functional avidity of the TCRs from donors A, B, and C, next to control TCRs, transduced into the Jurkat-76 cell line ($n = 2$ –5 from at least 2 independent transductions). (A–C) CD69 upregulation (percentage of CD69⁺ cells) upon stimulation with the Daudi cell line, normalized to the TCR LM1 within each experiment. (D–F) TCR downregulation measured by the MFI of TCRV δ 2 on Jurkat-76 cells coincubated with Daudi cells, relative to the MFI for TCRV δ 2 on the same transductants treated with PMA and ionomycin (PMA/iono). Data represent mean values \pm SD across experiments. (G and H) Functional avidity of the selected TCRs from donor A, next to control TCRs, transduced into primary human $\alpha\beta$ T cells. (G) IFN- γ production upon coincubation with Daudi cells in the presence of 100 μ M PAM. Data represent mean values \pm SD across 2 representative experiments. (H) Specific lysis of Daudi cells. Bars show mean values of 2 side-by-side biological replicates (see Supplemental Figure 21 for another representative experiment). A P value of less than 0.05 was considered significant. * $P < 0.05$, ** $P < 0.01$, and *** $P < 0.001$, by 2-way ANOVA with Tukey's post hoc test.

ers of the $\gamma\delta$ 2TCRs Cl5 and the nonfunctional TCR LM1 (15), as well as the low-reactive clone A2 and the high-reactive clone A3 identified here. As controls, we included an $\alpha\beta$ TCR and a V γ 4V δ 5 TCR. In contrast to previous reports with $\gamma\delta$ 2TCR tetramers (33–35) and in line with the more recent suggestions of low-affinity

interactions that are needed to sense target cells (6, 8), we found that $\gamma\delta$ 2TCR tetramers did not strongly stain the classical target Daudi, with only a slight but significant increase in MFI for the TCRs Cl5 and A3 compared with the control TCRs (Figure 3A). However, when we screened tumor cell lines other than Daudi,

such as HEK293T, $\gamma\delta$ TCR tetramers of the non-/low-reactive $\gamma\delta$ TCR LM1 and A2 had previously bound, whereas binding was further increased when using $\gamma\delta$ TCR tetramers were derived from the highly active clones, clone 5 and clone A3 (Figure 3B).

Since both Daudi and HEK293T cells are targets of the $\gamma\delta$ TCR, we addressed the question of whether raising the number of TCRs on a multimer to increase the overall avidity would result in a more pronounced staining of Daudi cells. The extracellular domains of the TCRs LM1, Cl5, A2, and A3 were coupled to streptavidin-conjugated fluorescent beads, which allowed at least 10^4 $\gamma\delta$ TCRs to be attached to the bead surface. $\gamma\delta$ TCR-coated beads were coincubated with the negative target cell line ML1 (6) and the positive target cell line Daudi. No bead conjugation to ML1 could be observed in any of the $\gamma\delta$ TCRs (Supplemental Figures 3 and 4), whereas bead conjugation to Daudi cells was dependent on the $\gamma\delta$ TCR CDR3 region (Figure 3C and Supplemental Figures 3 and 4). TCR LM1 cells showed some conjugation to Daudi cells, just like clone A2, whereas the “strong” TCRs Cl5 and A3 showed significantly higher conjugation, indicating that differences in functional avidity can be attributed to the unequal affinity of the $\gamma\delta$ TCR-ligand interaction. The pronounced differences observed on Daudi cells between the different $\gamma\delta$ TCRs were less evident, or even absent, when we used these fluorescent beads on HEK293T cells (Figure 3D). Importantly, bead staining could be blocked by an anti- $\gamma\delta$ TCR Ab (Supplemental Figure 5A), suggesting a TCR-specific interaction. Of note, we did not observe $\gamma\delta$ TCR tetramer staining or $\gamma\delta$ TCR bead staining for some of the tumor cells that have been reported to be targeted by the $\gamma\delta$ TCR (Table 2).

BTN2A1 was most recently suggested as a ligand for the $\gamma\delta$ TCR through its TCR γ chain (35, 36). First, we tested whether one of the proposed mutations in the TCR γ chain (His85Arg) (37), which has been suggested to interact with BTN2A1, could abrogate the binding of TCR multimers to tumor cells. Indeed, $\gamma_{\text{His85Arg}}\delta$ TCR tetramers and $\gamma_{\text{His85Arg}}\delta$ TCR-coated beads no longer bound to Daudi or to HEK293T cells (Figure 3, A–D). Next, we assessed the impact of CD277 KO and BTN2A1 KO on target cells in $\gamma\delta$ TCR tetramer binding. Although CD277 KO did not affect $\gamma\delta$ TCR tetramer binding, differences between $\gamma\delta$ TCR LM1 and Cl5 tetramers were still evident. KO of BTN2A1 completely abrogated $\gamma\delta$ TCR tetramer binding to both $\gamma\delta$ TCR LM1 and Cl5 tetramers, indicating that a high-affinity CDR3 TCR δ chain does not overcome the lack of binding through the TCR γ chain (Supplemental Figure 3A).

$\gamma\delta$ TCR target interaction does not depend on PAM, and CD277 binding does not require the $\gamma\delta$ TCR. $\gamma\delta$ TCRs have been reported to trigger T cell responses by sensing a joint spatial and conformational change in CD277 at the cell membrane of cancer cells (CD277) that is induced by a pAg-high state of the target cell (6, 8, 9, 11). Although we demonstrated that binding of tetramers to tumor cells heavily depended on BTN2A1, TCR binding could be further modulated through PAM-induced mobility changes of CD277. Therefore, we examined whether our $\gamma\delta$ TCR-coated beads fully mimicked the mode of action of a $\gamma\delta$ TCR in interacting with its target. First, we questioned whether the pAg level fine-tunes the activation state of the $\gamma\delta$ TCR cell by increasing TCR-ligand interaction affinity, and thus whether binding of the $\gamma\delta$ TCR-coated beads could be modulated by PAM treatment of

the target cells. Surprisingly, the addition of PAM did not change the intensity of the TCR tetramer or TCR-coated bead conjugation to either Daudi cells or HEK293T cells (Figure 3, A–D), indicating that while capturing unequal TCR affinities, TCR-coated beads do not sense the CD277 that induces $\gamma\delta$ TCR cell activation, and thus adding to the conflicting data on the possibility of a direct or indirect $\gamma\delta$ TCR-CD277 interaction (6–8, 13). Our observation would imply that, conversely, CD277 on a tumor cell does not need the $\gamma\delta$ TCR to bind to a T cell. Therefore, we next assessed whether the extracellular domains of CD277 interact with T cells in a $\gamma\delta$ TCR-independent manner. Tetramers of biotinylated CD277 ectodomains (CD277e) were incubated with Jurkat-76 cells, either nontransduced or transduced with $\gamma\delta$ TCR Cl5. In line with the assumption that CD277 is not the ligand for $\gamma\delta$ TCR, we observed strong CD277e tetramer and bead conjugation for both $\gamma\delta$ TCR⁺ and $\gamma\delta$ TCR-deficient Jurkat-76 cells, but not for Daudi cells (Figure 3, E and F). Binding of CD277e-coated beads could be specifically abrogated by anti-CD277 Abs and by the introduction of a mutation proposed by Willcox et al. (37), but not by anti- $\gamma\delta$ TCR Abs (Supplemental Figure 5B). Arg44Glu in CD277e abrogated the binding of this mutant to Jurkat-76 cells (Figure 3, E and F). These observations are in line with previous reports showing interaction of soluble CD277e constructs with T cell lines and activated primary T cells (38, 39).

In summary, our data show that (a) binding of $\gamma\delta$ TCR to BTN2A1 was dominated by the TCR γ chain and was further fine-tuned by the affinity of the CDR3 region of the TCR δ chain, (b) binding of the $\gamma\delta$ TCR to tumor cells was independent of CD277 and PAM, and (c) CD277 did not require $\gamma\delta$ TCR expression for interaction with T cells.

Conjugation enhancement depends on $\gamma\delta$ TCR affinity and PAM. Our data suggest that targeting a tumor cell with a $\gamma\delta$ TCR expressed on a T cell membrane depends at least on a 2-receptor, 3-ligand interaction rather than a single-receptor, single-ligand binding event. To gain more insight into the orchestration of BTN2A1 and BTN3A1 and the $\gamma\delta$ TCR, we assessed whether binding of the TCRs to tumor cells, as opposed to conjugation of the TCR beads, would change if we used a more dynamic model that would allow flexible membrane processes and assistance of additional cell membrane molecules. This was achieved by analyzing cell conjugation between $\gamma\delta$ TCR-expressing Jurkat/MA (JurMA) cells and HEK293FT cells as targets. Similar to $\gamma\delta$ TCR bead staining, cell-cell conjugate formation correlated with TCR affinity in the absence of PAM (Supplemental Figure 6), suggesting that TCR affinity governs conjugate formation between a T cell expressing the $\gamma\delta$ TCR and a tumor cell. However, whereas PAM treatment had no effect on $\gamma\delta$ TCR-coated bead binding, it increased the number of conjugates to the levels observed for the functional $\gamma\delta$ TCRs (Supplemental Figure 6), even in the case of the nonfunctional TCR LM1. Conjugation enhancement for a nonfunctional $\gamma\delta$ TCR suggested that PAM stabilized $\gamma\delta$ TCR-independent cell conjugates, which, in the case of a functional $\gamma\delta$ TCR, might lead to a reduction of the signaling threshold and increased $\gamma\delta$ TCR cell activation.

BTN2A1-BTN3A1 interaction is enhanced through a PAM-dependent increase in BTN3A1 surface recycling. BTN3A1 has been proposed to be regulated by RhoB through pAg accumulation (6).

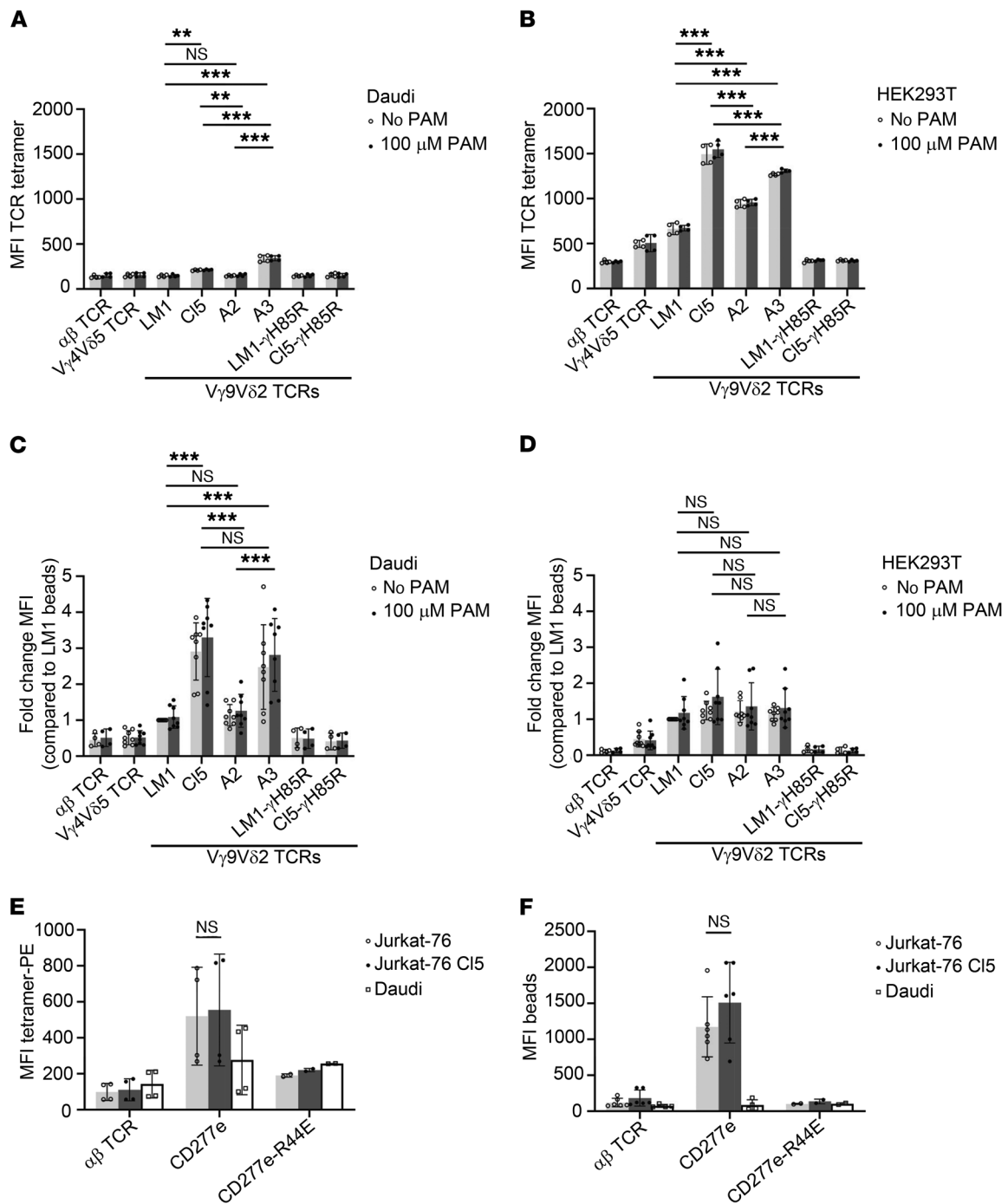


Figure 3. Staining of target and effector cells with $\gamma\delta$ TCR and CD277e multimers of increasing valency. (A and B) Results for staining of Daudi (A) and HEK293T (B) cells using fluorescent TCR tetramers (streptavidin-PE). Dots indicate the MFI values for individual FACS experiments ($n = 4$). (C and D) Results for staining of Daudi (C) and HEK293T (D) cells using fluorescent TCR beads (streptavidin-coated purple beads). Dots indicate the fold change in MFI values compared with LM1 (no PAM condition) in the same individual FACS experiment ($n = 4$, for $\alpha\beta$ TCR and $V\gamma 9V\delta 2$ -H85R-TCRs; $n = 8$ for all other TCRs). For the 100- μ M PAM condition, the cells were incubated for 1 to 2 hours with PAM before staining. (E) Results for staining of Jurkat-76 (untransduced or transduced with $V\gamma 9V\delta 2$ -TCR C15) and Daudi cells using fluorescent tetramers containing the $\alpha\beta$ TCR control, CD277e dimers (both $n = 4$), or the CD277e-R44E mutant ($n = 2$). Dots indicate the MFI values for individual FACS experiments. (F) Results for staining of Jurkat-76 and Daudi cells using fluorescent tetramers containing the $\alpha\beta$ TCR control, CD277e dimers ($n = 6$ for Jurkat-76, $n = 4$ for Daudi), or the CD277e-R44E mutant ($n = 2$ for all cells). Dots indicate the MFI values for individual FACS experiments. For the TCR tetramer and bead staining results shown in A–D, a 1-way ANOVA followed by Dunnett’s T3 multiple comparisons test was performed. A P value of less than 0.05 was considered significant. ** $P < 0.005$ and *** $P < 0.001$. For the CD277e tetramer and bead staining results shown in E and F, a 2 tailed, paired t test was performed. All statistical analysis was done using GraphPad Prism 8.3.0.

Table 2. TEG reactivity, soluble TCR tetramer and TCR bead binding, and RhoB relocalization per tumor

Tumor cell	TCR Cl5 TEG reactivity	TCR tetramer binding	TCR bead binding	RhoB relocalization
ML-1	-	-	-	-
AML 4 (primary blasts)	-	-	-	-
K562	+	+	+	+
HEK293T	+	+	+	-
Daudi	+	-	+	-
AML 1 (primary blasts)	+	-	-	+
AML 2 (primary blasts)	+	-	-	+
AML 7 (primary blasts)	+	-	-	+

TCR Cl5 TEG reactivity against a panel of tumor cell lines or patient-derived primary blasts was determined using INF- γ ELISA, tetramer and bead-binding assays were performed using TCR Cl5 as described in Figure 3, RhoB relocalization was determined by immunofluorescence staining for RhoB, and localization was analyzed using conformal microscopy as described previously (6).

However, RhoB relocalization alone did not fully explain the recognition of a tumor cell through a $\gamma 9\delta 2$ TCR (Table 2). To obtain additional insights into the regulation of BTN3A1 after pAg accumulation, we generated a stable cell line expressing FLAG-tagged BTN3A1 (Supplemental Figure 7), and to search for additional PAM-dependent interaction partners, we analyzed changes in the interactome of BTN3A1 with or without PAM treatment. Affinity purification coupled with mass spectrometry (AP-MS) from the cells expressing FLAG-BTN3A1 revealed a strong enrichment of BTN2A1 in the FLAG immunoprecipitates after PAM treatment (Figure 4A). This finding was confirmed by coimmunoprecipitating FLAG-BTN3A1 and HA-BTN2A1 (Figure 4C) and suggested that the accumulation of pAgs significantly enhanced the interaction or targeting to the same membrane domain of BTN3A1 and BTN2A1. By performing a gene ontology (GO) functional enrichment analysis of the proteins significantly enriched in the FLAG-BTN3A1 immunoprecipitates after PAM treatment, we found an enrichment of proteins targeted to cellular membranes, including the endoplasmic reticulum and the plasma membrane, and to biological processes associated with these compartments and vesicle trafficking (Figure 4B). This result is consistent with the reported subcellular localization of BTN3A1 (6) and suggests that the accumulation of intracellular pAgs enhances BTN3A1 targeting to cellular membranes. To test whether PAM enhances recruitment to the plasma membrane of CD277, we evaluated the CD277 surface expression on K562 cells treated with PAM for 5 hours after supplementing with anti-CD277-Alexa Fluor 647 Ab during the entire treatment. The PAM treatment resulted in a modest increase in CD277 surface levels compared with levels on untreated cells (Figure 4D). However, this result may have been influenced by the differential response of the different CD277 isoforms to the PAM treatment and by the inability to specifically evaluate the surface expression of endogenous BTN3A1. Nonetheless, this increase in CD277 surface levels was completely abolished once cells were treated with monensin, a known inhibitor of vesicle trafficking, suggesting that PAM influenced not necessarily the overall membrane expression but rather the recycling of surface BTN3A1. To further evaluate the contribution of the intracellular pool of CD277 to the $\gamma 9\delta 2$ TCR-dependent recognition, we evaluated the response of bulk $V\delta 2^+ \gamma \delta$ T cells to K562, preincubated with a control or anti-CD277-block-

ing Ab before or during PAM treatment. Although the anti-CD277-blocking Ab (clone 267-21f8.19) completely prevented target cell recognition when present throughout the PAM treatment, washing the Ab away before the treatment resulted in at least 50% recovery of target cell recognition by bulk $V\delta 2^+ \gamma \delta$ T cells (Figure 4E) and TEGs (Supplemental Figure 8). We confirmed this finding by preincubating target cells with inhibitors of vesicle trafficking such as brefeldin A or monensin, both of which were effective in preventing $\gamma 9\delta 2$ TCR Cl5 TEG-mediated recognition of the target cell (Figure 4F). These data suggest that the BTN2A1-BTN3A1 interaction or plasma membrane cotargeting is enhanced by PAM and is dependent on an increase in BTN3A1 surface recycling.

TCR affinity dictates the recruitment of a $\gamma 9\delta 2$ TCR toward the late IS. Next, we assessed whether formation of the late IS, a prerequisite for full T cell activation (40), depends on $\gamma 9\delta 2$ TCR affinity or PAM treatment. To address this question, we quantified accumulation of the $\gamma 9\delta 2$ TCRs to the contact area between tumor cells and T cells, relative to T cell membrane areas distant from the synapse (Figure 5, A and B). Synapse enrichment was significantly higher for the TCR CTE-Cl5 compared with the TCRs CTE-Cl3 and LM1 in the case of target cells that were not subjected to PAM treatment (Figure 5A). Even though T cells expressing a nonfunctional TCR LM1 showed an increase in conjugate formation upon PAM treatment (Supplemental Figure 6), this TCR did not enrich in the IS, even when tumor cells were treated with PAM (Figure 5A). PAM treatment did not further enhance polarization of the high-affinity CTE-Cl5 TCR to the IS and only significantly increased accumulation of the low-affinity TCR CTE-Cl3 in the synapse. This finding implies that TCR affinity dictates the recruitment of a $\gamma 9\delta 2$ TCR toward the late IS and that, in the case of a low-affinity TCR, PAM treatment can boost this recruitment by stabilizing the IS via CD277.

We investigated whether PAM might influence recruitment of a given $\gamma 9\delta 2$ TCR toward the IS through changes in CD277 clustering on the tumor cell. To explore this possibility, we treated HEK293 cells stably expressing the BTN3A1 isoform linked to EmGFP with either PAM or agonist and antagonist anti-CD277 Abs (clones 20.1 and 103.2, respectively) and analyzed the cells by confocal microscopy. The number and size of BNT3A1-EmGFP clusters in the cell membrane were similar in all conditions (Figure 5C), suggesting no

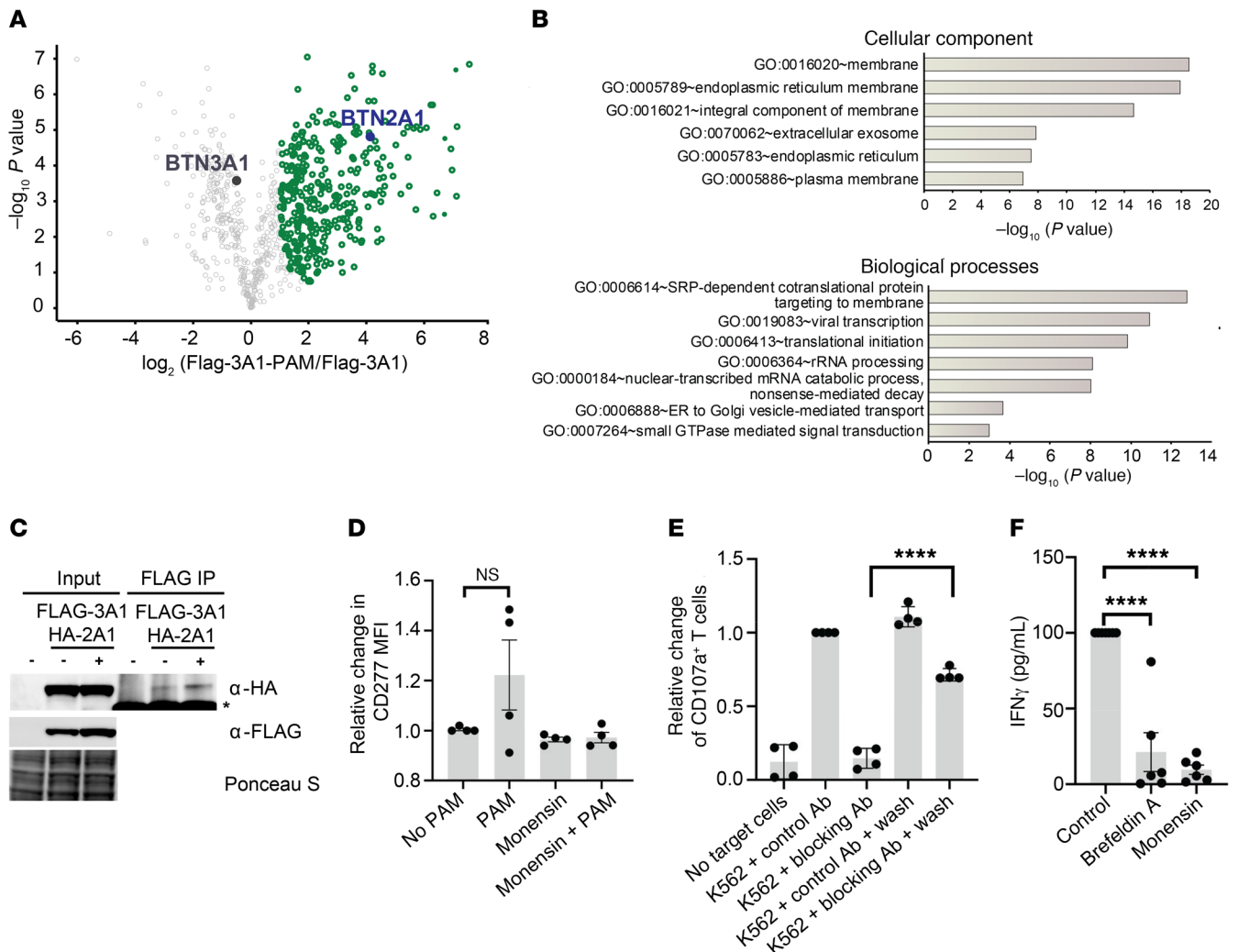


Figure 4. PAM treatment alters the interactome of BTN3A1 and the role of BTN3A1 turnover or the pAg-dependent V γ 9V δ 2 T cell activation. (A) Volcano plot shows the P values versus the difference in \log_2 protein intensities (proteins coenriched with BTN3A1 under the different treatment conditions). BTN2A1 coeluted with BTN3A1 after PAM treatment. The proteins significantly enriched after PAM treatment are highlighted in green, BTN2A1 in blue, and BTN3A1 in gray. (B) GO enrichment analysis of the proteins significantly coenriched with BTN3A1 after PAM treatment using the Database for Annotation, Visualization, and Integrated Discovery (DAVID). (C) Coimmunoprecipitation from HEK293T-CD277-KO cells transiently transfected with FLAG-BTN3A1 and HA-BTN2A1. PAM treatment enhanced the association between BTN3A1 and BTN2A1. The asterisk indicates the IgG heavy chain contaminant. Data are representative of 3 independent experiments. α , anti-FLAG. (D) Results for CD277 stainings of K562 cells cultured for 5 hours under different conditions in the presence of anti-CD277-AF647 Ab. Dots indicate the relative MFI values for individual FACS experiments. Data indicate mean values \pm SD. (E) Results for CD107a staining of V δ 2 T cells cocultured with K562 cells treated with 100 μ M PAM and preincubated for 45 minutes with a control Ab or a CD277-blocking Ab (clone 267-21f8.19) before starting the treatment or retained throughout the treatment. Dots indicate the relative changes in CD107a $^+$ T cells for individual FACS experiments. Data indicate mean values \pm SD. A P value of less than 0.05 was considered significant. **** P < 0.0001, by 2 tailed, unpaired t test. (F) IFN- γ release of TCR Cl5 TEGs cocultured with HEK293T cells pretreated with either monensin or brefeldin A and washed extensively before coculture. Data indicate mean values \pm SD. A P value of less than 0.05 was considered significant. **** P < 0.0001, by 1-way ANOVA, defining comparisons between the treatments and the control sample.

increase in CD277 cluster size induced by stimulation of the tumor cells with the activating Ab or PAM. However, the lateral resolution of the confocal microscope is limited to approximately 300 nm, and we may therefore have missed smaller clusters.

To examine CD277 organization at the nanoscale in the presence or absence of PAM stimulation, we performed direct stochastic optical reconstruction microscopy (dSTORM), a localization-based super-resolution imaging technique that provides approximately 20-nm resolution (41). CD277 molecules were labeled with AF647-CD277 Ab, and super-resolution images were

acquired at the basal membrane of HEK293T cells (Supplemental Figure 9). We found predominantly random CD277 distribution in both control and PAM-treated HEK293T cells, as Hopkins statistic values (H values) were centered around 0.5 (Figure 5D). By comparing the distribution of the localizations per cluster, we further confirmed that neither PAM treatment nor preincubation with agonist anti-CD277 20.1 Ab dramatically changed the spacing between CD277 molecules (Figure 5E and Supplemental Figure 9, A and B), whereas EGF treatment as a positive control resulted in enrichment of clusters of EGFR in the membrane of

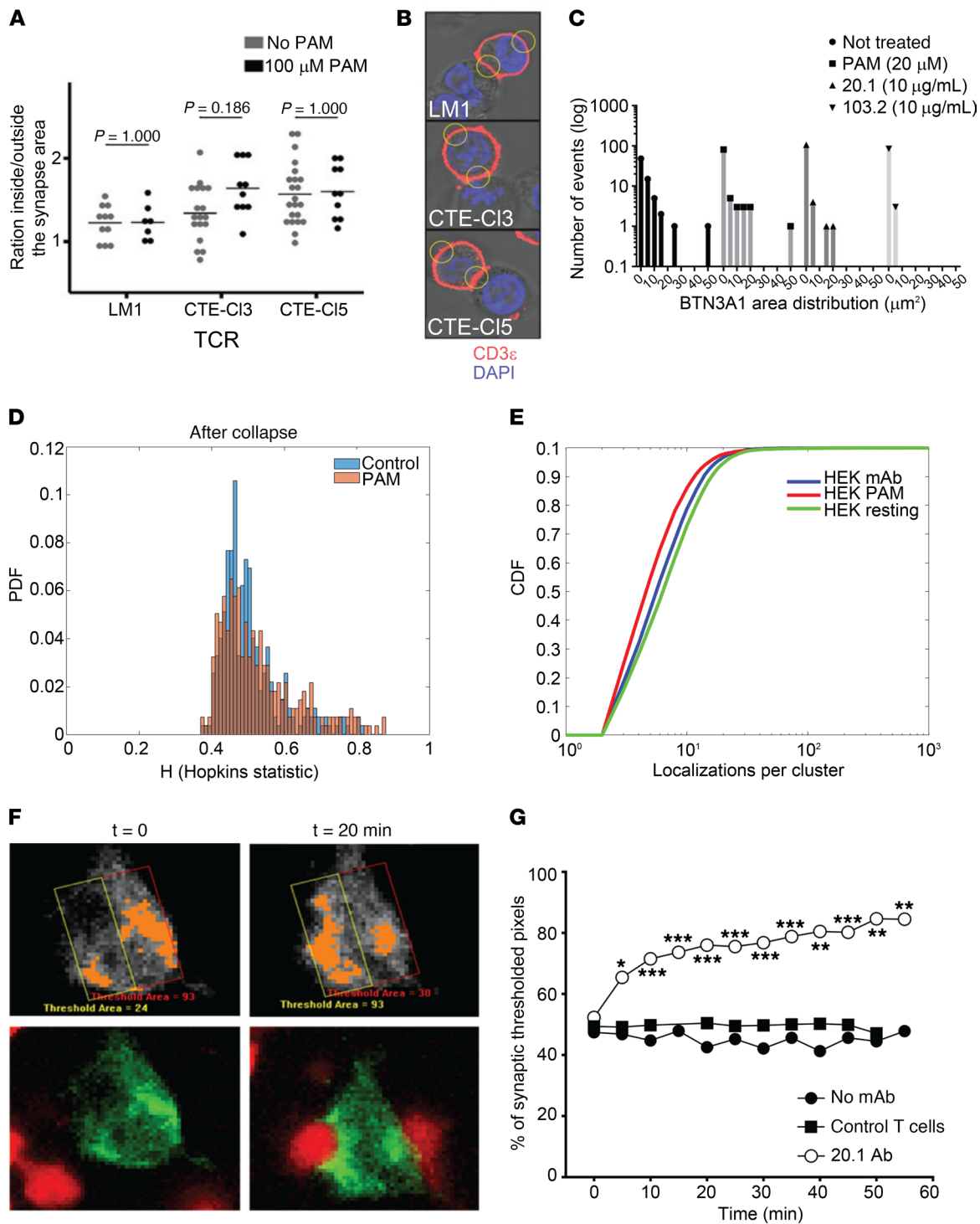


Figure 5. Target cell-surface distribution of BTN3A1 with and without $\gamma 9\delta 2$ T cell engagement. (A) Enrichment of the TCR in the IS was quantified as the ratio between CD3 ϵ -AF647 Ab signal intensity inside versus outside the synapse area. Analysis was done on at least 7 independent T cell–tumor cell conjugate images (representative images are shown in B). Significance was determined by 2-way ANOVA with Mann-Whitney *U* post hoc analysis. A *P* value of less than 0.05 was considered significant. Dots represent individual images, and bars represent the median values. (B) Representative images of the TEG–tumor cell conjugates and the regions of interest (ROIs) used for quantification. Original magnification, $\times 63$. (C) Topology of BTN3A1 molecules: the number of fluorescence events on the cell membrane (BTN3A1-EmGFP) was classified according to the cluster size based on confocal microscopic analysis of HEK293T cells stably expressing BTN3A1-EmGFP (*n* = 15). (D) Histogram comparing the Hopkins statistic (*H*) of BTN3A1 clustering for control and PAM-treated cells, imaged using dSTORM. A total of 275 ROIs across 10 cells were analyzed, and the probability distribution function (PDF) for *H* was plotted. (E) Cumulative distribution function (CDF) for the nearest neighbor distances (NNDs) for control-treated (HEK resting, green), PAM-treated (HEK PAM, red), and 20.1 Ab–treated (HEK mAb, blue) samples. The CDF plots presented are read as follows: the curve furthest to the right for high values of the CDF represent data that tend to have greater localizations per cluster compared with the curves on the left. (F) Example of BTN3A1-EmGFP polarization to the IS in the presence of 20.1 Ab. Threshold area boxes were used to calculate the percentage of synaptic threshold pixels at time point 0 (left) and 20 minutes after T cell conjugation (right, upper). The lower panel shows target cells (green) and T cells (red) at corresponding time points. Original magnification, $\times 20$. (G) Synaptic BTN3A1-GFP polarization measured on HEK293T cells following engagement with primary $\gamma 9\delta 2$ T cells with or without 20.1 Ab, NKT cells served as a control. *n* = 30 BTN3A1-GFP cells analyzed for each condition across 3 experiments. A *P* value of less than 0.041 was considered significant. **P* < 0.041, ***P* < 0.0041, and ****P* < 0.00041, by 2-way ANOVA with Bonferroni's post hoc correction for multiple comparisons.

HEK293T cells overexpressing EGFR (42) in the same experimental setting (Supplemental Figure 9C). These results are consistent with CD277 existing primarily as randomly distributed molecules in the plasma membrane of the isolated tumor cells in both pAg-low and pAg-high states. Importantly, when coincubating the same target cells with $\gamma 9\delta 2$ T cells, we detected a significant accumulation of BTN3A1-EmGFP within the IS in the presence of the activating 20.1 Ab. NKT cells targeting HEK293T cells but lacking a $\gamma 9\delta 2$ TCR did not induce BTN3A1-EmGFP accumulation at the IS. Accumulation of BTN3A1 molecules was significant 5 minutes after T cell conjugation (Figure 5, F and G).

Discussion

There has been a resurgence of interest in $\gamma 9\delta 2$ T cell therapies and medicinal products based on receptors extracted from $\gamma 9\delta 2$ T cells (1–3, 14). To overcome recent failures of clinical implementation, our data advocate for a substantial reduction of the high diversity usually observed in healthy $\gamma 9\delta 2$ T cell repertoires for experimental and, even more important, therapeutic purposes. In addition to functional diversity, we observed heterogeneity in $\gamma 9\delta 2$ TCR affinities, which, however, did not always correlate with the antitumor activity of the parental clone. In stark contrast to the current understanding of the mode of action, our data imply a 2-receptor, 3-ligand system, in which binding of the $\gamma 9\delta 2$ TCR to BTN2A1 is the major initiating event of the IS formation, and highlight the importance of the variable domain of the TCR γ chain between

the CDR2 and CDR3 regions for the first contact. Our data also emphasize the crucial role of the affinity mediated through the CDR3 region of the TCR δ chain, though the interaction of this domain with its binding partner has yet to be defined. In addition, CD277 was subsequently recruited to the IS for further stabilization in a pAg-dependent manner and ultimately ended up in close proximity to BTN2A1. This recruitment included the accumulation of CD277 from other membrane sites that were not in contact with the target, as well as the mobilization of intracellular CD277.

Both TCR CDR3 region diversity (15, 43) and variable expression of an array of other nonrearranged activating and inhibitory receptors (2, 30, 44) have been reported to contribute to functional heterogeneity within the $\gamma 9\delta 2$ T cell population. We provide formal evidence that this diversity also substantially affected the antitumor activity of $\gamma 9\delta 2$ T cells by highlighting the large variation in $\gamma 9\delta 2$ T cell antitumor reactivity on a clonal level, and further demonstrating unequal TCR affinities harbored within the diverse $\gamma 9\delta 2$ TCR repertoires. The reduced diversity within 1 donor and interindividual variations can most likely be attributed to a phenomenon called “repertoire focusing.” TCR repertoire focusing, or skewing toward a limited number of dominant clonotypes, has been described for both V $\delta 2^+$ and V $\delta 2^-$ subsets (22–24, 45). Major changes to the TCR $\delta 2$ repertoire happen postnatally, with a significant shift in dominant clonotypes according to their pAg reactivity (24), which is presumably antigen driven and strongly implies involvement of the $\gamma 9\delta 2$ TCR in constructing the functional landscape of $\gamma 9\delta 2$ T cells during ontogeny. Skewing of the repertoire toward a few dominant clonotypes implies a certain functional benefit in the conditions of selective pressure (46), however, although we observed highly focused repertoires, we did not find a correlation between clonotype frequency and antitumor functional potential of the respective T cell clone or its TCR in our experimental system. The impact of TCR affinity on maturation of the polyclonal $\gamma 9\delta 2$ T cell repertoire thus remains uncertain. The plausible explanation could be that clonotypic expansions, as a reflection of the microbial stimulation history of an individual after birth (24), either are not TCR driven or do not necessarily represent the functional hierarchy of the clonotypes when measuring in vitro tumor-induced activation.

Though the observed functional heterogeneity mediated through different $\gamma 9\delta 2$ TCRs emphasizes the possibly adaptive character of the $\gamma 9\delta 2$ T cell response, the fact that we did not always observe a direct correlation between functional avidity of the TCR and reactivity of its parental clone implies a significant “shaping” of individual $\gamma 9\delta 2$ T cell clones by other surface or cytoplasmic factors, which might be induced by microbial challenge, but may not be further involved in antitumor responses. Repertoire shifts (47) favoring expansion of potentially poorly or nontumor-reactive clones might explain a more than 15-year history of failure when using polyclonal and phenotypically diverse $\gamma 9\delta 2$ T cells to treat cancer (4), arguing for more precisely targeted applications to improve clinical outcomes when using $\gamma \delta$ T cells (14, 31, 44, 48–50).

Regardless of the role of the affinity of an individual $\gamma 9\delta 2$ TCR within the environment of its parental $\gamma \delta$ T cell, the TCR-mediated function is intriguing in the context of next-generation TCR and CAR-T therapies (3), since exploiting a $\gamma 9\delta 2$ TCR outside of its parental clone overcomes potentially unfavorable clonotypic

or phenotypic features (6, 15, 18, 51, 52). In the current study, we demonstrate intrinsic differences in $\gamma\delta$ TCR-mediated reactive potential that dictate the antitumor response of T cells engineered to express a defined $\gamma\delta$ TCR (TEGs). None of the newly isolated $\gamma\delta$ TCRs, however, exceeded the functional avidity mediated by the previously described model TCR clone 5 (15) in the TEG format, suggesting that a natural affinity ceiling is also relevant for the $\gamma\delta$ TCR, as reported for $\alpha\beta$ TCRs and Abs (53). The possible molecular determinants of $\gamma\delta$ TCR-mediated reactivity have been the focus of extensive studies, and “position 5” of the CDR3 δ (ImMunoGeneTics [IMGT] position 109), which is known to have highly conserved hydrophobic amino acids, was shown to determine the $\gamma\delta$ TCR-mediated response (54). In our data set, low or absent $\gamma\delta$ TCR-mediated reactivity was associated with amino acids A, G, R, or D at this position (TCRs A2 and C2, C7/C11, B2/B5, A7; see Table 1) rather than with the highly prevalent amino acids L, V, or W. In our study, the amino acid proline (TCR B1), reported earlier to confer nonreactivity to pAgs (54), mediated reactivity comparably to CDR3 δ sequences featuring L/V/W at this position.

When entering the clinic with medicinal products based on $\gamma\delta$ TCRs, it is, of course, of utmost importance to select for patients with a high likelihood of response and to thus be able to characterize the expression of the $\gamma\delta$ TCR target. However, the poor description of the mode of action of $\gamma\delta$ TCR-expressing T cells makes it challenging to develop tools that allow for robust detection of the target. To address this challenge, we developed $\gamma\delta$ TCR tetramers and $\gamma\delta$ TCR-coated beads as tools. Surprisingly, $\gamma\delta$ TCR-coated bead conjugation was independent of pAgs and did not fully mimic the currently understood mode of action, a finding that may fuel the ongoing debate as to whether CD277 directly interacts with the $\gamma\delta$ TCR (6–8, 13, 55). By using CD277-KO tumor cells and our staining tools, we demonstrate that binding of the $\gamma\delta$ TCR to its target can occur in the absence of CD277. In line with most recent reports (35, 36), our data indicate the crucial role of BTN2A1 as the first contact. However, we extended these findings by demonstrating that not only the variable framework region of the TCR γ chain but also the CDR3 region of the TCR δ chain is important in modulating the first contact between the $\gamma\delta$ TCR and its target. In addition, we demonstrate that this initiating event can substantially differ between different tumor targets, as implied by our observation that excellent tumor targets of a $\gamma\delta$ TCR like Daudi, which were barely stained with $\gamma\delta$ TCR tetramers, could bind to $\gamma\delta$ TCRs when coupled to beads. We also identified tumor cells such as primary AML blasts, which could not be stained at all with these tools, highlighting the question of whether such static tools can truly mimic the $\gamma\delta$ TCR-ligand interaction. When using $\gamma\delta$ TCR multimers alone, it is challenging to identify patients suitable for $\gamma\delta$ TCR-based immune therapies.

We therefore focused on the second key partner, CD277, and show that CD277e binding to a T cell did not depend on the $\gamma\delta$ TCR. Our data support a 2-receptor, 3-ligand model, in which the $\gamma\delta$ TCR mediates the first contact with BTN2A1 and a second ligand, followed by a third step involving CD277, which serves as an essential coactivating ligand and most likely interacts with another receptor to stabilize cell-cell vicinity and consequently reduce the $\gamma\delta$ TCR-mediated activation threshold. Unequal

$\gamma\delta$ TCR affinities further modulate the magnitude of the IS formation and T cell activation. The interaction partner of the CDR3 region of the TCR δ chain has yet to be defined (Figure 6).

We also elucidated the important role of cellular trafficking of CD277. After contact with the T cell, CD277 is recruited from the noninteracting cell membrane parts to the IS and relies on intracellular CD277, which then associates with more membrane-bound molecules, indicating a higher membrane recycling of CD277. In line with our previous observation, this second step is most likely coordinated by RhoB (6). The ability of a target cell to use the intracellular storage pools most likely depends on that cell's ability to activate RhoB-dependent pathways, so that RhoB is then located at the end of the process, next to the membrane, and induces changes in membrane mobility and conformation for CD277 (6). Juxtamembrane enrichment of RhoB could serve as part of an algorithm to assess the susceptibility of a tumor cell to $\gamma\delta$ TCR-based immune therapies (6). However, RhoB does not fully describe the target, as the most prominent $\gamma\delta$ TCR target, the Daudi cell, does not show the typical juxtamembrane enrichment of RhoB (6), implying that this mechanism is only part of the recognition process.

The final question is how to characterize the previously reported pAg-induced joint changes in membrane mobility and conformation of CD277 (6, 7, 9) and their role in $\gamma\delta$ TCR-induced T cell activation in the context of these new insights. In our experimental data set, PAM treatment did not induce changes in CD277 clustering when the surface distribution of CD277 was studied on HEK293T cells alone. However, we found increased membrane expression and active accumulation of the BTN3A1-EmGFP molecules in the contact area with the T cell, suggesting that prior formation of the early synapse through the $\gamma\delta$ TCR and its ligand was required. A PAM-induced decrease in membrane mobility of CD277 (6, 9) could facilitate the formation of heterogeneous clusters, including CD277 and the $\gamma\delta$ TCR ligand, allowing both molecules to interact with their cognate receptors on T cells within the IS. Candidate binding partners for CD277, one of which is the lymphotoxin β receptor, are currently being studied (56). Another intriguing and still-open question is whether an alternative model of dual $\gamma\delta$ TCR-mediated reactivity, as described recently for mouse V γ 7* and human V γ 4* TCRs (57), could explain the apparent “uncoupling” phenomena observed in this study. In this case, existing data on the expression of the CD277 receptor on cells of myeloid lineage and B cell lines (38) would argue for a multitude of potential binding partners for CD277. Therefore, to increase the likelihood of a positive signal in clinical studies, positive staining of the tumor with $\gamma\delta$ TCR tetramers or beads as well as identification of the characteristic distribution of RhoB (6) predictive of CD277J should assist with patient selection, as long as the crucial receptor-ligand interface has not been unambiguously identified.

In summary, we demonstrate a large variation in antitumor reactivity of primary $\gamma\delta$ TCR cell clones. Activity against cancer cells was not associated with clonal frequency of $\gamma\delta$ TCR cell clones within the repertoire, which resulted in only a small fraction of $\gamma\delta$ TCR cell clones that were strongly reactive against cancer cells. In addition, we propose a paradigm shift in the mode of action. Our data imply a 2-receptor, 3-ligand activation model for the description of the target-receptor interface, in which the interaction of the $\gamma\delta$ TCR with BTN2A1 is mediated by the TCR γ chain,

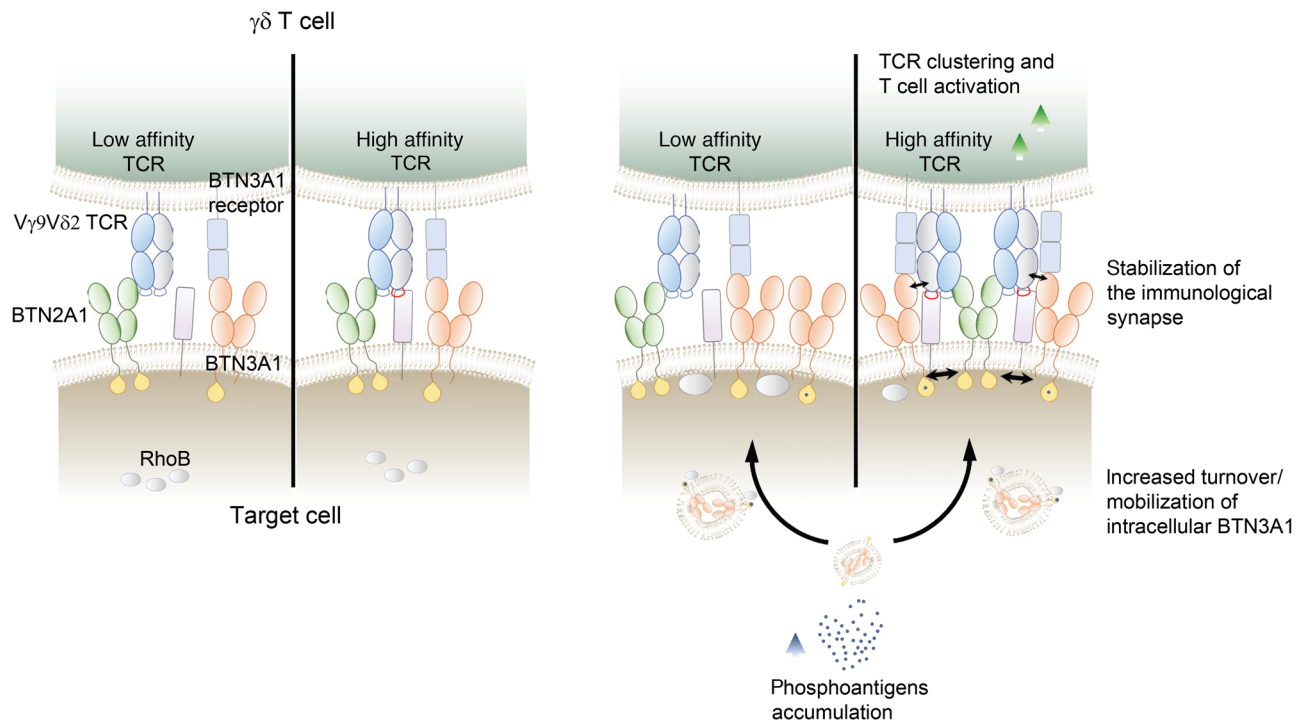


Figure 6. A 2-receptor, 3-ligand model for tumor cell recognition by $\gamma\delta 2$ TCR-expressing cells. While the region between CDR2 and CDR3 of the TCR γ chain (blue) directly binds to BTN2A1 on tumor cells as a major initiating event, the CDR3 region of the TCR δ chain most likely interacts with an additional, currently unknown ligand (purple) and modulates the strength of the interaction with the tumor cell. The interaction with the second ligand through the TCR δ chain critically depends on the affinity mediated via its CDR3 region. This first process is pAg independent (left). Full activation of $\gamma\delta 2$ TCR $^+$ T cells is only achieved, however, when BTN3A1 is recruited in a pAg-dependent manner through RhoB to the IS for further stabilization, where BTN3A1 comes into close proximity to BTN2A1. This recruitment of BTN3A1 includes the accumulation of BTN3A1 from other membrane sites that are not in contact with the target, as well as the mobilization of intracellular BTN3A1 (right). The ligand of BTN3A1 has not yet been identified.

as recently reported (35, 36). Subsequently, the CDR3 region of the TCR δ chain binds a second binding partner in a pAg-independent manner. However, the true interaction partner of the CDR3 region of the TCR δ chain has yet to be defined. The importance of this interaction was outlined by our observations that T cell conjugation, IS formation, and full T cell activation could be modulated by different affinities of the CDR3 region of the TCR δ chain. As a third event, CD277 was recruited to the IS and fixed via a RhoB-mediated mechanism. However, it is unlikely that CD277 interaction relied on the $\gamma\delta 2$ TCR and was mediating the pAg-dependent component via inside-out signaling, which had as a final event not only fixation of CD277 but also its close contact with BTN2A1, thereby providing the second essential coactivating signal. We believe our data pave the way for pursuit of the next ligand and have important implications for ongoing experimental and clinical efforts when exploring the use of $\gamma\delta 2$ T cells and their receptors for cancer immunotherapies.

Methods

Peripheral blood from anonymous healthy donors was purchased from the Dutch blood bank. $\gamma\delta 2$ T cells were single-cell sorted, expanded using a rapid expansion protocol (16), and tested functionally by overnight incubation with the tumor cell lines. Cytokine levels were subsequently measured in the supernatant using ELISA and/or Luminex assay. The sequences of the $\gamma\delta 2$ TCRs were determined by Sanger sequencing, ligated into retroviral vectors, transduced into

Jurkat-76 or human primary $\alpha\beta$ T cells, and, like the primary $\gamma\delta 2$ T cell clones, functionally tested.

The sequences of the extracellular domains of the model $\gamma\delta 2$ -TCRs (15, 28) and CD277 were ligated into pBullet vectors, expressed as soluble biotinylated proteins in Freestyle 293-F cells, and coupled with streptavidin-phycoerythrin (streptavidin-PE) to generate sTCR-CD277e tetramers and dextramers (58), or with streptavidin-conjugated microspheres. The fluorescent multimers were incubated with the target cells, and conjugation was measured by FACS. More detailed information on the methods, including HTS of the TCR δ chain, cell-cell conjugation assays, confocal and super-resolution imaging, videomicroscopy, and statistical analyses, can be found in the Supplemental Methods (59–66).

HTS data. The raw HTS data were deposited in the NCBI's Sequence Read Archive (SRA) database (SRA PRJNA631458).

Resources and reagents. Requests for resources and reagents should be directed to the corresponding author.

Study approval. AML blood samples were collected from the biobank of the UMC Utrecht in accordance with good clinical practices and Declaration of Helsinki principles. All patients provided consent prior to storage of their blood samples in the biobank (TCBio 16-088). This study was approved by TCBio 16-088, UMC Utrecht (Utrecht, Netherlands).

Author contributions

AV, DXB, ES, ZS, and JK designed the study. AV, DF, DXB, EVD, LK, FK, JS, ADH, LB, SN, TS, and ZS performed the experiments. AJ

and AV analyzed the HTS data. KAL, MJW, SLS, and DSL provided methodology and analyzed the super-resolution microscopy data. AJRH, DO, and JL provided reagents and intellectual input. ZS and JK supervised the project. AV, DXB, DSL, ZS, and JK wrote the manuscript. All authors provided critical review and discussions.

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Address correspondence to: Jürgen Kuball, Laboratory of Translational Immunology, University Medical Center Utrecht, Utrecht University, 3584 CX Utrecht, Netherlands. Phone: 31.88.755.7230; Email: j.h.e.kuball@umcutrecht.nl.

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