Innate Lymphoid Cell Activation and Sustained Depletion in Blood and Tissue of Children Infected with HIV from Birth Despite Antiretroviral Therapy

Highlights

- ILCs are depleted in children born with HIV-1 infection
- Only initiation of antiretroviral therapy at birth prevents ILC depletion
- Blood ILCs in HIV-1 infected children upregulate activation and cellular metabolism genes
- Tonsil-resident ILC3s and NK cells exhibit proliferation and innate immune signaling

Authors
Alveera Singh, Samuel W. Kazer, Julia Roider, ..., Philip Goulder, Alasdair Leslie, Henrik N. Kløverpris

Correspondence
henrik.kloverpris@ahri.org

In Brief
ILCs are dysregulated during HIV-1 infection in adults, but their fate in children is unknown. Singh et al. demonstrate that circulating and tonsil-resident ILCs are depleted in children infected with HIV-1 since birth. Transcriptionally, ILCs exhibit cell-type- and compartment-specific activity in HIV-1 infected children compared to healthy controls.

Singh et al., 2020, Cell Reports 32, 108153
September 15, 2020 © 2020 The Author(s).
https://doi.org/10.1016/j.celrep.2020.108153

Cell Press
Innate lymphoid cells (ILCs) are important for response to infection and for immune development in early life. HIV infection in adults depletes circulating ILCs, but the impact on children infected from birth remains unknown. We study vertically HIV-infected children from birth to adulthood and find severe and persistent depletion of all circulating ILCs that, unlike CD4+ T cells, are not restored by long-term antiretroviral therapy unless initiated at birth. Remaining ILCs upregulate genes associated with cellular activation and metabolic perturbation. Unlike HIV-infected adults, ILCs are also profoundly depleted in tonsils of vertically infected children. Transcriptional profiling of remaining ILCs reveals ongoing cell-type-specific activity despite antiretroviral therapy. Collectively, these data suggest an important and ongoing role for ILCs in lymphoid tissue of HIV-infected children from birth, where persistent depletion and sustained transcriptional activity are likely to have long-term immune consequences that merit further investigation.
We studied a total of 229 newborns (NBs), children, and adults and mapped the circulating and tissue-resident ILC and NK cell response to HIV infection from birth; we observed striking ILC depletion in both the blood and tonsil tissue of infected children. Moreover, we found distinct cell-type-specific transcriptional changes in activation and metabolism, suggesting a potentially important role for innate lymphocytes in response to HIV infection in early life.

**RESULTS**

**Helper ILCs in Blood Display a Distinct Transcriptional Profile Compared to NK Cells and Are Enriched in Children Compared to Adults**

To evaluate the role of ILCs in pediatric HIV infection, we first established a gating strategy to analyze ILCs and NK cells in blood based on distinct phenotype expression (Lim et al., 2017; Spits et al., 2013). We excluded lineage+ cells and used ILC- and NK-specific markers to simultaneously identify three different ILC (ILC1, ILC2, ILC pre-cursors [ILCPs]) (Lim et al., 2017) and two different NK cell populations (NK CD56-high and NK CD16-high) (Figure 1A). We performed RNA sequencing (RNA-seq) of sorted ILC2, ILCP, NK CD56-high, and NK CD16-high populations from 10 pediatric participants (median age 10.8 years, interquartile range [IQR] 6.4–11.9 years; Table S1). Principal component analysis (PCA) using 497 differentially expressed genes (DEGs) (false discovery rate [FDR]-corrected q < 0.01; see Method Details) demonstrated clean separation of these ILC subsets (Figure 1B; Table S2). Genes separating the subsets include granzyme B (GZMB), IFN-γ (IFNG), CD16 (FCGR3A), and KIR2DL1 expressed in NK populations; and high levels of KLRC1 (CD161), KLRC2, and KLRL1 (ST2), which binds IL-33 for activation in ILC2s (Figure 1C; Table S2). Thus, our flow cytometry panel successfully identifies the main ILC and NK cell subsets in pediatric blood, which also display the canonical gene signatures observed in adults.

Because the relative frequencies of many blood immune subsets change across the course of the normal lifespan (Prendergast et al., 2012; Shearer et al., 2003), we first studied ILC and NK levels in HIV-uninfected individuals spanning birth, childhood, and adulthood, in each case using samples from sub-Saharan African cohorts in Durban, South Africa (Table 1). Overall, among 138 HIV-uninfected individuals with an age range of 0–24 years, we found a strong reduction in the frequency of all ILC subsets with age (Figure 1D), whereas NK cell populations remained relatively stable (Figure 1E), consistent with a recent study (Vély et al., 2016). Together, these data define the circulating ILC populations present in children from sub-Saharan Africa and establish their normal frequencies in the absence of HIV infection.

**Depletion of All Circulating ILC Subsets in Treatment-Naïve HIV-Infected Children Irrespective of Disease Control**

Mortality among vertically HIV-infected children exceeds 50% by the age of 2 years in the absence of ART (Marston et al., 2011; Newell et al., 2004). Here, we studied a group of 26 untreated vertically transmitted children that survived to the age of minimum 5 years in two different groups: (1) PSPs (median...
CD4 = 37%, IQR 28%–44%, n = 15) and (2) pediatric progressors (PPs) (median CD4 = 15%, IQR 8%–28%, n = 11) who met World Health Organization (WHO) criteria prevailing at the time of the study to initiate antiretroviral treatment (Table 1; Figure 2A). Total ILC frequencies and all individual helper ILC subsets (ILC1, ILC2, and ILCPs) were significantly decreased compared to uninfected controls. However, there were no significant differences between PSPs and PPs (Figure 2B). Overall, the NK populations showed a similar pattern of depletion, though mostly found within the CD56<sup>high</sup> NK subset (p < 0.0002) (Figure 2C). Thus, all helper ILCs and NK cells are severely depleted in children infected with HIV at birth, even among the rare group of PSPs, who maintain normal-for-age CD4 levels (Figure 2A) despite being ART naive.

ILC Depletion in the Blood Is Sustained Despite Viral Suppression by ART but Can be Prevented by Immediate ART Treatment Initiation at Birth

We have previously reported that long-term ART in chronic adult infection is unable to restore circulating ILC levels (Kløverpris et al., 2016). However, HIV-infected children appear to possess a superior ability to restore adaptive immune function compared to adults (Lewis et al., 2012; Picat et al., 2013). Therefore, we next investigated the impact of long-term ART in pediatric HIV...
infection in a cohort of children (n = 38) treated for a median of 88 weeks (IQR 40–218) (Table 1). As in adult HIV infection, ART failed to restore total ILCs, ILC1, ILC2, ILCPs, and NK cells (Figure 3A). Longitudinal sampling of the PP cohort (n = 9) over three time points from a median of 12 weeks before treatment and again at 42 and 84 weeks after treatment initiation showed the same trend. Although a modest increase in both NK CD56high cells and ILCPs was observed, neither of which reached levels of uninfected controls (Figure 3B). This lack of ILC or NK cell reconstitution was confirmed in additional longitudinal cohorts undergoing viral suppression by ART or maintaining high CD4 levels in the absence of treatment over 84 weeks (Figure S1).

Our previous work demonstrated that early treatment of HIV-infected adults, prior to peak viremia, prevents significant loss of circulating ILCs (Kløverpris et al., 2016). To test this, we next examined the effect of early treatment initiation on circulating ILC subsets in a cohort of 27 in-utero-infected newborns (NBs), initiated on ART within minutes after birth (Adland et al., 2020). ART was initially prophylactic (nevirapine and zidovudine) and then increased to triple therapy treatment from a median of 7 days post-partum (range 0–18 days), with all individuals remaining undetectable for plasma HIV RNA at 21 months of age (Table 1). Strikingly, in these individuals, we found no difference in CD4+ T cell, ILC, or NK cell levels compared to age-matched HIV-uninfected children (Figure 3C). Thus, immediate ART initiation of in-utero-infected NBs preserves ILC and NK cell levels, which is consistent with that seen in adult horizontal HIV infection (Kløverpris et al., 2016).

Peripheral ILCs and NK Cells Are Transcriptionally Activated in the Blood of HIV-Infected Children

To investigate whether blood ILCs in children are modulated by HIV infection, we measured surface expression CD69 and FAS (CD95) and cytokine production (IL-2, IL-4, IL-5, IL-13) following mitogen stimulation on ILC2, ILC3, NK CD56high, and NK CD16high subsets. HIV had no impact on expression of CD69 and Fas on ILC2s and ILCPs, while NK CD56high and NK CD16high subsets displayed increased expression of Fas and CD69 (Munneke et al., 2014) (Figure S2A), indicating that they are stimulated by HIV infection. In the mitogen stimulation assay of ILC2s, we found no impact of HIV infection on cytokine production (Figure S2B). To further explore the impact of HIV on ILCs in circulation in infected children, we performed RNA-seq on ILC2, ILCP, NK CD56high, and NK CD16high subsets and CD4+ T cells from three ART-naive viremic PSpS, three ART-treated virally suppressed children, and four age-matched HIV-uninfected children (see Table S1 for replicate numbers and Method Details). Differential gene expression analyses on each subset between HIV-infected children (PSp+ ART+/–) and HIV-infected controls demonstrated >300 DEGs (FDR-corrected q < 0.1) in each of the CD4+ T cells, ILC2s, ILCPs, and NK CD16high cells and 61 DEGs in NK CD56high cells (Figure 4A; Table S3). Comparisons between both virally suppressed and viremic HIV-infected children and between virally suppressed HIV-infected children and HIV-uninfected controls revealed few transcriptional changes in all cell subsets (<57 and <49, respectively), suggesting that persistent viremia drives transcriptional changes in circulating ILCs (Figure 4B; Table S3).

Analysis of the DEGs observed in all subsets is consistent with the activation and loss of regulatory function of ILC subsets in chronically HIV-infected children. We note the following DEGs in each subset comparing HIV-infected children to healthy controls (Figure 4C; Table S3): (1) ILC2s: downregulation of inflammatory markers PTGS2, NFKBID, and TNF and upregulation of anti-apoptosis marker BIRC3, H4 clustered histone 3 (HIST1H4C), and SUB1, a transcript factor that mediates RNA polymerase binding; (2) ILCPs: downregulation of ILC3 lineage transcription factor RORC (Serafini et al., 2015) and upregulation of pro-survival markers (AXL, MET) and cytokine-induced transcription factor STAT4; (3) NK CD16high: downregulation of the mTORC-activating factor KLHL22 (Chen et al., 2018) and upregulation of TNFRSF9 (CD137), an activation marker of NK cells (Baessler et al., 2010) known to promote T cell expansion (Wilcox et al., 2002), and GK encoding glycerol kinase, an essential enzyme for glycerol conversion; and (4) NK CD56high: downregulation of canonical transcription factors involved in negative regulation of cellular proliferation and differentiation (DUSP6, FOS) and upregulation of MORN3, which recruits p53 and other transcription machinery (Sloan et al., 2016; Takahashi et al., 2007), and IFI30, an IFN-stimulated response gene. Moreover, genes involved in active metabolism like COX7A2, NDUFb6 (ILC2), NDUFa12 (ILCP), ACSL1 (NK CD16high), and DNAJ4 (NK CD56high) were upregulated in HIV-infected children, and then increased to triple therapy treatment from a median of 7 days post-partum (range 0–18 days), with all individuals remaining undetectable for plasma HIV RNA at 21 months of age (Table 1). Strikingly, in these individuals, we found no difference in CD4+ T cell, ILC, or NK cell levels compared to age-matched HIV-uninfected children (Figure 3C). Thus, immediate ART initiation of in-utero-infected NBs preserves ILC and NK cell levels, which is consistent with that seen in adult horizontal HIV infection (Kløverpris et al., 2016).

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suggesting active metabolism in these subsets in response to infection.

Gene set analysis using ingenuity pathway analysis (IPA) highlighted oncogenes ESR1, RET, and MYC as significantly enriched and upregulated upstream drivers potentially inducing the transcriptional changes in all ILC subsets (except for ESR1 in ILC2s and MYC in NK CD16<sup>high</sup> cells) between HIV-infected children and healthy controls (Figure 4D; Table S4). Consistent with a role for IFN in chronic HIV infection (Roff et al., 2014), IFN-γ is also significantly enriched, though lacking in ILC2s. These upstream drivers are corroborated by enrichment for genes associated with protein ubiquitination in all ILC subsets in downstream pathway analysis (Figure 4E; Table S4). Surprisingly, both ILC2s and ILCPs exhibit significant enrichment for pathways annotated for defects in cellular metabolism—mitochondrial dysfunction and unfolded protein response, respectively. To confirm these metabolic gene signatures, we performed gene set enrichment analysis (GSEA) on the DEGs from each ILC subset using the Gene Ontology (GO) and KEGG databases (see Method Details). In both ILC2s and ILCPs, several GO and KEGG terms encompassing cellular activation and metabolism were positively enriched (Figure 2C; Table S4). GSEA on NK CD16<sup>high</sup> DEGs implicated pathogen recognition receptor signaling and response to stress, while no terms were significant for enrichment from NK CD56<sup>high</sup> DEGs (Method Details). Together, these data demonstrate that peripheral ILC2s, ILCPs, NK CD16<sup>high</sup>, and NK CD56<sup>high</sup> subsets all express activation gene programming in HIV-infected children compared to HIV-uninfected controls. Moreover, enrichment results suggest differences in cellular metabolism in ILC2s and ILCPs.

Reduced ILC and NK Levels in Tonsils from HIV-Infected Children

While the role and function of helper ILCs in the blood are unknown, ILCs play a key role in human lymphoid tissue development (Koues et al., 2016; Vély et al., 2016) and in response to inflammation (Bernink et al., 2013). Using tonsils from children undergoing tonsillectomy as a source of secondary lymphoid tissue (Table S5) (Roider et al., 2019), we identified six different innate lymphocyte populations from lineage-negative cells (Figure S3A), which were dominated by NK cells.
and ILC3s (Figures S3B and S3C). We found differential expression of CD103 and CD69, described as surrogate markers for tissue residency (Masopust and Soerens, 2019; Skon et al., 2013) (Figure 5A). Overall, CD127– NK cells and NKp44+ ILC3s expressed higher levels of these markers of tissue residence compared to ILC1, ILC2, and CD127+ NK cells, consistent with mouse model experiments (Gasteiger et al., 2015) (Figure 5B). To test if tonsil-resident ILCs also were reduced by HIV infection, we compared the relative frequency of each of the tonsil NK and ILC subsets from 12 HIV-negative children to that of 4 ART-treated HIV-positive children in whom the ART initiation was unknown, of which 3 were virally suppressed (viral load <20 copies/ml plasma) with detectable antiretroviral drugs in plasma (Table S5). We found highly significant depletion for each ILC and NK subset (Figure 5C) but, surprisingly, no significant depletion of bulk CD4+ T cells or PD-1++CD69+ T-follicular helper-like cells in the same subjects (Figure S3D), in contrast to our previous work in adult tonsils where we found no HIV-associated ILC depletion (Kloverpris et al., 2016). Although the number of HIV-infected children studied here is small, these data suggest that HIV infection from birth has a more severe impact on tissue-resident ILCs than does infection in later life.

Transcriptional Profiling of NK and ILC3 Cells in Tonsils from HIV-Infected Children Reveals Subset-Specific Activation in Response to Infection

Next, to study the ILC and NK cell responses to HIV infection in pediatric tissue, we purified the dominant ILC (ILC3 NKp44–, ILC3 NKp44+) and NK cell subsets (NK CD127–) from pediatric tonsils, while insufficient cell numbers were available from ILC1 and ILC2 subsets (see Figure S3A), and performed transcriptional profiling directly ex vivo to characterize both the transcriptional differences between the subsets and their responses to HIV infection (Figure 6). In HIV-uninfected controls, a limited number of genes separated the two ILC3 subsets defined by NKp44 surface expression (238 DEGs, FDR-corrected q < 0.1) compared to >1,000 genes for both ILC3 subsets compared to NK CD127– cells (Figure 6A; Table S6). In the top 20 regulated genes between ILC3 subsets were NCR2, which encodes NKp44, and S1PR1 (CD69), which is associated with tissue residency (Skon et al., 2013); both are consistent with protein surface expression on ILC3 subsets (see Figure S3A). Comparison of the NK CD127– and ILC3 subsets showed strong differential expression of canonical genes known to be upregulated in ILC3s (ICAM1, ICOS, IL1R1, RORC, IL17RE, KIT, ICOS, CD83) whereas NK CD127– cells expressed genes...
involved in cytotoxicity (\textit{GZMA, GZMB, CCL5, GNLY, PRF1, IFNG}) together with expression of canonical NK cell surface molecules (\textit{FCGR3A, KLRF1, KLRG1}). In addition, we found ILC3-specific upregulation of genes associated with homing to lymphoid follicular zones (\textit{CXCR5, CXCR4, CCR6}) and regulation of adaptive immunity (\textit{ICOS, CD40L, IFNGR2}). These data are consistent with previous analysis in human tonsils (Björklund et al., 2016; Cella et al., 2019; Koues et al., 2016) and suggest that ILC3 and NK subsets have distinct functions in human lymphoid tissue as regulators of tissue homeostasis and killing potential, respectively.

To determine the response of each of these subsets to HIV infection in children, we compared the transcriptional response in ILC3 NKp44–, ILC3 NKp44+, and NK CD127– cells in tonsils from two virally suppressed ART-treated HIV-infected children with five age- and gender- (female) matched HIV-uninfected children (see Table S6). We found 195, 286, and 75 DEGs (FDR-corrected \(q < 0.1\)) for ILC3 NKp44–, ILC3 NKp44+, and NK CD127– cells, respectively (Figure 6B; Table S7). We note the following DEGs in each subset comparing HIV-infected children to healthy controls (Figure 6C; Table S7): (1) ILC3 NKp44–: upregulation of \textit{NF-\(\kappa\)B co-activator PRMT6} (Di Lorenzo et al., 2014), cell-cycle antigen \textit{PCNA}, and \textit{CD38}, which is associated with activation in T cells, but previously undescribed on ILCs; (2) ILC3 NKp44+: downregulation of \textit{NF-\(\kappa\)B inhibitor MAST2} (Xiong et al., 2004) and upregulation of cell-cycle-associated \textit{TUBB2A}, anti-inflammatory \textit{ANXA1} (Arcone et al., 1993), and lineage commitment protein \textit{ID1}, whose family member \textit{ID2} is known to regulate ILC differentiation in tissue (Zhong and Zhu, 2017); and (3) NK CD127–: upregulation of secreted pattern recognition receptor lectin \textit{CLEC18A} shown to be associated with hepatitis C infection and Notch signaling pathway genes \textit{JAG2} and \textit{NOD2}. Notch signaling has been shown to upregulate \textit{killer immunoglobulin-like receptor} (KIR) expression and drive maturation in NK cells (Felices et al., 2014), suggesting that this NK CD127– subset may play an extended effector role in the tonsil in HIV infection.

To examine the \textit{CD4+ T cell} response to HIV infection in pediatric tonsils, we sorted four different \textit{CD4+ T cell} populations based on \textit{CD103/PD-1} expression (Figure S4A) accounting for
the majority of CD4+ T cells with distinct gene expression to that of innate lymphocytes (Table S7). The PD-1++CD103− subset expressed high levels of the canonical T follicular helper (Tfh) cell genes \(\text{CXCR5} \) and \(\text{PDCD1}\) and low levels of \(\text{ITGAE}\) (CD103), reflecting the sorted phenotypes by flow (Figure S4B). Using samples collected from a separate set of children given limited sample availability (see Table S5), we compared gene expression in each CD4 T cell subset between HIV-infected (n = 3) and HIV-uninfected (n = 5) children (see Method Details for design). We found limited numbers of DEGs in all four CD4 T cell subsets (Figure S4C; Table S7), suggesting limited response from CD4+ T cells compared to that seen for ILCs and NK cells.

Gene set analysis distinguished the responses to HIV infection by subset, with unique potential upstream drivers and pathways significantly enriched in each (Figures 6D and 6E; Table S7). While the ILC3 NKp44− subset expressed DEGs associated with innate and cytokine-mediated immune signaling potentially induced by broad transcriptional activator \(\text{SP1}\) and epigenetic regulator \(\text{HDAC3}\), the enrichment on the DEGs in the ILC3 NKp44+ subset reflected general cellular activation and proliferation with potential activation by G-coupled protein receptors and nuclear transcription factor \(\text{HFN4A}\). Enrichment on the NK CD127− subset demonstrated a consistent role for NK cells in the tonsil as innate immune sentinels during HIV infection, with putative upstream driver \(\text{PTPRM}\), known to regulate cellular adhesion. GSEA using the KEGG and GO gene sets found significant enrichment only in the ILC3 NKp44− subset, supporting broad cellular activation and proliferation with significant GO terms encompassing protein complex assembly, signal transduction by \(\text{p53}\), and biogenesis (Figure S4D; Table S8).

In summary, the ILC and NK cells responses to pediatric HIV infection were cell-subset specific with distinct activation and proliferation programming. CD4+ T cells in the tonsil, on the other hand, did not demonstrate strong transcriptional differences between HIV-infected and HIV-uninfected children. While the ILC3 NKp44+ subset did not exhibit immune activation by flow cytometry, both the ILC3 NKp44− and NK CD127− subsets upregulated machinery associated with immune response and innate immune signaling. Together with the profound depletion of ILCs and NK cells observed in vertically infected children, these data suggest that HIV infection from birth has a persistent impact on innate lymphocytes within secondary lymphoid organs that may have important consequences for their downstream immune health.
DISCUSSION

The past decade has established ILCs as key players in orchestrating tissue homeostasis and repair (Vivier et al., 2018). While HIV infection is known to cause irreversible changes in some human tissues (Deeks et al., 2013), the impact on ILC number and function remains incompletely understood, especially following vertical HIV infection in blood and tissue during early life. We hypothesized that innate lymphocyte responses may be particularly important in pediatric HIV infection while the adaptive immune response is still developing.

Here, we studied 229 individuals spanning the time from birth to adulthood and show that circulating ILCs are dramatically reduced in vertically HIV-infected individuals and are not restored by successful long-term ART unless it is initiated at birth. This is similar to our finding in adult HIV infection, in which only ART started in Fiebig stages I–VI was able to preserve circulating ILC levels (Kløverpris et al., 2016). However, unlike adult infection, in which tissue-resident ILCs with tonsils were preserved, vertical HIV infection also caused severe depletion of all ILC subsets within these secondary lymphoid organs. Remaining tissue-resident ILCs displayed diverse responses to HIV infection that involves proliferation, activation, and potential differentiation, consistent with immune signatures observed in lymph nodes from adult non-human primates and human subjects (Mudd et al., 2018; Wang et al., 2020). These data showing persistent transcriptional activity suggest an ongoing functional role for lymph node ILCs in pediatric HIV infection. The severe depletion of these cells, therefore, seems likely to have long consequences to the lymph node function and immune health.

Using a gating strategy previously shown to identify circulating ILCs in adults (Kløverpris et al., 2016), we identify all known ILC subsets within our pediatric subsets and confirmed four of these subsets at the transcriptional level. Interestingly, we find that helper ILC levels, but not cytotoxic NK levels, are highly elevated in early life and decrease toward adolescence. This is consistent with recent data from cord blood and peripheral blood of pediatric in Caucasian populations (Vély et al., 2016) and suggests...
that helper ILCs may play a more dominant role in the immune response during early life while the adaptive immune response is maturing.

The cohorts spanning early life to adolescence show in each case that HIV infection reduces or depletes circulating helper ILCs and CD56<sup>high</sup> NK cells in the absence of early ART intervention. In our treatment-naïve cohorts, we stratified for disease control by comparing PSPs (Muenchhoff et al., 2016) and PPs but found no overall differences in ILC subsets between these otherwise clinically distinct groups. PSPs are characterized by normal-to-age CD4 levels and very low immune activation despite high viremia, in contrast to progressing children (Muenchhoff et al., 2016). This does not agree with the findings of Mudd et al. (2018), who found a direct correlation between ILCs and CD4 levels in individuals. Importantly, however, that association was observed in HIV-infected adults and in individuals with non-HIV-associated reduced CD4 counts. This study shows that ILC depletion in the context of vertical HIV infection, however, is clearly not directly related to CD4 counts. Indeed, our findings may question the mechanistic relationship between CD4 count and circulating ILCs, as it is clearly not a dependent relationship. Moreover, virally suppressed pediatric individuals followed longitudinally showed no reconstitution of helper ILCs or CD56<sup>high</sup> NK cells despite normalization of CD4 levels. Consistent with this, ILC levels in adolescents (14–18 years) remained low despite >4 years of treatment. Alternative mechanisms suggest HIV-induced cytokines to drive ILC depletion (Wang et al., 2020), although this was not investigated here.

While the importance of ILCs in tissue is now well established (Vivier et al., 2018), ILCs’ function in blood remains unknown. Our transcriptional profiling of blood ILCs showed robust transcriptional responses for helper ILCs and NK cell subsets consisting of DEGs associated with broad cellular activation. The fact that IFN-γ was predicted as an important upstream driver of this response (except in ILC2s) is consistent with a pervasive role for this cytokine in orchestrating peripheral immune responses during chronic HIV infection (Roff et al., 2014). In adult HIV infection, apoptotic signatures were detected in acute HIV infection and associated with ILC depletion (Kleverpris et al., 2016), consistent with recent work in showing HIV-induced cytokines can deplete ILC homeostasis in adult HIV infection (Wang et al., 2020). However, no such apoptotic signature was detected in the HIV-infected pediatric subjects studied here, who were not in the acute phase. Indeed, upregulation of the anti-apoptotic factor BIRC3 in ILC2s and AXL and MET in ILCPs was observed. We also observed significant upregulation of genes enriched for metabolic pathways in ILC2s and ILCPs. Recent work in healthy tissue has demonstrated that ILCs play an important role in regulating dietary and tissue metabolism, and changes in the cellular metabolism of ILCs can affect the immunoregulatory effects of these cells (O’Sullivan and Sun, 2017; Wilhelm et al., 2017). Thus, although this study lacks mechanistic detail, this gene modulation is consistent with published data and may suggest a role for ILCs in the immunometabolic effects of HIV infection. In the remaining ILCs sequenced from HIV-infected tonsils, we did indeed detect an upregulation of genes involved in metabolism. However, we do observe a clear signature of immune-related gene networks, including signaling and tissue repair. Surprisingly, we observed little impact from HIV infection on CD4<sup>+</sup> T cell subsets, including Tfh cells, in pediatric tonsils. Direct matched comparison of CD4<sup>+</sup> T cell subsets between blood and tonsils is needed to confirm this difference in transcriptional response by compartment. Although further work is needed, these data imply an ongoing role of ILCs within secondary lymphoid organs of HIV-infected children. Whether these are protective or detrimental to lymph node function in these individuals remains to be seen.

In conclusion, we demonstrate the impact of lifelong HIV infection on ILCs in both blood and lymphoid tissue. We used well-defined cohorts differentiated by relative natural disease control and time to treatment initiation at birth and early childhood. Despite their functional overlap to helper CD4<sup>+</sup> T cells, it is clear that ILCs in blood respond differently in both frequencies and function to HIV infection and to antiretroviral HIV treatment. Moreover, ILC responses at tissue effector sites point toward a role for these cells as important regulators of tissue homeostasis in chronic treated HIV infection. Properly functioning lymph nodes are crucial for the generation of optimal immune responses, and it is known that even treated HIV-infected children have impaired immune responses to both vaccination (Cagigi et al., 2012) and natural infection (Muenchhoff et al., 2019). ILCs are required for the formation of secondary lymphoid organs during development (van de Pavert et al., 2014) and their proper functioning (Bar-Ephraim and Mebius, 2016). Thus, the depletion of ILCs observed in children with lifelong HIV infection may contribute to suboptimal immunity in these individuals. Crucially, the consequences of HIV infection from birth in later life remain unknown. Understanding the impact of HIV-induced depletion of ILCs in lymph nodes may lead to interventions that improve immune function in this vulnerable and important population.

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Supplemental information can be found online at https://doi.org/10.1016/j.celrep.2020.108153.
ACKNOWLEDGMENTS

We wish to thank all the mother-child pairs, children, adolescents, and adults participating in this study. We also want to thank all the staff at the Human Pathogenesis Programme (HPP) and at the Africa Health Research Institute (AHRI), and associated medical and hospital clinical staff at thembalabantu clinic, Edendale, Mahatma Gandhi Memorial, and Stanger and Queen Nandi Memorial Hospitals in KZN. H.N.K. was supported by the Wellcome Trust (202485/Z/16/2) and the Maersk Foundation for Medical Improvements. A.L. is supported by the Wellcome Trust (210662/Z/18/2). A.K.S. was supported, in part, by the Searle Scholars Program, the Beckman Young Investigator Program, the National Institutes of Health (NIH) (SU24AI118672, 2R01HL095791, 2U19AI089992, 1R01HL134539, 1R01AI138546), a Sloan Fellowship in Chemistry, and the Bill and Melinda Gates Foundation. S.W.K. was supported by the Hugh Hampton Young Memorial Fellowship. P.G. was supported by the NIH (R01 AI133673) and the Wellcome Trust (WT104748MA). This work was supported through the Sub-Saharan African Network for TB/HIV Research Excellence (SANTHE), a Developing Excellence in Leadership, Training and Science (DELTAS) Africa Initiative (DEL-15-006). The DELTAS Africa Initiative is an independent funding scheme of the African Academy of Sciences (AAS)’s Alliance for Accelerating Excellence in Science in Africa (AESAf) and is supported by the New Partnership for Africa’s Development Planning and Coordinating Agency (NEPAD Agency) with funding from the Wellcome Trust (107752/Z/15/2) and the UK Government. The views expressed in this publication are those of the author(s) and not necessarily those of AAS, NEPAD Agency, the UK Government. H.N.K. and A.S. were supported by the Sub-Saharan African Network for TB/HIV Research Excellence (SANTHE).

AUTHOR CONTRIBUTIONS

A.S. and J.R. performed experiments. S.W.K. and K.K. performed transcriptional analysis. J.R., J.M., and M.M. contributed human samples. O.E.A., A.N., D.R., R.F., and A.A. contributed to experimental work. W.K. contributed surgical human tissue samples, F.K. and T.N. contributed sample collections. A.K.S. supervised data analysis. P.G. and A.L. provided clinical samples. S.W.K., A.L., and H.N.K. prepared the manuscript. H.N.K. conceptualized. S.W.K., A.K.S., P.G., A.L., and H.N.K. provided intellectual input.

DECLARATION OF INTERESTS

The authors declare no competing interests.

REFERENCES

Adland, E., Paioni, P., Thobakgale, C., Laker, L., Mori, L., Muenchhoff, M., Csala, A., Clapson, M., Flynn, J., Novelli, V., et al. (2018). Discordant Impact of HLA on Viral Replicative Capacity and Disease Progression in Pediatric and Adult HIV Infection. PLoS Pathog. 14, e1006954.

Adland, E., Millar, J., Bengu, N., Muenchhoff, M., Fillis, R., Sprenger, K., Ntiatnsana, V., Roider, J., Vieira, V., Govender, K., et al. (2020). Sex-specific innate immune selection of HIV-1 in utero is associated with increased female susceptibility to infection. Nat. Commun. 11, 1767.

Arcone, R., Arpaia, G., Ruoppolo, M., Malorni, A., Pucci, P., Marino, G., Ialenti, A., Di Rosa, M., and Ciliberto, G. (1993). Structural characterization of a biologically active human lipocortin 1 expressed in Escherichia coli. Eur. J. Biochem. 211, 347–355.

Ardain, A., Domingo-Gonzalez, R., Das, S., Kazer, S.W., Howard, N.C., Singh, A., Ahmed, M., Nhamoyebonde, S., Rangel-Moreno, J., Ongoro, P., et al. (2019). Group 3 innate lymphoid cells mediate early protective immunity against tuberculosis. Nature 570, 528–532.

Azzoni, L., Rutstein, R.M., Chehimi, J., Farabaugh, M.A., Nowmos, A., and Montaner, L.J. (2005). Dendritic and natural killer cell subsets associated with stable or declining CD4+ cell counts in treated HIV-1-infected children. J. Infect. Dis. 191, 1451–1459.

Baessler, T., Charton, J.E., Schmiedel, B.J., Grunefach, F., Krusch, M., Wacker, A., Ramnussen, H.G., and Salih, H.R. (2010). CD137 ligand mediates opposite effects in human and mouse NK cells and impairs NK-cell reactivity against human acute myeloid leukemia cells. Blood 115, 3058–3069.

Balian, W.M., Vu, B.A., Long, B.R., Loo, C.P., Michaelosien, J., Barbour, J.D., Lanier, L.L., Wiznia, A.A., Abadi, J., Fennelly, G.J., et al. (2007). Natural killer cells in perinatally HIV-infected children exhibit less degranulation compared to HIV-1-exposed uninfected children and their expression of KIR2DL3, NKp30, and NKp46 correlates with disease severity. J. Immunol. 179, 3362–3370.

Bar-Ephraim, Y.E., and Mebius, R.E. (2016). Innate lymphoid cells in secondary lymphoid organs. Immunol. Rev. 271, 185–199.

Bender, J.M., Li, F., Martelly, S., Byrt, E., Rouzier, V., Leo, M., Tobin, N., Panaraj, P.S., Adisetyo, H., Roline, A., et al. (2016). Maternal HIV infection influences the microbiome of HIV-uninfected infants. Sci. Transl. Med. 8, 343ra100.

Bennink, J.H., Peters, C.P., Munneke, M., te Velde, A.A., Meijer, S.L., Weijer, K., Hreghjvidsdottir, H.S., Heinsbroek, S.E., Legrand, N., Busken, C.J., et al. (2013). Human type 1 innate lymphoid cells accumulate in inflamed mucosal tissues. Nat. Immunol. 14, 221–229.

Björklund, A.K., Forkel, M., Picelli, S., Konya, V., Theorell, J., Friberg, D., Sandberg, R., and Mjöberg, J. (2016). The heterogeneity of human CD127+ innate lymphoid cells revealed by single-cell RNA sequencing. Nat. Immunol. 17, 451–460.

Cagigi, A., Cotugno, N., Giaquinto, C., Nicolosi, L., Bernardi, S., Rossi, P., Douagi, I., and Palmia, P. (2012). Immune reconstitution and vaccination outcome in HIV-1 infected children: present knowledge and future directions. Hum. Vaccin. Immunother. 8, 1784–1794.

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Cella, M., Gamini, R., Secca, C., Collins, P.L., Zhao, S., Peng, V., Robinette, M.L., Schettini, J., Zaitsev, K., Gordon, W., et al. (2019). Subsets of iLC3-ILC1-like cells generate a diversity spectrum of innate lymphoid cells in human mucosal tissues. Nat. Immunol. 20, 980–991.

Chahrour, A., and Silvestri, G. (2016). What pediatric nonprogressors and natural HIV hosts teach us about HIV. Sci. Transl. Med. 8, 358fs16.

Chen, J., Ou, Y., Yang, Y., Li, W., Xu, Y., Xie, Y., and Liu, Y. (2018). KLHL22 activates amino-acid-dependent mTORC1 signalling to promote tumorigenesis and ageing. Nature 557, 585–589.

Deeks, S.G., Tracy, R., and Douek, D.C. (2013). Systemic effects of inflammation on health during chronic HIV infection. Immunity 39, 633–645.

Di Lorenzo, A., Yang, Y., Macaluso, M., and Bedford, M.T. (2014). A gain-of-function mouse model identifies PRMT6 as a NF-κB coactivator. Nucleic Acids Res. 42, 8297–8309.

Felices, M., Ankarlo, D.E., Lenvik, T.R., Nelson, H.H., Blazar, B.R., Verneris, M.R., and Miller, J.S. (2014). Notch signaling at later stages of NK cell development enhances KIR expression and functional maturation. J. Immunol. 193, 3344–3354.

Fernandes, S.M., Pires, A.R., Matoso, P., Ferreira, C., Nunes-Cabaço, H., Correia, L., Valadas, E., Poças, J., Pacheco, P., Veiga-Fernandes, H., et al. (2018). HIV-2 infection is associated with preserved GALT homeostasis and epithelial integrity despite ongoing mucosal viral replication. Mucosal Immunol. 11, 236–248.

Gasteiger, G., Fan, X., Diky, S., Lee, S.Y., and Rudensky, A.Y. (2015). Tissue residency of innate lymphoid cells in lymphoid and nonlymphoid organs. Science 350, 981–985.

Gomez de Aguero, M., Galan-Vonarburg, S.C., Fuhrer, T., Rupp, S., Uchimura, Y., Li, H., Steinetz, A., Heikenwalder, M., Hapelmeier, S., Sauer, U., et al. (2018). The maternal microbiota drives early postnatal innate immune development. Science 351, 1296–1302.
Goulder, P.J., and Walker, B.D. (2012). HIV and HLA class I: an evolving relationship. Immunity 37, 426–440.

Goulder, P.J., Lewin, S.R., and Leitman, E.M. (2016). Paediatric HIV infection: the potential for cure. Nat. Rev. Immunol. 16, 259–271.

Gury-BenAri, M., Thaiss, C.A., Serafini, N., Winter, D.R., Giladi, A., Lara-Astiaso, D., Levy, M., Salame, T.M., Weiner, A., David, E., et al. (2016). The Spectrum and Regulatory Landscape of Intestinal Innate Lymphoid Cells Are Shaped by the Microbiome. Cell 166, 1231–1246.e1213.

Huot, N., Jacquelin, B., Garcia-Tellez, T., Rascle, P., Ploquin, M.J., Madec, Y., Reeves, R.K., Derreudre-Bosquet, N., and Muller-Trutwin, M. (2017). Natural killer cells migrate into and control simian immunodeficiency virus replication in lymph node follicles in African green monkeys. Nat. Med.

Klatt, N.R., Estes, J.D., Sun, X., Ortiz, A.M., Barber, J.S., Harris, L.D., Cervasi, B., Muenchhoff, M., Prendergast, A.J., and Goulder, P.J. (2012). Loss of mucosal CD103+ DCs and IL-17+ and IL-22+ lymphocytes is associated with mucosal damage in HIV infection. Mucosal Immunol.

Kraemer, B., Goeser, F., Lutz, P., Glaessner, A., Boesecke, C., Schwarze-Roosen, E., Bari, H., and Malmberg, K.J., Berning, J.H., Blom, B., Huisman, C., van Oers, M.H., Spits, H., Malmberg, K.J., and Hazenberg, M.D. (2014). Activated innate lymphoid cells are associated with a reduced susceptibility to graft-versus-host disease. Blood 124, 812–821.

Kremer, B., Goerlin, P., Lutz, P., Glässner, A., Boesecke, C., Schwarze-Zander, C., Kaczmarek, D., Nitschke, H.D., Branchi, V., Manekeller, S., et al. (2017). Compartment-specific distribution of human intestinal innate lymphoid cells is altered in HIV patients under effective therapy. PLoS Pathog. 13, e1006373.

Lewis, J., Walker, A.S., Castro, H., De Rossi, A., Gribb, D.M., Gianguito, C., Klein, N., and Gallard, R. (2012). Age and CD4 count at initiation of antiretroviral therapy in HIV-infected children: effects on long-term T-cell reconstitution. J. Infect. Dis. 205, 548–556.

Li, B., and Dewey, C.N. (2011). RSEM: accurate transcript quantification from RNA-Seq data with or without a reference genome. BMC Bioinformatics 12, 323.

Li, H., and Reeves, R.K. (2013). Functional perturbation of classical natural killer and innate lymphoid cells in the oral mucosa during SIV infection. Front. Immunol. 3, 417.

Lim, A.I., Li, Y., Lopez-Lastra, S., Stadhouder, R., Paul, F., Casrouge, A., Serafini, N., Piel, A., Bustamante, J., Surace, L., et al. (2017). Systemic Human ILC Precursors Provide a Substrate for Tissue ILC Differentiation. Cell 168, 1086–1100.e1010.

Love, M.I., Huber, W., and Anders, S. (2014). Moderated estimation of fold change and dispersion for RNA-seq data with DESeq2. Genome Biology 15. https://doi.org/10.1186/s13059-014-0550-8.

Mandl, J.N., Barry, A.P., Vanderford, T.H., Kozyr, N., Chavan, R., Klucking, S., Barratt, F.J., Coffman, R.L., Staphanos, S., and Feinberg, M.B. (2008). Divergent TLR7 and TLR9 signaling and type I interferon production distinguish pathogenic and nonpathogenic AIDS viruses infections. Nat. Med. 14, 1077–1087.

Marston, M., Becquet, R., Zaba, B., Moulton, L.H., Gray, G., Coovadia, H., Essex, M., Ekuoe, D.K., Jackson, D., Coutouco, A., et al. (2011). Net survival of perinatally and postnatally HIV-infected children: a pooled analysis of individual data from sub-Saharan Africa. Int. J. Epidemiol. 40, 385–396.

Masopust, D., and Soerens, A.G. (2019). Tissue-Resident T Cells and Other Resident Leukocytes. Annu. Rev. Immunol. 37, 521–546.

Meier, A., Chang, J.I., Chan, E.S., Pollard, R.B., Siddhu, H.K., Kulkarni, S., Wen, T.F., Lindsay, R.J., Orellana, L., Mildvan, D., et al. (2009). Sex differences in the Toll-like receptor-mediated response of placental dendritic cells to HIV-1. Nat. Med. 15, 955–959.

Mudd, J.C., Busman-Sahay, K., DiNapoli, S.R., Lai, S., Sheik, V., Lisco, A., Deleage, C., Richardson, B., Palesch, D.J., Pardiani, M., et al. (2018). Hallmarks of primate lentiviral immunodeficiency infection recapitulate loss of innate lymphoid cells. Nat. Commun. 9, 3967.

Muenchhoff, M., Prendergast, A.J., and Goulder, P.J. (2014). Immunity to HIV in Early Life. Front. Immunol. 5, 391.

Muenchhoff, M., Adland, E., Karimzaman, O., Crowther, C., Pace, M., Csala, A., Leitman, E., Moonsamy, A., McGregor, C., Hurst, J., et al. (2016). Nonprogressing HIV-infected children share fundamental immunological features of nonpathogenic SIV infection. Sci. Transl. Med. 8, 358ra125.

Newell, M.L., Coovadia, H., Cortina-Borja, M., Rollins, N., Gallard, P., and Da-bis, F.; GHent International AIDS Society (IAS) Working Group on HIV Infection in Women and Children (2004). Mortality of infected and uninfected infants born to HIV-infected mothers in Africa: a pooled analysis. Lancet 364, 1236–1243.

O’Sullivan, T.E., and Sun, J.C. (2017). Innate Lymphoid Cell Immunometabolism. J. Mol. Biol. 429, 3577–3586.

Pica, M.Q., Lewis, J., Musiime, V., Prendergast, A., Nathoo, K., Keziiriwa, A., Nahiriya Ntege, P., Gibb, D.M., Theibaut, R., Walker, A.S., et al.; ARROW Trial Team (2013). Predicting patterns of long-term CD4 reconstitution in HIV-infected children starting antiretroviral therapy in sub-Saharan Africa: a cohort-based modelling study. PLoS Med. 10, e1001542.

Prendergast, A.J., Klenerman, P., and Goulder, P.J. (2012). The impact of differential antiviral immunity in children and adults. Nat. Rev. Immunol. 12, 639–645.

Reeves, R.K., Rajakumar, P.A., Evans, T.I., Cononne, M., Gillis, J., Wong, F.E., Kuzmiczey, Y.V., Carville, A., and Johnson, R.P. (2011). Gut inflammation and indoleamine deoxyxigenase inhibit IL-17 production and promote cytotoxic potential in NKp44+ mucosal NK cells during SIV infection. Blood 118, 3321–3330.

Roff, S.R., Noon-Song, E.N., and Yamamoto, J.K. (2014). The Significance of interferon-γ in HIV-1 Pathogenesis, Therapy, and Prophylaxis. Front. Immunol. 4, 498.

Roider, J.M., Muenchhoff, M., and Goulder, P.J. (2016). Immune activation and paediatric HIV-1 disease outcome. Curr. Opin. HIV AIDS 11, 146–155.

Roider, J., Porterfield, J.Z., Ogongo, P., Muenchhoff, M., Adland, E., Groll, A., Morris, L., Moore, P.L., Ndung’u, T., Klevjerup, H., et al. (2019). Plasma IL-5 but Not CXCL13 Correlates With Neutralization Broad HIV-Infected Children. Front. Immunol. 10, 1497.

Serafini, N., Vossenhenric, C.A., and Di Santo, J.P. (2015). Transcriptional regulation of innate lymphoid cell fate. Nat. Rev. Immunol. 15, 415–428.

Shah, S.V., Manickam, C., Ram, D.R., and Reeves, R.K. (2017). Innate Lymphoid Cells in HIV/SIV Infections. Front. Immunol. 8, 1818.

Shearer, W.T., Rosenblatt, H.M., Gelman, R.S., Oyomopito, R., Piaegger, S., Steinh, E.R., Wara, D.W., Douglas, S.D., Luzuriaga, K., McFarland, E.J., et al.; Pediatric AIDS Clinical Trials Group (2003). Lymphocyte subsets in healthy children from birth through 18 years of age: the Pediatric AIDS Clinical Trials Group P1009 study. J. Allergy Clin. Immunol. 112, 973–980.
Skon, C.N., Lee, J.Y., Anderson, K.G., Masopust, D., Hogquist, K.A., and Jameson, S.C. (2013). Transcriptional downregulation of S1pr1 is required for the establishment of resident memory CD8+ T cells. Nat. Immunol. 14, 1285–1293.

Sloan, E., Orr, A., and Everett, R.D. (2016). MORC3, a Component of PML Nuclear Bodies, Has a Role in Restricting Herpes Simplex Virus 1 and Human Cytomegalovirus. J. Virol. 90, 8621–8633.

Slyker, J.A., Lohman-Payne, B., John-Stewart, G.C., Mbori-Ngacha, D., Tapia, K., Atzberger, A., Taylor, S., Rowland-Jones, S.L., and Blish, C.A. (2012). The impact of HIV-1 infection and exposure on natural killer (NK) cell phenotype in Kenyan infants during the first year of life. Front. Immunol. 3, 399.

Smith, C., Jaibert, E., de Almeida, V., Canniff, J., Lenz, L.L., Mussi-Pinhata, M.M., Cohen, R.A., Yu, G., Amaral, F.R., Pinto, J., et al. (2017). Altered Natural Killer Cell Function in HIV-Exposed Uninfected Infants. Front. Immunol. 8, 470.

Sonnenberg, G.F., Monticelli, L.A., Alenghat, T., Fung, T.C., Hutnick, N.A., Kunisawa, J., Shibata, N., Grunberg, S., Smha, R., Zahn, A.M., et al. (2012). Innate lymphoid cells promote anatomical containment of lymphoid-resident commensal bacteria. Science 336, 1321–1325.

Spits, H., Artis, D., Colonna, M., Diefenbach, A., Di Santo, J.P., Eberl, G., Koyasu, S., Locksley, R.M., McKenzie, A.N., Mebius, R.E., et al. (2013). Innate lymphoid cells—a proposal for uniform nomenclature. Nat. Rev. Immunol. 13, 145–149.

Subramanian, A., Tamayo, P., Mootha, V.K., Mukherjee, S., Ebert, B.L., Gillette, M.A., Paulovich, A., Pomeroy, S.L., Golub, T.R., Lander, E.S., and Mesirov, J.P. (2005). Gene set enrichment analysis: a knowledge-based approach for interpreting genome-wide expression profiles. Proc. Natl. Acad. Sci. USA 102, 15545–15550.

Takahashi, K., Yoshida, N., Murakami, N., Kawata, K., Ishizaki, H., Tanaka-Okamoto, M., Miyoshi, J., Zinn, A.R., Shime, H., and Inoue, N. (2007). Dynamic regulation of p53 subnuclear localization and senescence by MORC3. Mol. Biol. Cell 18, 1701–1709.

van de Pavert, S.A., Ferreira, M., Domingues, R.G., Ribeiro, H., Molenaar, R., Moreira-Santos, L., Almeida, F.F., Ibiza, S., Barbosa, I., Goverse, G., et al. (2014). Maternal retinoids control type 3 innate lymphoid cells and set the offspring immunity. Nature 508, 123–127.

Vély, F., Barlogie, V., Vallentin, B., Neven, B., Piperoglou, C., Ebbo, M., Perchet, T., Petit, M., Yessaad, N., Touzot, F., et al. (2016). Evidence of innate lymphoid cell redundancy in humans. Nat. Immunol. 17, 1291–1299.

Vivier, E., Artis, D., Colonna, M., Diefenbach, A., Di Santo, J.P., Eberl, G., Koyasu, S., Locksley, R.M., McKenzie, A.N., Mebius, R.E., et al. (2018). Innate Lymphoid Cells: 10 Years On. Cell 174, 1054–1066.

Wang, Y., Lifshitz, L., Geliaty, K., Vinton, C.L., Busman-Sahay, K., McCauley, S., Vangala, P., Kim, K., Derr, A., Jaiswal, S., et al. (2020). HIV-1-induced cytokines deplete homeostatic innate lymphoid cells and expand TCF7-dependent memory NK cells. Nat. Commun. 21, 274–286.

Wilcox, R.A., Tamada, K., Strome, S.E., and Chen, L. (2002). Signaling through NK cell-associated CD137 promotes both helper function for CD8+ cytolytic T cells and responsiveness to IL-2 but not cytolytic activity. J. Immunol. 169, 4230–4236.

Wilhelm, C., Kharabi Masouleh, S., and Kazakov, A. (2017). Metabolic Regulation of Innate Lymphoid Cell-Mediated Tissue Protection—Linking the Nutritional State to Barrier Immunity. Front. Immunol. 8, 1742.

Xiong, H., Li, H., Chen, Y., Zhao, J., and Unkeless, J.C. (2004). Interaction of TRAF6 with MAST205 regulates NF-kappaB activation and MAST205 stability. J. Biol. Chem. 279, 43675–43683.

Xu, H., Wang, X., Lackner, A.A., and Veazey, R.S. (2015). Type 3 innate lymphoid cell depletion is mediated by TLRs in lymphoid tissues of simian immunodeficiency virus-infected macaques. FASEB J. 29, 5072–5080.

Zhang, Z., Cheng, L., Zhao, J., Li, G., Zhang, L., Chen, W., Nie, W., Reszka-Blanco, N.J., Wang, F.S., and Su, L. (2015). Plasmacytoid dendritic cells promote HIV-1-induced group 3 innate lymphoid cell depletion. J. Clin. Invest. 125, 3692–3703.

Zhong, C., and Zhu, J. (2017). Transcriptional regulators dictate innate lymphoid cell fates. Protein Cell 8, 242–254.
## STAR METHODS

### KEY RESOURCES TABLE

| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| **Antibodies**      |        |            |
| Anti-CD1c AF488 (Lineage) | BioLegend | 301618; RRID: AB_439791 |
| Anti-CD14 FITC (Lineage)  | BD Bioscience | 555397; RRID: AB_395798 |
| Anti-CD19 FITC (Lineage)  | BD Bioscience | 560994; RRID: AB_10563406 |
| Anti-CD3 AF488 (Lineage)  | BioLegend | 300320; RRID: AB_493691 |
| Anti-CD4 FITC (Lineage)  | BioLegend | 317420; RRID: AB_571939 |
| Anti-TCRgd AF488 (Lineage)  | BioLegend | 331208; RRID: AB_1575108 |
| Anti-TCRab AF488 (Lineage)  | BioLegend | 306712; RRID: AB_528967 |
| Anti-CD34 FITC (Lineage)  | BioLegend | 343604; RRID: AB_1732005 |
| Anti-CD303 FITC (Lineage)  | BioLegend | 354208; RRID: AB_2561364 |
| Anti-CD19 FITC (Lineage)  | BD Bioscience | 560994; RRID: AB_10563406 |
| Anti-CD94 PerCp.Cy5.5  | BD Bioscience | 562361; RRID: AB_11152081 |
| Anti-CD117 BV421  | BioLegend | 313216; RRID: AB_11148721 |
| Anti-CD117 BV650  | BioLegend | 313221; RRID: AB_2562714 |
| Anti-CD161 BV605  | BioLegend | 339915; RRID: AB_11142679 |
| Anti-CD16 BV650  | BioLegend | 302042; RRID: AB_2563801 |
| Anti-CD56 BV711  | BioLegend | 318336; RRID: AB_2562417 |
| Anti-CD3 BV785  | BioLegend | 317330; RRID: AB_2563507 |
| Anti-CD294 AF647  | BD Bioscience | 558042; RRID: AB_2112699 |
| Anti-CD38 AF700  | BioLegend | 303516; RRID: AB_2072782 |
| Anti-CD95 PE-CF594  | BioLegend | 305634; RRID: AB_2564221 |
| Anti-CD127 Pe-Cy7  | Beckman Coulter | A14934; RRID: AB_2534372 |
| Anti-CD4 BUV496  | BD Bioscience | 564651; RRID: AB_2744422 |
| Anti-PD-1 BUV421  | BD Bioscience | 562516; RRID: AB_11153482 |
| Anti-CD103 BV605  | BioLegend | 350218; RRID: AB_2564283 |
| Anti-CD69 BV785  | BioLegend | 310932; RRID: AB_2563696 |
| Anti-CD3 PE-CF594  | BD Bioscience | 562280; RRID: AB_11153674 |
| Anti-CD366 (NKP44) PE-Cy5  | Beckman Coulter | A66903; RRID: AB_2857937 |
| Anti-CD8 BUV396  | BD Bioscience | 563795; RRID: AB_2722501 |
| Anti-CD19 BUV496  | BD Bioscience | 564655; RRID: AB_2744311 |
| Live/Dead Fixable Near-IR Dead Cell Stain Kit, 633nm  | Invitrogen | L10119 |
| Anti-IL-2 BV650  | BD Biosciences | 563467; RRID: AB_2738224 |
| Anti-IL-4 FITC  | BioLegend | 500807; RRID: AB_315126 |
| Anti-IL-5 APC  | BioLegend | 504305; RRID: AB_315329 |
| Anti-IL-13 BV421  | BioLegend | 561158; RRID: AB_10561838 |
| Anti-TNFα AFlour700  | BD Bioscience | 557996; RRID: AB_396978 |
| Anti-IFNγ PE-Cy7  | BD Bioscience | 557643; RRID: AB_396760 |
| Anti-CD294 (CTRH2) PE-CF594  | BD Bioscience | 563501; RRID: AB_2738244 |
| Anti-CD127 PE-Cy5  | Beckman Coulter | A64617; RRID: AB_2833010 |

### Biological Samples

| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| Human peripheral blood mononuclear cells (PBMCs)  | Human | Cohorts |
| Human tonsil mononuclear cells (TMCs)  | Human | Cohorts |

(See Table 1)
RESOURCE AVAILABILITY

Lead contact
Further information and requests for resources and reagents should be directed to and will be fulfilled by the Lead Contact (henrik.kloverpris@ahri.org).

Materials availability
This study did not generate new unique reagents.

Data and code availability
The RNA-seq datasets supporting the current study have not been deposited in a public repository because the subjects from which they were generated are at-risk children. The processed expression matrices are available upon request from the lead contact. Access to the raw data will be considered on a case-by-case basis with supporting IRB approval on the behalf of the requestor.

EXPERIMENTAL MODEL AND SUBJECT DETAILS

Peripheral blood (PB) samples from children were obtained from the Ithemabalabantu pediatric cohort in Durban, KwaZulu-Natal (KZN), South Africa (Muenchhoff et al., 2016) and from Stanger Hospital, Stanger, KwaZulu-Natal (KZN), South Africa (Roider et al., 2019). PB samples from newborn/infants were obtained from the Ucwaningo Lwabantwana (meaning learning from children) infant cohort from Edendale, Mahatma Gandhi Memorial, Stanger and Queen Nandi Memorial Hospitals in KZN (Adland et al., 2020). Tonsil tissue samples were obtained from pediatric patients undergoing routine tonsillectomy at Stanger Hospital, Stanger, KwaZulu-Natal (KZN), South Africa (Roider et al., 2019). Informed consent was obtained from all adult study participants; and for underage children and adolescents, informed consent was obtained from their guardians. All HIV infected individuals were infected via vertical transmission from maternal HIV infection. For non-adult participants, 4 age groups were defined: 1. Newborns, aged 3–45 h; 2. Infants, aged 2–60 months; Paediatrics, aged 5–18 years. Paediatric slow progressors (PSP) were defined by stable CD4 T cell percentage of total PBMCs > 20% and found to be clinically healthy, while untreated pediatric progressors (PP) were defined by CD4+ T cell percentage of total PBMCs < 20% or otherwise meeting requirements for treatment. The pediatric treated cohort have individuals on antiretroviral therapy (ART). All subjects are from black Sub-Saharan ethnicity. This study was approved by the respective institutional review boards and Biomedical Research Ethics Committee, University of KwaZulu-Natal (UKZN) in Durban, South Africa.
METHOD DETAILS

Cell isolation from human blood and tonsil
Peripheral blood mononuclear cells (PBMCs) were isolated by Histopaque 1077 (Sigma-Aldrich) density gradient centrifugation. Tonsil tissue samples was minced and digested with Collagenase D (0.5 mg/ml; Roche) and DNase I (20 μg/ml; Sigma-Aldrich) for 30 min in 37°C shaking incubator. Digested tissue was passed through 70 μm cell strainers. Lymphocytes from the tonsil were isolated by Histopaque 1077 (Sigma-Aldrich) density gradient centrifugation.

Flow cytometry analysis and cell sorting
For FACS analysis, different antibody panels for phenotype and intracellular cytokine staining (ICS) were used. A complete list of antibodies used with identifier and source information can be found in the Key Resources Table. All samples were surface stained at room temperature for minimum 20 min and near-infrared live/dead cell viability staining kit (Invitrogen). For experiments involving ICS, the cells were stimulated with PMA (5 ng/ml; Sigma) plus Ionomycin (1 μg/ml; Sigma) in the presence of Golgiplug and Golgistop (BD Biosciences) for 4 hr in 37°C incubator. Cells were stained with fluorochrome-conjugated monoclonal antibodies and subsequently fixed, permeabilized, and stained by BD Cytofix/Cytoperm Kit (BD Biosciences). Blocking with 20% goat serum for 20 min was done prior to intracellular antibody staining. After staining, cells were washed and fixed in 2% paraformaldehyde before acquisition on a 4 laser, 17 parameter BD FACSAura Fusion flow cytometer within 24 h of staining. Data were analyzed with FlowJo v.9.7.2 (TreeStar). For cell sorting experiments, cells were processed from cryopreserved, surface stained, kept on ice in PBS and sorted immediately after staining. All samples were surface stained at room temperature for a minimum 20 mins. Bulk populations were cell sorted to a purity 99% on the BD FACSAura Fusion flow cytometer.

RNA isolation, library construction, sequencing, and alignment
CD4+ T cells, IL2Cs, ILCPs, NK CD16high and NK CD56high cells from PBMCs and NKp44+ ILC3s, NKp44+ ILC3s, NK CD127 and 4 distinct CD4+ T cell subsets from TMCs (100 cell replicates) were FACS sorted directly into 50 μL of RLT Lysis Buffer (QIAGEN) supplemented with 1% v/v 2-mercaptoethanol. Briefly, 50 μL of mixed lysate from each sample was transferred to a skirted 96 well plate. Genetic material was pulled down and purified by mixing the lysate in each well with 2.2x volumes of Agencourt RNA-Clean XP SPRI beads (Beckman Coulter) and washing 3x with 75 μL of 80% ethanol. After drying, the SPRI beads were re-suspended in 4 μL of pre-reverse transcription (RT) mix, incubated for 3 min at 72°C, and placed on ice. Next, Smart-Seq2 Whole Transcriptome Amplification (WTA) was performed: 7 μL of RT mix was added to each well and RT was carried out; then, 14 μL of PCR mix was added to each well and PCR was performed. Thereafter a cDNA cleanup was performed using 0.6x and 0.8x volumes of Agencourt AMPure XP SPRI beads (Beckman Coulter) which was then quantified using a Qubit dsDNA HS Assay Kit (Life Technologies). Library size and quality were measured by Bioanalyzer using a High Sensitivity DNA Analysis Kit (Agilent Technologies). Sequencing libraries were prepared from WTA product using Nextera XT (Illumina). After library construction, a final AMPure XP SPRI clean-up (0.8 volumes) was conducted. Library concentration and size were measured with the KAPA Library Quantification kit (KAPA Biosystems) and a TapeStation (Agilent Technologies), respectively. Finally, samples were sequenced on a NextSeq500 (30 bp paired-end reads) to an average depth of 5 million reads. Reads were aligned to hg38 (Genencode v21) using RSEM and TopHat (Li and Dewey, 2011) and estimated counts and transcripts per million (TPM) matrices generated. Any samples with fewer than 5x10^5 or more than 6x10^5 aligned reads or fewer than 10,000 uniquely expressed genes were removed from subsequent analysis.

RNA-Seq Differential Expression Analysis
Differential expression analysis was performed using DESeq2 (v1.18.1) (Love et al., 2014). Expected counts from biological replicates for each cell type and participant were averaged prior to differential expression in order to prevent participant specific genes from generating false positives and reduce spurious heterogeneity from small (100-cell) populations. Small populations may show skewed expression based on the cell composition within; thus this replicate averaging approach is particularly important given our limited access to pediatric tissue sources and low frequency of these immune populations in order to remove further bias from small populations. See Tables S1 and S6 for replicate numbers. Tonsil analyses for the ILC3 and NK cell subsets were restricted only to those isolation sources and low frequency of these immune populations in order to remove further bias from small populations.

Gene set analysis was performed using Ingenuity Pathway Analysis (IPA; Winter 2019 Release, QIAGEN Inc.) and Gene Set Enrichment Analysis (GSEA) using the piano package in R (1.18.1). For IPA, DEGs whose FDR corrected q < 0.1 were used in the “Core” analysis with the log2FC and q values included in the analysis. To implement GSEA on our DESeq2 results, we used the log2FC of all genes whose FDR corrected q < 0.1 as t-value input into the runGSA function with setting the argument geneSetStat = “gsea.” We
chose to use the KEGG and GO databases (downloaded from MSigDB v7.0) (Subramanian et al., 2005) for GSEA analysis as these databases are well annotated for metabolic and cellular activation gene sets that are cell-type agnostic.

For the IPA enrichment on tonsil population comparisons, only 1-3 of the significantly enriched terms had non-N/A values for each population. Thus, we have omitted the z-scores from the manuscript.

**QUANTIFICATION AND STATISTICAL ANALYSIS**

Graphs were plotted using Prism 8.4.3 (GraphPad Inc.) Differences between groups were analyzed using Mann Whitney U-test or Dunn’s multiple comparisons test (two-sided) with specific test used stated in the figure legends otherwise. Data are presented as the medians values with boxes representing IQR range and range by error bars, with a p value < 0.05 considered statistically significant. The values of n refers to the number of participants used in study. In the other parts, it refers to the number of dependent experiments.