Macaque Paneth Cells Express Lymphoid Chemokine CXCL13 and Other Antimicrobial Peptides Not Previously Described as Expressed in Intestinal Crypts

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CXCL13 is a constitutively expressed chemokine that controls migration of immune cells to lymphoid follicles. Previously, we found CXCL13 mRNA levels increased in rhesus macaque spleen tissues during AIDS. This led us to examine the levels and locations of CXCL13 by detailed in situ methods in cynomolgus macaque lymphoid and intestinal tissues. Our results revealed that there were distinct localization patterns of CXCL13 mRNA compared to protein in germinal centers. These patterns shifted during the course of simian immunodeficiency virus (SIV) infection, with increased mRNA expression within and around follicles during AIDS compared to uninfected or acutely infected animals. Unexpectedly, CXCL13 expression was also found in abundance in Paneth cells in crypts throughout the small intestine. Therefore, we expanded our analyses to include chemokines and antimicrobial peptides (AMPs) not previously demonstrated to be expressed by Paneth cells in intestinal tissues. We examined the expression patterns of multiple chemokines, including CCL25, as well as α-defensin 6 (DEFA6), β-defensin 2 (BDEF2), rhesus θ-defensin 1 (RTD-1), and Reg3γ in situ in intestinal tissues. Of the 10 chemokines examined, CXCL13 was unique in its expression by Paneth cells. BDEF2, RTD-1, and Reg3γ were also expressed by Paneth cells. BDEF2 and RTD-1 previously have not been shown to be expressed by Paneth cells. These findings expand our understanding of mucosal immunology, innate antimicrobial defenses, homeostatic chemokine function, and host protective mechanisms against microbial translocation.
Animals and tissues. These studies were performed under the approval and guidance of the University of Pittsburgh Institutional Animal Care and Use Committee. Adult cynomolgus macaques (Macaca fascicularis) were used in this study and were inoculated intrarectally with the pathogenic SIV/DeltaB670 isolate (11). These animals and their clinicovirologic states have been described previously (12–14). Fixation and processing of tissues at necropsy were completed as described previously (15). Tissues used for visualization of Paneth cells were postfixed in 4% paraformaldehyde and then dehydrated in graded ethanols. These tissues were then stained by Lendrum’s procedure modified by extending tartrazine staining for a period of 5 h and 45 min.

Cloning of macaque DEFA6, BDEF2, and RTD-1 cDNAs. To generate α-defensin 6 (DEFA6), β-defensin 2 (BDEF2), and rhesus α-defensin 1 (RTD-1) cDNAs, RNA was purified from cryopreserved cynomolgus ileal tissue and cDNA synthesis was as described previously (16). Human Reg3γ cDNA was purchased from Open Biosystems (ThermoFisher Scientific, Lafayette, CO). The primers used for PCR amplification were designed based on the corresponding human sequences for α-defensin 5 and β-defensin 2 (GenBank accession no. AY659407 and AF401513, respectively). The resultant forward and reverse primers were rhDEFA5_R2_CF (5’-CTCACCATCTTGCTGCTATT-3’) and rhDEFA5_R2_CR (5’-CTAAGAGCTCGGCGACAGCA-3’) for DEFA6; SS_HBD2_F1 (5’-CATCAGCGTAGGTGGTGTG-3’) and SS_HBD2_R1 (5’-CTCACTTTTGTGAGGTCTT-3’) for BDEF2, and SS_RTD1_F1 (5’-GGAGAGCGGGACGAGGAC-3’) and SS_RTD1_R1 (5’-AAAAAGAGAAATTCCGTGCAACAGGTA-3’) for RTD-1. PCR products were ligated into the pGEM-T vector (Promega), and the DNAs were sequenced.

In situ hybridization and immunohistochemistry. In situ hybridization (ISH) for rhesus CXCL13, CCL25, α-defensin 6 (DEFA6), β-defensin 2 (BDEF2), RTD-1, and Reg3γ mRNAs was performed as described previously (15, 17). Immunohistochemistry (IHC) for CXCL13 protein was performed on tissues as previously described (18). CXCL13 protein was detected by heating tissues in the microwave in Dako target retrieval buffer (Dako, Carpenteria, CA) and then incubating them with a goat anti-CXCL13 polyclonal antibody (1:50) (R&D Systems; catalog no. AF801) for 1 h. A parallel tissue on the same slide was incubated with a goat IgG control antibody (1:500) (Dako, catalog no. AB108C). The primary antibodies were then detected with an anti-goat SuperPicture kit (Zymed Laboratories, South San Francisco, CA), according to the manufacturer’s instructions. All tissues were subsequently counterstained with hematoxylin (Fisher Chemicals, Fairlawn, NJ).

Simultaneous detection of CXCL13 protein and either DEFA6, BDEF2, DEFA1, RTD-1, or Reg3γ mRNAs was performed by conducting the previously described IHC procedure prior to completion of the ISH protocol. Briefly, tissues were pretreated in Dako target retrieval buffer, rinsed, and then incubated with goat anti-CXCL13 primary antibody for 1 h at room temperature. Tissues were then rinsed and incubated with anti-goat secondary antibody conjugated to peroxidase (SuperPicture kit; Invitrogen). Sections were rinsed and incubated with the substrate diamobenzidine (DAB) for 10 min. Upon completion, slides were rinsed briefly in water and then immediately placed in 0.1 M triethanolamine with 0.01% acetic anhydride to acetylate the tissue sections to reduce nonspecific riboprobe binding. The remaining ISH steps were followed as previously described.

Immunofluorescence and image capture. Two-color immunofluorescence detection of CXCL13 and DEFA6 was performed by incubating tissues simultaneously with anti–CXCL13 conjugated to donkey anti-goat Alexa Fluor 647 and anti–DEFA6 (1:1,000) (Sigma, catalog no. HPA019462) conjugated to donkey anti-rabbit Alexa Fluor 488. All tissue sections for immunofluorescence were treated with Autofluorescence Eliminator (Chemicon International) before being mounted with ProLong Gold antifade with DAPI (4’,6-diamidino-2-phenylindole) for nuclear clearing (Molecular Probes). Images of immunohistochemistry and immunofluorescence tissue sections were captured on a Nikon (Melville, NY) E600 microscope mounted with a SPOT charged-coupled device (CCD) camera (Diagnostic Instruments, Sterling Heights, MD). The MetaVue software package (Universal Imaging, West Chester, PA) was used to view captured images. Confocal microscopy images were captured on an Olympus Fluoview 500 microscope.

DLR assays. The HEK293T cell line was used for CXCL13 promoter activity analyses using a dual-luciferase reporter (DLR) assay (Promega). Cells (8 × 10^4) were plated in poly-L-lysine-coated 6-well culture plates (Corning) 24 h prior to transfection using PolyJet (SignaGen) per the manufacturer’s recommendations with 1 μg of total plasmid DNA per well. Cell lysates were harvested after in-well lysis with Promega passive lysis buffer (0.5 ml), frozen and thawed three times, and cleared by microcentrifugation at 4°C. The DLR assays were performed in white polystyrene 96-well plates (Costar) with 20 μl of lysate per well and read using a FLUOSTAR Optima with luminescence readings (firefly luciferase and then Renilla luciferase) taken from above individual wells. Cotransfection with plasmid pGL4.74 (Promega) encoding Renilla luciferase allowed normalization for transfection efficiency differences. Following normalization of firefly luciferase values to Renilla luciferase values, the ratios were multiplied by a constant of 10,000. The positive-control plasmid pGL2-Control (Promega), was included in all transfection experiments, as was the promoterless pGL2-Basic vector (Promega). Data analyses were performed using MARS analysis software (BMG Labtech), with further analyses using Prism software (GraphPad). Short-chain fatty acid (SCFA) stimulations with sodium butyrate, sodium acetate, sodium propionate, or sodium lactate (Sigma) were performed at 5 mM 24 h before harvesting of cell lysates. Trichostatin A and 5-azacytidine (Sigma; 5 μM) pretreatments were performed for 24 h prior to sodium butyrate stimulation.

RESULTS

CXCL13 mRNA and protein are differentially localized in macaque lymphoid tissues. To examine CXCL13 expression in lymphoid tissues and to define potential changes in CXCL13 expression during pathogenic SIV infection, we examined mRNA expression patterns in lymph node (LN) and spleen tissues of cynomolgus macaques by ISH. The tissues examined were collected at necropsy from cynomolgus macaques that were either uninfected or were infected intrarectally with the SIV/DeltaB670 pathogenic strain (11) and sacrificed during acute infection (2 weeks postinfection [p.i.]) or upon development of AIDS (14 to 62 weeks p.i.), which was defined by weight loss, loss of CD4^+ T lymphocytes, and/or development of opportunistic infections, such as Pneumocystis jirovecii pneumonia. These animals and their clinicovirologic states have been presented previously (12–14).

Cells producing CXCL13 mRNA localized to a ring around germinal centers (GCs) in LN tissue sections (Fig. 1A). Subjacent tissue sections were immunostained for CXCL13 protein, and, unexpectedly, CXCL13 protein signal was intensely localized to the center of the GCs (Fig. 1A). In these regions, there were networks of CXCL13 clearly visible after IHC and immunofluorescent staining for CXCL13 in LN (Fig. 1A). In LN paracortical,
there also were individual cells that stained for CXCL13. Parallel staining of subjacent tissue sections with control goat Ig did not yield specific signal (Fig. 1A, right). The mRNA and protein expression patterns of CXCL13 in macaque spleen tissues were similar to those in LNs (Fig. 1B). These data indicate that CXCL13 producer cells are localized in a microanatomic compartment distinct from and immediately adjacent to that harboring the highest levels of CXCL13 protein in the centers of GCs.

CXCL13 expression in LN and spleen tissues was also examined across SIV disease states by staining tissues from SIV/DeltaB670-infected animals that had been previously examined for inflammatory chemokine expression in lymphoid (14) and lung (13) tissues. We compared expression patterns in LNs from uninfected, acutely SIV-infected, and AIDS-developing macaques and found that CXCL13 mRNA expression was upregulated during acute infection and maintained at high levels during AIDS (Fig. 1A). In contrast, CXCL13 protein expression increased to a limited extent across disease states. Parallel analysis of spleen tissues from the same macaques yielded similar results (Fig. 1B).

**CXCL13 is expressed by Paneth cells.** Intestinal tissues are considered to be one of the largest lymphoid organs and are a site in which many CD4+ T lymphocytes are lost during HIV-1 and SIV infection (19). Therefore, we examined the microanatomic distribution of CXCL13 in different regions along the gastrointestinal (GI) tract. Using ISH and IHC, we observed mRNA and protein expression in lymphoid follicles throughout the intestines, including in duodenum, jejunum, ileum, and colon, again with mRNA+ producer cells distributed in a three-dimensional spherical shell surrounding high levels of CXCL13 protein within GCs (not shown). Interestingly, we observed intense expression of CXCL13 mRNA and protein in a subset of cells in the crypts of Lieberkühn (Fig. 2). Cells expressing CXCL13 in crypts were observed in duodenal, jejunal, and ileal tissues from all 16 macaques studied but were not observed in colon in any of these animals.

Paneth cells are specialized intestinal epithelial cells that produce AMPs and usually are found only in the small intestine (20). The regional distribution of CXCL13 expression in intestinal crypts we observed could be due to its expression by Paneth cells. To determine if Paneth cells were the source of CXCL13 in the intestinal crypts, we individually and simultaneously stained for DEFA6, which is an AMP uniquely expressed by Paneth cells. In macaque small intestine, we found Paneth cell secretory granules (Fig. 3A), DEFA6 protein (Fig. 3B), and DEFA6 mRNA (Fig. 4A) localized to intestinal crypts in patterns identical to those obtained for CXCL13. Similar to CXCL13, DEFA6 was expressed in jejunum and ileum, but not in colon. Simultaneous IHC for CXCL13 protein and ISH for DEFA6 mRNA demonstrated colocalization to the same cells in crypts (Fig. 3C). In addition, two-color immunofluorescence staining and confocal microscopy confirmed that CXCL13 and DEFA6 are coexpressed by Paneth cells in intracellular granules, with most CXCL13 signal overlapping a large proportion of the DEFA6 signal (Fig. 3H). These findings confirm that the source of CXCL13 in intestinal crypts is the Paneth cell and are to our knowledge the first to show the expression of a lymphoid homeostatic chemokine by this specialized intestinal epithelial cell.

**CXCL13 is uniquely expressed by Paneth cells among a subset of CC and CXC chemokines.** To determine whether other chemokines also were expressed strongly in intestinal crypts, we examined the localization of mRNA producer cells of multiple

![FIG 1 Localization of CXCL13 mRNA and protein in macaque lymph node and spleen. CXCL13 protein was detected by IHC (brown signal) or immunofluorescent staining (LN, AIDS, top row) in lymph node (A) or spleen (B) at the indicated SIV-related disease state. Parallel controls performed with control antibody or cognate sense riboprobes are presented to the far right. Original magnifications, ×200.](http://cvi.asm.org/Downloaded from http://cvi.asm.org)
chemokines along the macaque GI tract. These included CCL25, a ligand for CCR9 that directs the trafficking of intestinal homing T and B lymphocytes (Fig. 4D to F), as well as CXCL1, CXCL9, CXCL10, CXCL11, CXCL12, CCL19, CCL20, and CCL21. CCL25 is known to be expressed by small intestinal epithelium, and consistent with this, the mRNA expression patterns for CCL25 differed greatly from those for CXCL13, with CCL25 producer cells found uniformly across the epithelium of the small intestine and not in the GALT, Paneth cell-laden crypts, or colon. Cells producing the other chemokines examined were not localized to intestinal crypts and therefore were not in Paneth cells (not shown).

**Expression of other AMPs by Paneth cells.** Given the clear expression of CXCL13 by Paneth cells, suggesting it has antimicrobial functions in the intestine, we next examined whether other AMPs not previously recognized as expressed by Paneth cells are also produced by them. Although both DEFA5 and DEFA6 are AMPs produced by Paneth cells, β-defensins have not been localized to these specialized cells. Unexpectedly, β-defensin 2 (BDEF2) mRNA-producing cells also were localized to intestinal crypts (Fig. 4G), and simultaneous ISH for BDEF2 mRNA and IHC for CXCL13 protein revealed colocalization to the same cells in crypts (Fig. 3E). Similarly, rhesus β-defensin 1 (RTD-1) was expressed specifically in intestinal crypts (Fig. 4H) and Paneth cells (Fig. 3F). Macaques are not known to express the AMP Reg3γ as an AMP in Paneth cells (21, 22). By using ISH, we confirmed that Reg3γ was expressed in crypts throughout the small intestine in cynomolgus macaques but was absent from the colon (Fig. 3E).

In summary, these findings provide the first evidence that in macaques BDEF2 and RTD-1 are expressed by Paneth cells and expand our understanding of the repertoire of AMPs used by these cells to protect the crypts and shape the intestinal milieu.

To determine whether SIV infection impacted levels of expression of Paneth cell AMPs, including CXCL13, we examined tissues from animals in different stages of SIV infection. Admittedly these
intestinal tissues were fixed and cryopreserved without attention at the time to tissue orientation and interest in Paneth cells, so we were not able to reliably ensure a uniform representation of crypts and Paneth cells for quantitative analysis of tissue sections. However, at a qualitative level, differences in the intensities of the IHC signals for CXCL13 and DEFA6 did not appear to change, and the ISH signals appeared to increase slightly during AIDS. Multiple attempts to use a commercially available real-time reverse transcription (RT)-PCR assay for macaque cxcl13 mRNA or to develop an assay in house failed to meet quality control requirements, and therefore accurate measurement of cxcl13 mRNA in lymphoid and intestinal tissue RNA preparations was not achieved. In contrast, the real-time RT-PCR assay for DEFA6 was robust (not shown), and DEFA6 is known to be expressed uniquely by Paneth cells (e.g., Fig. 4A). Using this assay, we found that the levels of expression for this AMP increased ~10-fold during acute infection and AIDS in the ileum (Fig. 5). These changes were positively correlated with SIV plasma virus loads (PVLs) ($\log_{10}$ SIV PVLs versus ileum DEFA6 mRNA levels, $r^2 = 0.53$, $P = 0.04$). This increase in defensin expression is consistent with a recent report that SIV infection leads to induction of rhesus enteric defensin (RED) (19). We also developed real-time RT-PCR assays (SYBR green) for RTD-1 and BDEF2. The RTD-1 assay was not of sufficient robustness to allow quantitation, but the BDEF2 assay was robust. BDEF2 mRNA levels in homogenized macaque ileum did not change early or late after SIV infection (data not shown). Altogether, these data revealed that SIV infection impacts the expression of DEFA6 by Paneth cells.

The cxcl13 promoter is stimulated by the short-chain fatty acid sodium butyrate. To understand better the regulation of CXCL13 expression and its strong expression by Paneth cells, we cloned, sequenced, and functionally examined the cynomolgus (cyCXCL13p) and rhesus (rhCXCL13p) macaque cxcl13 promoters. rhCXCL13p and cyCXCL13p were 99% homologous to one another, and each was 94% homologous to the corresponding human sequences. Basal and cytokine-induced responses of the macaque cxcl13 promoters were measured by transient transfection of HEK293T cells with a reporter plasmid containing the cxcl13 promoter driving luciferase as a readout. Our results showed that neither of the proinflammatory cytokines tumor necrosis factor alpha (TNF-α) nor interleukin-1β (IL-1β) led to induction of luciferase activity from the rhesus or cynomolgus cxcl13 promoters (Table 1), although they potentely increased luciferase activity from the corresponding ccl20 promoters by up to 40-fold (data not shown; unpublished data). In contrast, we found that sodium butyrate, a short-chain fatty acid (SCFA) that is a product of bacterial metabolism and has histone deacetylase (HDAC) inhibitory activity, induced the macaque cxcl13 promoter by about 10-fold (Table 1). Sodium butyrate treatment of cells transfected with a promoterless pGL2-Basic plasmid, showed no change in the level of firefly luciferase activity compared to that in untreated cells, indicating that this intestinal short-chain fatty acid was not acting broadly on any DNA sequence upstream of the luciferase open reading frame (ORF). Treatment of transiently transfected cells with other SCFAs revealed that sodium propionate also stimulated the cxcl13 promoter by 2- to 3-fold, whereas sodium lactate and sodium acetate had only minimal effects (Table 1).

Sodium butyrate has histone deacetylase (HDAC) inhibitory activity (43), and to determine whether sodium butyrate's effects on the cxcl13 promoter were due to this effect, we compared its ability to modulate the cxcl13 promoter after pretreatment of cells

![FIG 4 Localization of antimicrobial peptide and CCL25 mRNAs in intestinal tissues.](image-url)

![FIG 5 Measurement of DEFA6 mRNA levels in macaque ileum. Real-time RT-PCR (TaqMan) was used to measure DEFA6 mRNA levels in homogenized ileum, which was normalized to the endogenous control β-GUS, and then all values were calibrated to one uninfected sample. Shown are the individual values and the means for each group. Statistically significant differences were observed relative to the uninfected group (Mann-Whitney test): *, $P < 0.05$; **, $P < 0.01$.](image-url)
with pharmacologic modulators of chromatin compaction. Pretreatment of cells with trichostatin A, a broad-spectrum inhibitor of HDACs, increased luciferase activity driven by the cxcl13 promoter by almost 2-fold, with no additional increase observed when cells were treated with sodium butyrate after trichostatin A pretreatment (Table 1, bottom). In contrast, pretreatment of cells with the DNA methyltransferase inhibitor 5-azacytidine did not stimulate the cxcl13 promoter and did not inhibit the ability of sodium butyrate to do so. Altogether, these findings suggest that pro-inflammatory cytokines are not likely to control the expression of CXCL13 by Paneth cells or stromal cells, whereas the bacterially produced SCFA sodium butyrate could contribute to CXCL13 expression by Paneth cells primarily through its HDAC inhibitory activity.

**DISCUSSION**

CXCL13 is a homeostatic chemokine that controls, in large part, the migration of B cells, follicular helper T (Tfh) cells, and DCs into the follicles of secondary lymphoid tissues. Because CXCL13...
is responsible for trafficking of B cells and T<sub>fh</sub> cells into the marginal zones of follicles, it is possible that modifications to its expression might contribute to the B cell dysfunction occurring in HIV-1 infection. Here we have examined the expression patterns of CXCL13 in lymphoid tissues, including intestine, and found that SIV infection causes a more disordered distribution of CXCL13 producer cells in LN and spleen and that CXCL13 is expressed by Paneth cells in the small intestine.

A limited number of studies have examined CXCL13 in the context of HIV-1 or SIV infection. CXCL13 levels are elevated in plasma during HIV-1 infection and correlate with levels of CXCL10 in advanced disease (23). In addition, CXCL13 production by B cells is increased in HIV-1-infected patients and there is increased migration toward CXCL13 by peripheral blood B cells from patients with advanced disease (24). CXCL13 protein expression during HIV-1-driven follicular hyperplasia is associated with the GC light zone and the follicular DC (FDC) network. These reports are consistent with our findings of an expansion and redistribution of CXCL13 producer cells in LNs and spleen.

During pathogenic SIV infection, we had previously identified CXCL13 as one of two chemokines with increases in mRNA expression in spleen tissues (10). We focused here on obtaining an improved understanding of the immunobiology of this chemokine in a robust macaque model for HIV/AIDS. We found CXCL13 protein was centrally located within GCs, whereas CXCL13 mRNA was generally localized in cells defining a shell in the mantle and marginal zones of follicles. This disparate microanatomic localization of CXCL13 mRNA and protein was unexpected and underscores the complexity of the in vivo immunobiology of chemokines within tissues. The mechanism by which CXCL13 is concentrated within GCs, compared to the spherical shell of producer cells and the adjacent paracortices, is not clear. There are a number of possible mechanisms, however. For example, the intense staining in the center of the sphere circumscribed by the CXCL13 producer cells could be a consequence of bidirectional diffusion away from the producer cells in both the central follicular and paracortical directions, resulting in an increased overall concentration within GCs and a reduced concentration in paracortices. Alternatively, CXCL13 release might be polarized and limited to the follicular surfaces of producer cells. This would lead to lower concentrations of CXCL13 outside the GCs and increased concentrations within the GCs. In addition, there might be an active transport mechanism that maintains the extracellular CXCL13 within the GC, perhaps piggybacking on cells migrating toward the GC or in microchannels. The latter would be consistent with the thin, cord-like structures we have observed (e.g., Fig. 1) within LNs and spleen. Furthermore, there might be differential expression of proteases that process CXCL13 in lymphoid tissue microcompartments, with a higher concentration of proteases in the surrounding paracortices relative to the inside the GCs. Such compartmentalization of proteases would be consistent with the ability of GCs to harbor immune complexes on FDC surfaces for long periods of time (25–27), and it is possible that FDCs also serve a depot function for CXCL13 as well. Regardless of the mechanism(s), the compartmentalization of CXCL13 within lymphoid tissue GCs is consistent with its role in bringing T<sub>fh</sub> cells, B cells, and DCs together.

In this study, we found there were also disease-specific changes in CXCL13 expression in lymphoid tissues, with the mRNA<sup>+</sup> producer cells expanding more broadly into the GCs and surrounding microcompartment as the course of infection progressed. Recently, increases in LN T<sub>fh</sub> cells in HIV-1 infected individuals were observed despite the overall reduction in the number of CD4<sup>+</sup> T cells that occurs during infection (28). Given that T<sub>fh</sub> cells express CXCR5, the CXCL13 receptor, as well as CXCL13 per se, the more distributed pattern of CXCL13 producer cells we observed could be driven by recruitment and/or expansion of T<sub>fh</sub> cells. In addition, a recent comparison between uninfected and SIV-infected rhesus macaque T<sub>fh</sub> cells revealed a 5-fold induction of CXCL13 expression in T<sub>fh</sub> cells that were SIV infected (29). B cell affinity maturation and isotype switching occur in GCs (30), and given the known B cell dysfunction that occurs during HIV-1 and SIV infection (31–34), it is conceivable that a chemokine that orchestrates the migration of these cells into follicles could contribute to humoral immune response abnormalities that arise during infection. Consistent with this notion, it has been argued recently that there is a decrease in memory B cell levels due to the distortion of the B cell population toward GC B cells and plasma cells during chronic HIV-1 infection (28).

A major finding from the studies presented here was the discovery that CXCL13 was expressed by Paneth cells. Paneth cells are specialized intestinal epithelial cells that express AMPs and proinflammatory cytokines that contribute to innate immunity (20). The AMPs secreted by Paneth cells have several overlapping roles in the intestine (35), including shaping the composition and limiting the number of commensals populating the small intestine, protecting the intestine and its stem cells from invading pathogens, and acting as paracrine signaling molecules (35). It has been argued that Paneth cells are responsible for the homeostatic environments surrounding the intestinal villi and crypts by regulating microbe infiltration (36). Given the expression of CXCL13 by Paneth cells, one of its major functions in the small intestine is likely as an AMP, which would not have been anticipated for an LN homeostatic chemokine. The same regional distribution of CXCL13 along the GI tract as other Paneth cell-expressed AMPs and the colocalization of CXCL13 and DEFA6 in Paneth cells support the interpretation that CXCL13 likely functions in vivo as an AMP. Furthermore, many chemokines have antimicrobial activity in in vitro assays, including CXCL13 (5). Among the microbes against which CXCL13 exhibited antimicrobial activity were the intestinal bacteria E. coli and S. aureus (5). Therefore, we posit that our findings here provide the strongest in vivo evidence, albeit indirect, that chemokines can function as AMPs. It was surprising, however, that the chemokine discovered as expressed by the AMP-producing Paneth cell was a chemokine previously relegated to that of a lymphoid tissue homeostatic chemokine.

In addition to CXCL13, we show here that BDEF2, RTD-1, and Reg3y are also expressed by macaque Paneth cells, this report being the first to demonstrate this for BDEF2 and RTD-1. β-Defensins are expressed by epithelial cells in multiple mucosal tissues, including skin, stomach, and colon. However, they have not been reported as expressed by Paneth cells. Interestingly, in addition to AMP activity, β-defensins also possess chemotactic activity. For example, BDEF2 is chemotactic for CCR6<sup>+</sup> cells (37), which as a regulator of Th17 migration (38) could participate in IL-17’s contributions to intestinal epithelial integrity. The θ-defensins are cyclic defensins originally isolated from leukocytes and bone marrow of rhesus macaques (39). Rhesus θ-defensins and repaired human retrocyclin have broad antimicrobial properties with inhibitory effects on HIV-1, influenza virus, and multiple
bacterial species. As with BDEF2, predominant expression in the GI tract by Paneth cells has not been reported. Finally, Reg3γ is an AMP expressed by murine Paneth cells (21), and our findings reveal the same for macaque Reg3γ. Of these AMPs, we were able to show that DEF6 was increased following SIV infection and these changes were positively correlated with plasma viral loads. Comparison of the sequence of our DEF6 cDNA and that of RED6 revealed they are the same gene, and therefore our finding of increased DEF6 by Paneth cells in cynomolgus macaques is consistent with findings by Zaragoza et al. (19) in rhesus macaques. It is not clear the mechanism(s) by which SIV infection leads to DEF6 upregulation, whereas, for example, BDEF2 is not upregulated. Nonetheless, increased AMP expression could have effects on the intestinal microbiome or could affect the levels of local inflammation. Since both of these effects could impact intestinal epithelial barrier integrity, extrapolating to HIV-1-infected individuals, outcomes might include increased microbial translocation and systemic immune activation (40). Altogether, these findings increase our understanding of the composition of the AMP repertoire produced by Paneth cells, the diversity of which likely reflects the diverse set of microbes targeted as an innate protective mechanism against microbial translocation in the intestinal crypts.

Little is known about the cxcl13 transcriptional promoter, and the limited expression of CXCL13 predominantly by GC-associated cells in lymphoid follicles and by Paneth cells indicates the cxcl13 promoter is tightly regulated. In our hands, even the potent inflammatory cytokines TNF-α and IL-1β failed to stimulate expression from the rhesus or cynomolgus macaque cxcl13 promoter, although cxcl20 promoter constructs were strongly induced by the same cytokines (not shown; unpublished data), indicating that HEK293T cells possess the corresponding, functionally active receptors. SCFAs, such as sodium butyrate, are produced by intestinal bacteria (41, 42), and we found that the macaque cxcl13 promoter was induced by this product of bacterial fermentation, most likely due to its HDAC inhibitor activity (43–45), since it did not stimulate the cxcl13 promoter above what was achieved with the broad-spectrum HDAC inhibitor trichostatin A (Table 1). The antimicrobial cathelicidin LL37 is also stimulated by sodium butyrate (43, 44, 46, 47). The upregulation of the cxcl13 promoter by microbial products, similar to LL37, is consistent with our interpretation that CXCL13 has AMP function in the intestine.

Overall, the findings presented here support the notion that chemokines function as AMPs. This has been demonstrated for many chemokines in vitro, but localization of CXCL13 to Paneth cells, which are specialized for production of AMPs, and transcriptional induction of the cxcl13 promoter by the intestinal bacterial product sodium butyrate provide strong support that at least a subset of chemokines provide antimicrobial function in vivo. That a lymphoid homeostatic chemokine is functioning in these dual capacities is intriguing, and it will be interesting to unravel the structural and biochemical aspects of these different functions of CXCL13.

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