NLRC5 promotes transcription of BTN3A1-3 genes and Vγ9Vδ2 T cell-mediated killing

HIGHLIGHTS
- BTN3A promoters contain a unique regulatory motif occupied by overexpressed NLRC5
- NLRC5 and BTN3A mRNA levels correlate in healthy and diseased cells
- NLRC5 overexpression increases susceptibility to Vγ9Vδ2 T-cell-mediated elimination

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Butyrophilins (BTNs) are an emerging family of molecules fulfilling immune and non-immune functions. Human BTNs comprise BTN1A1, BTN2A1, and BTN2A2, as well as BTN3A1, BTN3A2, and BTN3A3. The genes encoding these proteins are located in the extended major histocompatibility complex (MHC) locus on chromosome 6 (Abeler-Dorner et al., 2012; Arnett and Viney, 2014). Structurally, BTNs are membrane proteins sharing similarity with the B7 immunoglobulin superfamily of costimulatory/coinhibitory molecules (Arnett and Viney, 2014). In fact, BTN2A2 and BTN1A1 have been shown to act as co-inhibitory ligands hindering T cell activation and proliferation, and BTN2A2 knockout mice exhibit exacerbated T-cell-mediated autoimmunity (Sarter et al., 2016; Smith et al., 2010).

Human BTN3A1-3 proteins, which have no murine homologs, are composed of two extracellular immunoglobulin-like domains and a transmembrane region linked—in BTN3A1 and BTN3A3—to an intracellular B30.2 domain (Harly et al., 2012; Sandstrom et al., 2014; Vavassori et al., 2013). Despite this divergence in the intracellular portion, BTN3A1, BTN3A2, and BTN3A3 show >95% homology in the extracellular domain, suggesting that they are the products of recent duplications. Although not sufficient, BTN3A1 is necessary for the activation of Vγ9Vδ2 T cells (Riano et al., 2014; Sandstrom et al., 2014; Vantourout et al., 2018; Vavassori et al., 2013). Although the expression of BTN3A2 and BTN3A3 can support BTN3A1’s function, it recently became clear that BTN2A1 is the second critical molecule to stimulate Vγ9Vδ2 T cells (Karunakaran et al., 2020; Rigau et al., 2020; Vantourout et al., 2018). BTN2A1 presents the intracellular B30.2 domain and the two extracellular immunoglobulin-like domains (Abeler-Dorner et al., 2012; Arnett and Viney, 2014). Vγ9Vδ2 T cells are activated by phosphorylated metabolites, also called phosphoantigens (PAgs), that derive from a dysfunctional mevalonate pathway, such as isopentenyl pyrophosphate (IPP), or from microorganisms, as for instance Mycobacterium tuberculosis-derived (E)-4-hydroxy-3-methyl-but-2-enyl pyrophosphate (HMBPP) (Vantourout and Hayday, 2013). PAgs have been proposed to bind either the immunoglobulin-like domain or the B30.2 domain of BTN3A1, inducing a conformational change and/or stabilizing surface BTN3A1 to engage the γδ T cell receptor, whereas BTN2A1 interacts with the germline region of the Vγ9 chain (Gu et al., 2017; Harly et al., 2012; Karunakaran et al., 2020; Rigau et al., 2020; Sandstrom et al., 2014; Vavassori et al., 2013). Increased levels of PAgs from metabolically stressed, bacterial and/or infected donors’ blood cells, and in primary tumors. Moreover, forcing NLRC5 expression promoted Vγ9Vδ2 T-cell-mediated killing of tumor cells in a BTN3A-dependent manner. Altogether, these findings indicate that NLRC5 regulates the expression of BTN3A genes and hence open opportunities to modulate antimicrobial and anticancer immunity.
transformed, and infected cells are thus sensed by Vy9V62 T cells, leading to their activation, expansion, and participation in the immune response (De Libero et al., 2014). For instance, this subset of unconventional T cells is significantly expanded during M. tuberculosis infection (Cheng et al., 2018; Kabelitz et al., 1991). Furthermore, intratumoral Vy T cells emerged as the most significant favorable cancer-wide prognostic population, and their potential role in immunotherapy is being increasingly investigated (Bennyamine et al., 2016, 2017; Gentles et al., 2015; Le Page et al., 2012; Peedicayil et al., 2010; Zocchi et al., 2017).

The transcriptional regulation of BTN genes remains poorly characterized. Recently, it has been shown that Btn2a2 induction is regulated by the transcriptional regulator CIITA (class II major histocompatibility complex transactivator) (Sarter et al., 2016). This factor belongs to the nucleotide-binding oligomerization domain (NOD)-like receptor (NLR) family of proteins and, together with its closest homolog NLRCS (NLR family CARD domain containing 5), is known to control the transcription of MHC and related genes (Chelbi et al., 2017; Jongsma et al., 2019; Reith and Mach, 2001; Sarter et al., 2016). CIITA is the master transcriptional regulator of MHC class II genes, whereas we and others showed that NLRCS is an important transcriptional regulator of MHC class I genes, markedly in T lymphocytes (Chelbi et al., 2017; Meissner et al., 2010; Neerincx et al., 2013; Robbins et al., 2012; Staehli et al., 2012; Yao et al., 2012). These two NLRs, CIITA and NLRCS, are recruited to their respective target gene promoter by a multiprotein complex known as “enhanceosome” (Ludigs et al., 2015; Meissner et al., 2012a; Neerincx et al., 2012). This complex assembles on the promoter sequence called “SXY” module, which is composed of four individual elements (S, X1, X2, and Y) oriented and spaced in a specific manner (Anderson et al., 2017; Ludigs et al., 2015; Masternak et al., 2003; Meissner et al., 2012b; Neerincx et al., 2012 (Krawczyk et al., 2004)). Although we still do not know which factor recognizes the S-box, the X1-box is bound by the regulatory factor X (RFX) complex, the X2-box by cAMP-responsive element binding protein (CREB)/activating transcription factor (ATF) family members, and the Y-box by the nuclear transcription factor Y (NFY)-complex (Chelbi et al., 2017). Taken together, the finding that Btn2a2 is a target of CIITA and the localization of BTN genes close to the MHC locus suggest the existence of common evolutionary links. This prompted us to investigate the transcriptional regulation of BTN genes and to hypothesize that they represent a novel set of NLRCS or CIITA targets.

We found that BTN3A1-3 genes exhibited an atypical SXY module in their proximal promoter region. This regulatory motif presented a reverse complement Y-box at an altered spacing from the X-motif. Using chromatin immunoprecipitation and gene reporter assays, we demonstrated that overexpressed NLRCS occupies and transactivates this atypical module. We showed that forcing NLRCS expression led to increased levels of BTN3A mRNA and protein. Data mining in transcriptome datasets of M. tuberculosis-infected and uninfected individuals’ blood as well as of various cancers revealed a strong correlation between NLRCS and BTN3A1 and BTN3A2 gene expression. Furthermore, loss of both NLRCS copies was associated with significantly diminished expression of BTN3A1 and BTN3A2 in cancer cells. On a functional level, we observed that overexpression of NLRCS enhanced Vy9V62 T-cell-mediated elimination of target cells, which was mediated by BTN3A molecules as demonstrated using a loss-of-function approach. Altogether, these findings indicate NLRCS modulation as a possible way of targeting Vy9V62 T-cell-mediated immunity.

RESULTS

BTN3A genes have distinguishable S and X modules

To understand how far the mechanisms governing MHC gene transcription might be common to BTN genes, we screened their promoter sequences for the presence of SXY modules (Figures 1A and 1B) (Krawczyk et al., 2008; Ludigs et al., 2015). We confirmed the presence of an SXY consensus in the BTN2A2 promoter (Figure 1B) (Sarter et al., 2016). Previous findings showed that the S-motif of NLRCS-occupied promoters is unique and strictly required for NLRCS activity (Figure 1A) (Ludigs et al., 2015; Meissner et al., 2012a). The S-box sequence in both human and mouse BTN2A2 did not suggest regulation through NLRCS, but rather a role for CIITA (Figures 1B and S1A). Indeed, expression of murine Btn2a2 in various organs derived from Nlrc5-deficient mice was not significantly decreased (Figure S1A). We also noticed the presence of an X-motif in the promoters of BTN2A1, the pseudogene BTN2A3, BTN3A1, BTN3A2, and BTN3A3 genes (Figures 1A and 1B). Of note, 15 bp upstream of the X-box, the highly homologous promoter regions of the BTN3A genes, exhibited an S-box suggestive of NLRCS-mediated transcriptional regulation.

BTN3A1-3, BTN2A1, NLRCS, and CIITA transcripts were abundant in most of the tested immune organs and cells (Figure 1C). Among the main blood cell subsets, the mRNA profile of BTN3A1-3 was reminiscent
**A**

CIITA module

NLRC5 module

**B**

BTN2A1

BTN2A2

BTN2A3 (ps)

BTN3A1

BTN3A2

BTN3A3

**C**

- [Graphs showing relative mRNA levels for NLRC5, CIITA, and BTN2A1, BTN3A1, BTN3A2, BTN3A3 across different samples.]
of the one of NLRC5, whereas it largely differed from the one of CIITA (Figure 1D). Instead, BTN2A1 presented a distinct pattern, characterized by higher expression levels in monocytes (Figure 1D) (Rigau et al., 2020). We also observed a marked similarity between the expression pattern of NLRC5 and the BTN3A protein display (by using a monoclonal antibody anti-CD277, which recognizes all three BTN3A isoforms; Figures 1E and S1B). We next took advantage of bare lymphocyte syndrome (BLS) patient-derived and in-vitro-generated B cell lines lacking expression of CIITA or the RFX complex subunit RFX5 or RFXAP (Ludigs et al., 2015; Tarantelli et al., 2018). The absence of the RFX factors affected the expression of BTN3A1-3 and BTN2A1 transcripts, in particular following interferon (IFN) treatment (Figure S1C). This was not observed in the absence of CIITA, indicating that this NLR was not necessary for the regulation of these genes (Figure S1C). Taken together, these data support the involvement of the enhanceosome platform and prompted us to perform further analyses on NLRC5 in the regulation of BTN3A1-3 genes.

**NLRC5 overexpression induces transcription of BTN3A genes**

We next tested whether NLRC5 induced the transcription of BTN3A1-3 genes, first employing a primer pair that detects all three gene products. After 48 h, overexpression of NLRC5 in HEK293T cells led to increased transcript levels of BTN3A1-3 and HLA-B, the latter used here as a positive control (Figure 2A). To assess the specificity of the transcriptional effects of NLRC5, we overexpressed other NLR members. Although NOD1, NOD2, and NLRC3 had marginal effects on the expression of BTN3A1-3 and of HLA-B, two CIITA isoforms (CIITA I and CIITA III) induced the BTN3A1-3 and the HLA-B gene (Figure 2A). The induction in the levels of BTN3A1-3 and HLA-B transcripts by NLRC5 and CIITA I encoding plasmids was already detected 24 h following transfection (Figure S2A). Both NLRC5 and CIITA I induced the three BTN3A genes (Figure 2B). In contrast, only CIITA I moderately induced BTN2A1 mRNA (Figure 2B). Finally, we demonstrated that overexpression of NLRC5 and CIITA, but not of NLRC3, increased surface expression of BTN3A1-3 (Figure 2C). It is important to point out that although CIITA does not regulate MHC class I genes at the endogenous level, it is well established that its overexpression leads to their transactivation (Chang et al., 1996; Gobin et al., 1997; Ludigs et al., 2015; Martin et al., 1997; Robbins et al., 2012; Williams et al., 1998), as shown for HLA-B (Figure S2B). Conversely, the activity of NLRC5 maintains its specificity toward the HLA-B but not HLA-DRA genes even when overexpressed (Figure S2B). Therefore, these data encouraged us to further investigate whether BTN3A1-3 genes are regulated by mechanisms similar to the ones controlling MHC class I gene expression.

**BTN3A genes have an atypical SXY module**

In order to substantiate the hypothesis that BTN3A1-3 are direct targets of NLRC5, we cloned the proximal promoter region, containing the S- and X-motif, of BTN3A2—as a representative BTN3A promoter—into a luciferase reporter plasmid (referred to as “5X”). As we did not identify an “ATTGG” Y-box sequence nearby, we hypothesized that it could be dispensable for NLRC5-dependent transactivation of these genes (Figure 1B). However, we observed that these promoters were not transactivated by NLRC5 (Figure 3A). We therefore took a closer look at the promoters of BTN3A1-3 genes and identified the reverse complement of the “ATTGG” Y-box, “CCAA,” 13 bp downstream of the X-box (Figure 3B). As the latter constitutes the canonical NFY-binding site, we cloned an extended portion of the promoters, which included this motif, into a luciferase reporter plasmid (referred to as “SX-13bp” CCAAT”). This enabled transactivation by NLRC5 (Figure 3A). Moreover, scrambling of the “CCAA” sequence (referred to as “SX-13bp-CCTTT”) abrogated NLRC5-mediated transactivation, pinpointing the importance of this unique “Y” box (Figure 3A). NLRC5 was also able to transactivate the
``SX-13bp-CCAAT'' BTN3A1 promoter construct, which diverges from the ``SX-13bp-CCAAT'' BTN3A2 reporter by only 2 bp in the S-box, in a CCAAT-dependent manner (Figure 3C). Not surprisingly, similar results were observed when overexpressing CIITA I (Figures 3D and 3E). These results reveal the presence of an unconventional SXY module in the BTN3A promoters.

The S-box and the distance of the CCAAT-box are key for NLRC5-mediated transactivation

One intriguing feature of the BTN3A promoters is the fact that the spacing between the X-box and the CCAAT sequence (13 bp) is shorter than the usual one found at the MHC promoters between the X-box and the ATTGG (usually 17–18 bp). To assess whether the unusual orientation and distance were important, we generated a luciferase reporter plasmid in which the CCAAT sequence of BTN3A2 promoter was substituted by an ATTGG (referred to as ``SX-13bp-ATTGG'') at the 13 bp distance from the X-box. Interestingly, this promoter was transactivated neither by NLRC5 nor by CIITA (Figure 4A). We also moved the CCAAT sequence 18 bp downstream of the X-box (referred to as ``SX-18bp-CCAAT''). Again, NLRC5- and CIITA-mediated transactivation were impaired (Figure 4A). Yet, insertion of an ATTGG sequence at 18 bp distance from the X-box (referred to as ``SX-18bp-ATTGG''), which mirrors the classical organization of the HLA promoters, restored transactivation by NLRC5 and CIITA (Figure 4A). These results show that both orientation and distance of the NFY-binding site are crucial for transcriptional induction by these NLRs.

As the S-box found in the promoters of the BTN3A genes exhibits similarity to the one required for NLRC5-mediated transactivation, we next assessed its contribution to the transactivation of the BTN3A2 promoter. In line with results from the MHC class I gene promoter, scrambling the S-box sequence severely compromised NLRC5-mediated, but not CIITA-mediated, transactivation (Figure 4B). This underlines the importance of the S-box for NLRC5-mediated regulation.

Figure 2. NLRC5 overexpression increases BTN3A1-3 expression

(A and B) BTN3A1-3, BTN2A1, or HLA-B mRNA levels (relative to POLR2A mRNA) were measured by qRT-PCR 48 h following transfection of plasmids encoding the indicated NLR proteins or an empty vector (mock) in HEK293T. (C) HEK293T cells were co-transfected with vectors coding for the indicated NLRs or empty vector (mock) and a GFP-encoding plasmid to identify transfected cells. Graphs depict the quantification of HLA-A/B/C and BTN3A1-3 geometric MFI (gMFI) gating on GFP+ cells 48 h post-transfection. Histogram overlays show HLA-A/B/C and BTN3A1-3 expression for background (gray), mock- (blue), and NLRC5-transfected (pink) HEK293T cells. (A–C) Results are depicted as mean ± SD (n = 3 technical replicates) and are representative of at least two independent experiments. Statistical differences were determined by one-way ANOVA followed by comparison of the experimental conditions to the corresponding mock transfection and were corrected for multiple testing using the Dunnett method. Only statistically significant differences are illustrated. **p < 0.01; ***p < 0.001.

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BTN3A genes are direct targets of NLRC5

In order to prove that BTN3A genes are direct targets of NLRC5, we performed chromatin immunoprecipitation (ChIP) experiments. In the absence of an antibody against endogenous human NLRC5, we generated an HA-tagged version of wild-type NLRC5 (wt NLRC5) and of a mutant of the Walker A motif (mt NLRC5), which prevents NLRC5 nuclear translocation and transcriptional activity (Meissner et al., 2010). We then co-transfected HEK293T cells with plasmids coding for wt NLRC5 or mt NLRC5, and for the surface protein CD72, thus enabling the enrichment of transfected cells. ChIP using chromatin from wt NLRC5-transfected cells led to a substantial enrichment of the promoter region of HLA-B but also of BTN3A1 and BTN3A2 genes as compared with mt NLRC5-transfected cells (Figure 5). By contrast, the promoter of HOXC8, a gene not known to be controlled by NLRC5, was not enriched (Figure 5). Although we were unable to design primers specific for BTN3A3 promoter due to the high homology of the regulatory regions of these genes, these data demonstrate occupation of BTN3A1 and A2 promoters by NLRC5 (Figure 5).

BTN3A genes are co-regulated in health and disease

We next checked the correlation between NLRC5 and BTN3A1-3 expression in human pathological conditions in which Vγ9Vδ2 T cells are considered relevant. First, we analyzed transcriptomic data from a Gambia
Across cancers, we found a significant correlation between BTN3A and NLRC5, corroborating a generally weaker correlation between BTN2A and NLRC5 (Figures 6C and S3B). The correlation with HLA-B was generally good, with the exception of the dlbc dataset (Figure S3B) (Yoshihama et al., 2016; Staehli et al., 2012). In most cancers, we found a significant correlation between NLRC5 and BTN3A-3 transcript abundance (Figure 6C), whereas the correlation with NECTIN2 was low and mostly non-significant (Figure S3B). The correlation with HLA-B was generally good, with the exception of the dlbc dataset (Figure S3B). We therefore tested an independent cohort of DLBCL samples (GSE10846; Figure S3C), which corroborated a generally weaker correlation between NLRC5 and HLA-B than with the BTN3A genes. Across cancers, BTN3A-3 correlated better with NLRC5 than with CIITA expression, whereas the profile of BTN2A1 was quite independent from the transcript levels of both NLRs (Figure S3D).

We next investigated whether methylation of the NLRC5 promoter inversely correlated with the expression of the BTN3A genes. Data supported this hypothesis, including for HLA-B, whereas barely significant and variable results were observed for NECTIN2 (Figures 6D and S3E). We also interrogated whether structural alterations leading to NLRC5 copy number loss were concomitant with BTN3A gene expression decrease. We thus focused on breast (brca) and prostate (prad) cancer, the only tumor sets having more than five samples lacking both NLRC5 copies, and compared BTN3A expression of samples harboring loss of NLRC5 (homozygous deletion, HDEL) with samples having normal copy number (WT). Despite the small sample size, BTN3A1, BTN3A2, as well as HLA-B gene expression were reduced in the absence of NLRC5 (Figures S3E and S3F). As for most correlations tested in Figure 6, a similar trend was observed also for BTN3A3, although less robust (Figure 6E). We wondered whether the presence of a “C” instead of the conserved

*Figure 4. S-box sequence and CCAAT-box position are important for NLRC5 transactivation*

(A and B) Luciferase reporter assays were performed in HEK293T cells co-transfected with the parental pGL3 backbone or the indicated BTN3A2 promoter constructs, and a vector coding for NLRC5 or CIITA I, or an empty (mock) vector; mtSX indicates a scrambled S-box sequence (the CAACGTT sequence was substituted by CCAGAGT) (B). Data are expressed as fold transactivation as compared to the mock condition. Results represent mean ± SD of n = 3 technical replicates and are representative of at least three independent experiments. Statistical differences were determined by performing a two-way ANOVA followed by comparison of the indicated conditions and were corrected for multiple testing using the Holm-Sidak method. **p < 0.01; ***p < 0.001; ns: not significant.
“T” residue in the S-box of this gene’s promoter might underlie this observation (Figure S3G). We thus substituted this T with a C in the BTN3A2 SX-13bp-CCAAAT reporter plasmid. Supporting our hypothesis, this modification reduced NLRC5-mediated transactivation (Figure S3G). Taken together, these data underscore the robust correlation of NLRC5 with BTN3A1 and BTN3A2 transcript abundance in healthy and diseased conditions in which Vγ9Vδ2 T cells are considered relevant.

**NLRC5 overexpression confers susceptibility to γδ T-cell-mediated killing**

We thus overexpressed NLRC5 in Raji cells, a Burkitt lymphoma cell line known to trigger poor recognition by Vγ9Vδ2 T cells (Harly et al., 2012). As our results predicted, polyclonal NLRC5-transduced Raji cells presented increased levels of BTN3A1-3 surface expression as compared with mock-transduced control cells (Figure 7A). This was observed both in the absence and in the presence of zoledronate, a pharmacological inhibitor of the mevalonate pathway that increases IPP concentrations (Figure 7A). A similar trend was observed for HLA-A/B/C (Figure S4A). In a first set of experiments, killing of mock- or NLRC5-transduced polyclonal Raji cells was measured after 24 h of co-culture with in-vitro-expanded, primary Vγ9Vδ2 T cells in the presence of zoledronate. NLRC5 overexpression significantly enhanced the specific killing of Raji cells (Figure 7B). In these experiments, polyclonal populations of target cells with a transduction efficiency of roughly 68% were used. As this could have masked the effect of NLRC5 overexpression on Vγ9Vδ2 T-cell-mediated killing, we next generated subclones of mock- and NLRC5-transduced Raji cells. In line with data presented in Figure 2B, NLRC5-overexpressing subclones exhibited increased levels of BTN3A1-3 transcripts, but no difference in BTN2A1 mRNA (Figure 7C); in agreement, BTN3A1-3 surface expression was augmented (Figure 4B). Co-culture of Vγ9Vδ2 T cells with these subclones led to significantly increased IFNγ production (Figure 4C) and killing (Figure 7D). To demonstrate that this enhanced killing was mediated by the BTN3A molecules, we knocked out BTN3A1 by CRISPR/Cas9 in the NLRC5-overexpressing subclone 2. This loss-of-function strategy led to a decrease in the transcript levels of BTN3A1 and—due to the high homology of these two genes—of BTN3A2 (Figure S4D). As expected, these cells exhibited significantly reduced susceptibility to be killed by Vγ9Vδ2 T cells (Figure S4E). Therefore, NLRC5 overexpression not only increases the levels of BTN3A1-3 but also functionally promotes Vγ9Vδ2 T cell activation and killing.

**DISCUSSION**

The role of NLRC5 as a transcriptional regulator of MHC class I genes has been rapidly unveiled over the past years (Chelbi et al., 2017; Ludigs et al., 2015; Meissner et al., 2010, 2012a; Neerincx et al., 2012, 2013; Robbins et al., 2012; Staehli et al., 2012; Yao et al., 2012). Yet, our knowledge of its global transcriptional targets in humans remains limited. This work provides the first evidence for the transcriptional regulation of BTN3A1-3 genes by NLRC5, broadening our understanding of its role in humans.

Analysis of promoter sequences of BTN genes that are clustered in the adjacent MHC locus highlighted the presence of SXY modules. One of these modules, in the BTN2A2 promoter, is transactivated by CIITA (Sarter et al., 2016). In contrast, CIITA’s closest homolog NLRC5 did not contribute to the expression of Btn2a2 in the tested immune organs, adding further evidence for the role of CIITA in regulating this gene. We also identified an SXY module in the promoter of BTN2A1, and our results indicate that its expression is largely dependent on the RFX complex. However, BTN2A1 transcript levels, which were not increased by NLRC5...
Figure 6. BTN3A1-3 expression correlates with NLRC5 levels

(A and B) Pairwise correlation of NLRC5 and BTN3A1, BTN3A2, BTN3A3 expression are visualized. Analyses were performed using transcriptome datasets of the Gambia M. tuberculosis (TB) cohort study (GSE28623). (A) Scatterplots of gene expression are shown. The samples are divided in the groups “uninfected” (black), “latently infected” (red), and “active TB” (green). The table displays the Spearman’s correlation coefficient (R), and Bonferroni adjusted p-values.

| Gene   | R  | P_value | R  | P_value |
|--------|----|---------|----|---------|
| BTN3A1 | 0.79 | 1.53×10^{-2} | 0.83 | 1.71×10^{-2} |
| BTN3A2 | 0.53 | 6.78×10^{-2} | 0.58 | 1.96×10^{-2} |
| BTN3A3 | 0.23 | ns       | 0.23 | ns      |

(C and D) Correlation of NLRC5 mRNA and methylation levels with BTN3A1, BTN3A2, BTN3A3 expression.

(E) Differential expression of BTN3A1, BTN3A2, BTN3A3 in breast (brca) and prostate (prad) cancer cell lines. Fold change (fc) and p-values are indicated.
and moderately by CIITA, poorly correlated with these NLRs. Additional analyses are therefore needed to understand the transcriptional regulation of this gene. Further, data from BLS-derived B cell lines indicate that the induction of the BTN3A genes is also largely dependent on the presence of an enhanceosome, raising new questions on γδ T cell subsets in BLS patients.

Here, we identified an atypical SXY module in the promoters of BTN3A1-3 genes. Classical, non-classical, and selected MHC-related gene promoters contain—next to the S- and X-motif—an ATTTG Y-box (Krawczyk et al., 2008; Ludigs et al., 2015). Instead, the module of BTN3A1-3 promoters corresponds to the consensus sequence occupied by NLRC5 with regard to the S- and X-boxes but contains the reverse complement of the Y-box. This “CCAAAT” sequence corresponds to the canonical regulatory motif occupied by the trimeric NFY complex (Dolfi et al., 2012). Interestingly, the promoter of the MHC-class II-associated invariant chain (li, also called CD74) also contains an atypical SXY, with the CCAAT motif at a reduced distance from the X-box (Brown et al., 1991; Doyle et al., 1990; Zhu and Jones, 1990). This is in line with our observations in the BTN3A promoter, in which both orientation and spacing of the Y-box are crucial for NLRC5-mediated transactivation, presumably through formation of an alternative enhanceosome complex. Finally, we substantiate the importance of the S-box by showing that the substitution of a single conserved position affects NLRC5-mediated transactivation, possibly contributing to differences in the expression of individual BTN3A genes.

Importantly, overexpressed, but not endogenous, CIITA transactivates MHC class I genes, questioning its physiological contribution to BTN3A1-3 gene transcription (Chang et al., 1996; Gobin et al., 1997; Ludigs et al., 2015; Martin et al., 1997; Robbins et al., 2012; Williams et al., 1998). This is corroborated by the observation that CIITA deficiency did not reduce BTN3A expression in B cell lines. Although the question on the physiological contribution to BTN3A1-3 transcription is open also for NLRC5, robust correlative data support this possibility. BTN3A1-3 expression is abundant in T and NK cells, similar to the profile of NLRC5 (Neenncx et al., 2010; Staehli et al., 2012; Wu et al., 2009). The S-box in BTN3A1-3 promoters strongly resembles the one of NLRC5-transactivated genes (Ludigs et al., 2015; Meissner et al., 2012b). In the blood of both uninfected and M. tuberculosis-infected individuals, expression of NLRC5 strongly correlated with the profile of NLRC5. BTN3A1, which stimulates PAgS-mediated activation of Vγ9V2 T cells (Harly et al., 2012). Finally, we observed a robust coregulation between BTN3A1/BTN3A2 and NLRC5 expression in various cancers, and NLRC5 homozygous deletion was associated with a decrease in the abundance of BTN3A1 and BTN3A2 mRNA (Maertzdorf et al., 2011). Therefore, although we do not exclude a contribution of CIITA or a redundant function by these NLRs in the regulation of these genes, our results support a role for NLRC5 in regulating BTN3A gene transcription in normal as well as pathological conditions such as TB or cancer.

Our data demonstrate that forcing the expression of NLRC5 in cancerous target cells significantly promotes their killing by Vγ9V2 T cells, highlighting a novel functional link between NLRC5 expression and cytotoxicity by unconventional T cells. Because NLRC5 overexpression did not significantly impact BTN2A1 expression, and MHC-I and B2M are known to be dispensable and even hinder the activation of these unconventional T cells by engaging surface inhibitory receptors typical of NK cells (Bakker et al., 1998; Carena et al., 1997; Fisch et al., 1997; Halary et al., 1997; Morita et al., 1995), BTN3A molecules constitute the best candidates responsible for the NLRC5-driven enhanced Vγ9V2 T-cells-mediated killing. In accordance, CRISPR-mediated knockdown of BTN3A molecules nearly abrogated the killing enhancement induced by NLRC5 overexpressing cells. NLRC5 has already been linked to antitumor responses and patient prognosis (Farashi et al., 2019; Fernandez-Jimenez et al., 2019; Wang et al., 2019; Yoshihama et al., 2016). Although this has been attributed to its regulation of the MHC class I pathway, it will be important to
Figure 7. NLRC5 favors the activation of Vγ9Vδ2 T cells

(A) Polyclonal Raji cells were transduced with NLRC5-coding or mock vector and treated with zoledronate (+ZOL) for 24 h (bottom panel) or left untreated (top panel). BTN3A1-3 surface expression was analyzed by flow cytometry. Graphs depict a quantification of BTN3A1-3 geometric MFI (gMFI; top and bottom left), and the histogram overlays (bottom right) show BTN3A1-3 expression for unstained (US; black, gray), mock- (blue) and NLRC5-transfected polyclonal Raji cells (pink). Results represent six independent measurements.

(B) Specific killing of NLRC5- (pink) or mock-transduced polyclonal Raji cells (blue) was measured after 24 h of co-culture with Vγ9Vδ2 T cells at 3 effector-to-target ratios (1:1, 5:1, and 10:1) in the presence of ZOL. Depicted are the results for three independent healthy donors (HD) as mean ± SD (n = 3 technical replicates). The results are representative of four independent healthy donors and three independent experiments.

(C) BTN2A1, BTN3A1, BTN3A2, and BTN3A3 mRNA levels (relative to POLR2A mRNA) were assessed by qRT-PCR in the NLRC5- (red and violet) or mock-transduced Raji subclones (blue). The results depict mean ± SD (n = 3 technical replicates) and are representative of 2 independent experiments.

(D) Specific killing of the NLRC5- or mock-transduced Raji subclones was measured after 24 h of co-culture with Vγ9Vδ2 T cells at 3 effector-to-target ratios in the presence of ZOL. n.d. correspond to conditions where no specific lysis of the target cells by the Vγ9Vδ2 T cells was detected. Depicted are the results for three independent healthy donors (HD) as mean ± SD (n = 3 technical replicates). The results are representative of six independent healthy donors and three independent experiments. (A–D) Statistical differences between the condition with and without NLRC5 overexpression were calculated using paired Student’s t test (A) or post hoc Student’s t tests adjusted for multiple comparisons using the Holm-Sidak method (B–D). *p ≤ 0.05, **p ≤ 0.01, ***p ≤ 0.001. Only statistically significant differences are illustrated.
consider the potential contribution of the hereby discovered NLRC5/BTN3A regulation axis because altered expression of BTN3A genes has been associated with cancer and other diseases (Benyamine et al., 2017; Blazquez et al., 2018; Le Page et al., 2012; Peedicayil et al., 2010; Viken et al., 2009). Indeed, intratumoral γδ T cells have emerged as the most significant favorable cancer-wide prognostic infiltrating immune subset, and γδ T-cell-based clinical trials are increasingly performed, expanding our portfolio of immunotherapeutical approaches (Gentles et al., 2015). In addition to its role in the activation of γδ T cells, recent findings illustrate how BTN3A1 can play an inhibitory function on αβ T cells, by hindering the segregation of the phosphatase CD45 from the immune synapse (Payne et al., 2020). Importantly, this inhibition is relieved upon zoledronate treatment or by the use of CD277-specific agonistic antibodies, which enable the concomitant activation of VγVδ T cells. Our discoveries indicate therefore that targeting the NLRC5 axis, possibly in combination with such treatments, might represent an attractive anti-cancer strategy leveraging on both αβ and γδ T cells.

Our results suggest that, in humans, NLRC5 regulates more genes than previously thought and challenge its role as a transactivator of MHC class I and related genes only (Ludigs et al., 2015). BTN3A1 molecules have recently been shown to mediate Vy9Vy2 T cell activation in response to host- or microbial-derived metabolites, suggesting an involvement of NLRC5 in the γδ T-cell-mediated host immunity (Benyamine et al., 2016; Harly et al., 2012; Sandstrom et al., 2014; Vavassori et al., 2013). This observation is in agreement with previous findings that NLRC5 regulates non-classical MHC class I genes, such as murine H2-T10/H2-T22, which are recognized by a fraction of γδ T cells (Crowley et al., 2000). Therefore, regulation of BTN3A1 transcription by NLRC5 shows strong parallels with its established function, reinforcing its role as a modulator of conventional and unconventional T cell immunity.

Limitations of the study

Even if BTN3A molecules are induced by NLRC5 and required for Vγ9Vδ2 T cell activation, we cannot rule out the possibility that other NLRC5 targets, known or unknown, might contribute to the observed effect. In addition, further experiments are required to prove the pathophysiological relevance of these findings in cancer or infection.

Resource availability

Lead contact

Requests for further information and reagents should be directed to and will be fulfilled by the Lead Contact Greta Guarda (greta.guarda@irb.usi.ch).

Materials availability

Materials generated in this study will be made available upon reasonable request and may require a material transfer agreement.

Data and code availability

This study did not generate datasets or analyze codes.

METHODS

All methods can be found in the accompanying Transparent Methods supplemental file.

SUPPLEMENTAL INFORMATION

Supplemental Information can be found online at https://doi.org/10.1016/j.isci.2020.101900.

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AUTHOR CONTRIBUTIONS
A.T.D., J.S., A.Z., H.J.K., S.M.B., G.C., and S.T.C. performed the experiments; L.L., S.T.C., P.V.E., S.K., W.R., and S.M. shared reagents, help, and advice; I. K. performed the bioinformatic analyses; A.T.D., J.S., I.K., S.T.C., S.M., and G.G. designed the research, analyzed the data, and wrote the manuscript.

DECLARATION OF INTERESTS
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Supplemental Information

**NLRC5 promotes transcription of BTN3A1-3 genes and Vγ9Vδ2 T cell-mediated killing**

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Supplemental Information

Figure S1. BTN3A1, A2 and A3 mRNA is not decreased in CIITA-deficient cell lines (related to Figure 1). (A) Murine Btn2a2 promoter sequence and mRNA levels (relative to Polr2a mRNA) as assessed by quantitative RT-PCR (qRT-PCR) in lymph node (LN), spleen, thymus, and bone marrow (BM) of Nlrc5fl/fl and Nlrc5−/− mice. (B) Geometric mean fluorescence intensity (gMFI) of HLA-A/B/C and HLA-DR/DQ/DP as measured by flow cytometry in the indicated blood-derived cell subsets. (C) BTN3A1, A2 and A3, BTN2A1, HLA-B, and HLA-DRA mRNA levels (relative to POLR2A mRNA) were measured by qRT-PCR in B cell-derived cell lines not expressing CIITA (Rj2.2.5; black square, BLS-2; white square), RFX5 (SJO; black circle), or RFXAP (DA; white circle, 6.1.6; white circle with cross) or controls (SP-53; black triangle, Raji; white circle) treated or not with IFNγ for 16 hours. Results depict the mean ± SEM of n=3 mice per genotype and are representative of two independent experiments (A), the mean ± SEM of n=3 individual donors (B), or the mean of n=2 or n=3 cell lines, each represented as the average of n=3 independent measurements (C). (A) Statistical differences between genotypes were calculated using unpaired Student’s t-test, two-tailed, unequal variance; ns: non significant.
Figure S2. Overexpressed CIITA transactivates MHC class II and class I promoters (related to Figure 2). (A) BTN3A1-3 and HLA-B mRNA levels (relative to POLR2A mRNA) were measured by qRT-PCR at basal (t=0 h, untransfected) and 24 h and 48 h following transfection of plasmids encoding the indicated NLR proteins or an empty vector (mock) in HEK293T. Results are depicted as mean ± SD (n=3 technical replicates) and are representative of at least 2 independent experiments. Statistical differences were determined by two-way ANOVA followed by comparison of the experimental conditions to the corresponding mock transfection, and were corrected for multiple testing using the Holm-Sidak method. (B) Luciferase reporter assays were performed in HEK293T cells co-transfected with the parental pGL3 backbone, HLA-DRA, or HLA-B promoter constructs (reporters), and a vector coding for NLRC5, CIITA I, or an empty (mock) vector. Data are expressed as fold transactivation to the mock condition. Results represent mean ± SD of n=3 technical replicates and are representative of at least three independent experiments. Statistical differences were determined by performing a two-way ANOVA followed by comparison of the experimental conditions to the corresponding mock transfection and were corrected for multiple testing using the Dunnett method. *** p<0.001. Only statistically significant differences are illustrated.
Figure S3. NLRC5 mRNA correlates with HLA-B transcript abundance (related to Figure 6). (A) Pairwise correlation of NLRC5 and BTN2A1 and of CIITA and BTN3A1, BTN3A2, BTN3A3, BTN2A1 expression were analyzed using transcriptome datasets of Gambia M. tuberculosis (TB) cohort study (GSE28623). The samples are divided in the groups ‘uninfected’, ‘latently infected’, and ‘active TB’. The table displays the Spearman’s correlation coefficient (R), and Bonferroni adjusted p-values (P adj) for the 3 groups. (B, D-F) Data from The Cancer Genome Atlas (TCGA) provisional dataset collections were analyzed after adjustment for CD45 expression. (B) Spearman’s correlation coefficient (R) for NLRC5 and HLA-B or NECTIN2 mRNA expression across cancer
types (C). Data from GSE10846 lymphoma dataset were analyzed after adjustment for CD45 expression. Scatterplots for NLRC5 and BTN3A1, BTN3A2, BTN3A3, HLA-B, or NECTIN2 mRNA expression are shown. Spearman’s correlation coefficients (R) are indicated and significance was determined using the Bonferroni method. (D) Box plot depicting Spearman’s correlation coefficient (R) distribution for the indicated BTN genes and NLRC5, CIITA, HLA-B, or NECTIN2 mRNA expression across all cancers. Two group comparisons were performed using unpaired t-tests, two-tailed, unequal variance. (E) Spearman’s correlation coefficient for NLRC5 promoter methylation and HLA-B or NECTIN2 mRNA expression across cancer types. (F) HLA-B and NECTIN2 mRNA abundance is plotted according to NLRC5 copy number status. fc: fold change of expression in HDEL (n=14 and n=13 for brca and prad, respectively) over WT group (n=272 and n=361 for brca and prad, respectively). HDEL: homozygous deletion; WT: wild type. Two group comparisons were performed using unpaired t-tests, two-tailed, unequal variance, and p-values are indicated. (G) Luciferase reporter assays were performed in HEK293T cells co-transfected with the parental pGL3 backbone or the indicated BTN3A2 promoter constructs, and a vector coding for NLRC5 or an empty (mock) vector. In the “T>C SX-13bp-CCAAT” construct, the T position was mutated into a C (in red) to resemble the S-box sequence found in the BTN3A3 promoter. Data are expressed as fold transactivation as compared to the mock condition. Results represent mean ± SD of n=4 technical replicates and are representative of 2 independent experiments. Statistical differences were determined by performing a two-way ANOVA followed by comparison of the SX-13bp-CCAAT to the T>C SX-13bp-CCAAT condition and were corrected for multiple testing using the Holm-Sidak method. * p<0.05, **p<0.01, *** p<0.001, ns: not significant. (B, E) Grey bars indicate significant correlation (p<0.05 after Bonferroni correction), white bars non-significant ones.
Figure S4. NLRC5 overexpression increases BTN3A1-3 and HLA-A/B/C expression in Raji cells (related to Figure 7). (A) Polyclonal Raji cells were transduced with NLRC5 or empty vector (mock) and treated with zoledronate (ZOL) for 24 h. HLA-A/B/C surface expression was analyzed by flow cytometry. Graphs depict a quantification of HLA-A/B/C geometric MFI (gMFI; top) and histogram overlays (bottom) show HLA-A/B/C expression for unstained (US; black, grey), mock- (blue) and NLRC5-transduced polyclonal Raji cells (pink). Results represent 5 independent measurements. (B) Subclones were generated from Raji cells transduced with NLRC5 or empty vector (mock) and treated with ZOL for 24 h before BTN3A1-3 and HLA-A/B/C surface expression was analyzed by flow cytometry. Graphs depict a quantification of BTN3A1-3 and HLA-A/B/C gMFI (top) and histogram overlays (bottom) show BTN3A1-3 and HLA-A/B/C expression for unstained (US; black, grey), mock- (blue) and NLRC5-transduced Raji subclones (red, violet). (C) IFNγ production by Vγ9Vδ2 T cells was measured after 48 h of co-culture with the mock - or the two NLRC5-transduced subclones at an effector-to-target ratio of 10:1 in the presence of ZOL. Results are depicted as IFNγ fold induction in presence of NLRC5-transduced as compared to the mock-transduced subclones. (D, E) BTN3A1 was targeted by CRISPR/Cas9 in the NLRC5-transduced Raji subclone 2 (NLRC5 c2 sgBTN3A1). NLRC5 c2 and NLRC5 c2 sgBTN3A1 were assessed for BTN3A1, A2, and A3 mRNA levels (relative to POLR2A mRNA) by qRT-PCR (D) and for their susceptibility to Vγ9Vδ2 T cell-mediated killing after 24 h of co-culture at an effector-to-target ratio of 10:1 in the presence of ZOL (E). Results are depicted as mean ± SEM of n=3 independent measurements (B) as mean ± SEM of n=4 healthy donors (C), as mean ± SD of n=3 technical replicates (D), or as mean ± SEM of n=3 healthy donors (E). Results are representative of at least 2 independent experiments (A-E). Statistical differences between the condition with and without NLRC5 overexpression were calculated using paired Student’s t-test, two-tailed, unequal variance (A), by one-way ANOVA followed by comparison of the experimental conditions to the corresponding mock condition and were corrected for multiple testing using the Dunnett method (B, C), or by unpaired (D) or paired (E) Student’s t-test, two-tailed, unequal variance. *p<0.05, ***p<0.001. Only statistically significant differences are illustrated.
**Transparent Methods**

**Mice**

Sex- and age-matched 6- to 12-week-old \( Nlrc5^{+/+} \) and \( Nlrc5^{-/-} \) (Staehli et al., 2012) mice on a C57BL/6 (H2\(^b\)) background were housed at the animal facility of the University of Lausanne. All animal experimental protocols were approved by the Veterinary office regulations of the State of Vaud, Switzerland, and all methods were performed in accordance with the Swiss guidelines and regulations.

**Cell lines and transfections**

HEK293T cells were cultured using DMEM high glucose supplemented, whereas the B-cell lines SP-53, Raji, Rj2.2.5, BLS-2, SJO, DA, and 6.1.6 (previously described in (Ludigs et al., 2015; Tarantelli et al., 2018)) were cultured in RPMI 1640 medium (Life Technologies) containing at least 10 mM HEPES. Culture media were all supplemented with GlutaMAX (2 mM), sodium pyruvate (1 mM), 10% fetal calf serum (FCS), 100 U/ml penicillin and 100 \( \mu \)g/ml streptomycin. For HEK293T cells, 0.1 mM MEM Non-essential Amino Acids (NEAA) were added in some experiments. Cells were incubated at 37\(^\circ\)C in 5% CO\(_2\). For transfection, HEK293T cells were subconfluently seeded and transfected the following day using PEI reagent (1.5:1 up to 3:1 PEI: DNA ratio) and harvested at the indicated time point for analysis.

**Quantitative RT-PCR and sample preparation**

To assess expression in human immune tissues, cDNA included in the “Human immune system MTC™ panel” (TAKARA) were used (pool from at least 9 individuals). Peripheral blood mononuclear cells (PBMC) were isolated by density centrifugation over a Ficoll-Hypaque
gradient (LymphoPrep) from blood of healthy human donors. Total PBMCs were surface stained with CD4+ T, CD8+ T, NK, monocyte and B cell markers, and sorted by flow cytometry using FACSaria III sorter (BD Biosciences) as CD4+CD3+, CD8+CD3+, CD56+CD3- CD14+, and CD19+ cells, respectively. Immune organs were collected and processed from Nlrc5fl/fl and Nlrc5-/- mice. RNA from cell lines, PBMC subsets, and murine lymphoid tissues was extracted using TRizol® reagent (Ambion, Life Technologies) according to manufacturer's instructions. Annealing with random primers (Life Technologies) was performed at 70 °C for 5 min, followed by retrotranscription to cDNA with M-MLV RT, RNase H(−) point mutant (Promega), in presence of buffer, nucleotides (Roche Diagnostics), and RNasin® Plus RNase Inhibitor (Promega). Reaction was incubated at 40 °C for 10 min, 45 °C for 50 min and 70 °C for 15 min. cDNAs were diluted and quantitative PCR was performed using the LightCycler 480 SYBR Green I Master (Roche Diagnostics) on a LightCycler 480 machine (Roche Diagnostics). Expression was determined relative to POLR2A. Data were analyzed, and transcript abundance (gene/POLR2A) and s.d. were calculated using the LightCycler 480 software. Primers (Fwd; Rev) used were as follows: Polr2a: (5'-CCGGATGAATTGAAGCGGATGT-3'; 5'-CCTGCCCAGGATCCATTAGTC-3'); Btn2a2: (5'-TAGGGGTCTCTCCACACAGC-3'; 5'-TATGACCAGGCAACCATGAA-3'); POLR2A: (5'-CCGACCACATCAAGAGAGTCAGGCCTC-3'; 5'-GTATTTGATGCACCCTCCCTC-3'); BTN3A1-3: (5'-ATGAAAAAGCCCTGTCGGAGG-3'; 5'-TGTTATTTGGGTGTTGGGTGA-3'); HLA-B: (5'-CTACCCCTGCAGGAGATCA-3'; 5'-ACAGCCAGGCGCCAGACA-3'); HLA-DRA: (5'-GCCAACCTGGAAATCATGACA-3'; 5'-AGGGCTTGTGTGAGCACA-3'); BTN2A1: (5'-TGCTCGGCCAGAAGAAAGAA-3'; 5'-CCACATGATGGGCAGGCC-3'); BTN3A1: (5'-CTTCAGCTGCTCATGCTCA-3'; 5'-CAGATCAGCCTTCTCACC-3'); BTN3A2: (5'-
CAGTACTTGA\text{CTCGTGGAGAG}-3'; 5'-TCAGGCTGA\text{TATTGGATCGG-3'}; \text{BTN3A3} : (5'-TCGTGGAGAGAAG\text{TCTTTGG-3'}; 5'-ACATCCGCAGGTTGAAGA-3'); \text{NLRC5} : (5'-GTGCCCT\text{TCGGACCTGGAG-3'}; 5'-GA\text{GATTCCAGGTGCTTTTCC-3'}); \text{CIITA} : (5'-AGCCAAGTCCCTGAAGGATG-3'; 5'-TCTTAAGGTCCCGAACACAG-3').

\textit{Flow Cytometry}

For flow cytometry analysis, sorted PBMCs or cell lines were surface stained using antibodies against HLA-A/B/C (W6/32, eBioscience or BioLegend), HLA-DP/DQ/DR (Tu39, BioLegend) and BTN3A1-3/CD277 (BT3.1, BioLegend or Miltenyi Biotec). Data were acquired using a FACS Canto (Becton Dickinson) or a Gallios (Beckman Coulter) flow cytometer and analyzed with FlowJo software (LCC, Becton Dickinson).

\textit{Plasmids and constructs}

NOD1, NOD2, NLRC3 expression plasmid were obtained from the laboratory of the late J. Tschopp. CD72 expression plasmid was a kind gift from M. Thome-Miazz (UNIL, Switzerland). HLA-B170 (referred as p \textit{HLA-B}) luciferase reporter plasmid was kindly provided by P.J. van den Elsen (Leiden University, Netherlands). HLA-DRA luciferase reporter and CIITA I/III expression plasmids were kindly gifted by W. Reith (UNIGE, Switzerland). For overexpression experiments, a NLRC5 encoding plasmid previously described was used (Staehli et al., 2012). For ChIP assays, a plasmid encoding NLRC5 in frame with a N-terminal HA-tag, referred here as wt NLRC5, has been generated. ORF was amplified using the KAPA Hifi PCR kit (KAPA Biosystems) using NLRC5 containing plasmid as template. PCR primers were extended with KpnI or XhoI restriction sites for oriented cloning into the pCMV-HA backbone.
NLRC5 Walker A domain mutant (K234A), named as mt NLRC5, was generated by site directed point mutagenesis using the wt NLRC5 construct as template. The pHRSIN-CS-Luc-IRES-emGFP plasmid was a kind gift from A. Rodriguez (UAM, Spain). The pHRSIN-CS-IRES-mTagBFP2 was generated by removing the luciferase and substituting the emGFP for the mTagBFP2 in the pHRSIN-CS-Luc-IRES-emGFP plasmid using the Gibson assembly cloning method. The lentiviral construction pHRSIN-CS-HA-NLRC5-IRES-mTagBFP2 coding for NLRC5 was generated by Gibson assembly using the pHRSIN-CS-IRES-mTagBFP2 as recipient plasmid and the HA-NLRC5 insert PCR amplified from the wt NLRC5 plasmid. Lentiviral packaging plasmids pCMVDR8.74 and pMD2.G were a gift from D. Trono (EPFL, Switzerland). Luciferase reporter plasmids were created by replacing the MluI-BglIII fragment spanning the HLA-DRA SXY region in the pDRAprox plasmid (Krawczyk et al., 2004) with BTN3A promoter regions. Inserts were obtained either by PCR amplification using the GoTaq polymerase (Promega) or using annealed purchased DNA oligos (Microsynth AG). The pGL3 plasmid containing the remaining HLA-DRA core promoter (from −60 to +10) in the same reporter plasmid was used as negative control. All generated constructs were verified by sequencing. Sequences of primers used for cloning are available upon request.

**Luciferase reporter assay**

HEK293T cells were seeded into a 96-well plate and transfected the following day using PEI reagent with human NLRC5, or human CIITA I expression vectors, or an empty backbone as control (mock), and the indicated luciferase reporter constructs. The pRLTK (Renilla) luciferase reporter was included for normalization. Cells were harvested between 20 and 30 h post-transfection and cell lysates were analyzed using the Dual-Luciferase® Reporter Assay System
(Promega) following manufacturer’s instruction. Bioluminescence was measured using the Enspire™ Alpha2390 Multilabel Reader (PerkinElmer).

**Chromatin immunoprecipitation (ChIP)**

HEK293T cells were transiently co-transfected with plasmids encoding for HA-tagged human NLRC5 (wt NLRC5) or human NLRC5 Walker A mutant (mt NLRC5) and human CD72. After 48 h, cells were collected and stained with FITC-labeled α-CD72 (3F3; BioLegend) followed by incubation with α-FITC magnetic beads for MACS enrichment (Miltenyi Biotech). Chromatin was prepared from CD72+ cells as previously described (Masternak et al., 2003). Immunoprecipitation was performed using a ChIP grade anti-HA tag antibody (ab9110, Abcam). Analysis of specific DNA regions was performed by quantitative PCR. The amount of immunoprecipitated DNA was calculated from the standard curves generated with the input chromatin and fold enrichment was determined relative to the mt NLRC5 condition. The promoter of HOXC8, which is not a NLRC5 target, is used as negative control. Primers (Fwd; Rev) are listed hereafter: *BTN3A1*: (5’- GGGAGGTAGGGCAGGAATTT -3’; 5’- CACTGAGGAAGGCTGAAATGA-3’); *BTN3A2*: (5’- TGAGAAACATCACCTCTGAGCCA; 5’- CCATGAGAAACAGTAGTGCAGTG-3’); *HLA-B*: (5’- GTGTCGGGTCCTTCTGTCCTTCCA-3’; 5’- CCAATGGGAGTGGGAAGTG-3’); *HOXC8*: (5’- CTCAGGGCTACCAGAGAACC-3’; 5’- TTGGCGAGGAAAAACAGTAGTGCAGTG-3’).

**Generation of Raji target cells for cytotoxicity assays**

To produce lentiviral particles HEK293T cells were co-transfected with the pCMVDR8.74 and pMD2.G packaging vectors, the pHRSIN-CS-Luc-IRES-emGFP plasmid encoding for the
luciferase, and either the pHRSIN-CS-HA-NLRC5-IRES-mTagBFP2 construct expressing NLRC5 or the empty pHRSIN-CS-IRES-mTagBFP2 as control (mock). After transduction, Raji cells double positive for mBFP2 and luciferase expression were isolated using the MoFlo Astrios cell sorter (Beckman Coulter) and were used for the cytotoxicity assays as polyclonal population. To obtain Raji subclones, single-cell subcloning was performed using limiting dilution of the polyclonal Raji cells. In this context, culture medium was supplemented with 2-β-mercaptoethanol (0.05 mM).

CRISPR/Cas9-mediated gene disruption of BTN3A1 was conducted with a combination of two pre-designed chemically stabilized Alt-R CRISPR-Cas9 crRNAs (crRNA XT; IDT) and the Alt-R CRISPR Cas9 system of Integrated DNA Technologies. For duplex formation, the chemically stabilized crRNAs XT Hs.Cas9.BTN3A1.1.AL (5’-ACCAUCAGAAGUUCCCUCC-3’, IDT) and Hs.Cas9.BTN3A1.1.AP (5’-GAUGUGAAGGGUUACAAGGA-3’, IDT) were mixed in equimolar amounts with Alt-R CRISPR-Cas9 tracrRNA (IDT) to a final oligo concentration of 44 µM, heated to 95°C for 5 min and cooled down to RT for gRNA duplex formation (crRNA XT:tracrRNA). Alt-R S.p. Cas9 Nuclease V3 protein (IDT) was used at 36 µM. For ribonucleoprotein (RNP) assembly, the gRNA duplex was mixed with Alt-R S.p. Cas9 Nuclease V3 protein at equal volumes and incubated for 20 min at RT. RNP complexes were then stored on ice prior utilization. Electroporation of Raji cells was conducted with the Neon Transfection System (Invitrogen, Life Technologies, 1 pulse of 1,350 V for 30 ms) and in the presence of Alt-R Cas9 Electroporation Enhancer (10.8 µM; IDT). Cells were then transferred into pre-warmed complete medium and incubated at 37°C and 5% CO2.

*Expansion of human Vγ9Vδ2 T cells and cytotoxicity assays*
Human $\gamma\delta$ T cells were expanded from human PBMCs isolated from healthy donors using a Ficoll-Hypaque gradient. Cells were cultured in RPMI 1640 medium + GlutaMAX (Life Technologies) supplemented with 10% fetal calf serum (FCS), 50 U/ml penicillin, 50 μg/ml streptomycin, 10 mM HEPES buffer, 1 mM sodium pyruvate and 1x MEM non-essential amino acid solution. For $\gamma\delta$ T cell expansion, PBMCs were stimulated either with 2.5 μM zoledronate and 50 U/ml rIL-2 or with 1 μg/ml concanavalin A and 10 ng/ml rIL-2 and 10 ng/ml rIL-4. For zoledronate expanded $\gamma\delta$ T cells, fresh rIL-2 was added every second day over a culture period of 21 days. After 7 days, the expanded $\gamma\delta$ T cells were additionally split at a 1:2 ratio every second day. 14 days after stimulation, the purity of the expanded $\gamma\delta$ T cells was evaluated by flow cytometry and cells were MACS-enriched if required (TCR$\gamma/\delta+$ T Cell Isolation Kit, human, Miltenyi Biotech). For bioluminescence-based killing assays, 10,000 luciferase-expressing NLRC5- or mock-transduced Raji cells were co-cultured with either 10,000 (1:1), 50,000 (5:1) or 100,000 (10:1) expanded $\gamma\delta$ T cells in triplicates for 24 h at 37°C and 5% CO2 in the presence of 10 μM zoledronate. The proportion of living cells was measured as bioluminescence in relative light units (RLU) at an Infinite 200 PRO plate reader (Tecan) in the presence of the substrate D-firefly luciferin potassium salt (37.5 μg/ml, Biosynth). Specific killing was calculated relative to Raji cells treated either with medium alone (spontaneous death) or with 1% Triton X-100 (maximal death) with the following formula: % specific killing = 100 x (average spontaneous death RLU - test RLU) / (average spontaneous death RLU - average maximal death RLU). Quantification of IFN$\gamma$ production was performed by co-culturing 10,000 NLRC5- or mock-transduced Raji subclone cells with 100,000 $\gamma\delta$ T cells (10:1) for 48 h in the presence of 10 μM zoledronate, followed by standard enzyme linked immunosorbent assay (ELISA) from Invitrogen, Life Technologies.
**Bioinformatic analyses of Gambia cohort and tumor datasets**

We selected 32 TCGA cancer types from the TCGA provisional data set collections as available from cBioportal (June 2019). Genetic profiles for mRNA expression, methylation and copy number of selected genes were retrieved using the ‘cgdsr’ R/Bioconductor package. Adjustment for potential confounding factors (here CD45/PTPRC) was performed using the ‘removeBatchEffect’ function from the limma R/Bioconductor package using CD45 expression as covariate. For the independent diffuse large B-cell lymphoma (DLBCL) cohort, we downloaded gene expression data (GSE10846) from the NCBI GEO database using the ‘GEOquery’ R package, and corrected the data for CHOP/RCHOP treatment effects and for CD45 expression. Similarly, for the Gambia cohort, we downloaded the gene expression data (GSE28623) from GEO. For GSE10846 DLBCL, we used the 350 samples with ABC or GCB subtype and batch-corrected for chemotherapy. No batch correction or adjustment for CD45 was deemed to be needed. For continuous variables, statistical correlation and significance was assessed using Spearman’s correlation. For 2 group comparisons, the t-test (two-sided, unequal variance) was used to assess differences in means. Statistical analysis was performed using R version 3.5.2 and BioConductor 3.8 on Ubuntu/Linux. Cancer types are abbreviated according to the TCGA database (https://gdc.cancer.gov/resources-tcga-users/tcga-code-tables/tcga-study-abbreviations): dlbc: Lymphoid Neoplasm Diffuse Large B-cell Lymphoma; chol: Cholangiocarcinoma; thym: Thymoma; tget: Testicular Germ Cell Tumors; thca: Thyroid carcinoma; meso: Mesothelioma; paad: Pancreatic adenocarcinoma; kich: Kidney Chromophobe; lihc: Liver hepatocellular carcinoma; brea: Breast invasive carcinoma; prad: Prostate adenocarcinoma; kirc: Kidney renal clear cell carcinoma; laml: Acute Myeloid Leukemia; cesc:
Cervical squamous cell carcinoma and endocervical adenocarcinoma; lgg: Brain Lower Grade Glioma; ucs: Uterine Carcinosarcoma; gbm: Glioblastoma multiforme; acc: Adrenocortical carcinoma; luad: Lung adenocarcinoma; skcm: Skin Cutaneous Melanoma; ucec: Uterine Corpus Endometrial Carcinoma; kirp: Kidney renal papillary cell carcinoma; lusc: Lung squamous cell carcinoma; pcpg: Pheochromocytoma and Paraganglioma; esca: Esophageal carcinoma; sarc: Sarcoma; uvm: Uveal Melanoma; coad: Colon adenocarcinoma.

Ethics

Mice were treated in accordance with the Swiss Federal Veterinary Office guidelines. For expression analyses, PBMCs were isolated from blood donations obtained from the Blood Transfusion Center, Switzerland. For experiments using $V_\gamma 9V_\delta 2$ T cells, informed consent was obtained from the donors in accordance with the Declaration of Helsinki and Institutional Review Board approval from the University of Freiburg Ethics Committee (412/9). Human cell lines are established cell lines.

Statistical analysis

Statistical analyses were performed using either Prism software (GraphPad version 8.2.0) or R version 3.5.2 and BioConductor 3.8 on Ubuntu/Linux. For 2 group comparisons, t-tests (two-tail, unequal variance) were used to assess differences in means. In case of multiple testing, overall effects were determined by ANOVA and post hoc comparisons to the control condition were performed using either Dunnett’s or Holm-Sidak method. For continuous variables, statistical correlation and significance was assessed using Spearman’s correlation, Bonferroni method was
used to adjust for multiple correlations. Differences were considered significant when $P<0.05$ (*), very significant when $P<0.01$ (**) and highly significant when $P<0.001$ (***).