# C-type lectin receptor DCIR modulates immunity to tuberculosis by sustaining type I interferon signaling in dendritic cells

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Immune response against pathogens is a tightly regulated process that must ensure microbial control while preserving integrity of the infected organs. Tuberculosis (TB) is a paramount example of a chronic infection in which antimicrobial immunity is protective in the vast majority of infected individuals but can become detrimental if not finely tuned. Here, we report that C-type lectin dendritic cell (DC) immunoreceptor (DCIR), a key component in DC homeostasis, is required to modulate lung inflammation and bacterial burden in TB. DCIR is abundantly expressed in pulmonary lesions in Mycobacterium tuberculosis-infected nonhuman primates during both latent and active disease. In mice, we found that DCIR deficiency impairs STAT1-mediated type I IFN signaling in DCs, leading to increased production of IL-12 and increased differentiation of T lymphocytes toward Th1 during infection. As a consequence, DCIR-deficient mice control M. tuberculosis better than WT animals but also develop more inflammation characterized by an increased production of TNF and inducible NOS (iNOS) in the lungs. Altogether, our results reveal a pathway by which a C-type lectin modulates the equilibrium between infection-driven inflammation and pathogen's control through sustaining type I IFN signaling in DCs.

tuberculosis | inflammation | C-type lectin

L-12, IFN $\gamma$ -producing CD4<sup>+</sup> Th1 cells, and TNF are wellrecognized key players in immunity against *Mycobacterium tuberculosis*, and yet their uncontrolled production can result in progressive lung tissue destruction and tuberculosis (TB) disease (1, 2). Several host factors, including IL-10 and Treg cells, have been implicated in immune modulation in TB (2). Nevertheless, the balance between pro- and antiinflammatory reactions that dictates the magnitude of TB disease, from early bacterial clearance or latent infection to severe lung injury, remains incompletely understood (3).

In addition to T cells, dendritic cells (DCs) play a key part in antimycobacterial immunity, mostly through antigen processing and presentation, which together with the production of IL-12, drive the expansion of *M. tuberculosis*-specific Th1 cells (2). DCs express a wide range of so-called pattern-recognition receptors, including Toll-like receptors (TLRs), NOD-like receptors, and C-type lectin receptors (CLRs) (4), which are involved in recognition of exogenous or endogenous ligands, in antigen presentation and T-cell stimulation, as well as in gene expression rewiring and production of inflammatory mediators, such as cytokines and chemokines.

Among these CLRs, DC immunoreceptor (DCIR) (5) belongs to the Dectin 2 family (6). This lectin is mainly expressed by DCs and has a broad specificity for sugars, including mannose and fucose (7), of endogenous and exogenous origins (8, 9). DCIR is a rare case of a CLR that contains an immunoreceptor tyrosine-based inhibitory motif (ITIM) in its cytoplasmic tail and, as such, is generally believed to mediate inhibitory signals in DCs. In line with this concept, antibody-mediated DCIR triggering in human DCs inhibits TLR-mediated production of IL-1 $\beta$ , IL-6, TNF, IL-12, and IFN $\alpha$  (10–12), and mice deficient in the DCIR homolog Dcir1 (also known as Clec4a2) show an overexuberant expansion of DCs and develop more agingassociated or experimentally induced antibody- and T cellmediated autoimmune disorders than their WT counterpart (13–15). Intriguingly, the mechanisms responsible for these phenotypes still remain poorly understood.

# Significance

Tuberculosis (TB) is an immunopathology, mostly of the lung, due to an overexuberant immune response to the bacterial pathogen *Mycobacterium tuberculosis*. Here, we demonstrate in vitro and in vivo that dendritic cell (DC) immunoreceptor (DCIR), a C-type lectin receptor expressed by DCs, modulates immunity to TB by sustaining type I IFN signaling in DCs. These findings were generalized beyond TB, in a model of in vivo antigenpresentation assay unrelated to *M. tuberculosis*, suggesting that they may extend to other pathologies, such as viral infections or autoimmune disorders. Thus, modulating DCIR activity may help to develop type I IFN-targeting therapies for a large repertoire of inflammatory disorders, including TB.



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Here, using a combination of transcriptomic analysis and biochemical assays, we found that DCIR sustains type I IFN signaling by ensuring proper STAT1 phosphorylation in DCs. Infection of DCIR-KO mice along with in vitro assays revealed that DCIR impairs antimycobacterial Th1 response through type I IFNmediated down-modulation of IL-12. These findings were validated beyond TB, using an in vivo antigen-presentation assay based on the adoptive transfer of peptide-loaded DCs from WT and KO mice. In the case of TB, increased IL-12 production in DCIR-KO animals translates into reduced bacterial burden in the lungs of infected animals along with immunopathology. Considering that type I IFNs are generally believed to play a detrimental role during TB (16), our findings are in agreement that these cytokines lead toward susceptibility to M. tuberculosis infection, as evidenced by the lung bacterial burden in DCIR-KO animals. Nonetheless, we also show that DCIR, through supporting type I IFN signaling, may play a key role in the protection of host tissue integrity during a pertinent inflammatory infectious context.

# Results

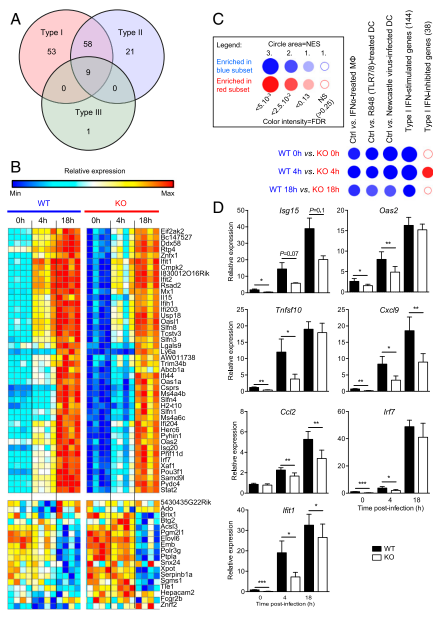
DCIR Expression Enables Sustained Type I IFN Signaling in DCs. Because DCIR polymorphisms were associated with several noninfectious inflammatory disorders in humans (17-19), we asked whether DCIR could play a part in modulating immunity in the context of a chronic inflammation of infectious nature, namely TB. We first found that DCIR is expressed at the periphery of lung granulomas in M. tuberculosis-infected nonhuman primates (NHPs), a model that closely mimics TB in humans, during both asymptomatic infection and active TB disease (SI Appendix, Fig. S1). To functionally explore the role of DCIR in antimycobacterial immunity, we sought to exploit a previously reported mouse model of DCIR (Dcir1/Clec4a2) deficiency (13, 20). Because DCIR is primarily expressed in DCs (5), in particular those from lungs and in bone marrow-derived DCs (SI Appendix, Fig. S2 A and B), we first evaluated the impact of this CLR in a global and unbiased manner through genomewide transcriptome analysis of bone marrow-derived DCs from WT and DCIR-KO animals before and during M. tuberculosis infection. Functional gene set enrichment analysis (GSEA) and interrogation of the Interferome database (21) revealed that a significant fraction of genes, which were down-regulated in DCIR-KO DCs compared with WT cells, was dependent on type I IFN and/or included genes involved in defense to viruses (SI Appendix, Fig. S3 and Fig. 1 A and B). Surprisingly, IFNstimulated genes (ISGs) modulated in KO DCs were already detected before infection [i.e., in resting cells (Fig. 1 C and D]. This finding strongly suggested that DCIR recognizes host endogenous ligand(s), which was already proposed in other contexts (20). In line with this possibility, we did not find evidence of any binding of DCIR to either whole mycobacterial cells (SI Appendix, Fig. S4A) or purified mycobacterial surface glycosylated components (SI Appendix, Fig. S4B), inferring that DCIR-mediated signaling likely relies on recognition of one or more, yet to be identified, endogenous ligand(s).

Sustained STAT1 Phosphorylation Requires DCIR Expression in DCs. We next explored if DCIR deficiency resulted in the impairment of production and/or of signaling of type I IFN. First, blocking ligand recognition by the type I IFN receptor subunit IFN $\alpha/\beta$  receptor 1 (IFNAR1) using the monoclonal antibody MAR1-5A3 resulted in a complete shutdown of ISG expression (Fig. 24), confirming that induction of these genes in infected DCs was solely due to type I IFN signaling. Second, the expression of mycobacterial receptors, type I IFN production or IFNAR expression in *M. tuberculosis*-infected DCs was unaltered in DCIR deficient DC (Fig. 2*B*–*D* and *SI Appendix*, Fig. S5). Third, further analysis of components of the JAK–STAT pathway, which mediates type I IFN signaling, revealed that JAK1 phosphorylation

was unaffected in DCIR-deficient DCs, compared with in WT cells (Fig. 2E). By contrast, even though there was no difference in protein levels, we found that STAT1 phosphorylation at Ser727 was diminished in KO DCs (Fig. 2F), implying a role for DCIR in regulating the activity of this transcription factor. For this reason, we analyzed the phosphorylation status of Src homology 2 domain tyrosine phosphatase 1 (SHP1) and 2 (SHP2), known to be involved in dephosphorylation of STATs (22, 23). We found that SHP2 phosphorylation was increased in DCIR-deficient cells by Western blotting (Fig. 2G) and by ELISA (Fig. 2H). This result suggested that the increase in the phosphorylated, active form of SHP2 might be responsible for STAT1 dephosphorylation. Noteworthy is that *M. tuberculosis* is known to stimulate TLR2, and our results could be recapitulated in DCs costimulated with a TLR2 agonist and IFN $\beta$  (Fig. 2 *E*, *F*, and *H*). In this condition, we found that neither STAT1 phosphorylation at Tyr701 (SI Appendix, Fig. S6A) nor SHP1 phosphorylation (SI Appendix, Fig. S6B) was modified in DCIR-KO cells compared with WT cells. Lastly, treatment of the cells with the recently reported highly specific SHP2 inhibitor GS-493 (24) resulted in restored STAT1 phosphorylation in DCIR-KO cells (Fig. 21), thus demonstrating that impaired STAT1 phosphorylation in DCIR-KO DCs was due to SHP2. Altogether, these findings are consistent with a scenario whereby DCIR deficiency leads to an increased phosphorylation of SHP2, which in turn decreases STAT1 phosphorylation and consequently type I IFN signaling. Not unexpectedly, because STAT1 and SHP2 are also involved in type II IFNy signaling, we found that the overall response of DCIR-KO DCs to IFN $\!\gamma$  was also altered, compared with their WT counterparts (SI Appendix, Fig. S6C).

### DCIR Deficiency Results in Enhanced Antimycobacterial Th1 Immunity.

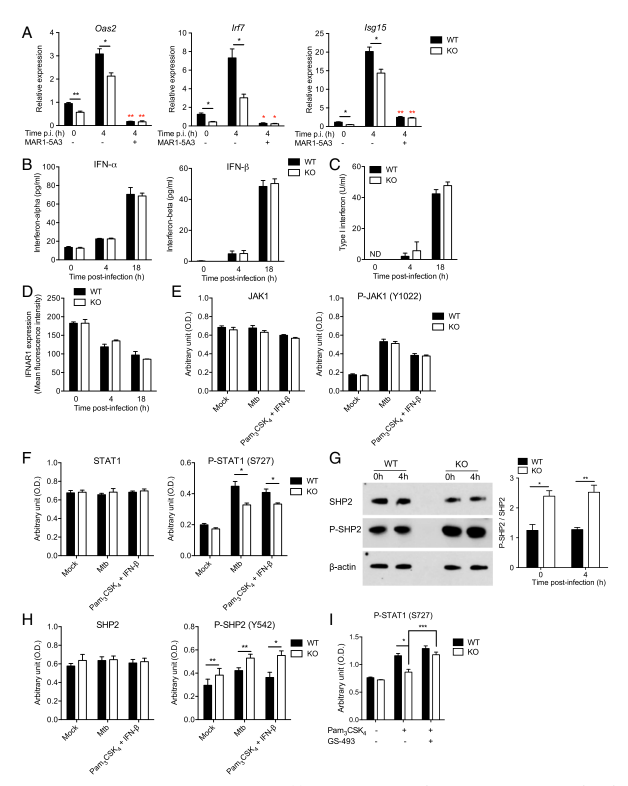
To explore the role of DCIR in immunity to *M. tuberculosis* in vivo, we infected DCIR-KO mice and found that they harbored significantly less bacteria in their lungs and spleen than WT animals (Fig. 3A). Indeed, this finding was associated with an increased cell infiltration in the lungs of KO mice (Fig. 3B), in particular with an increased influx of CD4<sup>+</sup> and CD8<sup>+</sup> T cells, B cells, macrophages, and DCs (Fig. 3 B and C and SI Appendix, Fig. S7). Th1 cells are critical in immunity to TB (2). Analysis of lung T cells from M. tuberculosis-infected animals revealed an increased proportion of IFNy-producing Th1 cells in the lungs of DCIR-KO mice, compared with WT animals (Fig. 4 A and B). Moreover, the overall production of IFNy was increased in the lungs of KO mice (Fig. 4C). Ex vivo restimulation of lung cells with purified protein derivative (PPD), a mixture of mycobacterial antigens, indicated that at least a fraction of these Th1 cells were *M. tuberculosis*-specific because they overproduced IFN $\gamma$ in a PPD-specific manner (Fig. 4D). In line with these results, given the role of the master regulator T-bet in IFNy production and Th1 lineage commitment (25), we confirmed an increase in CD3<sup>+</sup>T-bet<sup>+</sup> Th1 cells in pulmonary cell infiltrates of infected DCIR-KO mice (Fig. 4E). Importantly, we found that T cells from WT or KO mice can be equally activated by polyclonal stimuli (SI Appendix, Fig. S8), which support the notion that the observed phenotype relies on differential T stimulation by WT vs. DCIR-KO DCs. One of the functions of Th1 lymphocytes is to drive macrophage polarization toward a proinflammatory, antimicrobial M1-like profile (26). Gene expression analysis in the lungs of infected KO mice revealed a decreased expression of the M2 markers Arg1, Ym1, and Mrc1 and an increased expression of the M1 markers Nos2 and Tnf, compared with WT mice (Fig. 4F). Finally, IL-12-p70, which drives Th1 differentiation, was also produced more abundantly in the lungs of infected KO mice (Fig. 4G). Collectively, these findings indicated that inflammation and antimycobacterial Th1 response are augmented in DCIR-deficient animal.



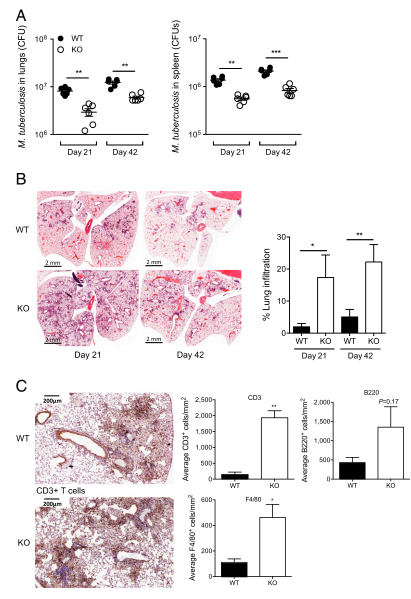
**Fig. 1.** DCIR expression results in sustained expression of ISGs in murine DCs. (*A*) Venn diagram of ISGs down-modulated in DCIR-deficient DCs, compared with in WT cells, at 0 or 4 h after *M. tuberculosis* infection. Data were generated using the Interferome online interface (21). (*B*) BubbleMap analysis, a high-throughput extension of GSEA, on the pairwise comparisons of DCs from WT vs. DCIR-deficient mice at different time points after *M. tuberculosis* infection. The gene sets shown, coming from independent experiments, include direct ISGs either up- or down-regulated in conventional DC subsets and genes regulated by type I IFN, by a TLR7/8 ligand known to induce type I IFN, or by a viral infection in macrophages or DCs. (C) Heat map of the genes belonging to the ISG gene set and found down-regulated in DCs from DCIR-KO compared with WT mice (*Top*) or belonging to the IFN-inhibited genes gene set and found up-regulated in DCs from DCIR-KO compared with WT mice (*Bottom*) upon *M. tuberculosis* infection. For each gene, relative gene expression values across DCs from WT and DCIR-KO mice infected with *M. tuberculosis* at 0, 4, and 18 h is shown and varies from low (blue) to high (red). (*D*) Expression profiles of selected genes from the gene set of down-modulated genes in DCIR-deficient DCs and other ISGs. Data are presented as means  $\pm$  SEM of four biological replicates. Statistical analysis was performed using Student's *t* test. \**P* < 0.05; \*\**P* < 0.01.

In the TB context, Th1 priming occurs in the lung-draining, mediastinal lymph nodes (LNs). Similar to the lungs, an increase in Th1 cells, IFN $\gamma$ , and IL-12 profiles was observed in the mediastinal LNs from KO mice compared with those from WT mice (*SI Appendix*, Fig. S9 *A–D*). Of note, IL-12–p70 overproduction in DCIR-deficient DCs was recapitulated in DCs stimulated with a TLR2 agonist (*SI Appendix*, Fig. S9*E*). Type I IFNs are known to inhibit IL-12–p70 in myeloid cells, including DCs (27), and in particular in *M. tuberculosis*-infected monocytes and macrophages (28, 29). Accordingly, stimulation by IFN $\beta$  resulted in a strong suppression of IL-12–p70 production by TLR2-activated

DCs (*SI Appendix*, Fig. S9*F*). Unlike that recently reported in *M. tuberculosis*-infected macrophages (29), IL-10 was not involved in IFN $\beta$ /DCIR-mediated IL-12 suppression because the expression of this cytokine was not modulated in DCIR-deficient DCs compared with WT cells (*SI Appendix*, Fig. S9*G*). Strikingly, treatment of TLR2-stimulated DCs with the SHP2 inhibitor GS-493 reduced IL-12–p70 production by DCIR-KO cells to WT levels (*SI Appendix*, Fig. S9*H*). All things considered, these findings indicated that overproduction of IL-12–p70, and consequently increased Th1 priming, by DCIR-KO DCs during *M. tuberculosis* infection in vivo is likely due to an impairment of



**Fig. 2.** DCIR expression leads to sustained type I IFN signaling in DCs. (*A*) Bone marrow-derived DCs from WT or KO mice were either left uninfected or infected (4 h) with *M. tuberculosis* H37Rv in the presence or absence of the IFNAR1-blocking monoclonal antibody MAR1-5A3, as indicated. Relative expression of the ISGs *Oas2, Irf7*, and *Isg15* was quantified by RT-qPCR. Red asterisks correspond to comparison analysis between MAR1-5A3–treated and –untreated cells. (*B–D*) WT or KO DCs were either left uninfected (0 h) or infected for 4 or 18 h with *M. tuberculosis*. IFN $\alpha$  and IFN $\beta$  (*B*) were quantified in the cell supernatants by ELISA or using the type I IFN-activity reporter cell line B16 Blue IFN $\alpha/\beta$  (*C*). The expression of IFNAR1 was quantified by FACS (*D*). (*E–I*) WT or KO DCs were either untreated or were treated with *M. tuberculosis* for 4 h, or were costimulated (Mock), were infected or were treated with the SHP2 inhibitor GS-493 (24). JAK1 (*E*), STAT1 (*F* and *I*), SHP-2 (*G* and *H*), and their phosphorylated forms were quantified in cell lysates by Western blot (WB) (G) or ELISA (*E*, *F*, *H*, and *I*). *G*, *Right* presents quantification of three independent WB analyses (*G*, *Left*) by densitometry. In *A–I*, data are presented as means ± SEM of at least three biological replicates and are representative of at least two independent experiments. Statistical analysis was performed using Student's *t* test. \**P* < 0.05, \*\**P* < 0.01.



**Fig. 3.** DCIR expression results in impaired bacterial clearance and controls pathology in *M. tuberculosis*-infected mice. (*A*–*C*) WT or KO mice were infected intranasally with ~1,000 cfu of *M. tuberculosis* H37Rv. Lungs and spleen were collected after 21 or 42 d of infection and analyzed for colony-forming unit (CFU) content (*A*), cell infiltration (*B*), and immune cell subsets at 42 d (*C*). In *A*, each symbol represents colony-forming units from one animal. Data are representative of three independent experiments. In *B* and *C*, data are presented as means  $\pm$  SEM of at least 25 fields chosen from at least 3 independent tissue slices and are representative of 2 independent experiments. Statistical analysis was performed using Student's *t* test. \**P* < 0.05, \*\**P* < 0.01. (*C, Top Left, Bottom Left, and Top Center*) CD3<sup>+</sup> T cells. (*C, Bottom Center and Right*) Quantification of other cell types (F4/80-expressing macrophages and B220-expressing B lymphocytes). Pictures are displayed in *SI Appendix,* Fig. S7*A*.

type I IFN signaling, which was also observed in the LNs of infected animals (*SI Appendix*, Fig. S91).

**DCIR Expression Results in Impaired Th1 Priming by DCs.** To generalize our findings beyond TB, we performed an in vivo antigen presentation assay based on adoptive transfer, as described previously (30). Briefly, OT-II antigen receptor transgenic T cells were purified and injected i.v. into C57BL/6J recipients. Two days later, mice were challenged in the hind footpads with ovalbumin (OVA) peptide-pulsed (or unpulsed) WT or DCIR-KO DCs. After 5 d of priming, popliteal LN cells were harvested from recipients, and cytokine secretion was assessed. In line with our findings in *M. tuberculosis*-infected mice, the OVA-specific Th1 response tended to increase in the recipients animals challenged with DCIR-KO DCs (Fig. 5A). More markedly, the production of IFN<sub>Y</sub> and IL-12–

p70 was strongly increased in animals challenged with DCIR-KO DCs, compared with WT DCs (Fig. 5 B and C), which correlated with a decreased expression of several ISGs (Fig. 5D).

# Discussion

In this study, we show that DCIR modulates immunity to *M. tuberculosis* through sustaining type I IFN signaling in DCs, which reduces IL-12–p70 production and Th1 expansion. As a consequence, DCIR-deficient mice control *M. tuberculosis* infection better than their WT counterparts but also develop more immunopathology in the lungs. These findings have several consequences for our understanding of DCIR signaling in immunity and of that of type I IFN in TB.

Because DCIR contains an ITIM in its cytoplasmic tail, it is thought to act as a negative regulator of immune cell signaling

+aCD3/CD28 Unstimulated В Day 42 Day 21 wт WT 1.8% 3.4% 10 WT 30-% IFN<sup>+</sup>CD4<sup>+</sup>T cells 🗆 ко % IFNy<sup>+</sup>CD4<sup>+</sup>T cells 8. 20 6 IFN-V-4 KO KO 10 1.4% 8.8% 10 2 C 0 . Mock +αCD3/CD28 Mock +αCD3/CD28 Day 21 Day 42 500 D WT 50-400-🗆 ко \*\*\* 400 40 WT IFN-y (pg/ml) IFN-y (pg/ml) 300 🗆 ко 300-30 200-200 20 100-100 10 ND Λ 0 n PPD Day 21 Day 42 PPD Unstimulated Unstimulated 80-% CD3<sup>+</sup>Tbet<sup>+</sup> cells 60 WT KO 40 Merge Merge 20 0 ŵт ĸо G 1,500-Mrc1 4-Arg1 5. Ym1 5 \* WT P=0.1 🗖 ко Relative expression IL-12p70 (pg/ml) 000 000 4 4 3. 3 3 2 2 0-0 . Day 42 Day 21 Day 21 Day 42 Tnf 8. Nos2 2.0-Relative expression 6 1.5 WT 4 1.0 🗖 ко =0.0.7 2 0.5

**Fig. 4.** DCIR expression leads to impaired Th1 immunity and M1-like macrophage polarization in *M. tuberculosis*-infected mice. Mice were infected as in Fig. 3. (*A* and *B*) IFN<sub>7</sub>-producing CD4<sup>+</sup> Th1 cells were quantified by intracellular cytokine FACS staining after either stimulation with a mixture of anti-CD3 and -CD28 antibodies or in the absence of stimulation (Mock), as indicated. A representative experiment among three at day 21 is depicted in *A. B* represents means  $\pm$  SEM of six biological replicates. (*C* and *D*) ELISA quantification of total IFN<sub>7</sub> in lung lysates of infected mice (*C*) or after ex vivo restimulation of total lung cells with PPD (*D*). ND, not detected. (*E*) Immunohistological analysis of expression of CD3 and the Th1-driving transcription factor T-bet in lungs of infected WT or KO mice. *E, Right* presents quantification of 404 and 313 WT or KO CD3<sup>+</sup> cells, respectively, and data are representative of *n* = 2 mice. Arrows show the presence of T-bet in the cell nuclei. (Scale bars: 10 µm.) (*F*) RT-qPCR quantification of 1L-12–p70 in lung lysates of infected mice. In *B–D, F,* and *G*, data are presented as means  $\pm$  SEM of at least three replicates and are representative of at least two independent experiments. Statistical analysis was performed using Student's *t* test. \**P* < 0.05; \*\**P* < 0.001; \*\*\*\**P* < 0.0001.

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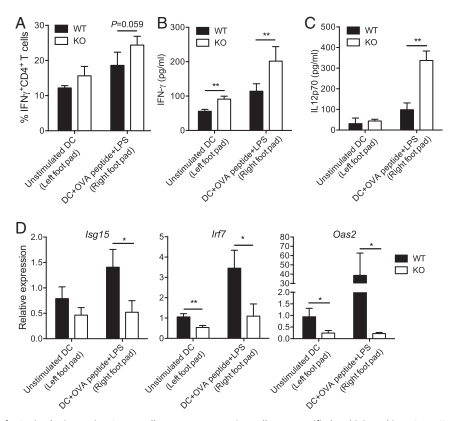
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**Fig. 5.** In vivo function of DCIR in Th1 immunity. OT-II T-cell receptor transgenic T cells were purified and injected into C57BL/6J recipients. Two days later, mice were challenged in the footpads with OVA peptide-pulsed or unpulsed WT or DCIR-deficient DCs. After 5 d of priming, popliteal LN cells were harvested from recipients. (*A*) Percentage of IFN $\gamma$ -producing CD4<sup>+</sup> OVA-specific (OT-II) Th1 cells adoptively transferred into C57BL/6J recipients and primed in vivo with peptide-pulsed or unpulsed DCs. (*B* and C) ELISA quantification of IFN $\gamma$  (*B*) and IL-12–p70 (*C*) in LN lysates from recipient mice. (*D*) RT-qPCR quantification of gene expression of the ISGs *Isg15, Irf7,* and *Oas2* in LN lysates from recipient mice. Data are presented as means ± SEM of five biological replicates. Statistical analysis was performed using Student's *t* test. \**P* < 0.05; \*\**P* < 0.01.

(4). However, our data clearly show that DCIR deficiency results in impaired type I IFN signaling in DCs, suggesting that this receptor activates, rather than inhibits, the IFNAR-associated JAK-STAT pathway. Based on these findings, we propose that one physiological role of DCIR is to sustain type I IFN signaling in DCs through interactions with self-glycosylated ligands that remain yet to be identified. The DCIR ITIM was shown to bind to unphosphorylated, but not phosphorylated, SHP2 (31). DCIR may therefore function as a molecular sink for unphosphorylated, inactive SHP2, thus limiting SHP2's capacity to deactivate STAT1. Our findings are reminiscent of several membrane receptors in which the ITIM activates, rather than inhibits, various signaling pathways (32). How ITIM-containing DCIR delivers an activating, rather than inhibitory, signal will need to be further dissected on the molecular level. In particular, whether DCIR delivers a signal on its own through its ITIM motif or requires a coreceptor will need to be addressed.

Another critical aspect worth highlighting is the notion that the levels of type I IFN produced during *M. tuberculosis* infection have important consequences for the host. Our results support the notion that type I IFNs are detrimental in TB (2) if one considers the lung bacterial burden as the main sign of TB disease. Nevertheless, we also show that DCIR, through sustaining type I IFN signaling, modulates lung immunopathology, which is a hallmark of TB in human, and therefore also contributes to protection. Further work is needed to understand the many roles of type I IFNs in TB, as suggested by a recent study showing that these cytokines can be protective or detrimental depending on the stage of *M. tuberculosis* infection (33) and on the *M. tuberculosis* strain (34). Interestingly, previous studies reported that DCIR-KO animals tend to develop

more autoimmune  $CD4^+$  T cell-associated (Th1- or Th17-associated) pathologies (14, 15). Based on our findings, it is likely that these phenotypes rely on an increased IL-12 production through impaired type I IFN signaling, which was not investigated in these cases.

In conclusion, our study identifies DCIR as a key molecular player not only in immunity to TB but more generally in tuning the balance between type I and type II IFN responses, which might be relevant in the context of a number of inflammatory diseases of noninfectious (e.g., autoimmune) or infectious (e.g., viral) nature. Identifying DCIR ligands with various pharmacological properties (agonist or antagonist) might help in developing type I IFN-targeting therapies for a range of inflammatory disorders, beyond TB.

# **Materials and Methods**

Ethics Statements for NHP Samples. The NHP study protocol was in compliance with the European Commission (EC) Directive 86/609/EEC, approved by the Biomedical Primate Research Centre ethical committee before the start of the study, and executed under Dutch law regulating animal experiments (Agreement DEC 579). For any particular animal, the endpoint was defined either by signs of severe disease (humane endpoint criteria, referring to animal condition by adverse body weight development, respiratory capacity, and animal behavior) or by protocol, which limited the follow-up time to 1 y postinfection.

**NHP Handling and Vaccination.** NHP materials were derived from animals that were used for vaccine research and development purposes. Specifically, healthy young adult female rhesus macaques (*Macaca mulatta*), all captive-bred for research purposes and of homogeneous breeding background, were vaccinated (or not) with a standard human dose of *M. bovis* Bacille Calmette et Guérin (bacillus Calmette–Guérin) Danish 1331 (Statens Serum

Institut) by intradermal injection. Nine months later, the animals were challenged with 500 cfu of *M. tuberculosis* strain Erdman K01 (prepared and provided under an agreement between the World Health Organization and Food and Drug Administration Center for Biologics Evaluation and Research with the assistance of Aeras), which was administered by intrabronchial instillation under sedation. At the endpoint, animals were sedated, euthanized, and submitted to macroscopic lung pathology scoring as described previously (35). Gross pathological findings were assessed and described by an experienced veterinary pathologist while blinded for treatment. Representative lung samples were collected and fixed in 10% neutral buffered formalin and embedded in paraffin for long-term storage. A blind analysis was performed to quantify immunohistochemical stainings of fixed samples.

Mice, Cells, and Bacteria. Dcir1(Clec4a2)<sup>-/-</sup> mice (20) were provided by the NIH-sponsored Mutant Mouse Regional Resource Center (MMRRC). The KO mouse strain was originally generated by the Consortium for Functional Glycomics (CFG) (http://www.functionalglycomics.org/static/consortium/resources/ DataCoreFdcir.shtml). The strain was then further backcrossed for more than 10 generations on a C57BL/6 background. Animals used in this study consisted in Dcir1(Clec4a2)<sup>+/+</sup> and Dcir1(Clec4a2)<sup>-/-</sup> mice originating from common Dcir1(Clec4a2)<sup>-/+</sup> breeding pairs. Details for genotyping are available on the CFG website. All mouse experiments were performed in animal facilities that met all legal requirements in France and by gualified personnel in such a way to minimize discomfort for the animals. All procedures, including mouse studies, were conducted in strict accordance with French laws and regulations, in compliance with the EC Directive 68/609/EEC guidelines and its implementation in France. All protocols were reviewed and approved by the Regional Ethical Committee (reference MP/ 04/26/07/03). All efforts were made to minimize suffering. The study did not involve humane endpoints. KO and WT (C57BL/6J) mice were housed and bred in a specific pathogen-free environment at the animal facility of the Transgenèse et archivage d'animaux modèle facility, Orléans, France. Animal care and experimentation were consistent with the French guidelines and were approved by the Ministry of Higher Education and Research (Agreement APAFIS 1269). In particular, all efforts were made to minimize animals discomfort and suffering.

Bone marrow-derived DCs were prepared from murine hind legs by flushing the uncovered and opened femurs and tibiae with Dulbecco's PBS (DPBS) (Gibco). Red blood cells were lysed by addition of red blood cell lysis buffer (Sigma) for 2 min. Subsequently, cells were separated with a cell strainer and cultivated in 6- or 24-well plates (Thermo Scientific) at  $2 \times 10^6$  or  $5 \times 10^5$  cells per well, respectively, in complete Roswell Park Memorial Institute (RPMI) 1640 medium (Gibco) supplemented with 10% (vol/vol) FCS (PAN-Biotech), 1% sodium pyruvate (Gibco), 0.1%  $\beta$ -mercaptoethanol (Gibco), and 20 ng/mL recombinant murine GM-CSF (Peprotech). Cells were allowed to differentiate for 5–7 d. The culture medium was replaced every 2–3 d, and the cells were used for stimulation studies between day 7 and day 9 after the start of differentiation. The complete DC culture medium was checked for LPS contamination using the HEK-Blue TLR4 reporter cell line (Invivogen) according to manufacturer's instructions, and was found free of LPS (*SI Appendix*, Fig. S10).

*M. tuberculosis* (H37Rv strain) was grown in Middlebrook 7H9 culture medium (Difco) supplemented with 10% albumin–dextrose–catalase (ADC) (Difco), 0.05% Tween-80 (Sigma), or on Middlebrook 7H11 agar medium (Difco) supplemented with 10% oleic acid–ADC (Difco).

**Mouse Infection and Cell Stimulation.** Six- to 10-wk-old female C57BL/6J or  $Dcir1^{-/-}$  mice were anesthetized in gas chambers containing isoflurane. Mice were infected intranasally with 1,000–1,500 cfu of *M. tuberculosis* in 25 µL of DPBS (Gibco). Mice were killed by cervical dislocation after 21 or 42 d. Lungs and spleens were harvested, homogenized, and plated onto 7H11 agar for colony counts. Bone marrow-derived DCs were stimulated at 37 °C for various periods of time with *M. tuberculosis* at a multiplicity of infection of three bacteria per cell or 1 µg/mL PAM<sub>3</sub>CSK<sub>4</sub> (Invivogen), 1,000 U/mL recombinant murine IFN $\beta$  (Biolegend), with or without prior incubation with 1 µg/mL of an IFNAR1 blocking antibody (clone MAR1-5A3; Biolegend) for 2 h at 37 °C or with the SHP2 inhibitor GS-493 (24) at 25 µM for 20 h at 37 °C.

**Microarray Analysis.** RNA from lungs, mediastinal, or popliteal LNs or from mouse bone marrow-derived DCs were extracted using the RNeasy mini kit (Qiagen). The amount and purity of RNA were quantified using a NanoDrop ND-1000 apparatus (Thermo Scientific) by measuring absorbance at 260/280 nm. Double-stranded cDNA was reverse-transcribed using the M-MLV Reverse Trancriptase kit (Invitrogen), according to the manufacturer's protocol. For microarray analysis, cDNA was labeled with Cy3–dCTP using the One-Color

DNA labeling kit (Roche NimbleGen). Gene expression was analyzed using SurePrint G3 Mouse gene expression 8\*60K microarray kit (Agilent). Hybridization was performed using 2  $\mu$ g Cy3–cDNA and the hybridization kit (Roche NimbleGen). Samples were incubated for 5 min at 65 °C and 5 min at 42 °C before loading for 17 h at 42 °C according to the manufacturer's protocol. Microarrays were washed and scanned using an MS200 microarray scanner (Roche NimbleGen). Microarrays were washed and scanned using an MS200 microarray scanner (Roche NimbleGen).

Agilent raw files extracted by using Feature Extraction software were processed through Bioconductor (version 3.1) in the R statistical environment (version 3.2.0) via the limma package (36). Raw expression values were background corrected in a "normexp" fashion and then quantile normalized (37). Quality of the hybridizations and evaluation of the sampling method were assessed by density plots, boxplots, principal component analyses, and hierarchical clusterings using the ade4 and cluster packages (38). Normalized expression values were then corrected for batch effects due to the cell sorts at different days, using the ComBat algorithm (39) through the sva package (40) and declaring two batches (two sorting days) and the genotype and postinfection time as covariates.

The ComBat-corrected expression values were then used to extract, via the limma package, the genes differentially expressed (false discovery rate, <0.05) between DCs from WT and  $Dcir1^{-/-}$  mice at each time point after *M. tuberculosis* infection. The JVenn plugin (41) was used to draw Venn diagrams showing the overlaps between lists of up- or down-regulated genes in the DCs from  $Dcir1^{-/-}$  compared with WT mice.

To assess the statistical enrichment of gene sets of interest in the pairwise comparisons between DCs from WT and  $Dcir1^{-/-}$  mice at each time point after *M. tuberculosis* infection, we used BubbleMap (42), a high-throughput extension of GSEA methodology (43). BubbleMap analysis was performed with 1,000 gene set-based permutations, and with "difference of classes" as a metric for ranking the genes because the data were expressed in log2 scale. The results are displayed as a BubbleMap, where each bubble is a GSEA result and summarizes the information from the corresponding enrichment plot. The color of the Bubble corresponds to the condition from the pairwise comparison in which the gene set is enriched. The bubble area is proportional to the GSEA normalized enrichment score. The intensity of the color corresponds to the statistical significance of the enrichment, derived by computing the multiple testing-adjusted permutation-based *P* value using the Benjamini–Yekutieli correction. Enrichments with a statistical significance above 0.05 are represented by empty circles.

Generation of the Type I ISG and IFN-Inhibited Gene Sets. To generate a list of genes directly activated or inhibited by type I IFN, we used data of one of our previous studies (44) where we compared the transcriptomic profiles of WT vs. *Ifnar1<sup>-/-</sup>* DC subsets coming from mixed bone marrow chimera mice, lethally irradiated and reconstituted with a mixture of CD45.1 WT and CD45.2 IFNAR<sup>-/-</sup> bone marrow, infected with murine cytomegalovirus. More specifically, we extracted, using the GeneSign module of the BubbleGUM software (42), the genes that were up- or down- regulated at least twofold in *Ifnar1<sup>-/-</sup>* compared with WT CD8 $\alpha^+$  or CD11b<sup>+</sup> conventional DCs (cDCs) from infected mice. We then selected the genes commonly found up- or down-regulated in both cDC subsets to generate the type I IFN inhibited or stimulated genes, respectively (*SI Appendix*, Table S1). We then added hundreds of public gene sets downloaded from MSigDB (45) to assess statistical enrichments of our gene sets of interest through BubbleMap (42).

The heat map was performed with Gene-E (https://www.broadinstitute. org/cancer/software/GENE-E), using the ComBat-corrected expression values of the genes belonging to the ISG or IFN-inhibited gene (IIG) gene sets that were found regulated in DCs from DCIR-KO mice upon *M. tuberculosis* infection. Specifically, using GSEA, we extracted the leading edges of the ISG gene set enriched in WT DCs at the three time points, selected the genes found in the overlap of the three leading edges to display their expression profiles in the heat map. For the IIGs, we focused on the leading edge of the IIG gene set enriched in DCIR-KO Dcs at 4 h after *M. tuberculosis* infection.

**Real-Time qPCR.** For real-time qPCR (RT-qPCR), specific PCR primers (*SI Appendix*, Table S2) were designed using QuantPrime. Real-time qPCR was performed with gene targeted primers using qPCR Mastermix plus SYBR Green (Eurogenetec), according to the manufacturer's protocol. All real-time qPCR reactions were carried out using a 7500 Real-Time PCR System and data were analyzed using the 7500 Software version 2.0.6 (Applied Biosystems). PCR array data were calculated by the comparative cycle threshold method, normalized with hypoxanthine-guanine phosphoribosyltransferase house-keeping gene, and expressed as mean fold change in experimental samples relative to control samples.

**ELISA and ISG Reporter Cell Line.** After stimulation, supernatants from lungs, from mediastinal or popliteal LNs, or from bone marrow-derived DCs were harvested, and IFN $\alpha$  and IFN $\beta$  (VeriKine ELISA kit), IL-12–p70, IFN $\gamma$ , and IL-10 (OptEIA kit; BD Biosciences) were detected by ELISA according to the manufacturer's instructions.

For IFN activity measurement, supernatants from DCs were added in sixwell plates containing the B16 Blue IFN $\alpha$ /b reporter cell line (Invivogen) for 24 h at 37 °C. This reporter cell line expresses the SEAP reporter gene under the control of the IFN $\alpha$ / $\beta$ -inducible ISG54 promoter enhanced by a multimeric ISRE. Levels of SEAP were monitored using the detection medium QUANTI-Blue (Invivogen) according to the manufacturer's instructions. For phosphorylation detection, DCs were lysed after stimulation in buffer containing phosphatase and protease inhibitors (Thermo Fisher). JAK1, phospho-JAK1, STAT1, phospho-STAT1 (Ser727), and phospho-STAT1 (Tyr701) were detected by ELISA (Raybiotech) according to the manufacturer's instructions. Colorimetric cell-based assays were performed for detection of SHP1, phospho-SHP1 (Tyr536), SHP2, and phospho-SHP2 (Tyr542) (AssaybioTech) according to the manufacturer's instructions.

Western Blotting. Cells in six-well plates were detached with trypsin-EDTA (Invitrogen), centrifuged and then lysed by adding 100  $\mu$ L of boiling 2× Laemmli buffer containing phosphatase inhibitors (2 mM orthovanadate, 5 mM NaF) onto cell pellets for 5 min; 20 µg of protein was subjected to electrophoresis in 10% SDS/PAGE gels and transferred onto a nitrocellulose membrane. Membranes were saturated with 3% BSA in TBS-T [50 mM Tris (pH 7.2), 150 mM NaCl, and 0.1% Tween 20) for 30 min and incubated with primary antibodies overnight at 4 °C. Antibodies used were: anti-SHP2 (clone M163; Abcam), anti-PY<sup>542</sup>-SHP2 [clone EP508(2)Y; Abcam], and anti- $\beta$  actin (clone 20-33; Sigma-Aldrich). Then, primary antibodies were revealed using a HRP-coupled secondary anti-mouse (Sigma) or anti-rabbit (Cell Signaling Technology) antibody for 1 h. Finally, HRP activity was revealed using an electrochemiluminescence kit (GE Healthcare Life Sciences) according to the manufacturer's instructions. All blots were normalized against actin expression and protein quantity was set to one in the control for quantification.

Binding Experiments. All binding experiments were performed in lectin buffer (50 mM Hepes, 5 mM MgCl<sub>2</sub>, 5 mM CaCl<sub>2</sub>) containing 0.5 mg/mL BSA (Sigma). Nonspecific binding to M. tuberculosis was prevented by incubating the bacteria for 30 min at room temperature in lectin buffer containing 2 mg/mL BSA. Bacteria were then incubated with 10  $\mu$ g/mL chimeric Dcir1-human (h)lgG1 fragment crystallizable (Fc) (20) at 37 °C for 1 h in the presence or absence of 20 mM EDTA (Euromedex). After washing in lectin buffer, bacteria were incubated with a PE-coupled polyclonal anti-hlgG (Fcy-specific; eBiosciences) at 5 µg/mL for 1 h at room temperature. Bacteria were then washed and fixed for 2 h at room temperature in PBS containing 4% paraformaldehyde (Polyscience) and analyzed by flow cytometry. For ligand binding experiments, different constituent of the mycomembrane, namely phosphatidylinositol dimannosides, arabinosylated lipoarabinomannan, mannosylated lipoarabinomannan, lipomannan, arabinogalactan, and trehalose dimycolate were coated onto 96well plates (Nunc Immuno Plates Maxisorp; Sigma) in water:ethanol (1:1, vol/vol) at different concentrations. The plates were dried and incubated with lectin buffer containing 2 mg/mL BSA at room temperature for 2 h (saturation step). The plates were then incubated with 10  $\mu$ g/mL of chimeric DCIR-hFc at room temperature during 2 h. After washing, binding was detected using a biotinylated polyclonal anti-human IgG (Fcy-specific; eBioscience) at 5 µg/mL for 1 h at room temperature, and streptavidin-HRP at 100 ng/mL for 30 min at room temperature. After washing, substrate solution (TMB substrate reagent set; BD Biosciences) was added for 30 min at room temperature and the reaction was stopped with H<sub>2</sub>SO<sub>4</sub>. Colorimetric analysis was performed at 450-570 nm using a spectrophotometer.

FACS Analysis. Mouse DCs or primary cells obtained from homogenized lungs or mediastinal or popliteal LNs were stained with fluorescein-conjugated monoclonal antibodies (1:100) directed against mouse Clec4a2 (320507; LS-Bio), CD11b (M1/70; eBiosciences), CD11c (N418; Biolegend), TLR-2 (6C2; eBiosciences), SIGNR1 (22D1; eBiosciences), Dectin 1 (bg1fpj; eBiosciences) CD86 (GL1; eBiosciences), I-A/I-E (M5/114.15.2; Biolegend), PD-L1 (10F.9G2; Biolegend), IFNAR1 (MAR1-5A3; eBiosciences), CD3 (17A2; Biolegend), CD4 (RM4-5; BD Biosciences), or F4/80 (BM8; Biolegend). For intracellular cell staining, cells from homogenized lung or mediastinal LN were stimulated during 4 h at 37 °C with a mix of purified anti-mouse CD3 (17A2; Biolegend) and CD28 (37.51; Biolegend) at 5 µg/mL or with PPD at 10 µg/mL. Cells were incubated in GolgiPlug (BD Biosciences) overnight. Subsequently, cells were resuspended in Fixation-Permeabilization solution (BD Cytofix/Cytoperm kit;

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BD Biosciences) and intracellular cytokine staining was carried out according to the manufacturer's protocol. Cells were stained with fluorescein-conjugated monoclonal antibodies (1:100) directed against mouse CD4 (RM4-5; BD Biosciences) and IFN $\gamma$  (XMG1.2; BD Biosciences). For mortality assessment, cells were stained with Annexin V (BD Biosciences) according to the manufacturer's instruction.

Histological Analysis. Paraffin-embedded tissue samples were sectioned and stained with hematoxylin and eosin for histomorphological analysis. Histopathological scoring of TB lesions in NHPs (SI Appendix, Table S3) was determined using a worksheet in which TB disease from lung biopsies was described (35). Histopathological scoring of TB lesions in WT and DCIR-KO mice was determined by measuring area of infiltrated lung tissue using the Panoramic viewer software (3DHISTECH). Immunohistochemical staining was performed on paraffin-embedded tissue sections, using polyclonal and monoclonal primary antibodies summarized in SI Appendix, Table S4. Immunostaining of paraffin sections was preceded by different antigen unmasking methods. After incubation with primary antibodies, sections were incubated with biotin-conjugated polyclonal anti-mouse, anti-rat or anti-rabbit Ig antibodies followed by the streptavidin-biotin-peroxidase complex (ABC) method (Vector Laboratories) and then were counter stained with hematoxylin. Slides were scanned with the Panoramic 250 Flash II (3DHISTECH). Virtual slides were automatically quantified for B and T lymphocytes, and macrophages distributions as described previously (46). For confocal microscopy, samples were stained with primary antibodies as described above and followed by antimouse IgG isotype-specific, anti-rat or anti-rabbit IgG antibodies labeled with Alexa 488 and Alexa555 (Molecular Probes). Samples were mounted with Prolong Antifade reagent (Molecular Probes) and examined using a 60×/1.4 NA objective of an Olympus FV1000 confocal microscope.

In Vivo Antigen-Presentation Assays. The assay was based on adoptive transfer, as described previously (30). OT-II T cells were purified from the spleen of C57BL/6J OT-II mice [a kind gift from S. Guerder, Center for Physiopathology of Toulouse–Purpan (CPTP), Toulouse, France] using mouse anti-CD4 microbeads (L3T4) positive selection and MACS separation columns (Miltenyi Biotec), according to the manufacturer's instructions. Purified T cells were injected i.v. into C57BL/6J mouse recipients. Two days later, OVA peptide (ISQAHAAHAEINEAGR)-pulsed-bone marrow DCs from C57BL/6J or Dcir1<sup>-/-</sup> mice were washed at least three times in RPMI 1640 and administered in DPBS at a dose of  $3.10^5$  cells in a volume of 25–40  $\mu$ L into the fore or hind footpads. The antigen-pulsed DCs were administered on one side, and the contralateral footpad served as the control. The control footpads were injected with DCs that had not been antigen-pulsed. At day 5, the draining popliteal LNs were removed and teased into a cell suspension. RNA was extracted to performed RT-gPCR as described in the experimental procedures of the manuscript. In parallel, cells were cultured and stimulated with antigens for 4 h in prewarmed RPMI-1640 supplemented with 10% FCS, 1% sodium pyruvate and 0.1% β-mercaptoethanol. Supernatants were collected to perform ELISA analysis and intracellular cell staining was performed.

**Statistics.** Statistical analyses were performed using GraphPad Prism (GraphPad Software). Data are presented as means  $\pm$  SEM of at least three independent experiments; *P* values (paired or unpaired Student's *t* test) are relative to the indicated control. Statistical significance was assumed when *P* < 0.05.

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