Peyer’s patch myeloid cells infection by *Listeria* signals through gp38+ stromal cells and locks intestinal villus invasion

Olivier Disson1,2, Camille Blériot1,2*, Jean-Marie Jacob3,4*, Nicolas Serafini5,6*, Sophie Dulauroy1,2, Grégoire Gessain1,2, Pierre Thouvenot1,2, Gérard Eberl6,7, James P. Di Santo5,6, Lucie Peduto3,4, and Marc Lecuit1,2,9

The foodborne pathogen *Listeria monocytogenes* (*Lm*) crosses the intestinal villus epithelium via goblet cells (GCs) upon the interaction of *Lm* surface protein InlA with its receptor E-cadherin. Here, we show that *Lm* infection accelerates intestinal villus epithelium renewal while decreasing the number of GCs expressing luminally accessible E-cadherin, thereby locking *Lm* portal of entry. This novel innate immune response to an enteropathogen is triggered by the infection of Peyer’s patch CX3CR1+ cells and the ensuing production of IL-23. It requires STAT3 phosphorylation in epithelial cells in response to IL-22 and IL-11 expressed by lamina propria gp38+ stromal cells. *Lm*-induced IFN-γ signaling and STAT1 phosphorylation in epithelial cells is also critical for *Lm*-associated intestinal epithelium response. GC depletion also leads to a decrease in colon mucus barrier thickness, thereby increasing host susceptibility to colitis. This study unveils a novel innate immune response to an enteropathogen, which implicates gp38+ stromal cells and locks intestinal villus invasion, but favors colitis.

**Introduction**

The intestinal epithelium directly interacts with the external environment and is constantly renewed by stem cells located in intestinal crypts (Barker, 2014). The maintenance of intestinal epithelial homeostasis is tightly regulated by pericryptal stromal cells (Stzepourginski et al., 2017) and the sensing of microbe-associated molecular patterns (MAMPs) from the microbiota (Rakoff-Nahoum et al., 2004; Sommer and Bäckhed, 2013).

Most invasive enteropathogens disrupt gut homeostasis: they can directly modulate epithelial proliferation by inhibiting cell cycle, as does *Shigella flexnerii* IpaB, which acts on Mad2L2 (Iwai et al., 2007), or promote it, as does the gastric carcinogen *Helicobacter pylori* (Mimuro et al., 2002). *Salmonella enterica, S. flexneri*, and attaching and effacing bacteria such as *Citrobacter rodentium*, which are all associated with enteritis, lead to epithelial barrier disruption and intestinal inflammation (Sansonetti et al., 1999; Collins et al., 2014). *C. rodentium*-associated epithelial damage leads to MyD88-dependent epithelial proliferation (Gibson et al., 2008). In a *Drosophila melanogaster* model of intestinal infection, epithelial proliferation is the consequence of the immune oxidative burst and is JAK/STAT dependent (Buchon et al., 2009). This phenotype is mediated by signaling from the damaged epithelium to stem cells and amplified by visceral muscles (Buchon et al., 2013). In noninfectious models of dextran sodium sulfate (DSS)–induced colitis and methotrexate-induced damage to stem cells, intestinal epithelium wound healing depends on intestinal epithelial cell STAT3 activation by IL-22 (Pickert et al., 2009; Aparicio-Domingo et al., 2015). Upon *C. rodentium* infection, IL-23 expression by CX3CR1+ cells triggers IL-22 expression by type 3 innate lymphoid cells (ILCs; Longman et al., 2014; Aychek et al., 2015). It has also been shown that IL-22 acts on enterocytes in a STAT3-dependent manner, inducing RegIIβ and RegIIγ expression (Zheng et al., 2008; Manta et al., 2013). Epithelial renewal upon infectious- and noninfectious-associated damages may therefore engage the same signaling.

The intestinal phase of listeriosis, a systemic infection caused by the foodborne pathogen *Listeria monocytogenes* (*Lm*), is mostly asymptomatic (Charlier et al., 2017), in line with the observation that *Lm* does not significantly alter the intestinal barrier integrity...
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immune cells responding to pathogen-elicited stimuli.

al., 2001; Disson et al., 2008). In humanized Ecad mice, Lm is rapidly transcytosed at the small intestinal level in an InlA–Ecad-dependent manner across goblet cell (GCs) expressing luminaly accessible Ecad and released into the lamina propria (LP; Fig. S1 A; Lecuit et al., 2001; Nikitas et al., 2011). Lm is also transferred, albeit at a lower efficiency, through M cells in an InlA-independent manner at the Peyer’s patch (PP) level, the only route of infection in nonhumanized mice (Jensen et al., 1998; Chiba et al., 2011; Gessain et al., 2015). We have shown by transcriptomic analysis that the global intestinal host response to Lm is InlA independent and triggered by invasion of PPs (Fig. S1 A; Lecuit et al., 2007). It requires the expression of listeriolysin O (LLO; Lecuit et al., 2007), a major Lm virulence factor involved in Lm escape from its phagocytic vacuole and survival in professional phagocytes (Hamon et al., 2012). We have also shown that Lm induces IL-22 and IFN-γ upon oral infection in humanized Ecad mice (Reynders et al., 2011). Whereas IFN-γ is required to control systemic Lm infection (Harty and Bevan, 1995), IL-22 is not (Graham et al., 2011).

Lm impact on intestinal epithelium homeostasis, although potentially critical for the outcome of the infection, has not been studied. We therefore investigated intestinal epithelium response to orally acquired listeriosis. We show here that Lm induces intestinal epithelial cell proliferation and depletion of GCs expressing accessible Ecad, leading to a complete blockade of Lm intestinal villus invasion. Intestinal epithelium proliferation and GC depletion are independent of Lm intestinal villus invasion, but strictly depend on infection of PP CX3CR1+ cells, which express IL-23 upon infection, leading to STAT3 activation in enterocytes. However, in contrast to host responses to intestinal epithelial damage, Lm-associated IL-23–dependent STAT3 phosphorylation involves not only IL-22 expression, but also induction of IL-11 by LP CD45+ gp38+ pericryptal stromal cells. Importantly, we show that IL-23/IL-22/IL-11–dependent intestinal epithelium response to Lm also critically requires IFN-γ–dependent STAT1 phosphorylation. We further demonstrate that this innate immune pathway leads to a decrease of mucus barrier thickness at the colon level, a known promoter of intestinal inflammation (Van der Sluis et al., 2006). Indeed, Lm-associated epithelial response leads to gut sensitization to colitis.

Altogether, these results identify a new innate immune pathway initiated at the PP level against infection, which relies on IL-22– and IL-11–dependent STAT3 activation in epithelial cells and the conjugated action of IFN-γ. This study unveils a new layer of regulation of the intestinal epithelium in response to invasive pathogens and the unexpected role of LP CD45+ gp38+ CD34+ stromal cells as innate immune cells responding to pathogen-elicited stimuli.

Results

Lm infection leads to intestinal epithelium proliferation

We first investigated intestinal epithelium proliferation upon Lm oral inoculation by quantifying BrdU incorporation in KIE16P humanized mouse intestinal epithelium. Whereas only cells located in intestinal crypts incorporated BrdU at steady-state (Barker et al., 2008), oral infection with two genetically distant WT Lm strains (EGD and EGDe) induced a significant increase in BrdU+ epithelial cells (Fig. 1A and Fig. S1 B). Increase in enterocyte BrdU incorporation was noticeable as early as day 2 post infection (pi). As BrdU was injected i.p. and incorporated in dividing cells 16 h before tissue sampling, this indicates that proliferation begins in the first day pi. Proliferation peaked between day 3 and 4 pi and returned to basal level at day 6 pi (Fig. 1B). In line with these results, more Ki67+ cycling cells were counted in crypts and at crypt-villus junctions upon Lm oral infection (Fig. 1C and Fig. S1 C; Whitfield et al., 2006; Cuylen et al., 2016). No leakage of the epithelial barrier was detected in a biotin barrier assay (Fig. S1 J).

Lm-associated intestinal epithelium proliferation is independent of villus epithelial cell invasion, but requires PP infection

We next investigated the bacterial determinants of intestinal epithelial proliferation. Since Lm crosses the villus intestinal epithelial barrier in an InlA-dependent manner (Fig. S1 A), we investigated if cell proliferation was dependent of InlA-mediated villus invasion. Enterocyte proliferation was detected in all intestinal crypts observed, irrespective of local Lm villus invasion, while only 10% of villi are infected (Fig. S1 G). This argues against local proliferation in response to InlA-dependent infection of villi. To directly investigate the role of InlA-dependent villus invasion in Lm-associated intestinal epithelium proliferation, KIE16P humanized mice were infected with an isogenic Lm ΔinlA mutant, which is unable to invade intestinal villi, but crosses via PPs. As expected, intestinal bacterial load was lower upon infection with Lm ΔinlA as compared with WT bacteria (Fig. S1 H). Nevertheless, Lm-associated intestinal epithelium proliferation was comparable in both conditions (Fig. 1, D and E). As previously shown, Lm invades via M cells in an InlA-independent manner the PP (Lecuit et al., 2001; Chiba et al., 2011; Nikitas et al., 2011), where host response to Lm is initiated (Fig. 1 F and Fig. S1 A; Lecuit et al., 2007). To investigate the role of PPs in Lm-associated intestinal epithelium proliferation, we injected Lm ΔinlA in ligated loops containing or not containing a PP. Lm-associated intestinal epithelium proliferation was strictly dependent of the presence of PPs (Fig. 1 G), demonstrating the critical role of PPs in Lm-associated intestinal epithelium proliferation and the absence of contribution of InlA-dependent invasion in this phenomenon. Of note, proliferation was higher in villi near to PPs, suggesting a gradient of induction from PP to LP (Fig. S1 I). We confirmed that proliferation is InlA-independent in nonhumanized C57BL/6J mice, in which Lm entry occurs only through PP M cells (Fig. S1 J).

As intestinal proliferation is regulated by the microbiota (Rakoff-Nahoum et al., 2004), we investigated whether Lm-associated intestinal epithelium proliferation was indirect, as a

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Figure 1. Lm-induced epithelial cell proliferation in an InlA-independent but PP-dependent manner. (A) Fluorescent imaging of sections of the ileum obtained from KIE16P mice 4 d after oral infection with $5 \times 10^9$ Lm. 16 h before euthanasia, mice were injected with BrdU. Sections are stained for proliferating cells (BrdU), epithelial cells (Ecad), and nuclei (Hoechst). Bars, 20 µm. (B) Quantification over time of epithelial proliferation. Fold increase is measured by dividing the length between the first and last BrdU$^+$ epithelial cells by the length of the villi, normalized by the proliferation from the noninfected mice. n = 9–39.
consequence of an interaction with the microbiota. As in conventional mice, proliferation was induced in germ-free mice mono-associated with Lm, demonstrating the direct impact of Lm on proliferation, rather than via interactions with the microbiota (Fig. 1 H).

**Lm-associated intestinal epithelium proliferation depends on epithelial STAT3 activation and leads to a blockade of Lm intestinal villus invasion via GCs**

STAT3 activation leads to epithelial cell proliferation upon DSS-induced colitis and methotrexate-induced damage to stem cells (Pickert et al., 2009; Aparicio-Domingo et al., 2015). We generated and infected nonhumanized C57BL/6 Stat3fl/fl vil-Cre-ERT2 mice, in which Stat3 is excised in intestinal epithelial cells (IEC) expressing Villin by hydroxytamoxifen. Proliferation was totally impaired in STAT3IEC-KO mice as compared with STAT3IEC-WT littermates, demonstrating the critical role of epithelial STAT3 in Lm-associated intestinal epithelium proliferation, independently of InlA-dependent entry (Fig. 2, A and B). GCs expressing accessible Ecad are the main sites of InlA-dependent Lm translocation across the intestinal epithelium (Nikitas et al., 2011). Since STAT3-dependent epithelial proliferation may be linked to the turnover and differentiation of GCs (Gonneaud et al., 2016), we investigated the impact of Lm-associated intestinal epithelium proliferation on GCs. We labeled Muc2, the major component of mucin synthesized by GCs, with an anti-Muc2C3 antibody (Johansson et al., 2008). We also stained glycosylated mucins with wheat germ agglutinin (WGA), which specifically binds to sialic acid and N-acetyl-glucosaminyl carbohydrate residues on mature, modified mucins (Fischer et al., 1984). The number of WGA+ Muc2+ GCs was significantly decreased in intestinal villi from infected mice as early as 1 d pi and recovered 14 d pi (Fig. 2, C and D; and Fig. S2 A). The decrease in WGA+ GC number was STAT3-dependent, as excision of STAT3 in Stat3fl/fl vil-Cre-ERT2 mice prevented Lm-induced decrease in GC number (Fig. 2 E and Fig. S2 B). As for proliferation, WGA+ GC number regulation was InlA independent (Fig. S2 C). We next investigated the expression of Muc2 in GCs. Unlike the clear decrease of WGA+ GCs, the number of total Muc2+ GCs only slightly decreased in intestine (Fig. 2, D and F). Data from literature indicate that GCs that harbor less glycosylated mucins are not fully mature (Asker et al., 1998; Nowarski et al., 2015). We therefore investigated if the decrease in WGA+ GCs may correlate with other markers of GC differentiation. We have shown previously that WGA+ GCs express luminaly accessible Ecad, which is targetted by Lm (Nikitas et al., 2011). We observed a dramatic STAT3-dependent decrease of WGA+ GCs expressing accessible Ecad upon Lm infection (Fig. 2, G and H).

We next investigated the impact of decreased accessible Ecad on Lm InlA-dependent villus invasion. We generated humanized Ecad KIE16P STAT3IEC-KO and STAT3IEC-WT mice in which InlA-dependent entry occurs. To this end, we inoculated these mice with WT or Lm ΔInlA at 5 × 10^6 CFUs to activate STAT3. As expected, proliferation was inhibited in KIE16P STAT3IEC-KO compared with KIE16P STAT3IEC-WT (Fig. S2 D). 3 d later, we reinoculated with the same bacteria at higher dose of 5 × 10^8 CFUs for 4 d. Strikingly, InlA-dependent Lm entry was fully inhibited in KIE16P STAT3IEC-WT mice, whereas it was not in KIE16P STAT3IEC-KO, showing that Lm infection blocks subsequent InlA-dependent entry in a STAT3-dependent manner (Fig. 2 I, middle). In sharp contrast, no impact was detected on translocation of Lm ΔInlA, suggesting that STAT3 activation inhibits specifically InlA-dependent entry that occurs via GCs (Fig. 2 I, right). As a control, we infected nonhumanized C57BL/6 Stat3fl/fl vil-Cre-ERT2 and STAT3IEC-WT mice (Fig. 2 I, left) and confirmed that STAT3 has no impact on InlA-independent invasion, whereas proliferation and the number of WGA+ GCs are modulated in these mice.

Together, these results indicate that STAT3 activation, which is induced by Lm in an InlA-independent manner, leads to a complete blockade of InlA-dependent villus invasion by depleting specifically GCs expressing luminaly accessible Ecad.

**Infection of PP CX3CR1 initiates Lm-associated intestinal epithelium response**

We next investigated how STAT3 is activated in the epithelium upon Lm infection in an InlA-independent but PP-dependent manner. We observed that Lm is rapidly captured mostly by CX3CR1+ cells in PPs upon oral infection with 5 × 10^8 CFUs, whereas no bacteria was detected in PPs with 5 × 10^7 CFUs, which fits with the inoculum required for enterocyte proliferation (Fig. 3 A and Fig. S2, E and G). We next investigated the role of PP CX3CR1 in Lm-associated intestinal epithelium response. When compared with Cx3cr1GFP/+ littermates, Lm-associated epithelial cell proliferation in Cx3cr1GFP/GFP mice was totally abolished (Fig. 3 B), even though Lm were present in both CX3CR1GFP/+ and CX3CR1GFP/GFP cells (Fig. 3 A and Fig. S2 F). This indicates that CX3CR1 expression is critically needed for Lm-associated intestinal epithelium proliferation. Proliferation of epithelial cells and WGA+ cell decrease required expression of LLO, which is encoded by hly and is necessary for bacterial escape from phagocytic vacuole (data

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Figure 2. *Lm* triggers a STAT3-dependent negative feedback loop blocking its access to intestinal villi via GCs. (A) Confocal imaging of sections of the ileum obtained from STAT3IEC-WT and STAT3IEC-KO mice treated as in Fig. 1A with $5 \times 10^9$ *Lm* and isogenic mutants. Bars, 20 µm. (B) Quantification of epithelial proliferation in STAT3IEC-WT and STAT3IEC-KO littermate mice 4 d after oral inoculation. Each dot corresponds to one quantified villus, except the red dots, which correspond to the mean of each mouse. *n* = 3–5 mice (PBS) or 6–7 mice (*Lm*)/condition. Data are pooled from three independent experiments. A one-way ANOVA test followed by a Sidak’s multiple comparisons was performed on the mean of each mouse. (C) Quantification of WGA* GCs/villous at different time points after inoculation. *n* = 8–25 independent villi. Data are represented as mean ± SEM. Mann-Whitney test.
not shown). LLO is also required for *Lm*-associated intestinal epithelium proliferation in germ-free mice as well as in C57BL/6 mice, where entry occurs only via PPs (Fig. 1 H and Fig. S1 I). This suggests that *Lm* access to PP CX3CR1*+* cytoplasm is needed for signaling. We next investigated if PP CX3CR1*+* cells are involved in signaling upstream of STAT3. IL-23 and its downstream effector IL-22 are expressed in a CX3CR1-dependent manner upon *C. rodentium* infection (Longman et al., 2014; Aychek et al., 2015), and IL-22 is a potent inducer of epithelial proliferation (Pickert et al., 2009; Lindemans et al., 2015). We investigated if, as do colonic CX3CR1*+* cells upon *C. rodentium* infection, PP CX3CR1*+* cells express IL-23 upon *Lm* infection. At steady-state, most CX3CR1*+* cells were CX3CR1*high* in PPs and villi, as previously reported (Varol et al., 2009; Bonnardel et al., 2015). However, upon 24 h of *Lm* oral infection, the number of CX3CR1*high* mononuclear phagocytes (MPs) was significantly increased in PPs compared with noninfected tissue and villi from infected mice (Fig. 3 C). Upon infection, CX3CR1*+* and especially PP CX3CR1*int* MPs, express IL23 (Fig. 3 D and S2 H). We next showed that induction of *Il23p19* expression is CX3CR1-dependent in CX3CR1*GFP/GFP* mice (Fig. 3 E). These data demonstrate that CX3CR1 expression in PPs is necessary for IL-23 production in response to *Lm* infection.

We next investigated the role of IL-23 on *Lm*-associated intestinal epithelial proliferation. IL-23 pathway disruption in *Il23p19−/−* mice fully impaired *Lm*-associated enterocyte proliferation (Fig. 3 F), indicating that IL-23, while being produced at the PP level, exerts its effect at the villus level. Surprisingly, enterocyte proliferation in response to *Lm* was still observed in *Il22−/−* mice (Fig. 3 F). We therefore investigated whether other factors contributing to *Lm*-associated intestinal epithelial proliferation were involved and induced by IL-23. *Lm*-associated enterocyte proliferation was not abolished in *Rag2−/−* mice, devoid of B and T cells, and was partially observed in *Rag2−/−Il17ra−/−* mice, which lack IL-17-induced signaling (Fig. 3 G). However, *Lm*-associated enterocyte proliferation was absent in *Rag2−/−Il17ra−/−Il11ra−/−* mice when treated with an anti–IL-22 antibody (Fig. 3 G), indicating that both IL-22 and IL-17 are involved in *Lm*-associated epithelial proliferation.

**IL-11 expressed by gp38+ stromal cells is involved in *Lm*-associated epithelial response**

To decipher the complementary roles of IL-22 and IL-17 in *Lm*-associated epithelial response, we investigated the contribution of another pathway which is induced by *Lm* infection. In vitro stimulation of gp38+ stromal cells by recombinant IL-23 did not induce expression of IL-11 (Fig. 4 D), consistent with the observation that gp38*+* stromal cells do not express IL-23 receptor (Fig. S3 C). These results suggest that another factor bridges IL-23 signaling with IL-11 expression. Since IL-17 plays a role in *Lm*-associated enterocyte proliferation (Fig. 3 G), we investigated its contribution to gp38*+* stromal cells’ IL-11 production. IL-17 was sufficient to increase IL-11 release by gp38*+* stromal cells (Fig. 4 D), indicating a role for IL-17 in IL-11 production.

We next investigated in vivo whether IL-11, IL-17, and IL-22 complement IL-23 inhibition to induce *Lm*-associated enterocyte proliferation. Treatment of *Lm*-infected *Il22p19−/−* mice with recombinant IL-11, IL-17, or IL-22 fully restored *Lm*-associated epithelial proliferation, as well as a decrease in WGA*+* GCs (Fig. 4, E and F). Together, these results show that intestinal gp38*+* stromal cells are the major producer of IL-11 after *Lm* infection, which acts in concert with IL-22 downstream of IL-23 and IL-17 to induce *Lm*-associated intestinal epithelium proliferation. However, injection of recombinant IL-11 or IL-22 does not induce enterocyte proliferation or a decrease in WGA*+* GCs in uninfected mice (Fig. 4, E and F), indicating that STAT3 activation is necessary, but not sufficient, to induce epithelial proliferation and requires another pathway which is induced by *Lm* infection.
Figure 3. IL-23 produced by CX3CR1 is involved in Lm-associated epithelial proliferation. (A) Confocal imaging of a vibratome section from a PP infected by $5 \times 10^9$ Lm 24 h after oral inoculation. Bars, 20 µm. (B) Quantification of epithelial proliferation in Cx3cr1<sup>GFP/GFP</sup> and Cx3cr1<sup>GFP/+</sup> cohoused control mice 4 d after oral inoculation. **, P < 0.01; ***, P < 0.001. Each dot corresponds to one quantified villus, except the red dots, which correspond to the mean of each mouse. n = 4 mice/condition. Data are pooled from three independent experiments. A one-way ANOVA test followed by a Sidak’s multiple comparisons was performed on the mean of each mouse. (C) Cell number per organ counted by flow cytometry. All PP versus entire intestinal tissue. Data are representative of ...
IFN-γ production by natural killer (NK) cells is required for Lm-associated epithelial response

IFN-γ is a major cytokine controlling Lm infection in vivo (Harty and Bevan, 1995). IFN-γ is also involved in tissue homeostasis, in particular in the intestine and liver (Smith et al., 2000; Brooling et al., 2005). We investigated enterocyte proliferation in Ifnγ−/− mice orally inoculated with Lm. As expected, IFN-γ depletion led to an increased Lm systemic burden (Fig. S4 A), yet no enterocyte proliferation was detectable, demonstrating the critical role of IFN-γ in Lm-associated enterocyte proliferation (Fig. 5 A). As previously shown, IFN-γ was mostly produced by NK cells and to a lower degree by T cells (Fig. 5 B and Fig. S4 B; Andersson et al., 1998; Reynders et al., 2011). Lm-associated enterocyte proliferation was still observed in Rag2−/− mice, which lack T and B cells, but was fully abolished in Rag2−/−γC−/− mice, which also lack ILCs and NK cells (Fig. 5 C). To investigate if group 1 ILCs (including ILC1 and NK cells) are the cells involved in Lm-associated enterocyte proliferation, we depleted all ILC1 cells in Rag2−/− mice injected with an antibody against NK1.1, a condition where ILC2s and ILC3s are preserved, but ILC1s are totally absent. Lm-associated epithelial proliferation was completely abrogated, as in Rag2−/−γC−/− mice, indicating the essential role of ILC1s in Lm-associated enterocyte proliferation (Fig. 5 C).

We next investigated the signal triggering IFN-γ production by NK cells in response to Lm oral infection. It has been shown that CX3CR1+ cells produce IL-12 during sepsis (Ishida et al., 2008), leading to IFN-γ production by NK cells. Accordingly, CX3CR1− MPs in PPs, but not in villi, expressed IL-12 in response to Lm infection (Fig. 5 D), and expression of both IFN-γ and IL-12 was CX3CR1-dependent (Fig. S4, C and D). IL-23 is a potent inducer of IFN-γ in Helicobacter hepaticus infection in mice (Buonocore et al., 2010). However, in our model, IFN-γ was induced after Lm infection in both WT and Il23p19−/− mice, indicating that in this context, IFN-γ production is IL-23 independent (Fig. S4 E).

IFN-γ can act both on hematopoietic and nonhematopoietic cells. To decipher if myeloid hematopoietic (radio-sensitive) or nonmyeloid nonhematopoietic (radio-resistant) cells are involved in IFN-γ-dependent Lm-associated enterocyte proliferation, we analyzed bone marrow chimera mice for which particular in the intestine and liver (Smith et al., 2000; Brooling and Bevan, 1995). IFN-γ is also involved in tissue homeostasis, in particular in the intestine and liver (Smith et al., 2000; Brooling and Bevan, 1995). IFN-γ is also involved in tissue homeostasis, in particular in the intestine and liver (Smith et al., 2000; Brooling and Bevan, 1995). IFN-γ can act both on hematopoietic and nonhematopoietic cells. To decipher if myeloid hematopoietic (radio-sensitive) or nonmyeloid nonhematopoietic (radio-resistant) cells are involved in IFN-γ-dependent Lm-associated enterocyte proliferation, we analyzed bone marrow chimera mice for which particular in the intestine and liver (Smith et al., 2000; Brooling and Bevan, 1995). IFN-γ is also involved in tissue homeostasis, in particular in the intestine and liver (Smith et al., 2000; Brooling and Bevan, 1995). IFN-γ can act both on hematopoietic and nonhematopoietic cells. To decipher if myeloid hematopoietic (radio-sensitive) or nonmyeloid nonhematopoietic (radio-resistant) cells are involved in IFN-γ-dependent Lm-associated enterocyte proliferation, we analyzed bone marrow chimera mice for which particular in the intestine and liver (Smith et al., 2000; Brooling and Bevan, 1995). IFN-γ is also involved in tissue homeostasis, in particular in the intestine and liver (Smith et al., 2000; Brooling and Bevan, 1995).

We have previously shown that Lm also targets the cecum and colon (Disson et al., 2008), and we investigated if Lm also targets the cecum and colon (Disson et al., 2008). We observed an increase of proliferation in the colon (Fig. S5 A), as well as a decrease in the number of WGA+ GCs upon Lm infection, while the number of Muc2+ GCs was not modified (Fig. S5, B and C).

Depletion of mucus sensitizes mice to colitis (Van der Sluis et al., 2006). We therefore investigated the impact of Lm-associated WGA+ GC depletion on luminal mucus content and its consequences on susceptibility to colitis. The distance between the microflora and the epithelium, which reflects mucus thickness, was strongly decreased in a STAT3-dependent manner in the ileum and cecum (Fig. S5, D–G). At the colon level, the thickness of the highly organized inner mucus layer, measured by a Muc2+ staining, was reduced upon Lm infection (Fig. 7, A and B). Thus, Lm-associated decrease of the differentiation of the GCs inhibits Lm entry, while leading to a significant decrease in the physical context, IFN-γ production is IL-23 independent (Fig. 5 E).

STAT3 and STAT1 activations lead to intestinal enterocyte proliferation

Both IL-22 and IL-11, known to activate STAT3 and IFN-γ, which activates STAT1, are required in vivo for Lm-associated enterocyte proliferation. IL-22/IL-11 and IFN-γ are expressed 24–48 h pi (data not shown; Reynders et al., 2011). To determine whether IL-22, IL-11, and IFN-γ act in an interdependent and direct manner on epithelial cells to induce proliferation, we used intestinal organoids. We first determined by Western blotting that STAT3 is phosphorylated upon IL-22 or IL-11, but not IL-23, treatment of intestinal organoids and that STAT1 is phosphorylated upon IFN-γ treatment, indicating a direct effect of these cytokines on epithelial cells (Fig. S4, F and G). As expected, induction of proliferation in organoids did not occur with IL-23 (data not shown). It also did not occur with IL-22 or IFN-γ alone, indicating that these cytokines are not sufficient to induce proliferation (Fig. 6 A and Fig. S4 I for the method). In contrast, proliferation was observed when both a STAT3 inducer (IL-22 or IL-11) and a STAT1 inducer (IFN-γ) were simultaneously added to intestinal organoids (Fig. 6, A–C). STAT3 excision by the addition of hydroxytamoxifen to Stat3fl/fl Villin-cre intestinal organoids blocked proliferation induced by IL-22 and IFN-γ, indicating the critical requirement for STAT3 in proliferation (Fig. 6 D and Fig. S4 H). STAT1 inhibitor fludarabine (Frank et al., 1999) also led to a decrease in proliferation, indicating that both STAT3 and STAT1 activators are required to induce epithelial cell proliferation (Fig. 6 E).

Lm infection increases susceptibility to colitis

We have previously shown that Lm also targets the cecum and colon (Disson et al., 2008). We investigated if Lm regulates epithelial homeostasis at the cecum and colon level. We observed an increase of proliferation in the colon (Fig. S5 A), as well as a decrease in the number of WGA+ GCs upon Lm infection, while the number of Muc2+ GCs was not modified (Fig. S5, B and C). Depletion of mucus sensitizes mice to colitis (Van der Sluis et al., 2006). We therefore investigated the impact of Lm-associated WGA+ GC depletion on luminal mucus content and its consequences on susceptibility to colitis. The distance between the microflora and the epithelium, which reflects mucus thickness, was strongly decreased in a STAT3-dependent manner in the ileum and cecum (Fig. S5, D–G). At the colon level, the thickness of the highly organized inner mucus layer, measured by a Muc2+ staining, was reduced upon Lm infection (Fig. 7, A and B). Thus, Lm-associated decrease of the differentiation of the GCs inhibits Lm entry, while leading to a significant decrease in the physical context, IFN-γ production is IL-23 independent (Fig. 5 E).

Two independent experiments with three mice per group. Data are represented as mean ± SEM. A one-way ANOVA test followed by a Sidak’s multiple comparisons was performed on the mean of each mouse. n = 4–9 mice/condition. Data are pooled from four independent experiments. A one-way ANOVA test followed by a Sidak’s multiple comparisons was performed on the mean of each mouse.
separation between the epithelium and the luminal content, a well-established risk factor for colitis. To investigate whether Lm-associated decrease in mucus layer increases susceptibility to colitis, we fed mice with DSS for seven consecutive days, 7 d after Lm ΔinlA infection, when the IL-23 and IFN-γ pathways are no more induced (our observation; Reynders et al., 2011). Mice were sacrificed 14 d pi, and the colitis was evaluated. Infection with Lm ΔinlA alone, which does not invade epithelial cells, did not
induce gut lesions 14 d pi, and pathology was similar to noninfected control (Fig. 7 C, top and second from bottom; and Fig. 7 D). As previously described, DSS treatment of noninfected mice induced colitis (Okayasu et al., 1990), as quantified by colonic inflammation and epithelial and mesenchymal lesions (Fig. 7 C, second from top; and Fig. 7 D). Very interestingly, as compared with controls treated by DSS only or with an Lm Δhly strain (Fig. S5 H), with which Lm-associated intestinal response is abolished, animals exposed to Lm ΔinlA before DSS, in which the mucus thickness was reduced, developed significantly more severe colitis (Fig. 7 C, bottom; and Fig. 7 D). Altogether, this indicates that Lm significantly increases susceptibility to colitis.

Discussion
In contrast to other foodborne infections, the intestinal phase of listeriosis is clinically silent in most patients (Charlier et al., 2017). In line with this observation, the crossing of the intestinal barrier by Lm is not associated with epithelial damage, in contrast to other enteropathogens such as S. enterica, S. flexnerii,
and C. rodentium, a model for attaching/effacing pathogens in mice inducing epithelial lesions (Collins et al., 2014). Here, we have shown that upon Lm infection, PP CX3CRI+ cells trigger both IL-23/IL-22/IL-11 and IFN-γ pathways, which act in combination to accelerate epithelial cell renewal and decrease the number of WGA+ GCs expressing accessible Ecad. This leads to a complete blockade of intestinal villus invasion by Lm. Unexpectedly, this response, which critically requires IL-23, is partially IL-22–dependent and requires the expression of IL-11 by intestinal gp38+ stromal cells (Fig. 8).

The number of GCs and their differentiation are modulated upon infection with other enteropathogens. Indeed, C. rodentium leads to a depletion of GCs and a decrease in the thickness of the mucus layer during the first phase of the infection, which is reversed during the clearance of the infection (Gustafsson et al., 2013). C. rodentium and Salmonella typhimurium modulate GC number in an IFN-γ-dependent manner, and IFN-γ is expressed by CD4+ T+ cells in the case of C. rodentium infection (Songhet et al., 2011; Chan et al., 2013; Klose et al., 2013). Lm translocation across intestinal epithelium at the villus level occurs in an InlA-dependent manner via luminally accessible Ecad on GCs (Nikitas et al., 2011). Decrease of WGA+ GC number expressing luminally accessible Ecad in response to Lm therefore selectively blocks InlA-dependent translocation of Lm at the villus level and constitutes a novel and specific innate immune mechanism against an enteropathogen (Fig. S1 A and Fig. 8).

This pathway, which is not initiated upon villus infection, is triggered at the PP level. This is in agreement with previously published results showing that host response to Lm at the intestinal level is fully dependent on Lm PP infection (Lecuit et al., 2013).
Our results further illustrate that invasion of villi via GCs in an InlA-dependent manner is silent. This study underlines on one hand the relative unresponsiveness of the villus LP to pro-inflammatory stimuli and on the other hand the critical role of PP in danger detection and propagation of defense signals to the intestinal epithelium.

IL-22 induces epithelial proliferation through STAT3 activation during DSS-induced colitis (Pickert et al., 2009) and is expressed downstream of IL-23 in response to C. rodentium infection in the colon (Aychek et al., 2015). However, although epithelial proliferation is fully IL-23 and STAT3 dependent upon Lm infection, IL-22 only partially relays IL-23 signaling at the intestinal level. We have shown here that IL-17, which is produced by γδ T cells upon Lm infection (Hamada et al., 2008; Sheridan et al., 2013), induces Lm-associated enterocyte proliferation together with IL-22. IL-17 in turn triggers IL-11 expression by LP gp38⁺CD34⁺ stromal cells, which activates STAT3 in epithelial cells (Fig. 8). IL-11 is a known activator of STAT3, involved in gastrointestinal tumorigenesis in human and in mouse (Putoczki et al., 2013). IL-11 is also induced upon C. rodentium, by a so-far unknown
mechanism and is involved in protection against TLR4-mediated colitis (Gibson et al., 2010). Whereas inhibition of IL-11 does not exacerbate Lm burden in systemic listeriosis (Opal et al., 2000), the role of IL-11 during Lm oral infection had so far not been investigated. Several cell types have been described to express IL-11 in the intestine (Putoczki and Ernst, 2010). We show here that LP gp38+CD34+ stromal cells, a mesenchymal cell subset in close contact with the intestinal crypts, are the major intestinal source of IL-11 upon Lm infection. Several evidences indicate that mesenchymal stromal cells are involved in mucosal immune responses, notably via IL-33 (Owens and Simmons, 2013; Mahapatro et al., 2016). Here, we have uncovered that gp38+CD34+ stromal cells are part of a host response pathway involving CX3CR1-derived IL-23 and γδ T-derived IL-17, which leads to intestinal epithelial cells proliferation and depletion of GCs expressing accessible Ecad in a STAT3-dependent manner in response to an enteric infection. The partially redundant and complementary roles of IL-11 and IL-22 upon Lm infection remain to be characterized in more detail. We have shown that IL-11 effect is compensated by IL-22 in Il11ra−/− mice, but is unmasked when IL-22 is also absent. The induction of STAT3 via the respective receptors of IL-11 and IL-22 may allow the fine tuning in time and space of host tissue responses to infection (Ernst et al., 2014).

IL-22 and IL-11 are not sufficient to induce proliferation and GC depletion in noninfected mice, indicating that a cofactor induced by Lm infection is necessary. IFN-γ is expressed upon Lm infection (Anderson et al., 1998), and both IL-22 and IFN-γ are induced in an InlA-independent manner 24 h pi and peak 48 h pi upon Lm oral infection (Reynders et al., 2011), indicating that they may act together rather than sequentially. We could recapitulate ex vivo stem cells proliferation by simultaneously treating with IL-22 and IFN-γ intestinal organoids. These results indicate that IL-22/IL-11–dependent STAT3-induced proliferation is not sufficient to induce proliferation, which indeed requires STAT1 phosphorylation, in response to IFN-γ receptor engagement. In line with this finding, IL-22–dependent STAT3-induced proliferation is involved in vivo in tissue repair in a graft-versus-host tissue damage model (Lindemans et al., 2015) and DSS-induced colitis (Pickert et al., 2009; Geng et al., 2018). IL-22 and IFN-γ are also induced upon the protozoan Tritrichomonas musculis infection, which increases susceptibility to colorectal cancer induced in Apcmmin/+ mice (Chudnovskiy et al., 2016).

WGA+ GC depletion and the decrease of luminally accessible Ecad constitute a specific and very efficient short-term host defense response to Lm infection, by locking its portal of entry (Fig. 8 and Fig. S5 I). But this response is also potentially detri-
nal for the host by reducing the thickness of the protective mucosal barrier all along the gut and especially in the colon where the mucus layer is highly organized. Decrease of mucus layer thickness has been shown to be associated with the development of colitis (Van der Sluis et al., 2006). In line with these results, we show here that Lm increases host susceptibility to colitis. This provides support to previous report on the association between Lm presence in the gut and the development of inflammatory bowel disease (IBD; Liu et al., 1995; Hugot et al., 2003; Miranda-Bautista et al., 2014). Decrease of mucus layer thickness has also been shown to be associated with colorectal cancer (Velchich et al., 2002) and Crohn’s disease (Niv, 2016). IL-23, IL-22, and IL-11 can promote tumor growth in a mouse model of colorectal tumorigenesis (Grivennikov et al., 2012; Putoczki et al., 2013) and are associated with IBD (Eken et al., 2014). Chronic Lm intestinal carriage is more likely in Western countries where refrigeration favors Lm food contamination and human exposure (Swaminathan and Gerner-Smidt, 2007) and may be associated with gallbladder infection (Begley et al., 2009; Charlier et al., 2014). This study suggests that chronic Lm intestinal carriage may favor both IBD and tumorigenesis, through both decreased mucus layer thickness and production of cytokines such as IL-23 and IL-11, in conjunction with host factor predisposing for these conditions. This hypothesis is in line with the highest incidence of IBD and colon cancer in Western countries (Ng et al., 2013; Kuipers et al., 2015). Assessing the biological and medical relevance of this correlation will require further epidemiological and experimental studies.

Materials and methods

Mice and infection

All the procedures were in agreement with the guidelines of the European Commission for the handling of laboratory animals, directive 86/609/EEC. They were approved by the ethical committees No. 59 under the number 2010-0020 and by the CET EA/CEEA No. 89 under the numbers HA0017, 2015-0014, and 2017-0057. Knock-in E16P mouse line was from our laboratory. Cx3cr1GFP mice (Jung et al., 2000; RRID: MGI:2670353) were provided by the Cryopréservation, Distribution, Typage et Archivage animal Orléans, Orléans, France. Il23p19−/− mice (Ghilardi et al., 2004) were provided by Genentech through B. Ryffel (CDTA Orléans). Rag2−/−, Rag2−/−γc−/−, Rag2−/−Il17ra−/−, and Rag2−/− mice were provided by Genentech through B. Ryffel (CDTA Orléans). Rag2−/−, Rag2−/−γc−/−, Rag2−/−Il17ra−/−, and Rag2−/−Il15−/− mice were provided by the laboratory of J. Di Santo. Il22−/− mice were provided by R. Flavell (Yale University School of Medicine, New Haven, CT) via C. Leclerc (Institut Pasteur, Paris, France; Zenewicz et al., 2007). Il11ra−/− mice were provided by T. Putoczki (University of Melbourne, Parkville, Australia; Nandurkar et al., 1997). Stat3fl/fl Villin-cre mice are from Stat3fl/fl mice (Takeda et al., 1999) crossed with Villin-Cre (Cre-Ert2) mice (El Marjou et al., 2004). C57BL/6J were mixed with 300 µl of CaCO3 (50 mg/ml–1) before mid-log growth phase. For intragastric gavage, 200 µl of bacteria in PBS were mixed with 300 µl of CaCO3 (50 mg/ml–1) before injection in overnight starved animals. For BrdU assays, 150 µl of BrdU-labeling reagent solution (10373283; Life Technologies) were injected i.p. 16 h before sacrifice. To monitor bacterial burden, organs were removed from infected killed animals and homogenized. Before homogenization, intestines were rinsed twice with DMEM and incubated for 2 h at room temperature in DMEM supplemented with gentamicin (100 µg/ml; Sigma-Aldrich). Serial dilutions of cell suspensions in PBS were plated on brain–heart infusion agar plates. After 24 h of incubation at 37°C, CFUs were counted.

For in vivo IL-22 blocking experiments, mice were injected i.p. either with PBS (control) or with 0.3 mg purified anti–IL-22 (AM22.1 provided by J.C. Renaud and L. Dumoutier, Université Catholique de Louvain, Brussels, Belgium) at days −3 and 0 of infection. To deplete NK1.1+ cells, control and Lm-infected Rag2−/−Il15−/− mice were injected i.p. with 100 µg of anti-NK1.1 antibody (PK136; Biolegend) in 0.1 ml PBS i.d before Lm injection.

For in vivo cytokine treatment, mice were injected with 5 ng/g of mouse recombinant IL-22 (R&D; 582-ML-CF) or mouse recombiant IL-11 (R&D; 418-ML-CF) or mouse recombiant IL-17 (R&D; 421-ML-CF) at day −1, 0, 1, 2, and 3 after injection.

Hydroxytamoxifen treatment

Hydroxytamoxifen was dissolved in corn oil at 10 mg/ml. Stat3fl/fl Villin-cre mice were treated for five consecutive days with 1 mg/d, as previously described (Indra et al., 1999), 7 d before infection.

Bone marrow engraftment

Total bone marrow cells were removed from legs (femora and tibiae) of donor WT mice and intravenously injected into 5-wk-old lethally irradiated (1,250 rad) host mice (one donor for four recipient mice) to generate bone marrow chimera. After 10–12 wk of bone marrow reconstitution, chimeras were used to perform experiments.

Biotin penetration experiment

Biotin was used as a molecule to address the integrity of intestinal epithelium as described previously (Tsai et al., 2013). In brief, 2 mg/ml of EZ-link Sulfo-NHS-Biotin (Pierce) in PBS was slowly injected into the lumen of ileum loop via the open end adjacent to cecum immediately after removal of the entire ileum. After 3 min, the loop was opened and followed by PBS wash and 4% paraformaldehyde (PFA) fixation.

Ligated loops

8–12-wk-old mice were fasted for 16 h before surgery. 30 min after injection of the buprenorphine (0.1 mg/kg body weight), deep anesthesia was induced with a mix of ketamine (100

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Infection of Peyer’s patch locks villus invasion
mg/kg body weight; Imalgene 1000; Merial), and xylazine chlo-chorhydrate (10 mg/kg body weight; Rompun; Bayer). Skin was cleaned with Vetadine, a laparotomy was performed, the small intestine was exposed, and ileal loops of 1.5-cm long containing or not containing one FP were prepared. 100 µl of inoculum (10⁵ CFUs) was injected into each loop. Loop was placed back in the peritoneal cavity. PBS was injected subcutaneously to prevent dehydration and BrdU was injected intraperitoneally. Peritoneal membrane and skin were sutured, and animals were returned to their warmed cage. Water was allowed, but not food. 24 h pi, mice were killed. Intestinal loops were harvested, washed in PBS, and fixed in 4% PFA overnight. Tissue staining is described below.

**Quantitative real-time PCR (qRT-PCR)**

We performed RNA isolation and qRT-PCR as previously described (Dulauroy et al., 2012). In brief, we extracted total RNA from FACs-sorted cells using RNeasy Micro kit (Qiagen) and assessed the quality of total RNA using the 2100 Bioanalyzer system (Agilent Technologies). Transcription was performed using Superscript III reverse transcription (Invitrogen). All procedures were performed according to the manufacturers’ protocols. We performed qRT-PCR using RT² qPCR primer sets (SA Biosciences) and RT² SYBR-green master mix (SA Biosciences) on a PTC-200 thermocycler equipped with a Chromo4 detector (Bio-Rad Laboratories) and analyzed data using Opticon Monitor software (Bio-Rad Laboratories). We normalized Cd values to the mean Ct values obtained for the housekeeping genes Hsp90ab1, Hprt, and Gapdh.

**Cell isolation, flow cytometry analysis, and sorting**

Isolation of intestinal myeloid and stromal cells were performed as previously described (Satoh-Takayama et al., 2008; Szepourginski et al., 2015). Cells were blocked and stained with indicated antibodies: CD3 (clone 145-2C11; ebioscience); CD19 (clone ID3; BD); NKp46 (clone 29A1.4; ebioscience); IFN-γ (clone XMG1.2; BD); NKI.1 (clone PK136; Biolegend); and CD31 (clone 1D3; BD). Syrian hamster antibody to gp38 was a gift from A. Farr (University of Washington, Seattle, WA).

Cells were analyzed with an LSRFortessa flow cytometer (BD) or sorted with a FACSAria machine (BD). Flow cytometry analysis was done with the FlowJo software (Tree Star). Sorted stromal CD45+ gp38+ cells were grown in DMEM containing 10% fetal calf serum at 150,000 cells/well in 24-well plates coated with rat tail collagen (10 µg/cm²). Cells were treated 4 d after sorting with the mouse recombinant IL-17 (R&D; 421-ML) or mouse recombinant IL-23 (R&D; 1887-ML) for 24 h.

**Ex vivo organoid culture**

Intestinal organoids were obtained as previously described (Sato et al., 2009). Organoids were treated as follows: mouse recombinant IL-11 (R&D; 418-ML) at 50 ng/ml; mouse recombinant IL-22 (R&D; 582-ML) at 50 ng/ml; IFN-γ (ebiosciences 34-8311-85) at 50 ng/ml for 24 h followed by 16 h of BrdU staining (1/100 dilution of the BrdU labeling reagent; 10373283; Life Technologies). For STAT inhibition, fludarabine was added at 50 mM 24 h before cytokine treatment in WT organoid. For STAT3 depletion, Stat3fl/fl Villin-cre organoids were treated with hydroxytamoxifen 24 h before cytokine treatment.

**Immunofluorescence staining**

Tissues and organoids were labeled as previously described (Disson et al., 2009; Mahe et al., 2013). Intestine or colon was removed from cervical dislocated animals, flushed, and fixed in 4% PFA overnight. Tissues were then washed in PBS and (i) embedded in agarose for 150-µm-thick vibratome section or (ii) cryoprotected in sucrose 30%, embedded in optimal cutting temperature compound and nitrogen frozen for 8-µm-thin cryostat section. Organoids were fixed for 30 min in PFA 4% and washed in PBS. For BrdU staining, samples were incubated sequentially in HCl 1N for 10 min at 4°C, HCl 2N for 10 min at room temperature, HCl 2N for 20 min at 37°C, buffered with borate buffer (0.1 M) for 12 min at room temperature, and washed in PBS. Tissues were permeablized and blocked with 3% BSA in PBS containing 0.4% (thick sections) or 1% (thick sections and organoids) Triton X-100 for at least 1 h. Tissues were then probed with the appropriate primary and secondary antibodies at least 1 h at room temperature or overnight at 4°C. Sections were then mounted in fluorescent mounting medium (Fluoromount G; Beckman Coulter) and dried at 37°C for 30 min. For staining of accessible Ecad, whole mount intestine tissues were labeled with Ecad-2 in PBS BSA 3% as previously described (Nikitas et al., 2011). R2 anti-Lm rabbit polyclonal antibody was provided by P. Cossart (Institut Pasteur, Paris, France). Anti-Ki67 was obtained from Abcam. Anti BrdU clone MoBU-1 and anti-Ecad clone Ecad-2, Alexa Fluor 647 Phalloidin, Alexa Fluor 647, Alexa Fluor 488 WGA, Alexa Fluor 546 and 647 goat anti-rabbit, Alexa Fluor 488 and 647 goat anti-rat, and Hoechst 33342 are from Life Technologies.

For I65 staining, tissues were fixed in Carnoy solution for 24 h and then cut, deparaffinized, and treated as previously described (Canny et al., 2006). For Muc2 staining, tissues were fixed in Carnoy solution for 24 h and then cut and deparaffinized. Sections were boiled in 0.01 M citric acid buffer, pH 6, to retrieve the antigens and were permeablized, blocked, and stained with the anti-MUC2C3, a gift from G. Hansson, Institute of Biomedicine, Gothenburg, Sweden, at 4°C overnight (Johansson et al., 2008).

TUNEL assay was performed according to manufacturer instructions (click-iT Plus TUNEL Assay; Molecular Probes).

Sections were imaged with an inverted Axioobserver or LSM710 confocal Zeiss microscope (20× or 40× oil objectives). BrdU fold increase was calculated by dividing the distance between basal and upper BrdU+ cells by the length of the villus. When indicated, this ratio was normalized by noninfected animals (PBS) of the experiment to normalize for inter-experiments variations. For each experiment, two to three infected animals and one to two control noninfected animals were tested. Experiments were done in duplicate or triplicate. 5–10 villi were counted for each animal. For organoid, BrdU quantification was obtained by measuring the number of BrdU pixels from three-dimensional acquisition with Icy (de Chaumont et al., 2012), divided by the number of crypts (script in Fig. S4 I). All acquisitions and quantifications were done in a blinded manner.
Immunoblot
Tissue was lysed in lysis buffer containing radioimmunoprecipitation assay 1× (Sigma-Aldrich), anti-phosphatase, and anti-protease (Roche). Protein concentration from total protein extracts was quantified using the BCA Protein Assay kit (Thermo Fisher Scientific). 20 µg of protein for each sample was boiled for 5 min in NuPage buffer (Invitrogen) with reducing agent (Invitrogen) and radioimmunoprecipitation assay 1×. Samples were then subjected to SDS-PAGE and transferred onto polyvinylidene fluoride membranes (Millipore). Membranes were activated by absolute ethanol, followed by electro-transfer and saturation with PBS supplemented with 0.1% Tween 20 (Sigma-Aldrich) and 5% non-fat milk. Membranes were incubated with primary antibodies: rabbit anti-phospho-Stat3 (Tyr705; EP2147Y; ab76315; Abcam), mouse anti-phospho-Stat1 (Tyr701; M135; ab29045; Abcam), and mouse anti-β-Actin (clone AC-15; A1978; Sigma-Aldrich). Membranes were washed three times in PBS supplemented with 0.1% Tween 20 (Sigma-Aldrich), incubated with mouse (NA931; GE Healthcare) or rabbit (NA934; GE Healthcare) secondary antibody coupled to peroxidase, washed three times in PBS supplemented with 0.1% Tween 20 (Sigma-Aldrich), and revealed by chemiluminescence with ECL Western blotting detection system (GE Healthcare). Images were acquired and quantified with the PXi4 GeneSys software version 1.3.9.0.

ELISA
IFN-γ, IL-12, and IL-11 were measured with ELISA kits from Abcam from supernatant of intestinal tissue or in vitro cells in culture. Assays were performed according to the manufacturer’s instructions.

Histological analysis of DSS-treated mice
DSS was added to drinking water at a concentration of 2.5% 36–50 kDa DSS (MP Biomedicals) for seven consecutive days (Chassaing et al., 2014) after Lm infection. Histopathological examination of colon was done after fixation of the tissues in PFA 4% and embedding in paraffin. Tissue sections were cut and stained with hematoxylin and eosin. All slides were coded and evaluated in a blinded manner as previously described (Chassaing et al., 2014). In brief, each section was assigned four scores based on the degree of epithelial damage and inflammatory infiltration into the mucosa, submucosa, and muscularis/serosa (0–3). Each of the four scores was multiplied by 1 if the change was focal, 2 if it was patchy, and 3 if it was diffuse. The four individual scores per colon were added, resulting in a total scoring range of 0–36 per mouse.

Statistical analysis
Student’s t tests, nonparametric Mann-Whitney, or one-way ANOVA tests were routinely used for statistical analysis. Figure legends show the minimum number of mice used in the experiment and the statistical test used. Statistical significance is expressed as follows: *, P < 0.05; **, P < 0.01; ***, P < 0.001; ****, P < 0.0001.

Online supplemental material
Fig. S1 shows that Lm-induced epithelial cell proliferation is Lm strain independent, is not associated to barrier damage, is dose-dependent, and is initiated at the PP level, but not at the villus level. Fig. S2 illustrates STAT3’s role on WGA+ GC number decrease and shows that the phenotype is reversible and LLO dependent, but InLA independent. It shows that infected cells are PP CX3CR1+ myeloid cells, which express IL-23 upon infection. Fig. S3 defines the studied gp38+ CD45+ CD31− stromal cells population. Fig. S4 shows that IFN-γ is expressed by sNK cells in a CX-3CR1-dependent, but IL-23-independent, manner. It also shows that STAT-3 phosphorylation in ex vivo organoid is IL-11 or IL-22 dependent, while STAT1 phosphorylation is IFN-γ dependent. It illustrates the script used to quantify BrdU incorporation in organoids. Fig. S5 shows that Lm-induced epithelial cell proliferation is observed in colon, where the mucus layer is smaller. It also illustrates the model we propose.

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