RhoB Mediates Phosphoantigen Recognition by Vγ9Vδ2 T Cell Receptor

Highlights
- Identification of SNPs near RhoB is associated with poor Vγ9Vδ2 T cell activation
- RhoB activity and distribution in tumor cells modulate Vγ9Vδ2 T cell activation
- Relocalization of RhoB induces membrane immobility of BTN3A1
- Tumor recognition by a Vγ9Vδ2 TCR depends on BTN3A1 conformation

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In Brief
Sebestyen et al. show that Vγ9Vδ2 TCR activation is modulated by the GTPase activity of RhoB in tumor cells, and by the relocalization of RhoB to BTN3A1. Subsequently, a phosphoantigen-induced conformational change in BTN3A1 leads to its recognition by Vγ9Vδ2 TCRs.
RhoB Mediates Phosphoantigen Recognition by Vγ9Vδ2 T Cell Receptor

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http://dx.doi.org/10.1016/j.celrep.2016.04.081

SUMMARY

Human Vγ9Vδ2 T cells respond to tumor cells by sensing elevated levels of phosphorylated intermediates of the dysregulated mevalonate pathway, which is translated into activating signals by the ubiquitously expressed butyrophilin A1 (BTN3A1) through yet unknown mechanisms. Here, we developed an unbiased, genome-wide screening method that identified RhoB as a critical mediator of Vγ9Vδ2 TCR activation in tumor cells. Our results show that Vγ9Vδ2 TCR activation is modulated by the GTPase activity of RhoB and its redistribution to BTN3A1. This is associated with cytoskeletal changes that directly stabilize BTN3A1 in the membrane, and the subsequent dissociation of RhoB from BTN3A1. Furthermore, phosphoantigen accumulation induces a conformational change in BTN3A1, rendering its extracellular domains recognizable by Vγ9Vδ2 TCRs. These complementary events provide further evidence for inside-out signaling as an essential step in the recognition of tumor cells by a Vγ9Vδ2 TCR.

INTRODUCTION

γδT cells are unconventional T cells with strong reactivity toward a broad spectrum of tumors of diverse tissue origin. γδT cells combine potent anti-tumor effector functions with the recognition of broadly expressed tumor-associated molecules, and these features have put γδT cells in the spotlight for clinical application in cancer immunotherapy. Activation of γδT cells involves the sensing of metabolic changes in cancer cells that result in the expression of generic stress molecules. These molecules are upregulated upon transformation or distress (Bonneville et al., 2010; Vantourout and Hayday, 2013). However, progress in the clinical application of γδT cells for cancer treatment is hampered by conflicting published data from various labs that describe contradicting molecular requirements for γδT cell activation (Scheper et al., 2014; Vavassori et al., 2013; Sandstrom et al., 2014) as well as by a lack of prognostic markers to assess which patients may benefit from such therapy.

Vγ9Vδ2 T cells, the major γδT cell subset in human peripheral blood, express γδT cell receptors (TCR) composed of Vγ9 and Vδ2 chains and are specifically activated by intermediates of the mammalian mevalonate pathway (Sober et al., 2003; Constant et al., 1994), such as isopentenyl pyrophosphate (IPP), or by the microbial 2-C-methyl-D-erythritol 4-phosphate (MEP) pathway (Morita et al., 2007). Intracellular phosphoantigen (pAg) levels accumulate in tumor cells due to dysregulation of the mevalonate pathway or upon microbial infection, allowing the targeting of transformed or infected cells by Vγ9Vδ2 T cells. Similarly, intracellular phosphoantigen levels can be pharmacologically increased by treating cells with mevalonate pathway inhibitors such as aminobisphosphonates (ABPs), thereby sensitizing cells toward recognition by Vγ9Vδ2 T cells. Although the involvement of the Vγ9Vδ2 TCR in detecting elevated phosphoantigen levels was demonstrated as early as the 1990s (Bukowski et al., 1998; Davodeau et al., 1993; Wang et al., 2010), the molecular determinants required for activation of Vγ9Vδ2 TCRs on target cells have long remained elusive.

Although recent breakthrough studies have identified the membrane-expressed butyrophilin BTN3A1 (CD277) as a key molecule in phosphoantigen-induced activation of Vγ9Vδ2 T cells, (Harly et al., 2012; Sandstrom et al., 2014; Vavassori et al., 2013), they describe conflicting molecular roles for BTN3A1 in the Vγ9Vδ2 T cell activation process. While Vavassori et al. (2013) suggest that BTN3A1 as antigen presenting molecule triggers Vγ9Vδ2 T cell activation, data from Sandstrom et al. (2014) and Harly et al. (2012) support an inside-out signaling mechanism for BTN3A1, where phosphonatigens bind to its intracellular region and in which model immobilization of BTN3A1 at the cell surface may contribute to an extracellular...
cue for recognition by V_{γ9}V_{δ2} TCRs. Since BTN3A1 is expressed on both transformed and healthy human cells (Compte et al., 2004), additional molecules are likely involved in mediating selective recognition of tumor targets by V_{γ9}V_{δ2} T cells, including: e.g., the cytoskeletal adaptor protein Periplakin (Rhodes et al., 2015). However, it is unclear how these different proposed molecules and mechanisms are linked. Here, we developed an unbiased, genome-wide screening method, SNP-Associated Computational Pathway Hunt Including short hairpin RNA (shRNA) Evaluation (SAPPHIRE), and identified important molecular requirements for V_{γ9}V_{δ2} T cell-mediated recognition of cancer (stem) cells. We propose a two-component mechanism occurring in a tumor cell, which allows V_{γ9}V_{δ2} T cells to specifically sense cellular transformation.

RESULTS

Identification of Genetic Loci Associated with V_{γ9}V_{δ2} TCR-Mediated Target Cell Recognition by SNP-Associated Computational Pathway Hunt Including shRNA Evaluation

We hypothesized that differences in the genetic backgrounds of tumor cells affect recognition of these cells by V_{γ9}V_{δ2} TCR-engineered T cells. Therefore, we utilized the library of cell lines from the Centre d’Étude du Polymorphisme Humain (CEPH), which contains a large collection of EBV-transformed B cell lines (EBV-LCLs) obtained from several family pedigrees (Dausslet et al., 1990) and genotyped for millions of SNPs (International HapMap Consortium, 2003). CD4^{+} αβ T cells engineered to express one defined V_{γ9}V_{δ2} TCR were utilized for the functional screening in order to eliminate fluctuations in recognition by a diverse γδ TCR repertoire (Gründler et al., 2012) and varying expression of natural killer (NK) receptors (Scheper et al., 2013; Correia et al., 2013). The recognition phenotypes of EBV-LCLs by V_{γ9}V_{δ2} TCR T cells were assessed by means of interferon (IFN) γ production and showed an activating phenotype for 33 EBV-LCLs, while 7 were non-activating (Figures S1A and S1B). Importantly, in the zygosity analysis, EBV-LCLs with a non-activating phenotype represent more power than those with an activating phenotype analyzed with SAPPHIRE. Hypothetical zygoses for candidate genetic loci were deduced using classical Mendelian inheritance patterns within CEPH family trios, where the influence of candidate alleles on V_{γ9}V_{δ2} TCR-mediated recognition was assumed to be dominant. The resulting recognition phenotypes combined with family pedigrees of the CEPH cell lines overcame the need to screen large numbers of LCL lines (Spaapen et al., 2008), and allowed the precise prediction of zygoses of candidate loci for 12 CEPH individuals (Figure 1A). The hypothetical loci zygoses were then correlated with available genotype information of SNPs within the study’s population (Spaapen et al., 2008), resulting in the identification of 17 SNPs whose genotypes correlated perfectly (100%) with predicted zygoses (Figure 1B). Since none of these 17 SNPs, nor their proxy SNPs within high linkage disequilibrium (r^2 > 0.8), directly affected genes by causing changes in protein coding sequences, we speculated that rather than playing direct roles, the SNPs we identified could represent surrogate markers for genetic regions associated with susceptibility to V_{γ9}V_{δ2} TCR^{+} T cell recognition. In addition, we queried the genomic vicinity of the 17 SNPs for neighboring candidate genes (Figure 1C).

To test the relevance of these genes for V_{γ9}V_{δ2} TCR^{+} T cell-mediated target recognition, we knocked down all 17 SNP-adjacent genes in the V_{γ9}V_{δ2} TCR^{+} T cell-activating EBV-LCL line 48. We then assessed the effect of knockdown on activation of V_{γ9}V_{δ2} TCR^{+} T cells by measuring IFN-γ production, which was reduced upon the knockdown of three genes (RAB4A, RHOB and UBE3C) (Figure 1B). To ensure that potential knockdown effects pointed to genes that selectively affect V_{γ9}V_{δ2} TCR-dependent activation, the three knockdown variants of EBV-LCL line 48 were pulsed with Wilms’ tumor 1 (WT1) peptide and tested for recognition with T cells engineered to express the cognate WT1-specific αβ TCR (Kuball et al., 2007). The selective knockdown of the small GTPase RhoB significantly affected the activation of V_{γ9}V_{δ2} TCR- but not αβ TCR-engineered T cells (Figure 1B). Similar data were observed after partial knockdown of RhoB in the prototypic V_{γ9}V_{δ2} T cell target cell line Daudi (Figure S2B), as well as after CRISPR/Cas-mediated partial RhoB knockout in the renal carcinoma cell line MZ1851RC (Figure 2A). Interestingly, even complete knockout of RhoB in 293 HEK cells resulted only in partial depletion of target cell-mediated V_{γ9}V_{δ2} TCR T cell activation (Figure S2A). In addition, knockout of either RhoA or RhoC genes in 293 HEK cells did not significantly influence their ability to activate V_{γ9}V_{δ2} TCR^{+} T cells (Figures 2B and S2C), emphasizing that RhoB modulates the recognition of tumor cells by a defined V_{γ9}V_{δ2} TCR in a non-redundant role.

Target Cell Recognition by V_{γ9}V_{δ2} TCRs Depends on Rho GTPase Activity

Genomic sequencing of RhoB and associated promoter regions did not identify genetic variations associated with our marker SNP (data not shown), suggesting that RhoB is indirectly regulated by other gene loci influencing expression or activity (Pombo and Dillon, 2015). Our primary observations, however, confirmed that neither RhoB RNA nor protein levels in various recognized and two non-recognized tumor cell lines correlated with their ability to activate V_{γ9}V_{δ2} TCR^{+} T cells (Figures 2C, S2D, and S2E). Therefore, we assessed the impact of RhoB GTPase activity by pretreating different tumor cell lines with either the Rho GTPase activator calpeptin or inhibitor C3 transferase (Aktories et al., 1992; Schoenwaelder and Burridge, 1999). Pretreatment with calpeptin significantly sensitized EBV-LCL 93 cells for recognition by V_{γ9}V_{δ2} TCR^{+} T cells while, conversely, inhibition of Rho GTPase with C3 transferase resulted in significantly reduced activation of V_{γ9}V_{δ2} TCR^{+} T cells by LCL 48 cells (Figure 2D). Importantly, modulation of Rho GTPase activity did not affect the recognition of WT1 peptide-pulsed EBV-LCL 48 cells by WT1 αβ TCR-transduced T cells. To specify that enzymatic activity of RhoB regulates tumor cell recognition by V_{γ9}V_{δ2} TCR^{+} T cells, we transfected HEK293 cells with wild-type or dominant-negative form corresponding to the GDP-bound state (RhoB-ΔN), or with the constitutively active form corresponding to the GTP-bound form (RhoB-CA) of RhoB (Kamon et al., 2006) and used them as target cells for V_{γ9}V_{δ2} TCR^{+} T cells (Figure 2E). HEK cells overexpressing RhoB-CA mutants were able to trigger significantly stronger V_{γ9}V_{δ2} TCR-specific responses than cells expressing wild-type
RhoB, while RhoB-DN-transfected HEK cells showed a significantly reduced ability to stimulate Vγ9Vδ2 TCR+ T cells compared to cells expressing wild-type RhoB. Thus, these data suggest that the biochemical activity of RhoB GTPases is a critical factor modulating the recognition of cancer cells by Vγ9Vδ2 TCR+ T cells.

The Intracellular Distribution of RhoB in Cancer Stem Cells Marks Susceptibility for Vγ9Vδ2 TCR-Mediated Recognition

RhoB proteins have been shown to translocate within cellular compartments upon activation (Kranenburg et al., 1997). Therefore, we hypothesized an impact on subcellular localization of RhoB in recognized versus non-recognized cells. Additionally, ABP treatment of cells have been demonstrated to increase the GTP-bound forms of Rho GTPases (Dunford et al., 2006) that could provide a possible link between dysregulation of mevalonate pathway in tumor cells and RhoB cellular compartmentalization. This hypothesis was supported by confocal imaging, which revealed that distribution of RhoB patches that were primarily reported to locate to membranes of intracellular vesicles such as endosomes (Adamson et al., 1992; Michaelson et al., 2001), segregated strongly with the susceptibility of different target tumor cell lines to Vγ9Vδ2 TCR-mediated recognition. RhoB was selectively excluded from nuclear areas in cells that are able to activate Vγ9Vδ2 TCR+ T cells (Figures 3A, 3B, S3A, S3B).

Figure 1. SNP-Associated Computational Pathway Hunt Including shRNA Evaluation to Identify Genetic Loci Associated with Vγ9Vδ2 TCR Stimulation

(A) The recognition of CEPH EBV-LCL lines by Vγ9Vδ2 TCR+ T cells provided the basis for deducing hypothetical zygosities of candidate loci in each cell line (black, recognized; white, not recognized; square, male; circle, female; +/-, heterozygous; -/-, homozygous negative; +/-?, undetermined). Members of two CEPH families are shown as examples. For CEPH ID numbers of cell lines, see Figure S1A.

(B) Genetic association analysis revealed 17 SNPs of which genotypes correlated 100% (r² = 1) with predicted zygosities of cell lines. Locations and nearest neighboring genes of SNPs are indicated. The effect of knocking down candidate genes on recognition of EBV-LCL 48 by T cells transduced with either Vγ9Vδ2 TCR clone G115 or an HLA-A*0201-restricted WT1 126–134-specific abTCR are indicated by black circles (significant effect on T cell activation) and white circles (no effect). For testing recognition by WT1 abTCR+ T cells, the EBV-LCL 48 line was pulsed with WT1 126–134 peptide.

(C) Associating SNPs resulting from association analysis with candidate genes. The genetic region of the SNPs neighboring RhoB is shown as an example. Each bar represents one SNP, and r² values represent correlation between predicted zygosities and SNP genotypes.
Figure 2. RhoB Activity Correlates with Target Cell Recognition by Vγ9Vδ2 TCR+ T Cells

(A) RhoB was knocked out in the renal cancer cell line MZ1851RC using the CRISPR/Cas system. MZ1851RC cells were treated with either pamidronate or HLA-A*0201-restricted WT1126–134 peptide, and the effect on target cell recognition by Vγ9Vδ2 TCR+ and WT1 αβTCR+ T cells, respectively, was determined by measuring IFN-γ production. A guide RNA targeting an irrelevant sequence was used as control for knockout, while medium or irrelevant peptide loaded tumor cells were used as controls for T cell stimulation. Data show mean ± SEM of three independent experiments, in duplicate samples. Level of knockout was determined by intracellular flow cytometry.

(B) The effect of knockout of RhoA, B, and C in 293 HEK cells on recognition by Vγ9Vδ2 TCR+ T cells was assessed by measuring IFN-γ production. A guide RNA targeting an irrelevant sequence was used as control. Figure shows IFN-γ production normalized to irrelevant knockout samples of three independent experiments, in duplicate samples.

(C) RNA expression of RhoB was measured by qPCR in either non-recognized (black bars) or recognized (white bars) EBV-LCLs and tumor cell lines. Data are representative of two repeated experiments.

(D) The non-recognized EBV-LCL line 93 or the recognized EBV-LCL 48 were pretreated with either calpeptin or C3 transferase in combination with pamidronate or IPP, and the effect on stimulation of Vγ9Vδ2 TCR+ T cells was assessed by measuring IFN-γ. The effect of Rho-modulating compounds on recognition of WT1126–134 peptide-pulsed EBV-LCL 48 cells by WT1 αβTCR+ T cells was measured in parallel. Data show mean ± SEM of at least three independent experiments.

(E) HEK293 cells were transfected with dominant-negative (RhoB-DN), constitutively active (RhoB-CA), or wild-type RhoB (RhoB-WT), and the effect of activity variants on target cell recognition by Vγ9Vδ2 TCR+ T cells in the presence of pamidronate was determined by measuring IFN-γ. Figure shows IFN-γ production normalized to wild-type RhoB samples of three independent experiments.

The significance of data has been analyzed by Mann-Whitney test in (A), (B), and (D) and by Kruskal-Wallis test and Dunn’s multiple comparison test in (E).
and S3E). Crucially, re-localization of RhoB from the nucleus, or from the nuclear membrane to extranuclear sites was induced by ABP as well as by soluble phosphoantigen IPP in cell lines (Figures 3C and 3D), emphasizing that this process is dependent on accumulation of intracellular phosphoantigen. Expressing transgenic RhoB-GFP in either recognized or non-recognized tumor cell lines followed intracellular distribution of the endogenous RhoB (Figure S3B), suggesting post-translational regulation of RhoB. The homogenous intracellular distribution of other GTPases such as RhoA did not change upon ABP treatment (Figure S3C), supporting the observation that RhoA knock down did not impact V$_y$9Vl2 TCR-mediated recognition (Figure 2B). We found that RhoB was excluded from the nucleus in human, but not in mouse dendritic cells selectively treated with ABP, even though RhoB protein sequences are identical in both species (Figures 3E and S3D). This is consistent with our previous observations that human, but not mouse monocyte-derived dendritic cells are able to activate V$_y$9Vl2 TCR$^+$ T cells (Marcu-Malina et al., 2011). In order to test whether redistribution of RhoB occurs upon ABP treatment in leukemic cancer stem cells also, we sorted leukemic blasts, cancer stem cells and healthy stem cells from the very same donor based on flow markers (Terwijn et al., 2014), and we quantified distribution of RhoB. Indeed, in primary blasts from an acute myeloid leukemia patient/donor, RhoB localization correlated with the recognition by V$_y$9Vl2 TCR-engineered T cells (Figures 3B and S3A) including leukemic stem cells, but not healthy stem cells from the same donor (Figure 3F). This supports our previous observation that V$_y$9Vl2 TCR-mediated recognition selectively targets the leukemic but not the healthy stem cell pool (Marcu-Malina et al., 2011). Taken together, the intracellular distribution of the small GTPase RhoB strongly correlates with recognition of diverse tumor targets, including cancer stem cells, and exclusion of RhoB from the nucleus is a signature of tumor cells that are susceptible for targeting by V$_y$9Vl2 TCR$^+$ T cells.

**RhoB Regulates Membrane Mobility of BTN3A1 on Cancer Cells**

Decreased mobility of the transmembrane protein BTN3A1 on target tumor cells has recently been identified as a critical determinant of V$_y$9Vl2 T cell activation (Harly et al., 2012; Sandstrom et al., 2014; Vavassori et al., 2013). However, the mechanisms linking intracellular accumulation of phosphoantigens to changes in BTN3A1 membrane mobility are unclear. Therefore, the impact of ABP and RhoB, an important player in the cytoskeletal reorganization and formation of actin stress fibers (Prendergast, 2001), on mobility of BTN3A1 was tested. Consistent with previous reports (Harly et al., 2012; Sandstrom et al., 2014), treating 293 HEK cells with the ABP zoledronate resulted in decreased BTN3A1 membrane mobility. Strikingly, treatment with calpeptin induced immobilization of BTN3A1 to similar levels as those of ABP treatment alone, whereas C3-transferase counteracted this ABP effect (Figure 4A). This points to the possibility that Rho GTPase activity acts on BTN3A1 membrane immobilization upstream of the mevalonate pathway. Selective depletion of RhoB by CRISPR/Cas inhibited the ABP–induced immobilization of BTN3A1 to levels comparable to those of medium controls, suggesting that ABP-mediated changes in BTN3A1 mobility depend on RhoB.

To next assess a role for RhoB-induced cytoskeletal rearrangements in mediating the observed changes in BTN3A1 mobility, the relation between BTN3 molecules and F-actin was investigated by colocalization experiments. HEK293 cells were stained with fluorescently labeled anti-BTN3 and phalloidin, and colocalization measurements in mediating the observed changes in BTN3A1 mobility, the relationship between BTN3 and F-actin was investigated by colocalization experiments. HEK293 cells were stained with fluorescently labeled anti-BTN3 and phalloidin, and colocalization coefficients were determined in response to treatment with ABP and calpeptin. In cells in a culture medium, a variable but considerable colocalization between BTN3 and F-actin was observed, and was markedly reduced by ABP treatment (Figure 4B). Strikingly, and similar to its effect on BTN3A1 membrane mobility, calpeptin reduced colocalization between BTN3 and F-actin to comparable levels observed with ABP treatment. This reduction suggests that both phosphoantigen accumulation and Rho activation may induce the formation of membrane domains surrounded by cytoskeleton, where BTN3 molecules could be trapped and immobilized, as recently described for FcεRI (Andrews et al., 2008). Importantly, C3-transferase prevented ABP-induced BTN3-actin segregation, indicating again the crucial involvement of active Rho in this process. Together, these data suggest that RhoB activity contributes to target recognition by V$_y$9Vl2 TCR$^+$ T cells by modulating BTN3A1 membrane mobility through cytoskeletal rearrangements.

**RhoB Interacts with BTN3A1 Homodimers in Cancer Cells Recognized by V$_y$9Vl2 TCRs**

Given the strong requirement for RhoB activity in the membrane immobilization of BTN3A1, we questioned whether regulation of BTN3A1 involved direct interactions with RhoB. Using in situ proximity ligation assay (PLA), RhoB and BTN3 were observed to be in close proximity in recognized EBV-LCL 48 cells only when pretreated with the ABP (Figure 5A). Importantly, PLA signals were typically excluded from the nuclear area and distributed close to the plasma membrane, in line with our data that RhoB is involved in V$_y$9Vl2 TCR$^+$ T cell recognition by regulating membrane-expressed BTN3A1.

To determine whether BTN3A1 exists as a homodimer when expressed in a cellular context, as suggested from crystalization studies (Palakodeti et al., 2012), and to study RhoB-BTN3A1 interactions at even higher resolution, we examined close interactions by utilizing fluorescence resonance energy transfer (FRET). Flow cytometry FRET measurements were performed on ABP-sensitive HEK293 cells (Harly et al., 2012), by either overexpressing FRET compatible fusion proteins or labeling endogenous proteins with antibodies coupled to FRET-compatible fluorochromes. These experiments showed that BTN3A1 molecules are expressed as homodimers on the cell surface of cancer cells (Figure 5B); however, the pairing of BTN3A1 molecules was insensitive to ABP-induced phosphoantigen accumulation. Close association between RhoB and BTN3A1 was undetectable in ABP-untreated HEK cells but increased markedly after treating cells with the ABP (Figure 5C). We used Biolayer Interferometry (BLI) to formally define a possible docking site for RhoB on the intracellular domain of BTN3A1. RhoB binding was detected with recombinant full-length BTN3A1 intracellular domain (BFI) (Figure 5D, left, and Table S1) yet was significantly reduced when using a recombinant BTN3A1 B30.2 domain, lacking the N-terminal region connector to the transmembrane domain (Figure 5E left). These data indicate an important role for the
Phosphoantigen Accumulation Associates with Conformational Changes of BTN3A1 Dimers

Phosphoantigen accumulation associates with conformational changes of BTN3A1 dimers. This interaction is regulated by phosphoantigens. The crystal structures of the extracellular domain of BTN3A1 in complex with the functionally well-characterized BTN3-specific antibodies 20.1 and 103.2 have recently been resolved and have revealed that both antibodies bind to different epitopes on the membrane-distal Ig-V domains of BTN3A1 dimers (Harly et al., 2012). To study BTN3A1 conformational changes in response to increased phosphoantigen levels, surface membranes of either unstimulated or ABP-stimulated HEK293 cells were labeled with the fluorescent lipid conjugate BODIPY FL (donor), and subsequently stained with acceptor dye-labeled BTN3-specific antibodies on ice, in order to prevent conformational changes that could be driven by these antibodies under stimulatory conditions.
physiological circumstances (Gáspár et al., 2001). Without ABP stimulation, potent FRET efficiencies between stained membrane and both antibodies were observed (Figure 6B), suggesting that the BTN3 Ig-V domain is in close proximity to the cell membrane. Strikingly, however, treatment of cells with ABP resulted in a marked reduction in FRET signals (Figure 6B), demonstrating that intracellular phosphoantigen accumulation associates with a conformational change of BTN3 molecules.

Figure 5. RhoB Interacts with BTN3 Molecules and Dissociates after Phosphoantigen Treatment
(A) EBV-LCL 48 cells were treated either with medium or ABP pamidronate, loaded onto poly-L-lysine-coated coverslips, and permeabilized. The interaction between RhoB and BTN3 was subsequently assessed by Duolink PLA using anti-RhoB and anti-CD277 antibodies. Duolink PLA without antibodies against RhoB and BTN3 served as negative control (red, PLA signal; blue, nucleus [DAPI]; dotted line, cell membrane). Figures are representative of two independent experiments.

(B) HEK293 cells were treated with either medium or pamidronate and co-stained with equal amount of anti-CD277-PE (donor) antibodies, and FRET efficiency in cells was measured as described in the Supplemental Information. Data shown are mean ± SEM of three independent experiments, in triplicate samples, where Mann-Whitney test was used to analyze statistical significance.

(C) HEK293 cells were pretreated either with medium or pamidronate, trypsinized, permeabilized, and stained with anti-RhoB-Alexa Fluor 488 (FRET donor) and anti-CD277-DyLight 680 (FRET acceptor) antibodies. FRET efficiency was subsequently measured by flow cytometry as described in the Supplemental Information. Data show mean ± SEM of three independent experiments, in triplicate samples, where Mann-Whitney test was used to analyze statistical significance.

(D) Concentration dependent binding of the full-length BTN3A1 intracellular domain (BFI) with RhoGTPase in the presence or absence of the phosphoantigen cHDMAPP. Binding of BFI to RhoGTPase was measured using Biolayer Interferometry (BLI) either in the absence of cHDMAPP (left panel) or presence of cHDMAPP (1:1) (right panel). Concentrations of BTN3A1 BFI shown in the upper panel are 6.25, 12.5, 25, 50, and 100 μM shown in gray. The kinetics fitting curves are shown as black. In the lower panel, concentrations of BTN3A1 BFI shown are 3.75, 7.5, 15, 30, and 60 μM shown in gray. Data show one representative experiment.

(E) Same experimental setup but with recombinant BTN3A1 B30.2 domain, lacking the N-terminal region connector to the transmembrane domain. In the left panel, the interaction was measured without cHDMAPP. Concentrations of BTN3A1 B30.2 shown were 12.5, 25, 50, 100, and 200 μM shown in gray. The kinetics fitting curves are shown as black. In the lower panel, the interaction was measured with cHDMAPP (1:1). Concentrations of B30.2 domain shown are 3.75, 7.5, 15, 30, and 60 μM shown in gray. Data show one representative experiment.
This change involves a pronounced distancing of the Ig-V domain from the cell membrane. Importantly, Rho-inhibitor C3-transferase treatment (Figure S2F) of cells did not prevent ABP-driven conformational changes of the BTN3A1 homodimer (Figure 6B) indicating that this conformational change is independent of the enzymatic activity of RhoB. These data provide support for the hypothesis that increases in intracellular phosphoantigen levels can induce extracellular changes in BTN3A1 dimers that may act as, or contribute to a molecular signature recognized by Vγ9Vδ2 TCRs.

**DISCUSSION**

The translation of tumor-reactive Vγ9Vδ2 T cells to clinical applications has proven challenging, due to a limited understanding of the molecular mechanisms by which Vγ9Vδ2 T cells recognize their targets. Important progress has been made in this area by recent elegant reports identifying BTN3A1 as a phosphoantigen sensor that plays a key role in the activation of Vγ9Vδ2 T cells (Vavassori et al., 2013; Harly et al., 2012; Sandstrom et al., 2014). These studies suggest, however, nearly opposing mechanisms for the exact role of BTN3A1 in this process. Importantly, these findings, based on the model of Vγ9Vδ2 TCR-engineered CD4+ αβT cells, are most likely transferrable to the recognition of Vγ9Vδ2 T cells, as both populations have comparable activity against defined target cells as we recently demonstrated (Marcu-Malina et al., 2011). However, we cannot exclude that the modulating effect of RhoB for the recognition of tumor cells by Vγ9Vδ2 T cells can be partially overcome by utilizing Vγ9Vδ2 TCRs of higher affinity, or additional co-stimulatory molecules (Gründer et al., 2012). Together, these data suggest that RhoB is a key actor in the cellular mechanisms that couple phosphoantigen accumulation to BTN3A1-mediated triggering of Vγ9Vδ2 T cell responses.

The intrinsic GTPase activity of RhoB that cycles between biologically active GTP-bound and inactive GDP-bound conformations enables the differential association with downstream signaling pathways. As such, RhoB functions as a molecular switch in diverse cellular processes, ranging from gene transcription to the regulation of cytoskeletal changes and vesicle transport (Prendergast, 2001). Our data provide several lines of evidence that support a direct link between the mevalonate pathway-dependent changes in RhoB activity in tumor cells, cytoskeletal reorganization, and tumor cell-induced activation of Vγ9Vδ2 TCR+ T cells. However, the biological activity of RhoB seems to be partially genetically predetermined, as suggested by our initial screening strategy, we could not identify a direct genetic link of the surrogate marker SNP with RhoB activity. It is possible that differential SNPs found in recognized tumor cells, with the frequency of around 70% in the normal population, act as enhancer of a distant genomic region through chromatin looping (Pombo and Dillon, 2015), and indirectly influence RhoB activity. The importance of RhoB expression in the studied cancer cell lines was further emphasized by the loss in proliferative activity of complete RhoB knockdowns. Although we encountered a genetic block in a small subset of patients who were largely non-responsive to Vγ9Vδ2 TCR+ T cell-mediated tumor control, our results identified in RhoB a crucial link connecting intracellular changes to the recognition of target cells by Vγ9Vδ2 TCR+ T cells (Gober et al., 2003). These changes are
induced by the accumulation of phosphoantigens occurring in the majority of cancer cells.

Our data suggest that pAgs act differently in separate cell compartments and support a two-component model. A dysregulated mevalonate pathway leads to the activation and relocation of RhoB to the membrane-proximal connector region of the BTN3A1 cytoplasmic tail, through a process requiring GTPase activity (Figure 7). B30.2 domains have been reported to function as scaffold modules by spatiotemporally sequestering signaling proteins to distinct subcellular locations (Perfetto et al., 2013). This interaction favors an enhanced submembranous cytoskeletal arrangement, which leads to the observed immobilization of BTN3A1 at the cell surface, similar to its reported role in modulating membrane structures such as focal adhesions (Allal et al., 2002; Vega et al., 2012). Moreover, our data suggest a sequential, second cascade of events in the transmembrane compartment, which is necessary for recognition by a Vγ9Vδ2 TCR+ T cell: RhoB dissociates from immobilized BTN3A1 most likely after phosphoantigen binding to B30.2 and the conformational change in the extracellular region (ER) of BTN3A1 leading to Vγ9Vδ2 TCR T cell activation. Consistent with our observations, other recent data also suggest that additional factors are likely to be involved in the

**Figure 7. Two-Component Mechanism in Tumor Cells that Leads to Activation of Vγ9Vδ2 TCR T Cells**

Non-activating phase: no accumulation of phosphoantigens (pAg, yellow stars), which retains GDP-bound RhoB (red circles+GDP) from extranuclear areas.

Activating phase component I: accumulation of phosphoantigens is followed by more GTP-bound RhoB formation (red circles+GTP). GTP-bound RhoB undergoes subcellular re-compartementalization (black arrows) accumulating at extranuclear areas and facilitates spatial redistribution of BTN3A1 by promoting cytoskeletal trapping (green lines) in the plasma membrane binding to the B30.2 domain proximal connector region (CR) of BTN3A1 (blue hexagon). Activating phase component II: GTP-bound RhoB dissociates (black arrow) from while pAg binds to the B30.2 domain of BTN3A1, which triggers a conformational change of the extracellular region (ER) of BTN3A1 leading to Vγ9Vδ2 TCR T cell activation.
recognition of tumor cells by Vδ9Vα2 TCR+ T cells. In particular, transfer of human BTN3A1 alone to rodent cells, which are not recognized by Vδ9Vα2 T cells (Morita et al., 2007), was insufficient to sensitize cells toward recognition by Vδ9Vα2 T cells (Sandstrom et al., 2014). Instead, this sensitization required co-transfer of a substantial proportion of the human chromosome 6 (Riaño et al., 2014; Vavassori et al., 2013), suggesting that the combined presence of BTN3A1 and additional human genes located on this chromosome is essential for facilitating target cell recognition by Vδ9Vα2 T cells. The human RhoB gene locates to chromosome 2, and moreover, protein sequences of RhoB are fully conserved between human and rodents. Thus, other human-specific genes that mediate activation of Vδ9Vα2 T cells are yet to be identified. These might include GTPase activating proteins (GAPs) and guanine nucleotide exchange factors (GEFs), both of which coordinate actions that regulate the balance between the active, GTP-bound state and inactive, GDP-bound state of Rho GTPases (Rosman et al., 2005; Moon and Zheng, 2003). However, none of the known regulators of Rho GTPase signaling locate to the region on chromosome 6 required to transfer Vδ9Vα2 T cell susceptibility to rodent cells. Identification of these missing factors linking RhoB and BTN3A1 to target cell recognition by Vδ9Vα2 T cells is therefore an important challenge for further elucidating the molecular activation mechanisms of Vδ9Vα2 TCR+ T cells.

The identification of RhoB as a mechanistic link between transformation-associated dysregulation of the mevalonate pathway and BTN3A1-dependent activation of Vδ9Vα2 TCR+ T cells has important implications for cancer immunotherapeutic concepts using Vδ9Vα2 T cells or their TCRs. Our observations strongly suggest the potential of Vδ9Vα2 T cell-mediated immune interventions to target selectively the leukemic but not the healthy stem cell fraction. Thereby, the divergent intracellular distribution of RhoB among recognized and non-recognized tumor cells may be used as a biological surrogate marker for the identification of patients susceptible to Vδ9Vα2 TCR+ T cell-mediated tumor cell killing. Second, although we have not been able to provide a direct genetic link between the marker SNP and the function of RhoB, our data strongly argue for a genetic predisposition, which most likely acts via post-translational regulation of RhoB, and partially explains why some patients do not benefit from Vδ9Vα2 TCR+ T cell-mediated tumor control. Third, our data put RhoB forward as a key therapeutic target for improving immunotherapy using Vδ9Vα2 TCR+ T cells. In contrast to BTN3A1, which is ubiquitously expressed on healthy as well as malignant cells, the data presented here and elsewhere (Prendergast, 2001) demonstrate that RhoB possesses transformation-selective characteristics, such as spatial distribution, and could therefore represent a more selective therapeutic target compared to BTN3A1. Moreover, the mobilization of Vδ9Vα2 T cells in cancer patients using in vivo or ex vivo stimulation protocols with ABPs such as zoledronate (Fournié et al., 2013; Fisher et al., 2014; Schepers et al., 2013) is likely limited by the poor pharmacokinetic profile of ABPs (Lin, 1996) and their toxicity to T cells upon prolonged exposure (Wang et al., 2011). Agents modulating RhoB GTPase activity could raise attractive opportunities to complement currently pursued Vδ9Vα2 TCR+ T cell-based therapies. In summary, with RhoB we provide a key molecule that regulates the ability of cancer cells to activate Vδ9Vα2 TCR+ T cells, by orchestrating BTN3A1 in the plasma membrane. This knowledge will enable the pre-selection of patients susceptible to Vδ9Vα2 T cell based therapies and will enhance the efficacy of such therapies.

**EXPERIMENTAL PROCEDURES**

### Retroviral Transduction of TCRs

The Vδ9Vα2 TCR genes were transduced into αβ T cells using retroviral supernatant from Phoenix-Ampho packaging cells that were transfected with gag-pol, env, and pBulret retroviral constructs containing TCR genes.

### Functional T Cell Assays

IFN-γ ELISPOT was performed as 15,000 T cells and 50,000 target cells were cocultured in nitrocellulose plates, and the number of spots was quantified using ELISPOT Analysis Software.

### Zygosity/SNP Correlation Analysis

Recognition of CEPH EBV-LCL lines by Vδ9Vα2 TCR+ T cells was defined in IFN-γ ELISpots. Hypothetical zygosities for candidate genetic loci were deduced using classical Mendelian inheritance patterns within CEPH family pedigrees, where the influence of candidate alleles on Vδ9Vα2 TCR-mediated recognition was assumed to be dominant. Correlations of predicted zygosities with HapMap SNP genotypes of CEPH individuals were calculated with the software tool ssSNPer.

### Confocal Microscopy and Data Analysis

Cells were allowed to attach onto poly-L-lysine precoated coverslips and were permeabilized and stained with anti-RhoB antibody. Images were acquired using a Zeiss confocal laser scanning microscope LSM 700. Ratios between nuclear and extranuclear signal of RhoB was determined using ImageJ software, using DAPI staining to mark nuclei.

### Statistical Analysis

All experiments were repeated at least three times unless otherwise indicated. All data were shown as mean ± SEM. Statistical significance was analyzed by either Mann-Whitney or Kruskal-Wallis test and Dunn’s multiple comparison test.

All patients or their legal guardians provided written informed consent.

### SUPPLEMENTAL INFORMATION

Supplemental Information includes Supplemental Experimental Procedures, four figures, and one table and can be found with this article online at http://dx.doi.org/10.1016/j.celrep.2016.04.081.

### AUTHOR CONTRIBUTIONS

Z.S., W.S., and J.K. designed the study and wrote the manuscript. Z.S., R.O., and T.M. designed and performed the computational SNP analysis. S.G. and E.J.A. designed and performed in vitro protein expression and purification and BLI assays. Z.R. and Z.S. generated T cells and performed EBV-LCL screening. A.C., A.V., Z.R., and Z.S. designed and performed Rho activity assays. M.S., M.v.N., and A.V. performed miRNA analysis. R.J.L. designed and produced CRISPR/Cas constructs and viruses. C.M.P., E.S., and Z.S. designed and performed Rho activity assays. O.C., A.V., and Z.S. performed confocal analysis. G.J.S. designed and carried out FACS sorting of leukemia stem cells. D.O. produced and purified CD277 antibodies.

### ACKNOWLEDGMENTS

Funding for this study was provided by ZonMW 43400003 and VIDI-ZonMW 917.11.337, KWF. UU 2010-4689, UU 2013-6426, UU 2014-6780, and UU 2015-7601, Vrienden van het UMCU, ACR 10-0736, and 15-0049 to J.K.
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