ILC1s in tissue inflammation and infection

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ILC1s in Tissue Inflammation and Infection

Anja Fuchs*

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Innate lymphoid cells (ILCs) are innate immune cells that provide an early source of cytokines to initiate and tailor the immune response to the type of the encountered pathogen or insult. The group 1 ILCs are comprised of conventional natural killer (cNK) cells and subsets of “unconventional NK cells,” termed ILC1s. Although cNK cells and ILC1s share many features, such as certain phenotypic markers and the ability to produce IFN-γ upon activation, it is now becoming apparent that these two subsets develop from different progenitors and show unique tissue distribution and functional characteristics. Recent studies have aimed at elucidating the individual contributions of cNK cells and ILC1s during protective host responses as well as during chronic inflammation. This review provides an overview of the current knowledge of the developmental origins as well as of the phenotypic and functional characteristics of ILC1s.

Keywords: ILC1, innate lymphoid cells, inflammation, host response, ILC development, antimicrobial defense

INTRODUCTION

Innate lymphoid cells (ILCs) are a recently identified group of cells of the innate immune system that modulate immune functions prior to the generation of an adaptive immune response. Unlike T cells and B cells, ILCs lack antigen-specific receptors; instead, their function is mediated by cytokines released by antigen-presenting cells or damaged tissue cells in response to infection or tissue damage. Activated ILCs secrete cytokines that tailor the immune response to the encountered pathogen or insult. Based on their secreted cytokine profile, ILCs are regarded as the innate equivalent of adaptive T cells: group 1 ILCs are equivalent to Th1 T cells, group 2 ILCs to Th2 T cells, and group 3 ILCs to Th17 and Th22 cells (1–4). The group 1 ILCs are comprised of conventional natural killer (cNK) cells and several groups of “unconventional NK cells,” termed ILC1s, which differ from conventional NK cells in their phenotypes, locations, functions, and/or transcription factor dependence. Within group 1 ILCs, cNK cells appear to be more potent in mediating cytotoxicity (“killer ILCs”), while the primary role of ILC1s is the production of pro-inflammatory cytokines (“helper-like ILCs”) (2). This review will briefly touch on the key features of conventional NK cells as the prototype of group 1 ILCs and will then discuss in more detail the known characteristics of the “unconventional NK cells,” referred to here as ILC1s. Several subsets of ILC1s have been identified, which are located predominantly within non-lymphoid tissues, such as the intestinal mucosa, liver, salivary gland, and in the female reproductive tract. For some of the ILC1 subsets, a clear lineage relationship to ILC1s or cNK cells has not firmly been established. The paucity of specific surface markers and transcription factors for ILC1s has made studying these cells difficult. The ILC field in general is a relatively young field of study, and within this field, group 1 ILCs are the least-well studied ILCs compared to ILC2s and ILC3s. This review aims at giving an overview of the current knowledge on the phenotype and
developmental origin of ILC1 subsets, as well as their roles during antimicrobial immune defense and chronic inflammation.

HETEROGENEITY WITHIN GROUP 1 ILCs: LOCATION, PHENOTYPE, AND FUNCTIONS

The unifying characteristic of group 1 ILCs is their ability to produce IFN-γ. Apart from this common attribute, significant differences exist between the known ILC1 subsets. The following sections summarize the current knowledge of the tissue location, surface marker expression, and functional characteristics of the main group 1 ILC subsets.

Conventional NK Cells

Conventional NK cells are predominantly found as circulating in the blood and within secondary lymphoid tissues such as the lymph nodes and spleen. However, cNK cells are also present in some non-lymphoid tissues such as the liver and the lung. Resting mature cNK cells are identified as CD3ε− NKp46+ DX5 (CD49b)+ lymphocytes in the mouse (Table 1) and as CD3ε− CD56ε− NKp46+ NKp44− cells in humans (Table 2) (5, 6). Mature cNK cells express the transcription factors T-bet and Eomes, which are important in mediating their functions. cNK cells can be activated through cytokines, such as IL-12, IL-15, and IL-18, which are important in mediating their functions. cNK cells can undergo an expansion and contraction phase during microbial infections, followed by the generation of a pool of memory cells. These memory cNK cells possess enhanced cytolytic and cytokine responses during secondary exposure to certain microbial pathogens, a finding that suggests immunological memory is not confined to adaptive lymphocytes (14).

ILC1s

Unconventional NK cells (ILC1s) are currently defined as tissue-resident NK-like cells that do not develop from conventional NK cell precursors (NKP) and are not typically found in blood or lymphoid organs. Subsets of ILC1s have been identified in a variety of non-lymphoid tissues, including small intestinal mucosa, liver, salivary glands, and the female reproductive tract (Tables 1 and 2). In mice, these pools of tissue-resident ILC1s do not recirculate and appear to be maintained predominantly via local self-renewal rather than through replenishment from blood-derived ILC1s or their precursors (15–17). Apart from these common attributes, significant differences exist between

### Table 1 | Surface marker and transcription factor expression by murine group 1 ILCs.

<table>
<thead>
<tr>
<th>NK1.1</th>
<th>NKp46</th>
<th>DX5</th>
<th>TRAIL</th>
<th>CD49a</th>
<th>CD160</th>
<th>CD127</th>
<th>CD11b</th>
<th>CD69</th>
<th>CXCR6</th>
<th>T-bet</th>
<th>Eomes</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Splenic cNK</td>
<td>+</td>
<td>+</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>+</td>
<td>-</td>
<td>-</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>LP ILC1</td>
<td>+</td>
<td>+</td>
<td>-</td>
<td>n.d.</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>-</td>
<td>±</td>
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<td>ile ILC1</td>
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<td>+</td>
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<tr>
<td>Liver ILC1</td>
<td>+</td>
<td>+</td>
<td>-</td>
<td>+</td>
<td>+</td>
<td>±</td>
<td>-</td>
<td>+</td>
<td>±</td>
<td>-</td>
<td>±</td>
<td>+</td>
</tr>
<tr>
<td>Saliary ILC1</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>n.d.</td>
<td>-</td>
<td>+</td>
<td>+</td>
<td>n.d.</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>-</td>
</tr>
</tbody>
</table>

± heterogenous expression; n.d., not determined.

### Table 2 | Surface marker and transcription factor expression by human group 1 ILCs.

<table>
<thead>
<tr>
<th>CD56</th>
<th>NKp46</th>
<th>NKp44</th>
<th>CD103</th>
<th>CD49a</th>
<th>CD160</th>
<th>CD127</th>
<th>CD11b</th>
<th>CD69</th>
<th>CXCR6</th>
<th>T-bet</th>
<th>Eomes</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Blood cNK (CD56++)</td>
<td>+</td>
<td>+</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>+</td>
<td>±</td>
<td>+</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>Blood cNK (CD56++)</td>
<td>+</td>
<td>+</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>+</td>
<td>±</td>
<td>∓</td>
<td>-</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>LP ILC1</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>n.d.</td>
<td>+</td>
<td>n.d.</td>
<td>+</td>
<td>n.d.</td>
<td>±</td>
<td>-</td>
<td>n.d.</td>
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</tr>
<tr>
<td>ile ILC1</td>
<td>±</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
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<td>-</td>
<td>+</td>
<td>±</td>
<td>+</td>
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<td>+</td>
</tr>
<tr>
<td>Liver ILC1</td>
<td>+</td>
<td>+</td>
<td>-</td>
<td>±</td>
<td>n.d.</td>
<td>n.d.</td>
<td>n.d.</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>±</td>
<td>+</td>
</tr>
</tbody>
</table>

± heterogenous expression; n.d., not determined.
the individual ILC1 subsets in regards to their phenotype and known functions. Furthermore, for some of these subsets, which appear to be ILC1s based on their characteristic cytokine production, a clear lineage relationship to other ILC1s or cNK cells has not firmly been established. For the purpose of this review, I will focus on some of the better-characterized members of the ILC1 lineage, which includes the subsets of ILC1s in the intestine and liver.

**Intestinal ILC1s**

Intestinal ILCs that express typical NK cell receptors – NKP46 and NK1.1 in the mouse; NKP46 and/or CD56 in humans – have in the past collectively been considered as cNK cells due to their NK cell-like phenotype and their ability to produce IFN-γ. However, evidence is now emerging that the intestinal mucosa contains several types of group 1 ILCs that are distinct from cNK cells. Two main subsets of intestinal ILC1s were recently identified in mice and humans: intraepithelial ILC1s (ieILC1s) and lamina propria-resident ILC1s (LP ILC1s) (18–20). ieILC1s are found in the intestinal epithelium and resemble cNK cells in their expression of canonical NK cell receptors. However, unlike resting cNK cells, ieILC1s express surface markers typical of intraepithelial T cells, such as the integrin CD49a, the activation marker CD69, the HVEM receptor CD160, and in humans, the integrin CD103 (19). To date, it is unclear whether this surface marker pattern is a result of tissue factors encountered in the intestinal epithelium, or is a characteristic of a separate lineage of cells. Like cNK cells, ieILC1s express the transcription factors T-bet and Eomes; however, studies in mice provided evidence that ieILC1s do not develop from a conventional NKP (discussed below in Section Development). Like most other ILC1s, ieILC1s are potent producers of IFN-γ, which they release in response to IL-12 and IL-15, and are likely to contribute not only to immune protection but also to chronic inflammation of the intestinal mucosa.

Murine LP ILC1s share characteristics with cNK cells and ieILC1s, including surface expression of NKP46 and NK1.1. However, unlike the latter two subsets, mouse LP ILC1s express high levels of the IL-7 receptor alpha chain (CD127). Furthermore, LP ILC1s are positive for CD27, Thy1, and c-kit, which are not typically expressed on mature cNK cells, and they are negative for most Ly49 receptors. Unlike cNK cells and ieILC1s, LP ILC1s only express T-bet but not Eomes (20). In human intestinal lamina propria, a CD127+ ILC subset with ILC1-like functions has recently been described. These cells lack CD56 and NKP46 but express CD161, a marker also found on some human cNK cells, ILC2s, and ILC3s (18, 21–23). These ILC1s are Eomes negative; subsets of these cells are T-bet positive and produce IFN-γ (18, 24).

**Liver ILC1s**

The liver contains cNK cells as well as ILC1s, the two of which can be distinguished by their different expression of surface markers and transcription factors. In mice, liver ILC1s are CD3ε− NK1.1+ DX5− cells with high expression of both the integrin CD49a (VLA-1) and the cytotoxicity-inducing ligand Trail. In contrast, cNK cells in the liver are DX5+ CD49a− Trail−. Both cNK and ILC1s in the liver express T-bet; however, like LP ILC1s, liver ILC1s do not express Eomes. Originally believed to represent immature cNK cells (25), liver-resident ILC1s are now recognized as a separate lineage of group 1 ILCs with different progenitor origin, cell trafficking, and functional capacities than cNK cells (26–28). Liver ILC1s are potent producers not only of IFN-γ but also of additional cytokines such as TNF-α, IL-2, and GM-CSF. Liver-resident ILC1s lack the typical cytotoxic machinery – perforin and granzymes A and B – that are common to cNK cells. In contrast, they express high levels of granzyme C and display potent TRAIL-mediated cytotoxicity (17, 29, 30). Human liver ILC1s have been identified as CD56hi CD16+ cells capable of producing IFN-γ upon activation (31). While the majority of these cells are CD49a Eomes+, a minor subset expresses CD49a− and lacks Eomes, and may thus represent the human counterpart of murine liver CD49a+ ILC1s (32). In mice, liver ILC1s are capable of generating immunologic memory to hapten and viral antigens and are capable of mediating robust recall responses upon rechallenge with the same antigen (16, 33), a feature that has not yet been demonstrated for any of the other known ILC1 subsets.

**Salivary Gland ILC1s**

In mice, a unique subset of NK-like cells resides in salivary glands (34–36). While these cells closely resemble cNK cells in both surface receptor and transcription factor expression, they also share several features with unconventional NK cells, and in particular, with liver ILC1s. Most notably, both salivary gland ILC1s and liver ILC1s express TRAIL and CD49a (34, 35). Unlike liver ILC1s; however, salivary gland ILC1s express DX5 and Eomes. Salivary gland ILC1s are similar to liver ILC1s in their ability to induce TRAIL-mediated cytotoxicity; however, in contrast to most other ILC1s, salivary gland ILC1s are poor producers of IFN-γ and thus do not entirely fit the classical definition of group 1 ILCs (34, 36).

**ILC1s in Other Tissues**

NK-like cells are also found in other non-lymphoid organs, such as the uterus, kidney, and skin of mice and humans (17, 37–41). These cells vary in their phenotype, transcription factor expression, and functions, which may in part be dictated by their specific tissue microenvironment. Common features of these NK-like cells are their relatively poor cytotoxic potential but potent production of cytokines and growth factors. In the pregnant uterus, non-classical NK cells are thought to play important roles during fetal implantation and vascular remodeling at the decidua (40–42). These uterine NK cells share features with cNK cells, such as the expression of certain activating and inhibitory NK cell receptors, and the expression of perforin and granzymes. However, their developmental origin is still debated (40, 41). In the uterus of virgin mice, several distinct cell populations with NK-like (NK1.1+ NKP46+ T-bet+) phenotype were recently described: DX5+ CD49a− cells that likely represent cNK cells as well as a DX5− CD49a+ putative ILC1 population that contains both Eomes+ and Eomes− subsets (17, 37). Further studies are needed to investigate the lineage relationships of these distinct uterine group 1 ILC subsets and their individual functions at this tissue site. DX5− CD49a+ ILC1s with similarity to liver ILC1s were also described in murine skin and kidneys (17, 38). In the murine lung, cNK cells have been studied extensively (5); however, it is currently unknown whether lung tissue also contains distinct
ILC1 subsets. In human lung, a putative ILC1 subset has recently been identified (43); however, further studies are required to confirm its specific phenotype and to investigate its functions.

**ILC Plasticity**

Further complicating the identification of ILC1s, a subset of group 3 ILCs in murine intestinal tissues, termed NCR$^+$ ILC3s, was recently shown to convert to IFN-γ-producing, T-bet-expressing cells during microbial infections (20, 21, 44). Conversion of these ILC3s is accompanied by their downregulation of their specific transcription factor RORγt, resulting in a cell population that resembles ILC1s in their phenotype and cellular functions. Due to their different lineage relationship compared to ILC1s, these converted ILC3s, termed “ex-RORγt ILCs,” are not typically regarded as a “true” ILC1 subset. Additional ILC plasticity was recently identified for a subset of human tonsil ILC1s, which in culture with ILC3-polarizing cytokines acquired RORγt expression and switched their cytokine profile from predominantly IFN-γ to the ILC3-specific cytokine IL-22 (24). This conversion was not seen in mice, as LP ILC1s were unable to convert to ILC3s; however, “ex-RORγt ILCs” were capable of reverting back to RORγt-expressing ILC3s (24).

**ILC1 Signature Genes**

Transcriptional profiling of ILC1s and cNK cells from liver, small intestine, and spleen recently demonstrated that the gene expression by these tissue ILC1s clearly differs from that of cNK cells. Principle-component analysis from this study placed the tested ILC1 subsets closer to each other than to cNK cells from the same tissues. However, in addition to shared gene signature, each ILC1 subset possesses a tissue-specific transcriptional signature (43). In contrast, liver and spleen cNK cell transcriptional profiles were indistinguishable, indicating that cNK cells from different tissues represent a relatively homogeneous population. Despite the differences between individual ILC1 subsets, common signature genes were identified that distinguish ILC1s from cNK cells; among these are higher transcript levels for CD127, the chemokine receptor CXCR6, and, interestingly, for some T cell receptor (TCR) chains, although no TCR chain protein expression was detected either intracellularly or on the cell surface. The latter finding is intriguing as it suggests common transcriptional programs between ILC1s and T cells.

**DEVELOPMENT**

All ILCs develop from a common lymphoid progenitor (CLP), which further differentiates into a shared ILC precursor, termed α-lymphoid precursor (αLP) (or common ILC progenitor, CILP). This shared precursor can be identified by its expression of the integrin α4β7 and of CD127. Further differentiation of αLP generates at least two different precursors with more restricted lineage repopulation capacity: a common helper ILC precursor (CHILP) that can give rise to ILC1s and all other ILC subsets except cNK cells, and an NKP, which can differentiate into cNK cells but no other ILCs (2, 46–48) (Figure 1). Differentiation of these precursors from CLP requires the acquisition of the transcription factors Id2, NFIL3, and Tox; mice deficient in any of these factors show greatly reduced numbers of all ILCs (49–55). Additionally, the Runx family of transcription factors and in particular, Runx3 is required for normal development of group 1 and group 3 ILCs (56, 57). The development of mature ILC1s and cNKs from CHILPs and NKP, respectively, depends on a variety of additional transcription factors and cytokines, many of which have only very recently been elucidated. The following sections discuss the individual requirements for the differentiation of cNK cells and ILC1s from these early precursors.
Development of cNK Cells

Direct precursors to cNK cells (NKp) were originally defined as lineage-negative (Lin−) cells that express the IL-2 receptor beta chain (CD122) but none of the typical NK cell receptors. This precursor population, however, was later found to include cells with T cell differentiation potential and was subsequently refined to include only Lin−CD122+ cells that co-expressed CD244, CD27, and CD127. The generation of NKp from CLPs is dependent on the transcription factors NFIL3, Id2, and Tox, which are also involved in the development of CHILPs for ILC development (1, 2). Unlike other ILCs, however, development of cNK cells does not include a precursor that expresses the transcription factor PLZF. As recent studies demonstrated, PLZF is specifically expressed during ILC differentiation from CHILPs, allowing a further refinement of the differences between NKp and ILC1 precursors. PLZF lineage-tracing experiments revealed that true NKp have the phenotype Lin−CD122+CD244+CD27+CD127+PLZF−α4β7+, while ILC1 precursors are characterized by additional expression of PLZF and α4β7 (58).

Further differentiation of NKp gives rise to immature cNK cells, a process that is accompanied by gradual acquisition of the NK cell receptors NK1.1 and NKp46 and the transcription factors T-bet and Eomes. Immature cNK cells can be identified as DX5−CD27+CD11b− cells, which are T-bet+ but Eomes− (23, 25, 59). With these marker characteristics, immature cNK cells have striking similarity to ILC1s, making it difficult to differentiate between the two populations. Maturation of cNK cells is dependent on the cytokine IL-15 and is accompanied by the acquisition of DX5, upregulation of CD11b, and loss of CD27 expression. The cytokine IL-7, which is crucial for the development of group 2 and 3 ILCs, is not required for mature cNK cell or ILC1 development (20, 26, 51). cNK cell maturation is dependent on Eomes and T-bet, as mature cNK cells are absent in Eomes-deficient mice and reduced in numbers and with less mature phenotype in T-bet-deficient mice (25, 60).

Development of ILC1s

It has been proposed that ILC1s, like group 2 and 3 ILCs, develop from a CHILP that further differentiates into an ILC precursor (ILCp) in a PLZF-dependent process (48, 58). Lineage tracing for PLZF expression revealed that all ILCs, with the exception of cNK cells and a subset of ILC3s (LTi-like ILC3s), develop from a PLZF-positive precursor (48). Among the investigated group 1 ILCs, liver ILC1s and intestinal iILC1s, but not splenic or liver cNK cells, were lineage traced for PLZF expression, and thus, the former two are likely to arise from the same ILC precursor (48). Other ILC1s such as LP ILC1s or salivary gland ILC1s were not interrogated in this study (48). Interestingly, PLZF-deficient bone marrow can still generate normal numbers of iILC1s (48), indicating the existence of alternative pathways for ILC1 differentiation, which may involve branching from either PLZF−CHILP or NKp. Within ILCp, the highest ILC1-generating capacity was found in a CD122+ subset that lacks the chemokine receptor CXCR5, suggesting that these CD122+ CXCR5+ ILCp may represent the direct precursors to ILC1s (61).

Differentiation of liver ILC1s and intestinal ILC1s (both LP ILC1s and iILC1s) further depends on the transcription factor T-bet (17, 19, 20, 25, 62). In contrast to cNK cells, which require Eomes for their final maturation, liver ILC1s and presumably all other Eomes− ILC1s develop independently of this transcription factor (25). Interestingly, there is remarkable diversity among ILC1 subsets in relation to their dependence on other transcription factors and cytokines. Although NFIL3 is required for the development of most group 1, 2, and 3 ILCs, including cNK cells and intestinal iILC1s, mice deficient in this transcription factor have normal numbers of salivary gland, uterine, and kidney-resident ILC1s (17, 34, 38). In regards to liver ILC1s, the precise role of NFIL3 remains to be determined, as currently conflicting data exists showing either dependence (49, 55) or independence of liver ILC1s on this transcription factor (17, 63). Similarly, dependence on the transcription factor GATA-3 appears to vary between the individual ILC1 subsets. GATA-3 deficiency severely reduces numbers of intestinal LP ILC1s but has no effect on iILC1 or cNK cell numbers (64). As development of all CD127+ group 2 and group 3 ILCs is dependent on GATA-3, it has been suggested that only CD127+ ILC1s but not CD127− ILC1s require GATA-3 for their development (20, 64).

Development of most group 1 ILCs, including cNK cells, is dependent on the cytokine IL-15. We recently found that in contrast to all other ILC1s, intestinal iILC1s were only partially dependent on IL-15, as their numbers were reduced but not absent in mice deficient in the IL-15 receptor alpha chain (19). Although some ILC1 subsets express the IL-7 receptor CD127, IL-7 appears dispensable for ILC1 development, as mice deficient in IL-7 or its receptor CD127 have normal numbers of intestinal LP and liver ILC1s (20, 26, 51).

The differences in transcription factor and cytokine requirements within ILC1s may be a result of different lineage relationships between these individual subsets. Further studies are needed to identify the individual precursors and their location (bone marrow vs. development in peripheral tissues) for each of these ILC1 subsets and elucidate potential pathways of conversion within group 1 ILCs and potentially between other ILCs.

ILC1s in Antimicrobial Responses

The prototypic function of group 1 ILCs is potent expression of IFN-γ upon activation with cytokines or surface receptor crosslinking. IFN-γ plays important roles in the immune defense to intracellular pathogens, and cNK cells have been recognized for their critical functions in the immune defense against a variety of viral and bacterial pathogens (7, 9). In most of these studies, all cells with NK-like surface phenotype and ability for IFN-γ production were regarded as cNK cells. However, it appears likely that in some of the earlier studies, ILC1s may have contributed to IFN-γ production but were not recognized as a separate lineage. Thus, specific roles of ILC1s, as compared to cNK cells, during immune defenses to pathogens are only now being investigated.

The identification of specific markers for ILC1s, such as CD49a expression in liver or expression of CD127 and lack of Eomes in LP ILC1s, has made it possible to more accurately assess specific host protective roles of ILC1s compared to cNK cells. However, as marker expression can change during cellular activation, results
still need to be interpreted with caution. Recent studies have focused on the potential roles of intestinal ILC1s during protective responses to intestinal pathogens. New findings in this respect suggest that following oral infection of mice with the intracellular pathogen *Toxoplasma gondii*, ILC1s produce the majority of IFN-γ, as well as TNF-α, while cNK cells and ILC3s contribute to a lesser extent (20). In this particular infection model, Diefenbach and colleagues identified ILC1s as the cell type responsible for controlling infection, which they accomplish by their rapid attraction of inflammatory monocytes to the site of infection. Unfortunately, due to the lack of specific ILC1-knockout mice or depleting antibodies, it is currently impossible to unequivocally attribute a particular function to either ILC1s or cNK cells. In the mentioned study on host responses to *T. gondii* infection. However, it should be noted that T-bet also plays roles in cNK cell maturation and IFN-γ production (60); thus, the failure of T-bet-deficient mice to control parasite burden may still partially be due to defects in cNK cell functions. Additionally, T-bet is also required for the development of NCR+ ILC3s and their conversion into IFN-γ-producing cells (65).

During acute infection with the intestinal pathogen *Clostridium difficile*, ILC1s represent the major innate cell subset to respond with IFN-γ production in the LP, epithelium, and MLN of infected mice (66). Consistent with this finding, Pamer and colleagues further demonstrated that mice lacking ILC1s as well as mice deficient in IFN-γ were more susceptible to lethal *C. difficile* infection. For this study, T-bet-deficient mice on a Rag-deficient background were used to deduce a specific role for ILC1s during the immune defense, and adoptive transfer of T-bet+ CD127+ LP ILC1s to these mice imparted partial protection from lethal infection. However, as ILC1 transfer only provided partial rescue, the authors speculate that the contribution of other innate cells, such as cNK cells, neutrophils and monocytes, is likely required for protective IFN-γ responses to *C. difficile* (66).

IFN-γ is also a major factor in the immune defense to intestinal infections with *Salmonella enterica* serovar *Typhimurium*. A recent study demonstrated that a group of lamina propria ILCs was shown to provide the majority of IFN-γ during acute infection. However, in this study, it was concluded that converted ILC3s, rather than true ILC1s, were responsible for the majority of the produced IFN-γ (65). As illustrated in the examples above, there is emerging evidence that intestinal ILC1s contribute to the immune defense to microbial pathogens. However, to date, an accurate assessment of the individual roles of ILC1s, cNK cells, and converted ILC3 remains impossible. Further exploration of novel markers and transcription factors involved in ILC1 development and functions may aid in further elucidating their specific roles during host protective responses.

**ILC1s IN TISSUE INFLAMMATION AND AUTOIMMUNITY**

The production of pro-inflammatory cytokines by group 1 ILCs has important functions during antimicrobial immune responses. However, exaggerated or prolonged cytokine responses can also lead to chronic inflammation and autoimmunity. Several studies demonstrated that inflamed intestinal tissues from patients with Crohn's disease harbor larger numbers of ILC1s, suggesting a role for ILC1s in inflammatory pathology (18, 19). We recently investigated the role of ILC1s in a mouse model of colitis induced by anti-CD40 injection into Rag-deficient mice. In this model, IFN-γ is known as the major factor driving wasting disease and systemic inflammation (67). We find that in this model of inflammatory bowel disease, intestinal iILC1s contribute to intestinal pathology through production of IFN-γ (19). Similarly, mice with a human immune system show accumulation of IFN-γ-producing human ILC1s in the inflamed intestine upon challenge with the colitis-inducing agent dextran sodium sulfate (18, 19).

A recent study by Victorino et al. demonstrated that ILC1s contribute to organ dysfunction seen in a mouse model of ischemic kidney injury. The authors of this study identified NK1.1+ non-T cells as the major culprit that mediate kidney dysfunction following ischemia–reperfusion injury, and found that depletion with anti-NK1.1, which depletes both cNK cells and ILC1s in kidneys, ameliorated disease, whereas anti-asialo-GM1 treatment, which preferentially depletes cNK cells, did not protect from disease (38). The exact mechanism for this ILC1-mediated tissue damage awaits further investigation.

In contrast to the potential detrimental roles of ILC1s described above, ILC1s in the salivary gland appear to play tissue-protective functions during chronic infection. ILC1s at this tissue site show low ability for IFN-γ production; however, they can induce cytotoxicity through their expression of TRAIL. Two recent studies demonstrated that during chronic infection with murine cytomegalovirus, ILC1s protected from autoimmunity by regulating both innate and adaptive immune responses in the salivary gland. In these studies, salivary gland NK1.1+ cells were shown to preserve gland functions by limiting eosinophil infiltration and by preventing T cell-mediated autoimmunity through TRAIL-mediated cytotoxicity toward activated CD4 T cells (35, 68). Although individual roles of cNK cells vs. ILC1s were not investigated in these studies, a contribution of salivary gland-resident ILC1s appears likely, in particular in relation to the observed TRAIL-mediated regulatory roles.

**OPEN QUESTIONS**

The ILC1 field is still in its infancy, and many questions remain as to their specific developmental pathways, and to the lineage relationship between the to-date characterized ILC1s subsets. For instance, the exact molecular processes that mediate lineage specification of ILC1 vs. cNK cells during the branch point from μLP to either CHILP or NKP are insufficiently understood. Furthermore, for several of the putative ILC1 subsets, a lineage relationship to ILC1s awaits confirmation. Examples of those subsets include salivary gland ILC1s, uterine ILCs, and subsets of dermal ILCs. Additional NK-like cells with non-typical phenotype and functions have been described, such as CD127+ blood and splenic ILC1s and GATA-3-dependent thymic NK cells (13, 45, 69, 70).
Furthermore, to date, the extent of ILC plasticity is not clear. As discussed above, ILC3s can convert to ILC1-like cells during inflammatory conditions. A recent study now also provided evidence that a subset of human ILC1s can gain ILC3-like phenotype and function (24). Further studies will be required to define whether this ILC1 subset represents a precursor population that differentiates into ILC3s, or a mature ILC1 population that undergoes conversion to ILC3s, and to investigate the mechanism and functional relevance of this ILC1 to ILC3 conversion.

The roles of ILC1s vs. cNK cells during protective host responses and chronic inflammation are only now starting to be investigated and there is little evidence of their beneficial or pathogenic functions during these processes. Group 2 and 3 ILCs play important roles both as initiators of local inflammation, as well as in the resolution of inflammation and restoration of tissue integrity (71). Further studies are needed to elucidate the specific contributions of ILC1s in these processes, and to investigate potential crosstalk between ILC1s with other ILCs and with adaptive immune cells during the initiation and resolution of inflammation.

AUTHOR CONTRIBUTIONS

AF researched the literature and wrote the article.

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REFERENCES


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