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Notch signaling in group 3 innate lymphoid cells modulates their plasticity


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Abstract

The Notch signaling pathway is conserved through evolution and controls processes including cell fate determination, differentiation and proliferation. It influences fate choice during hematopoiesis. Its role is well documented for the development and differentiation of lymphocytes in contrast to innate lymphoid cell (ILC) subsets. ILC are lymphoid cells without rearranged receptors that fulfill effector and regulatory functions in innate immunity and tissue remodeling. Type 3 innate lymphoid cells (ILC3s) reinforce the epithelial barrier and maintain the homeostasis with the intestinal microbiota. Here, we demonstrated that the population of natural cytotoxic receptor–positive (NCR\(^+\)) (ILC3s) in mice is composed of two subsets with distinct developmental requirements. A major subset depended on the activation of Notch2 in NCR\(^-\) ILC3 precursors in the lamina propria of the small intestine to stimulate expression of Tbx21, Rorc and AhR, respectively encoding T-bet, ROR\(\gamma\)t and AHR transcription factors. Notch signaling contributed to the transition of NCR\(^-\) cells into NCR\(^+\) cells in a cell-autonomous manner. In the absence of Notch signaling, NCR\(^-\) ILC3s lost the gene expression profile of NCR\(^+\) ILC3s but adopted a transcriptional profile uniquely resembling that of NCR\(^+\) ILC3s proving the absence NCR\(^+\) precursors. A second subset of NCR\(^+\) ILC3s did not depend on Notch for their development or for the increased expression of Tbx21, Rorc, and AhR; however, their production of cytokines and their cell surface abundance of the natural cytotoxic receptors were decreased. Together, our data suggest that Notch is a regulator of the plasticity of ILC3s by controlling NCR\(^+\) cell fate.
**Introduction**

Innate lymphoid cells (ILCs) promote inflammatory responses and tissue homeostasis through the rapid secretion of effector cytokines at mucosal barriers and secondary lymphoid organs. These cells differentiate from a common lymphoid progenitor (CLP); however, unlike T cells or B cells, they are devoid of rearranged antigen-specific receptors. All ILCs require the transcriptional repressor Id2 for their development (1-3). ILCs can be subdivided into three groups, which follow different developmental pathways and produce different cytokines (4). The transcriptional profiles of distinct ILC subsets in different tissues were analyzed in depth as part of the immunological genome project (5).

Group 1 ILCs consist of Eomes-positive conventional natural killer (cNK) cells and Eomes-negative ILC1s, secrete the inflammatory cytokine interferon-γ (IFN-γ), and require the transcription factor T-bet for proper development. ILC2s produce the cytokines interleukin-5 (IL-5) and IL-13 and are characterized by the expression of the transcription factors GATA-3 and Retinoic acid receptor related Orphan Receptor alpha (RORα). The ILC3 group is heterogeneously composed of the fetal subset of lymphoid tissue–inducer (LTi) cells (6) and the adult ILC3 subsets, which are mainly found at mucosal surfaces. All ILC3s depend on the transcription factor RORγt (which is encoded by Rorc) for their development and function, and they produce IL-17 and IL-22 (7). In mice, adult ILC3s can be subdivided based on the cell surface expression of the natural cytotoxic receptor 1, Natural Killer protein 46 (NKp46); mostly found present on the surface of NK cells. The NKp46⁺ (NCR⁺) subset is characterized by the cell-surface expression of the CC chemokine receptor CCR6 and produces both IL-22 and IFN-γ. The NKp46⁻ (NCR⁻) subset is composed of various sub-fractions that differently express
combinations of CD4 and CCR6 on the cell surface. Adult NKp46⁺ CCR6⁺ cells consist of both CD4⁺ and CD4⁻ cells, which share several features with LTi cells and are also called LTi-like cells. The lineage relationship between adult ILC3 subsets and the molecular mechanisms that drive their development and effector functions have been under intense investigation (8-15). It was proposed that CCR6⁻ and CCR6⁺ ILC3s arise from distinct progenitors (9, 12, 16, 17). Within the CCR6⁻ population, T-bet–dependent ILC3s were identified as a source of NCR⁺ cell precursors (11-13). ILC3s exhibit plasticity in vitro and in vivo, and environmental cues, such as microbiota and IL-7, stabilize these cells by maintaining RORγt abundance (9). Plasticity phenomenon is recognized during T cell differentiation where the phenotypic and functional characteristics of a given population are dependent on the milieu in which they operate. Several studies support the idea that Notch signaling is important for the generation of the NCR⁺ ILC3 subset (10, 16, 18). It was also proposed that expression of the genes encoding T-bet and AhR by the NCR⁻ ILC3 subset is Notch-dependent (12, 13); however, the potential role of the Notch pathway in adult ILC3s has not been directly analyzed in vivo (10-13, 19).

The Notch pathway is involved in several developmental processes in different tissues. Upon receptor-ligand interaction, the Notch intracellular domain (NICD) translocates into the nucleus where it acts as a transcriptional co-factor together with the recombining binding protein suppressor of hairless Kappa (RBP-Jκ). In T lymphocytes, the canonical Notch signaling pathway drives the expression of different target genes such as Hes1, Tcf7 and Dtx1, respectively encoding the transcription factor HES1, TCF1 and the DTX1 protein, all important for T cell development. Notch1 is especially implicated in various stages of T cell development by
repressing B cell fate, whereas Notch2 signaling is essential for the generation of marginal zone B cells and for the development of CD11b⁺ dendritic cells (DCs) in the spleen and intestine (20).

Here, we analyzed the distribution of adult ILC3 subsets in the lamina propria of mice deficient in Notch signaling through knockout of the genes encoding either Notch2 (Il7r<sup>Crepo</sup>No<sub>otch2<sup>flo/flo</sup></sub> mice) or RBP-Jk (Il7r<sup>Crepo</sup>Rbp<sub>j<sup>flo/flo</sup></sub> mice) under control of the IL7r promoter to ensure deletion at an early stage of lymphoid cell development or of mice that exhibit enhanced Notch signaling through expression of the constitutively active Notch intracellular domain (NICD) under the control of the same promoter (Il7r<sup>Crepo</sup> NICD mice). We showed that the Notch pathway was essential for the generation of NCR⁺ ILC3s. Competitive bone marrow reconstitution experiments indicated that the action of the Notch pathway was direct and cell-intrinsic. Mice that exhibited constitutively active Notch signaling in the earliest lymphoid progenitors showed an increased number not only of NCR⁺ cells, but also of NCR⁻ ILC3s, which is compatible with a role for Notch in the differentiation of NCR⁺ cells from an NCR⁻ ILC3 subset. In contrast, mice that exhibited constitutively active Notch signaling at the later developmental NKp46⁺ stage retained NCR⁺ ILC3s that were indistinguishable from those of their littermate controls. We found that Notch acted on NCR⁻ precursor cells by inducing expression of the genes encoding the transcription factors T-bet, AhR, Gata3, thus enabling their differentiation into NCR⁺ cells. A Notch-independent subset of NCR⁺ ILC3s (around 20% of total NCR⁺ ILC3s) was also identified, and these cells expressed Tbx21 and Tcf7 independently of the Notch signaling pathway.
Results

Notch signaling regulates intestinal NCR$^+$ ILC3s

To assess roles for the Notch pathway in ILC3 homeostasis, we generated $IIL7\text{Cre}^{+/+} Rbpj^{0/\text{fl}}$ mice, which have defective canonical Notch signaling in all lymphoid cells because of their lack of the transcriptional co-activator RBP-Jk. We quantified the $Rbpj$ floxed alleles and determined that less than 1% of IL-7R$^+$ CLPs retained one $Rbpj$ copy (fig. S1, A and B). In contrast, the $IIL7\text{Cre}^{+/+}$ deletion in lymphoid –primed multipotent progenitors (LMPP) was restricted to a few $Rbpj$ alleles. The Notch signaling pathway is essential for T cell development, which was blocked at the transitional stage between double-negative 1 (DN1) and DN2 cells (fig. S1, C and D) in $IIL7\text{Cre}^{+/+} Rbpj^{0/\text{fl}}$ mice; however, a few progenitors (less than 1%) escaped deletion and generated detectable peripheral T cells (fig. S1, C to F). When $Rbpj$ was deleted at the hematopoietic stem cell stage in $\text{Vav}^{\text{Cre}^{+/+}} Rbpj^{0/\text{fl}}$ mice, T cells were almost undetectable in the peripheral organs (fig. S1G).

We next assessed the effect of loss of Notch signaling on the composition of ILC subsets in the lamina propria of adult mice. cNK cells and ILC1s were defined by flow cytometric analysis as NK1.1$^+$NKp46$^+$IL-7R$^-\alpha^-$ and IL-7R$^+\alpha^+$ cells, respectively (Fig. 1A), whereas NCR$^+$ and NCR$^-$ ILC3s were defined as RORγ$^+\text{IL-7R}\alpha^+\text{NK1.1}^+\text{NKp46}^+$ cells and RORγ$^+\text{IL-7R}\alpha^+\text{NK1.1}^-\text{NKp46}^-$ cells, respectively, (Fig. 1A). ILC1s and cNK cells represented less than 2% of the total number of lin$^-$ intestinal lamina propria cells, and their percentages and absolute numbers were similar in $Rbpj$-deficient and $Rbpj$-sufficient mice. We concluded that ILC1s were not affected, in terms of their total numbers or proportions, by the loss of Notch signaling (Fig. 1B).
We found that there was a 5- to 10-fold decrease in the number and percentage of NCR$^+$ ILC3s in Rbpj-deficient mice (0.8 x 10$^4$ cells, representing 1% of lamina propria lineage negative (Lin$^-$) cells) compared to those in Rbpj-sufficient or control Il7r$^{+/+}$ mice (~4 x 10$^4$ cells, representing 8%) (Fig. 1C). In contrast, the NCR$^-$ ILC3 subset was apparently unaffected by loss of Rbpj (~9 to 10 x 10$^4$ cells, representing 15 to 20%) (Fig. 1D), which suggests that neither the development nor the maintenance of this subset required Notch signaling in vivo. The phenotypic effect of Sca1 increase and Thy1 decrease caused by the loss of Notch signaling is common to most ILC subsets with the exception of cNK cells (Fig. 1, D and E). The increase of Ly49CIFH marker, exclusively found in cNK, is compatible with the idea that NK cell precursors branch off from a common differentiation pathway before the other ILC lineages (16, 17). In deficient Notch signaling ILC3s, Thy1 quantities were decreased whereas only NCR$^+$ ILC3s showed a substantial increase in the abundance of cKit, a receptor tyrosine kinase that respond to stem cell factor for proliferation and differentiation of hematopoietic precursors (Fig. 1F). An important molecular sensor detecting “induced self” on cells in danger, the active receptor NKG2D, was substantially less abundant in Rbpj-deficient cells; however, the difference in abundance was statistically significantly different only for the ILC1 and NCR$^+$ ILC3 subsets (Fig. 1, E and F). A comparison of the mean fluorescence intensity (MFI) of NKp46 showed that it was decreased in Rbpj-deficient NCR$^-$ ILC3s (fig. S1H). Moreover, NCR$^+$ ILC3s that were NKG2D$^+$ had substantially more NKp46 than did the NKG2D$^-$ subset (fig. S1I). The increased amounts of Sca1 were statistically significant in both the NCR$^+$ and NCR$^-$ ILC3 subsets from the Rbpj-deficient mice (Fig. 1F).
We previously reported that Notch2 is found early in ILC development (19). In contrast to T cell development, which is mainly dependent on Notch1, ILC3 differentiation could thus be Notch2-dependent. We generated Il7r<sup>Cre/+</sup>Notch2<sup>fl/fl</sup> mice and found that the number of NCR<sup>+</sup> ILC3s was markedly reduced compared to those in Il7r<sup>Cre/+</sup>Notch2<sup>fl/+</sup> mice (fig. S2A). Contrary to our findings with Il7r<sup>Cre/+</sup>Rbpj<sup>fl/+</sup> mice, the decrease in the number of NCR<sup>+</sup> ILC3s was statistically significantly different in the Il7r<sup>Cre/+</sup>Notch2<sup>fl/+</sup> mice (four times less than the number of cells in Il7r<sup>Cre/+</sup>Rbpj<sup>fl/+</sup> mice), suggesting that Notch2 haplodeficiency, which results in the reduced cell-surface abundance of Notch2, was sufficient to reduce the absolute numbers of these ILCs. The absence of both Notch2 alleles (in Il7r<sup>Cre/+</sup>Notch2<sup>ff</sup> mice) resulted in there being similar numbers of NCR<sup>+</sup> and NCR<sup>-</sup> cells to those found in the Il7r<sup>Cre/+</sup>Rbpj mice. Similar to what was observed in the Il7r<sup>Cre/+</sup>Rbpj<sup>ff</sup> mice, no changes were found in the numbers of NCR<sup>-</sup> ILC3s or in the percentages or numbers of ILC1s after Notch2 deletion (fig. S2A).

A residual population of NCR<sup>+</sup> ILC3s was still detectable in both Il7r<sup>Cre/+</sup>Notch2<sup>ff</sup> mice and Il7r<sup>Cre/+</sup>Rbpj<sup>ff</sup> mice, which might reflect incomplete Cre-mediated deletion in lymphoid progenitors. We therefore analyzed ILC3s in Vav<sup>Cre/+Rbpj</sup> mice in which peripheral T cells are undetectable, indicating complete inactivation of Rbpj. A small population of NCR<sup>+</sup> ILC3s was also present in these mice, which is suggestive of a Notch-independent pathway for the generation of this subset (fig. S2B). Overall, the intestinal ILC subsets in Vav<sup>Cre/+Rbpj</sup> mice mirrored those found in Il7r<sup>Cre/+Rbpj</sup> mice, with reduced numbers of NCR<sup>+</sup> cells, unchanged numbers of ILC1 and cNK subsets, and a small decrease in the number of NCR<sup>-</sup> ILC3s (Fig. 1B and fig. S2B). An increase in the abundance of Sca1 was maintained in the ILC1 and ILC3 subsets in Vav<sup>Cre/+Rbpj</sup> mice (fig. S2C); however, an increase in the abundance of Thy1 was
observed only for the ILC1 subset (fig. S2C). Together, these results suggest that Notch signaling through Notch2 is essential for the development of appropriate numbers of NCR⁺ cells, whereas a minor subset of NCR⁺ ILC3s were derived through a Notch-independent pathway.

Loss of Notch signaling changes the secretive capacities of intestinal ILC3s

We next assessed whether loss of the Notch signaling pathway affected cytokine production by ILC3s and ILC1s. The abundances of mRNAs for Il17a, Il17f, Il22, Csf2, and Ifng were measured by quantitative RT-PCR analysis of mRNA extracted from ex vivo unstimulated sorted ILCs from the lamina propria (Fig. 2A). ILC1s did not have any Th17 cytokine family transcripts, whereas the abundance of Ifng mRNA was not affected by the loss of Notch signaling. NCR⁺ and NCR⁻ ILC3s from the Il7rCre/+Rbpjfl/fl mice had similar amounts of Il22 and Il17a mRNAs to NCR⁺ and NCR⁻ ILC3s from control mice (Fig. 2A). The deficiency in Rbpj resulted in increased amounts of Csf2 mRNA in NCR⁺ ILC3s and decreased amounts in NCR⁻ ILC3s, whereas Il17f mRNA was more abundant in the NCR⁻ ILC3s from Il7rCre/+Rbpjfl/fl mice than in the NCR⁻ ILC3s from control mice (Fig. 2A).

Under steady-state conditions, we observed differences for Il17f and Csf2 transcripts, that are both significantly increased in NCR⁺ subset (Fig. 2A). IFNγ secretion was tested after Phorbol Myristate Acetate (PMA) activation, whereas the amounts of IL-17A and IL-22 proteins produced were then tested by treating intestinal ILCs with IL-23. No differences in cytokine production were observed in ILC1s from either strain of mice (Fig. 2B), whereas Rbpj-deficient NCR⁺ ILC3s produced substantially more IL-22 than did NCR⁺ ILC3s from their littermate controls. NCR⁻ ILC3s were the major producers of IL-17A irrespective of their genotype (Fig. 2,
B and C). When we used a combination of IL-23 and PMA to activate intestinal lamina propria ILC3s (fig. S3A), we found that the NCR\(^-\) ILC3 subset of the Rbpj-deficient mice produced more IL-22 than did the Rbpj-haploinsufficient mice. The abundance of Il1r1 mRNA was substantially decreased in NCR\(^-\) subsets compared to that in NCR\(^+\) subsets (fig. S3B). In conclusion, these data suggest that IL-22 production from ILC3s is increased by loss of the Notch signaling pathway.

**Notch signals are cell-intrinsic during the differentiation of NCR\(^-\) cells into NCR\(^+\) ILC3s**

To determine whether the decrease in the number of cells in the NCR\(^+\) ILC3 subset of IL7r\(^{Cre/+}\)Rbpj\(^{fl/fl}\) mice was a result of a direct or an indirect effect of the loss of Notch signaling, we analyzed competitively mixed bone marrow chimeric mice. The extent of reconstitution of ILCs and T cells were similar in both types of chimeras (Fig. 3). Wild-type competitor cells are more efficient to reconstitute than both IL7r\(^{Cre/+}\)Rbpj\(^{fl/+}\) and IL7r\(^{Cre/+}\)Rbpj\(^{fl/fl}\) donor cells since they are coming from mice deficient in one allele of IL7Ra (Fig. 3, A to D). This could be explained by the better proliferation and survival of both wild-type progenitor and mature cells that depend on IL-7R signaling. Hence, the reconstitution experiments were analyzed by first determining the overall percentages of the cells of interest and then by evaluating the ratio of reconstituted cells between donor cells (CD45.1\(^+\)CD45.2\(^-\)) and competitor cells (CD45.1\(^-\)CD45.2\(^+\)). Similar to what was found in the Rbpj\(^{fl/fl}\) mice, intestinal ILC3 subsets in the chimeric mice were differentially affected by the loss of Notch signaling (Fig. 3, A and B). Rbpj-deficient or Rbpj-sufficient hematopoietic progenitors made similar contributions to the CD4\(^+\)CCR6\(^-\) and CD4\(^-\)CCR6\(^+\) cells within the NCR\(^-\) ILC3 population; however, the size of the NCR\(^+\) ILC3 population was statistically significantly reduced among the lymphoid cells derived
from Rbpj-deficient progenitors as compared to that of Rbpj-sufficient progenitors (Fig. 3B). As expected, Rbpj-deficient progenitors were unable to reconstitute T cells (Fig. 3, C and D). Together, these results suggest that the mechanism for the decrease in the size of the NCR⁺ population after inactivation of the Notch signaling pathway is cell-autonomous.

**Intestinal NCR⁺ ILC3s are derived from NCR⁻ precursors in a Notch signaling–dependent manner**

Intestinal NCR⁻ ILC3s contain precursors of NCR⁺ ILC3s (11-13, 19). The NCR⁻ ILC3 subset is heterogeneous and can be subdivided based on the cell-surface expression of CD4 and CCR6. The CCR6⁺ fraction is composed of CD4⁺ and CD4⁻ cells, whereas the CCR6⁻ fraction is diverse and contains precursors that are potentially dependent on the Notch signaling pathway (12, 13). We purified all three intestinal ILC3 subsets from Il7rCre⁺Rbpjflo/⁺ mice and Il7rCre⁺Rbpjflo/⁻ mice (NCR⁺, CCR6⁻NCR⁻, and CD4⁺CCR6⁻NCR⁻) and analyzed the role of Notch signaling in their survival and differentiation (Fig. 4, A and B). The cell-surface abundance of IL-7Rα was reduced on a fraction of CCR6⁻ NCR⁻ cells isolated from the Il7rCre⁺Rbpjflo/⁻ mice (Fig. 4A). The cells were then cultured for 4 days on control OP9 stromal cells or on OP9 cells that were transduced to express the Notch ligand Delta-like 4 (OP9-Dl4 cells) (Fig. 4, C and D). The CD4⁺CCR6⁻NCR⁻ ILC3 subset was stable and insensitive to Notch signaling. Substantial differences were observed in the maintenance of NCR⁺ cells under both conditions. Less NCR⁺ cells were maintained either in the absence of Notch ligands (that is, when cultured on OP9 cells) or in cells from Il7rCre⁺Rbpjflo/⁻ mice compared to their respective controls (Fig. 4D). Conversely, we found more frequently NCR⁻ cells after culture of sorted NCR⁺ cells on OP9 cells (Fig. 4D), which suggested a role for Notch signaling in the maintenance of NKp46, but not RORγt or IL-
7Rα, in ILCs. When CCR6−NCR− cells were cultured, most of them (~60%) maintained their phenotype (Fig. 4D). Moreover, NCR+ cells obtained from Notch signaling–competent precursors were 1.5 times more abundant in the presence of the Notch ligand than in its absence (Fig. 4D). The few NCR+ cells that were obtained from Notch signaling–deficient precursors were insensitive to the presence of Notch ligands (Fig. 4D).

We then analyzed the abundances in the NCR− and NCR+ subsets of various transcripts that are direct targets of the Notch pathway or are important for the ILC developmental program. In contrast to Tbx21, the genes Dtx1, Hes1, and Tcf7 could be considered to be Notch-inducible during ILC development. We found that mRNAs for these three genes were more abundant in the NCR− cells than in the NCR+ cells, and were substantially decreased in the absence of Notch signaling, which suggests that the Notch pathway is active at the NCR− stage (Fig. 4E). In contrast, the abundances of Ild2, IL7r, and Gata3 mRNAs were not statistically significantly different between the NCR− and NCR+ cells in the context of loss of Notch signaling (Fig. 4E and fig. S4). Tbx21 mRNA was more abundant in NCR+ cells than in the NCR− ILC3s (Fig. 4E). In both cell populations, Tbx21 mRNA was almost undetectable in the absence of Notch signaling. We also performed intracellular staining of ILC1s, NCR− cells, and NCR+ cells (Fig. 4F). Whereas T-bet was similarly low in abundance in all ILC3 subsets, confirming its essential role in the generation of NCR+ cells, RORγt protein was substantially reduced in cells from the Il7rCre+/Rbpjfl/fl mice (Fig. 4G). These results suggest that two distinct pools of NCR− ILC3s differentiate in the lamina propria through Rbpj-dependent and Rbpj-independent pathways, and that the Notch signaling pathway operates in NCR− ILC3s by increasing not only the abundance of RORγt, but also the expression of Tcf7 and Tbx21.
Notch signaling directly stimulates the development of NCR⁻ ILC3 precursors into NCR⁺ ILC3s

To complement our analyses of the role of the Notch signaling pathway on ILC3s, we generated mice with constitutive expression of the Notch intracellular domain (NICD) in lymphoid cells (Il7r<sup>Cre/+</sup> NICD mice) (Fig. 5A). The percentages and total numbers of CCR6⁻NCR⁻ cells and NCR⁺ ILC3s were statistically significantly increased in Il7r<sup>Cre/+</sup> NICD mice compared to those in the control mice, whereas the number of CCR6⁺NCR⁻ ILC3s remained unchanged (Fig. 5B). These results indicated that the Notch pathway was active in the CCR6⁻NCR⁻ cells. There was no increase in the mRNAs of pro-apoptotic genes in the absence of Notch signaling (fig. S5). Thus, we concluded that the absence of a large fraction of NCR⁺ ILC3s in Notch signaling–deficient mice was not a result of increased apoptosis, but rather to a low rate of differentiation. We did not detect the decreased expression of any gene encoding pro-survival factors in the absence of Notch signaling (fig. S5). On the contrary, the only difference detected in the absence of Notch signaling was the increase of the pro-survival gene Bcl2 in NCR⁻ precursors (fig. S5). Hence, the increased numbers of ILC3s in the IL-7r<sup>Cre</sup>-Rosa26<sup>NICD/+</sup> mice with a constitutive activation of the Notch signaling pathway was likely not because of the increased survival of these cells.

We further tested whether Notch signaling also affected developing NCR⁺ ILC3s by analyzing ILCs in the lamina propria of Ncr1<sup>Cre/+</sup> NICD mice, that constitutively express the active form of Notch in all cells expressing Ncr1 (Nkp46). We found that all three ILC3 subsets were unchanged in percentages and numbers (Fig. 5, C and D). All subsets of ILC3s also had similar
amounts of RORγt, NKp46, and IL-7Rα (fig. S6A). We concluded that the Notch signaling pathway had no effect on the survival of NCR+ ILC3s. The MFI of NKp46 was increased in NCR+ ILC3s expressing NICD, which is consistent with a role for Notch signaling in regulating the cell-surface abundance of NKp46 (Fig. 5E), whereas the abundance of RORγt was unchanged in all ILC3 subsets (Fig. 5F). Furthermore, the total number of intestinal ILC1s was increased in the NICD-expressing mice (Fig. 5G). Because NICD had no effect on the ratio of NCR+ ILC3s among total ILCs or the abundance of RORγt, we propose that constitutive activation of the Notch signaling pathway at the NKp46-positive stage increased the size of the ILC1 and cNK subsets independently of a potential ILC3-ILC1 plasticity. Moreover, flow cytometric analysis of the cell surface amounts of both IL-7Rα and Dx5 enabled the identification of ILC1s among total NKp46−NK1.1+ cells, and tended to show that percentage of ILC1s was decreased among group 1 ILCs (fig. S6B). Together, these results support our conclusion that Notch has a direct role on the differentiation of NCR− precursors into NCR+ cells in the intestine, rather than modulating the survival or plasticity of NKp46+ ILC3s.

**In the absence of Notch signaling, the population of CCR6−NCR− cells is devoid of Notch-dependent precursors of NCR+ cells**

We hypothesized that the maturation of NCR− ILC3 precursors into NCR+ ILC3s is blocked in the absence of canonical Notch signaling. Thus, we investigated the effect of loss of the Notch signaling pathway on the transcriptional profiles of different NCR− ILC3 subsets. All NCR− ILC3 subsets (CD4+CCR6+, CD4−CCR6+, and CD4−CCR6−) were present at comparable percentages and numbers in the lamina propria of Rbpj-deficient and Rbpj-haploinsufficient mice (Fig. 6, A and B). We then analyzed the transcriptional profiles, in Notch signaling–competent
and deficient cells, of a set of transcripts that includes transcription factors, chemokines, and cytokine receptors expressed in ILCs, as well as components of the Notch signaling pathway. We also analyzed ILC1s, CD4\(^+\)CCR6\(^+\) ILC3s, and CD4\(^-\)CCR6\(^+\) ILC3s and compared them to the NCR\(^+\) and CCR6\(^-\)NCR\(^-\) ILC3 subsets. Genes with low expression overall [less than 20-fold of that of Tbp (<3 mRNA molecules per cell) (22)] were removed from the analysis, and the abundances of all mRNAs were normalized to the average mRNA abundances of the housekeeping genes Actb, Gapdh, and Hprt.

Clustering of the samples and transcripts enabled the definition of gene signatures that characterized each subpopulation (Fig. 6C). ILC1s were separated from ILC3 subsets, whereas CD4\(^+\)CCR6\(^+\) ILC3s clustered together with CD4\(^+\)CCR6\(^+\) ILC3s, confirming their close relationship. Only for CCR6\(^-\)NCR\(^-\) ILCs did cells from Il7r\(^{Cre/+}\)Rbpj\(^{fl/fl}\) mice not cluster with cells from Il7r\(^{Cre/+}\)Rbpj\(^{fl/+}\) mice, consistent with their dependence on Notch signaling. CCR6\(^-\)NCR\(^-\) cells from Il7r\(^{Cre/+}\)Rbpj\(^{fl/+}\) mice clustered with NCR\(^+\) ILC3s, whereas cells from Il7r\(^{Cre/+}\)Rbpj\(^{fl/fl}\) mice clustered with other NCR\(^-\) ILC3 subsets. Target genes were identified as either decreased in expression (Fig. 6D, top) or increased in expression (Fig. 6D, bottom) with at least a 1.5-fold change in NCR\(^-\) cells, CCR6\(^-\)NCR\(^-\) ILC3s, and CCR6\(^-\)NCR\(^-\) ILC3s from Il7r\(^{Cre/+}\)Rbpj\(^{fl/fl}\) mice compared to haploinsufficient mice. CCR6\(^-\)NCR\(^-\) ILC3s were the cells that showed the largest number of genes that were decreased in expression (20 genes) compared to those in CCR6\(^+\)NCR\(^-\) and NCR\(^+\) cells, whereas increased gene expression was mainly found in NCR\(^+\) ILC3s (29 genes) and CCR6\(^-\)NCR\(^-\) ILC3s (23 genes).
Accordingly, most of the genes that were decreased in expression in the CCR6⁻NCR⁻ subset after Notch signaling deletion were found in NCR⁺ ILC3 clusters (cluster III, 6/10 transcripts), including Runx1, Bcl11b, Icos, and Dtx1, and in ILC1 clusters (cluster I, 4/19 transcripts), including Zbtb16, Ets1, and Nfil3. On the other hand, increased gene expression was found in NCR⁻ ILC3 clusters (clusters II and IV), including genes encoding cytokines (Il-17f and Ltb) or cytokine and chemokine receptors (Ccr6, Ccr8, Il-17rb, and Cxcr5). These results suggest that the transcriptional profile of CCR6⁻NCR⁻ cells from Il7r\textsuperscript{Cre/+}Rbpj\textsuperscript{fl/+} mice was similar to that of NCR⁺ ILC3s in haplosufficient mice and was completely different by inactivation of Notch signaling. Conversely, many of the transcripts that were increased in abundance in CCR6⁻NCR⁻ cells from Il7r\textsuperscript{Cre/+}Rbpj\textsuperscript{fl/fl} mice were shared with other NCR⁻ ILC3 subsets. The expression of Id2, Gata3, AhR, and Nfil3, which are essential for the generation of helper-like ILCs, although present in all ILC subsets (1-3, 10, 23), were substantially decreased by the loss of Notch signaling in CCR6⁻NCR⁻ cells (fig. S7). Nfil3 and the Notch-dependent gene Gata3 were also reduced in expression, whereas the expression of Tox, which is important for ILC development (24), was not affected (Fig. 6D and fig. S7). Zbtb16 (which encodes the PLZF protein) is transiently expressed by most ILCs in a Notch-dependent manner (16). Accordingly, we found that the expression of Zbtb16 was decreased in all ILC3 subsets from Il7r\textsuperscript{Cre/+}Rbpj\textsuperscript{fl/fl} mice, statistically significantly decreased in ILC1s (fig. S7).

Finally, we were interested in the expression patterns of Rorc and Tbx21, both of which are required for the development of NCR⁺ ILC3s (3, 25, 26). Rorc expression in ILC3 subsets from Il7r\textsuperscript{Cre/+}Rbpj\textsuperscript{fl/fl} mice was substantially reduced compared to that in ILC3 subsets from Il7r\textsuperscript{Cre/+}Rbpj\textsuperscript{fl/+} mice cells. In comparison, T-bet was highly expressed in ILC1s, and, although it
was also found in *Rbpj*-competent and deficient ILC3s, quantities of the T-bet protein did not exhibit a corresponding reduction in protein abundance than observed for its encoding Tbx21 gene after Notch signaling deletion (Fig. 4, E-G). This apparent discrepancy might be explained by the low abundance of T-bet protein in ILC3s, which prevented its proper quantification (Fig. 4F). These results suggest that CCR6 NCR− cells in which Notch signaling was lost could not give rise to NCR+ ILC3s such that, in contrast to Notch signaling–competent cells, they had more genes with a similar transcriptional pattern to that found in CCR6+CD4+ and CCR6+CD4− NCR− cells. Hence, the Notch signaling pathway appeared to activate a complete transcriptional program that drove progression of the cells toward an NCR+ fate.

**Discussion**

Intestinal NCR− ILC3s contain the putative NCR+ precursors previously suggested to be Notch-dependent (10-13). Intestinal ILC3s are both Notch1+ and Notch2+, whereas fibroblastic stromal cells in lymph nodes (27) and in intestinal Paneth cells (28) produce the Notch ligands. ILC3s are more abundant in the small intestine than in the colon (9), and a study showed a lower ratio of NCR+ cells to NCR− cells in the colon compared to in the small intestine (29).

In this study, we demonstrated that the intestinal lamina propria is composed of at least two distinct pools of NCR+ ILC3s that differ in their dependency on the Notch pathway. Both a major Notch-dependent and a minor (20%) Notch-independent subsets were found in mice in which the *Notch2* or *Rbpj* loci were ablated in hematopoietic progenitor cells (under either the *Il7r* or *Vav* promoters). These results suggest that the reduction in the number of NCR+ ILC3s in *Rbpj*-deficient mice was Notch2-dependent, independent of bystander T cells, and cell-
autonomous, which was confirmed by analysis of competitive mixed bone marrow chimeras. On the other hand, CCR6⁺NCR⁻ ILC3s, which were largely composed of LTi-like cells, were only mildly affected by the absence of Notch signaling. Thus, we propose a model for the Notch-dependent and Notch-independent differentiation of NCR⁺ ILC3s (Fig. 7). We further obtained evidence for the segregation of Notch-dependent and -independent NCR⁺ subsets in vitro. The analysis of 93 transcripts showed that CCR6⁻NCR⁻ ILC3 precursors that clustered with NCR⁺ cells in haplosufficient mice were more similar to LTi-like cells in the absence of Notch signaling. Hence, the specific effect of loss of Notch signaling in CCR6⁻NCR⁻ cells resulted in the absence of a subset that resembles NCR⁺ cells.

The progression from NCR⁻ toward NCR⁺ cells is partly controlled by AhR (10) and T-bet (11-13). Accordingly, the Notch-independent NCR⁺ subset that we identified still had Tbx21 and AhR mRNAs, albeit at reduced amounts compared to those in the Notch-dependent cell subsets. Previous studies with Tbx21⁻/⁻ mice revealed that the few T-bet–independent NCR⁺ cells showed a reduction in the amount of RORγt (11). Consistent with this finding, we observed substantial decreases in Rorc mRNA and RORγt protein in both NCR⁻CCR6⁻ cells and NCR⁺ cells in Notch signaling deficient mice, which suggests a role for Notch in activating the expression of both Tbx21 and Rorc. Whereas Dtx1 expression was undetectable, Hes1, Tcf7, Zbtb16, Nfil3, and Gata3 transcripts were only partially reduced in abundance in the CCR6⁻NCR⁻ precursors, which suggests that their expression is regulated in part through an alternative pathway. Because our analyses were performed on populations rather than on single cells, the decreased expression of these genes may rather reflect the absence of the Notch-dependent precursors. Nonetheless, NCR⁺ cells exhibited decreased amounts of Hes1, Tcf7, Tbx21, and Rorc transcripts in Notch
signaling–deficient ILCs, which suggests that the Notch-independent generation of NCR$^+$ ILCs is compatible with reduced amounts of these transcripts.

Gata3 inhibits ROR$\gamma$t expression in ILC3s (14), and a 50% reduction in ROR$\gamma$t abundance is sufficient to promote the development of T-bet$^+$ NCR$^+$ ILC3s (14), presumably through derepression of Gata3 expression. Because the expression of both Gata3 and Tbx21 was reduced in NCR$^-$ cells, the decrease in ROR$\gamma$t abundance appeared to be attributable to the loss of Notch signaling. In Tbx21 haploinsufficient mice, NCR$^+$ ILC3s are decreased in number, and T-bet directly stimulates the expression of Notch2, but not Notch1, during the development of NKp46$^+$ ILCs (13). In our analysis, when Notch signaling was inactivated, both Notch1 and Notch2 transcripts were increased in abundance in NCR$^+$ ILC3s, although the abundance of T-bet was maintained. These results suggest that the Notch signaling pathway also regulates receptor abundance through a negative feedback loop.

It was proposed that Notch presence in ILC3s depends on AhR (10). Although decreased in abundance among the NCR$^-$ precursors, AhR transcripts were still present in Notch signaling–deficient cell populations, which could explain the maintenance of an AhR-dependent, IL-22–secreting CD4$^+$CCR6$^+$ population in normal numbers. It thus appears that the coordinated actions of ROR$\gamma$t, T-bet, Gata3, AhR, and Notch on the same target cells maintains the equilibrium between the NCR$^-$ and NCR$^+$ ILC3 subsets. When one of these co-regulators is perturbed by environmental cues or pathologic conditions, the equilibrium shifts toward another subset, thus explaining the plasticity of ILC3s.
Experiments with NCR+ cells in vitro showed that NKp46+ ILC3s were more efficiently maintained in the presence than absence of Notch ligands, which suggests that Notch signaling is important for the cell-surface expression of NKp46 or for maintaining it. Conversely, in Notch signaling–deficient conditions, the NK cell markers NKp46 and NKG2D were minimally abundant on the surface of NCR- cells. Other cell-surface markers, such as Thy1, cKit, and Sca1, also changed their abundances on ILC3 populations when the Notch signaling pathway was lost, which implies that they are partially regulated by the Notch pathway. Transcripts of genes encoding pro- and anti-apoptotic factors in ILC3s were similarly abundant in the presence or absence of Notch signaling, which suggests that NCR+ cells do not undergo apoptosis in the absence of Notch. CCR6+NCR- ILC3s (LTi-like cells) appeared as a stable population, which was in comparison to NCR+ ILC3s, which lost RORγt and produced IFN-γ, entering a state known as “ex-ILC3” (9, 12). However, neither enhancement nor abrogation of the Notch signaling pathway alone was sufficient to drive the conversion of ILC3s to ILC1s. The increase in the number of NK cells in mice expressing NICD under the control of the Ncr1 regulatory sequences was probably because of a specific action of Notch on ILC1s. Indeed, this observation did not correlate with the transition from ILC3s to ILC1s, because the number of NCR+ ILC3s was stable, the abundance of RORγt ILC3s was normal, and the ILC1 subset decreased as a percentage of type I ILCs.

In mice in which NICD was expressed at the earliest developmental stage of IL-7R precursor, more NCR- cells received a Notch signal and consequently both the CCR6+NCR- and NCR+ populations were increased in number. In culture, NCR- cells that originated from NCR+ cells were detected, and they were more frequent in the absence of Notch signaling, which suggests
that NCR\(^+\) cells may revert to NCR\(^-\) cells in the absence of Notch signaling. However, no differences in NCR\(^-\) cell numbers were found in vivo in the absence of Notch signaling. Hence, these results suggest that *Ncr1* is a direct target of Notch in ILC3s and is required to maintain the NCR\(^+\) identity of the Notch-dependent cell subset. As observed in vitro, the cell-surface expression of NKp46 could be transient on ILC3s, which implies that the NCR\(^-\) cell population could contain some “ex-NCR\(^+\)” cells. Only a fate map mouse model tracing the NKp46\(^+\) cells could determine if the reversion of NCR\(^+\) cells to NCR\(^-\) ILC3s is a common event in the intestine. However, we consider that process to be unlikely because a decrease in *Nkp46* expression in the absence of Notch signaling should have revealed an enrichment, rather than a reduction, of the NCR\(^+\) transcriptional signature among the CCR6 NCR\(^-\) population in Notch signaling-deficient mice.

In comparison to previous studies, the mouse model reported here exhibits an inactivation of the Notch signaling pathway that is restricted to the lymphoid compartment and does not affect the Notch2-dependant DC subsets that deliver regulatory cytokines to ILCs. Under steady-state conditions, ILC3s support intestinal homeostasis through such diverse mechanisms as IL-22 secretion to control intestinal microbiota (30) or GM-CSF secretion, which preserves the tolerogenic function of intestinal DCs (31). We report that Notch signaling–deficient ILC3 subsets have quantitative differences in their secretory capacities that could reinforce or challenge their steady-state functions. Indeed, in Notch signaling–deficient ILC3 subsets, the production of IL-17F, IL-22, and Csf2 was increased compared to that of their Notch signaling–sufficient counterparts, whereas the production of IFN-\(\gamma\) was unaffected. Notch signaling–deficient NCR\(^+\) ILC3s that also had lower amounts of cell-surface NCR markers may have
different functions during infection and inflammation. In conclusion, our study reveals that the Notch2 canonical signaling pathway regulates both the plasticity and function of adult ILC3s.

**Materials and Methods**

**Mice**

Il7r<sup>Cre</sup>Rosa26<sup>YFP</sup> mice [kindly provided by H-R Rodewald, German cancer research center Heidelberg, (32)] were crossed with Rbp<sup>jtm1Hon</sup> mice (33) (Il7r<sup>Cre/+</sup>Rbp<sup>jfl/+</sup>Rosa26<sup>YFP</sup>), Notch2 mice (Il7r<sup>Cre/+</sup>Notch2<sup>fl/+</sup>Rosa26<sup>YFP</sup> (34), or NICD mice (Il7r<sup>Cre/+</sup>Rosa26<sup>NICD/+</sup>) (35). Vav<sup>Cre</sup> and Ncr1<sup>Cre</sup>Rosa26<sup>NICD/+</sup> mice were kindly provided by C. Vosshenrich. All animal experiments were approved by the Pasteur Institute Safety Committee in accordance with French Agriculture ministry and the EU guidelines.

**Cell preparation**

Fetal liver, bone marrow, thymic lobes, spleens, and lamina propria lymphocytes (LPLs) were harvested, dissociated, and resuspended in Hanks’ balanced-salt solution (HBSS) supplemented with 1% fetal calf serum (FCS, Gibco). Fetal liver, bone marrow, and spleen cells were depleted of Lineage<sup>+</sup> (lin<sup>−</sup>) cells by staining with biotinylated-conjugated antibodies specific for lineage markers (for bone marrow cells, these include: CD3ε, CD5, CD8α, CD11c, CD19, Ter119, Gr-1, NK1.1, TCRβ, and TCRδ; for spleen cells, these include: CD3ε, CD5, CD8α, CD19, Ter119, Gr-1, TCRβ, and TCRδ), followed by incubation with streptavidin microbeads (Miltenyi Biotec). Depletion was performed on LS+MACS columns (Miltenyi Biotec) from which the negative fraction was recovered. To isolate LPLs, the small bowel was flushed with phosphate-buffered saline (PBS) and the conjunctive tissue and Peyer’s patches were carefully removed. The
intestine was opened and cut into 1-cm pieces. To eliminate epithelial cells and intra-epithelial lymphocytes, these fragments were incubated at 37°C in 50 ml of RPMI 1640 (Gibco) containing 10% FCS and 10 mM Hepes buffer under strong agitation for 30 min, which was followed by vortex treatment for 4 min. For LPL isolation, the remaining fragments were incubated in identical medium to which was added type VIII collagenase (0.5 mg/ml, Sigma) and were shaken for 30 min at 37°C. To complete digestion, the suspension was repeatedly passed through a 10-ml syringe for 5 min and then filtered through a 40-μm cell strainer (BD Bioscience) and collected by centrifugation. The cell pellet was re-suspended in 44% Percoll (GE Healthcare), laid over 67% Percoll, and centrifuged at 600g for 20 min at 20°C. Cells at the interface were collected, washed in HBSS containing 1% FCS, and recovered.

**Flow cytometry and cell sorting**

Flow cytometry data were acquired with a FACS Canto II or LSR Fortessa flow cytometer (Becton Dickinson) and analyzed with FlowJo software (TreeStar). Dead cells were eliminated by exclusion with propidium iodide. Cells were stained intracellularly after permeabilization and fixation with Foxp3 Permeabilization/Fixation Concentrate and Diluent (eBioscience). Cells were then incubated for 5 min with DAPI (4,6-diamidino-2-phenylindol; Invitrogen). LPL cells were purified with a FACS Aria III (Becton Dickinson). Cells were recovered in tubes or in 96-well qPCR plates for gene expression analysis.

**Antibodies**

All antibodies were from BD Biosciences, eBioscience, Biolegend, Cell Signaling Technologies, or R&D Systems. Antibodies were biotinylated or conjugated to fluorochromes (FITC, PE,
PECy5, PerCPc5.5, PECy7, APC, AlexaFluor647, APC-Cy7, Pacific Blue, BV421, eFluor450, V500, BV605, BV655, BV700, BV786) and were specific for the following mouse antigens: Ly76 (TER-119), Gr-1 (RB6-8C5), CD11c (HL3), CD3ε (145-2C11), CD5 (53-7.3), CD19 (6D5), NK1.1 (PK136), IL-7Rα (A7R34), c-Kit (2B8), Sca-1 (D7), RORγt (AFKJS-9), α4β7 (DATK32), Flt3 (A2F10), Ly6D (48-H4), CD8 (53-6.7), TCRβ (H57-597), TCRδ (GL3), CCR6 (29-2L17), CD4 (GK1.5), CD25 (PC61), CD44 (IM7), Thy1.2 (53-2.1), NKp46 (29A1.4), GATA-3 (L50-823), ICOS (C398.4A), Ly49 CIFH (14B11), IFN-γ (XMG1.2), IL-22 (1H8PWSR), T-bet (eBio4B10), CD27 (LG.3A10), T1/ST2 (DIH9), CD45.1 (A20), CD45.2 (104), CD23 (B3B4), IL-17A (TC11.18H10.1), CD49a (HMa1), CD49b (DX5), NKG2D (CX5), and CD21 (7E9). Antibody against human Ki67 (B56) was obtained from BD Pharmingen.

Cell culture

All experiments were performed in 96-wells plates at 37°C and 5% CO₂ and in culture medium consisting of OptiMEM, 10% (vol/vol) FCS, penicillin (100 U/ml), streptomycin (100 μg/ml), and 50 μM 2-mercaptoethanol (Gibco). OP9 and OP9-DL4 stromal cells were seeded into 96-well plates. The culture medium was supplemented with saturating amounts of c-Kit ligand, Flt3 ligand, IL-2, and IL-7. The presence of ILC3 subsets was assayed by flow cytometry analyses for the expression of NKp46, RORγt, CCR6 and CD4 proteins among cultured cells after 4 days on OP9 or OP9-DL4 cells.

Chimeric reconstitution

Recipient Ly5.1 mice were sub-lethally irradiated (800rad) before injection. Fetal liver progenitors (Lin⁻IL-7Rα⁻cKit⁺Sca-1⁺) cells were sorted from donor Ly5.2 or competitor Ly5.1 ×
Ly5.2 E15 embryos and then mixed at 1:1 ratio for reinjection. Additionally, protective congenic bone marrow cells (1 x 10^6 T cell–depleted Ly5.1 bone marrow cells) in PBS were also injected at the same time into the retro-orbital vein. Reconstituted mice were analyzed after 8 weeks. Cells from the recipient mice were CD45.1⁺CD45.2⁻, whereas competitor cells (WT B6) were CD45.1⁺CD45.2⁺, and Rbpj-deficient or Rbpj-sufficient donor cells were CD45.1⁻CD45.2⁺.

RT-PCR analysis

Cells were sorted in RLT buffer (Qiagen) containing 2-mercaptoethanol (Sigma) and were frozen at -80°C. RNA was obtained with an RNeasy Micro Kit (Qiagen), and cDNA was obtained with the PrimeScript RT Reagent Kit (Takara). A 7300 Real-Time PCR System (Applied Biosystem) and Taqman technology (Applied Biosystem) or SYBRGreen Technology (Qiagen) were used for quantitative reverse transcriptase polymerase chain reaction (RT-PCR) analysis. A bilateral unpaired Student’s t-test was used for statistical analysis. The following primers were from SA Biosciences: Il7f (PPM05398E), Il22 (PPM481A), and Gapdh (PPM02946E). The following primers were from Applied Biosystems: Il1r1 (Mm_00434237_m1), Il23r (Mm_00519943_m1), Csf2 (Mm_01290062_m1), Ifng (Mm_01168134_m1), Casp3 (Mm_01195085_m1), Casp9 (Mm_00516563_m1), Bcl2 (Mm_00477631_m1), Bcl2cl1 (Mm_00427783_m1), Bcl2cl11 (Mm_00432359_m1), Birc3 (Mm_01168413_m1), Bik (Mm_00476123_m1), Hprt (Mm_00446968_m1), Actb (Mm_02619580_g1).

Multiplex RT-PCR analysis

Total RNA from the appropriate cell populations was extracted as previously described (19) and mixed in 10 µl of CellsDirect One-Step qRT-PCR Kit (Life Technologies), containing a mixture
of diluted primers (0.05× final concentration). See table S1 for sequences. Pre-amplified cDNA (22 cycles) was obtained according to the manufacturer’s instructions and was diluted 1:5 in TE buffer (pH 8, Ambion). The sample mixture was as follows: diluted cDNA (2.9 µl), Sample Loading Reagent (0.32 µl, Fluidigm), and either Taqman Universal PCR Master Mix (3.5 µl, Applied Biosystems) or Solaris qPCR Low ROX Master Mix (3.5 µl, GE Dharmacon). The assay mixture was as follows: Assay Loading Reagent (Fluidigm) and either Taqman (Applied Biosystem) or Solaris (GE Dharmacon). A 96.96 dynamic array integrated fluidic circuit (IFC, Fluidigm) was primed with control line fluid, and the chip was loaded with assays (either Taqman or Solaris) and samples with an HX IFC controller (Fluidigm). The experiments were run on a Biomark HD (Fluidigm) for 40 cycles. Samples that did not express at least one of three housekeeping genes (Actb, Gapdh, or Hprt) and genes that were present in less than 5% of the wells (except for Dtx1, Pax5, Ebf1 and Rag2) were removed from the analysis. Gene expression was assessed by the 2−ΔCt method.

Statistical analysis

Statistical analysis was performed with the Mann Whitney nonparametric test where appropriate. These tests were performed with Prism software (GraphPad). Graphs containing error bars show means ± SEM. Statistical significance is represented as follows: *P < 0.05, **P < 0.01, ***P < 0.001.

Supplementary Materials

Fig. S1. Rbpj is efficiently deleted in IL-7Rα+ CLPs by Cre recombinase.
Fig. S2. Notch2 signaling is specifically required for both NCR+ and NCR− ILC3s.
Fig. S3. Analysis of cytokine secretion and receptors in Notch signaling–competent (Ii7rCre+/Rbpjfl/fl) and deficient (Ii7rCre+/Rbpjfl/fl) mice.
Fig. S4. Expression of Gata3 and Il7ra in ILC3 subsets in the presence or absence of Notch signaling.

Fig. S5. The expression of genes encoding pro-apoptotic proteins in ILC3 subsets is unchanged by the loss of Notch signaling.

Fig. S6. Analysis of ILC3 and ILC1 subsets in Ncr1Cre/+Rosa26NIcD/+ mice.

Fig. S7. Analysis of transcription factors expressed from ILC3 and ILC1 subsets in Notch signaling–competent (Il7rCre/+Rbpjfl/+ ) and -deficient (Il7rCre/+Rbpjfl/fl ) mice.

Table S1. P values for the differential expression of 92 genes between Il7rCre/+Rbpjfl/+ and Il7rCre/+Rbpjfl/fl ILC3 subsets (respectively, NCR+, CCR6+ NCR- and CCR6- NCR- subsets).

References and Notes


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**Author contributions:** S.C. performed most of the experiments and analyzed data; T.P. performed and analyzed experiments and prepared figures; M.P. performed and analyzed experiments; T.V. performed experiments with the Ncr1CreNICD mice; E.G.B. performed the mouse genotyping; D.G.G. supervised experiments with lamina propria cells and contributed to the writing of the manuscript; C.V. generated the Ncr1Cre mice; J.P.D. and A.C. contributed to the writing of the manuscript; and R.G. directed the research, designed experiments, analyzed
data, and wrote the manuscript. **Competing interests:** The authors declare that they have no competing interests.
Fig. 1. Loss of Notch signaling alters the homeostasis of intestinal ILC3 subsets. (A) Flow cytometric analysis of lamina propria lymphocytes (LPLs) from C57Bl/6 mice. Lin− cells were defined according to the presence of RORγt, IL-7Rα, and NKp46 into cNK cells (RORγt− NKp46−NK1.1+IL-7Rα−), ILC1s (RORγt−NKp46+NK1.1+IL-7Rα+), NCR+ ILC3s (RORγt−NKp46+IL-7Rα+), and NCR− ILC3s (RORγt+NKp46+NK1.1+IL-7Rα−). (B) Percentages (top) and total numbers (bottom) of cNK cells and ILC1s from Il-7rCre/+Rbpjfl/fl, Il-7rCre/+Rbpjfl/fl, and Il-7rCre/+Rbpjfl/fl mice. (C) Percentages (top) and total numbers (bottom) of NCR+ ILC3s and NCR− ILC3s from Il-7rCre/+Rbpjfl/fl, Il-7rCre/+Rbpjfl/fl, and Il-7rCre/+Rbpjfl/fl mice. (D) Flow cytometric analysis of the amounts of Icos, Sca-1, CD25, Thy1, c-Kit, and Ly49 CIFH in cNK cells and ILC1s from Il-7rCre/+Rbpjfl/fl mice (black) and Il-7rCre/+Rbpjfl/fl (red) mice compared to those in total Lin− cells (gryy). (E) Flow cytometric analysis of the amounts of Icos, Sca-1, CD25, Thy1, c-Kit, and Ly49 CIFH in NCR+ ILC3s and NCR− ILC3s from Il-7rCre/+Rbpjfl/fl mice (black) and Il-7rCre/+Rbpjfl/fl (red) mice compared to those in total Lin− cells (gray). (F) Mean fluorescence intensities (MFIs) of Thy1, c-Kit, NKG2D, and Sca1 for ILC1s, NCR+ ILC3s, and NCR− ILC3s from Il-7rCre/+Rbpjfl/fl mice (white) and Il-7rCre/+Rbpjfl/fl (black) mice. a.u., arbitrary units. Data are representative of at least three independent experiments with at least three mice of each genotype analyzed per experiment. Data in (B), (C), and (F) are means ± SEM of three independent experiments, with three mice of each genotype analyzed per experiment. Data points in (B) and (C) represent individual mice. *P < 0.05; **P < 0.01; ***P < 0.001 by Mann Whitney test.
Fig. 2. Changes in the cytokine profiles of ILC3 subsets from mice deficient in Notch signaling. (A) Quantitative RT-PCR analysis of the relative abundances of Il17f, Il22, Il17a, Csf2, and Ifng mRNAs in ILC1, NCR⁺ ILC3, and NCR⁻ ILC3 subsets (as defined in Fig. 1A) from Il-7r<sup>Cre/+ Rbpj<sup>fl/+ mice (white) and Il-7r<sup>Cre/+ Rbpj<sup>fl/fl mice (black). The abundances of the indicated mRNAs are presented relative to the average abundances of mRNAs of housekeeping genes. Data are means ± SEM of three independent experiments, with three mice of each genotype analyzed per experiment. *P < 0.05, **P < 0.01, ***P < 0.001 by Mann Whitney test. (B) cNK-ILC1, NCR⁺ ILC3, and NCR⁻ ILC3 subsets isolated from Il-7r<sup>Cre/+ Rbpj<sup>fl/+ mice (black) and Il-7r<sup>Cre/+ Rbpj<sup>fl/fl mice (red) were treated with PMA (left) or IL-23 (middle and right) and then were analyzed by flow cytometry to determine the relative abundances of IFN-γ (left), IL-22 (middle), and IL-17A (right). Data are representative of two independent experiments. (C) Analysis of the percentages of the indicated subsets of cells from Il-7r<sup>Cre/+ Rbpj<sup>fl/+ (white bars) and Il-7r<sup>Cre/+ Rbpj<sup>fl/fl mice (black bars) that were positive for IL-17A (top) or IL-22 (bottom) in response to treatment with IL-23. Data are means ± SEM of two independent experiments, with two mice of each genotype analyzed per experiment. Analysis by Mann Whitney test showed that there were no statistically significant differences among the groups.
Fig. 3. The disruption of ILC3 homeostasis in Notch signaling–deficient mice is cell-
intrinsc. (A to D) A total of 2000 fetal liver LSK cells from embryonic day 15.5 (E15.5) CD45.2+CD45.1- donor mice (either Il7r<sup>Cre/+Rbpj<sub>fl/fl</sub></sup> or Il7r<sup>Cre/+Rbpj<sub>fl/+</sub></sup>) and from CD45.2+CD45.1+ competitor B6 wild-type (WT) mice were injected at a 1:1 ratio into lethally irradiated CD45.2-CD45.1+ recipient mice, as indicated. The mice were analyzed 8 weeks after reconstitution. (A) Flow cytometric analysis of the ILC3s in the lamina propria of the indicated mice after reconstitution. Lamina propria ILC3s (LPL ILC3s) were assigned, based on their cell-surface expression of CD4, CCR6, and NKp46, into CD4+CCR6+NCR- cells (pink), CD4+CCR6-NKR- cells (orange), and CD4’NKp46’NKR+ cells (blue). (B) Percentages of donor ILC3 subsets from the lamina propria of both type of reconstituted mice (Il7r<sup>Cre/+Rbpj<sub>fl/+</sub></sup>: WT in white or Il7r<sup>Cre/+Rbpj<sub>fl/fl</sub></sup>: WT in black). (C) Donor:competitor ratios of ILC3 subsets [as defined in (A)] after reconstitution from reconstituted mice with Il7r<sup>Cre/+Rbpj<sub>fl/+</sub></sup>: WT (white) or Il7r<sup>Cre/+Rbpj<sub>fl/fl</sub></sup>: WT (black). (D) Flow cytometric analysis of CD4+ T cell (defined as CD3+CD4+) and CD8+ T cell (defined as CD3+CD8+) subsets in the spleen of the indicated mice after reconstitution. (E) Percentages of donor CD4+ T cell and CD8+ T cell subsets from the lamina propria of both type of reconstituted mice (Il7r<sup>Cre/+Rbpj<sub>fl/+</sub></sup>: WT in white or Il7r<sup>Cre/+Rbpj<sub>fl/fl</sub></sup>: WT in black). (F) Donor:competitor ratios of CD4+ T cell and CD8+ T cell subsets [as defined in (C)] from reconstituted mice with Il7r<sup>Cre/+Rbpj<sub>fl/+</sub></sup>: WT (white) or Il7r<sup>Cre/+Rbpj<sub>fl/fl</sub></sup>: WT (black). Data are representative of two independent experiments, with four or five mice of each group analyzed per experiment. Data in (B) and (D) are means ± SEM. *P < 0.05, **P < 0.01, ***P < 0.001 by Mann Whitney test.
Fig. 4. Notch signaling contributes to the maintenance of NCR⁺ ILC3s and the conversion of NCR⁻ ILC3s into NCR⁺ ILC3s. (A) Sorting strategy of NCR⁺ ILC3s (in blue; Lin⁻NK1.1⁻ KLRG1⁻IL-7Rα⁺Thy1.2⁻NKp46⁺), CCR6⁻ NCR⁻ ILC3s (in orange; Lin⁻NK1.1⁻KLRG1⁻IL-7Rα⁺Thy1.2⁻NKp46⁻CCR6⁻), and CCR6⁺ NCR⁻ ILC3s (in pink; Lin⁻NK1.1⁻KLRG1⁻IL-7Rα⁺Thy1.2⁻NKp46⁻CCR6⁺) from Il-7rCre/+ Rbpjfl/+ mice (top) and Il-7rCre/+ Rbpjfl/fl mice (bottom). (B) Purity sorting test. After sorting, sorted cell populations were analyzed by flow cytometry a second time to check cell populations purity. All sorted cell population purity were superior to 99%. Data in (A) and (B) are representative of at least 3 experiments. (C) Flow cytometric analysis of the indicated ILC3 subsets isolated from Il-7rCre/+ Rbpjfl/+ mice (top) and Il-7rCre/+ Rbpjfl/fl mice (bottom) and cultured for 4 days on OP9-DL4 cells. Data are representative of four independent experiments. (D) Analysis of the indicated ILC3 subsets after culture on OP9-DL4 cells (white) or OP9 cells (black) expressed as a percentage of CD45⁺ cells. Data are means ± SEM of four independent experiments. (E) Quantitative RT-PCR analysis of NCR⁺ ILC3s and NCR⁻ ILC3s isolated from Il-7rCre/+ Rbpjfl/+ mice (white) and Il-7rCre/+ Rbpjfl/fl mice (black). The abundances of the indicated mRNAs are presented relative to the average of those of the appropriate housekeeping genes. N.D., not detected. Data are means ± SEM of two independent experiments, with four mice of each genotype analyzed per experiment. (F) Flow cytometric analysis of the relative amounts of T-bet in ILC1s, NCR⁺ ILC3s, and NCR⁻ ILC3s isolated from Il-7rCre/+ Rbpjfl/+ mice (black) and Il-7rCre/+ Rbpjfl/fl mice (red) compared to those in total Lin⁻ cells (gray). Data are representative of three independent experiments. (G) Comparison of the MFIs of RORγt and T-bet in the indicated sorted cells isolated from Il-7rCre/+ Rbpjfl/+ mice (white) and Il-7rCre/+ Rbpjfl/fl mice (black). Data are means ± SEM of three independent
experiments, with two mice of each genotype analyzed per experiment. *$P < 0.05$, **$P < 0.01$, ***$P < 0.001$ by Mann Whitney test.
A. 

B. 

C. 

D. 

E. 

F. 

G.
Fig. 5. Notch signaling favors the differentiation of ILC3s from lamina propria ILC precursors. (A) Flow cytometric analysis of cNK-ILC1s, NCR⁺ ILC3s, and NCR⁻ ILC3s in the lamina propria of Il-7rCre⁺Rosa26⁺/+ and Il-7rCre⁺Rosa26NICD/+ mice. Data are representative of three independent experiments. (B) Percentages (top) and total numbers (bottom) of cNK cells, ILC1s, NCR⁺ ILC3s, and NCR⁻ ILC3s in the lamina propria of Il-7rCre⁺Rosa26⁺/+ mice (white) and Il-7rCre⁺Rosa26NICD/+ mice (black). Data are means ± SEM of three independent experiments, with three mice of each genotype analyzed per experiment. (C) Flow cytometric analysis of the indicated ILC3 subsets assigned according to their cell-surface expression of NKp46 and CCR6: NCR⁺ ILC3s (NKp46⁺CCR6⁻; blue), CCR6⁻ NCR⁻ ILC3s (NKp46⁻CCR6⁻; orange), and CCR6⁺ NCR⁻ ILC3s (NKp46⁻CCR6⁺; pink). Data are representative of two independent experiments. (D) Percentages (left) and total numbers (right) of the indicated ILC3 subsets from Ncr1Cre⁺Rosa26⁺/+ mice (white) and Ncr1Cre⁺Rosa26NICD/+ mice (black). (E) NKp46 MFIs of NCR⁺ ILC3s from Ncr1Cre⁺Rosa26⁺/+ mice (white) and Ncr1Cre⁺Rosa26NICD/+ mice (black). (F) RORγt MFIs of the indicated ILC3 subsets from Ncr1Cre⁺Rosa26⁺/+ mice (white) and Ncr1Cre⁺Rosa26NICD/+ mice (black). (G) Total numbers of cNK-ILC1s (NK1.1⁺NKp46⁺) from Ncr1Cre⁺Rosa26⁺/+ mice (white) and Ncr1Cre⁺Rosa26NICD/+ mice (black). Data in (D) to (G) are means ± SEM of two independent experiments, with two mice of each genotype analyzed per experiment. *P < 0.05, **P < 0.01, ***P < 0.001 by Mann Whitney test.
Fig. 6. The decrease in the number of NCR⁺ ILC3s in Notch signaling–deficient mice results from loss of the Notch-dependent RORγt⁺ subset. (A) Flow cytometric analysis of NCR⁻ ILC3 subsets from Il-7r⁺/⁺Rbpj⁺/⁺ mice. All CD4⁺NCR⁻ ILC3s are CCR6⁺ (dark green), whereas CD4⁺NCR⁻ ILC3s are either CCR6⁺ (light green) or CCR6⁻ (orange). (B) Percentages (top) and total numbers (bottom) of the indicated NCR⁻ ILC3 subsets from Il-7rCre⁺/⁺Rbpjfl/fl mice, Il-7rCre⁺/⁺Rbpjfl/fl mice, and Il-7r⁺/⁺Rbpjfl⁺/⁺ mice. Each data point represents an individual mouse. Data are means ± SEM of at least 3 independent experiments, with 2 mice of each genotype analyzed per experiment. *P < 0.05, **P < 0.01, ***P < 0.001 by Mann Whitney test. (C) Single cell multiplex mRNA abundances of cNK-ILC1s, NCR⁺ ILC3s, and NCR⁻ ILC3s in the lamina propria of Il-7rCre⁺/⁺Rbpjfl⁺/⁺ mice and Il-7rCre⁺/⁺Rbpjflfl mice. Results are presented as relative to the average expression of housekeeping genes. After hierarchical clustering, data were median-centered and scaled for visualization (red: high expression, blue: low expression, gray: no expression). (D) Venn diagrams of the genes that were decreased in expression (top) and the genes that were increased in expression (bottom) in the indicated subsets of cells isolated from Il-7rCre⁺/⁺Rbpjflfl mice compared to those in the same subsets isolated from Il-7rCre⁺/⁺Rbpjfl⁺/⁺ mice. Genes with at least a 1.5-fold change in expression are listed, whereas those with at least a two-fold change in expression are in bold. Genes that are in colors are either increased or decreased in expression in other cell populations. Data are representative of at least two independent experiments, with four mice of each genotype analyzed per experiment. P values from statistical analysis of the data in (D) are summarized in table S1.
**Fig. 7. Schematic representation of the differentiation of intestinal NCR$^+$ ILC3s.** The recapitulative model shows that whereas 80% of the NCR$^+$ cells were derived from $Dtx1^+$ T-bet$^+$ precursors, the Notch-independent pool was derived from a $Dtx1^-$ T-bet$^+$ precursor and represents 20% of the total number of NCR$^+$ cells. The absence of Notch signaling is associated with decreased amounts of NKG2D, RORγ$t$, and Thy$^1$, as well as of transcription factors, such as Zbtb16, Tcf7, Gata3, Hes1, Nfil3 and AhR. Whereas the genes encoding most of these factors are Notch targets, the factors were decreased in abundance, but were not absent, in Notch-independent precursors. Similar to their NCR$^-$ precursors, the Notch-independent NCR$^+$ cells showed decreased amounts of similar proteins (including NKG2D, RORγ$t$, and Thy$^1$) and transcription factors (including Tcf7 and Hes1). They also exhibited specific increases in cKit and Sca1 abundance. The transcripts for Notch1 and Notch2 were also both increased in abundance in Notch independent NCR$^+$ cells. The Notch-independent NCR$^+$ cells expressed more Csf2 transcripts under steady-state conditions and secreted more IL-22 before and after activation than did Notch dependent NCR$^+$ cells. The CCR6$^+$ subset was added to the scheme as an early progenitor of Notch-dependent NCR$^+$ cells because a previous study showed that CD4$^-$ CCR6$^+$ ILC3s can differentiate into NCR$^+$ ILC3s in a T-bet–dependent manner after culture in the presence of Notch ligands (that is, on OP9-D11 cells) (13).