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Pro-inflammatory role of monocyte-derived CX3CR1<sup>int</sup> macrophages in *Helicobacter hepaticus*-induced colitis

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Running title: Myeloid cells in *H. hepaticus* colitis
ABSTRACT

Cells of the monocyte-macrophage lineage play important roles in the pathogenesis of inflammatory bowel diseases, but they are also present in the normal healthy intestine, where they are critical for maintaining homeostasis. It has been unclear whether the pro-inflammatory roles of intestinal macrophages reflect altered behaviour of the existing resident cells, or if they involve recruitment of a distinct cell type. Here we have explored these ideas using the model of colitis induced by Helicobacter hepaticus (Hh) in the context of neutralisation or deletion of interleukin 10 (IL-10). Granulocytes and monocytes made up most of the inflammatory myeloid infiltrates found in the colon of Hh-infected colitic mice, rising to a peak within 2 weeks of Hh inoculation, but taking several months to resolve completely. The inflammatory response was dependent on the combined presence of Hh and absence of IL-10, and was accompanied by increased production of inflammatory mediators such as IL-1β, TNFα, IL-6 and IL-23p19 by infiltrating myeloid cells, mostly relatively immature cells of the macrophage lineage that express intermediate levels of CX3CR1. In contrast, the population of mature CX3CR1^hi macrophages did not expand as markedly during colitis, and these cells made little contribution to inflammatory mediator production. Taking into account their numerical dominance in the myeloid compartment, we conclude that newly recruited monocytes are the main source of pro-inflammatory mediators in colitis induced in the absence of IL-10 signalling, and that altered behaviour of mature macrophages is not a major component of this pathology.
INTRODUCTION

Inflammatory bowel diseases (IBD), comprising Crohn’s disease and ulcerative colitis, are growing health problems in the developed world. Although recent advances have helped elucidate some of the mechanisms underlying IBD, treatment remains unsatisfactory and individual regimes are seldom effective for all patients. Thus, a better understanding of the processes and cells involved in the pathogenesis of IBD could help develop new targets for therapy.

Macrophages (mø) have received considerable attention in recent years because of their potential roles in both steady-state and inflamed intestine (1-3). Resident møs are abundant in the healthy intestine, where they are involved in clearance of apoptotic cells and play a crucial role in maintaining homeostasis, ingesting and killing commensal bacteria that cross the epithelial barrier (reviewed in (2)). In contrast to other tissues, these processes do not provoke overt inflammation in the intestine, due to powerful control mechanisms that prevent local mø from producing pro-inflammatory mediators in response to stimuli such as TLR ligands (4, 5). However, findings from both IBD and experimental models of the disease have demonstrated that møs are also crucial components of the inflammatory infiltrate (4, 6-9), raising the possibility of exploiting them as therapeutic targets.

We and others have shown recently that intestinal mø originate from Ly6C\textsuperscript{hi} blood monocytes that continuously enter the colonic mucosa and differentiate locally through a series of intermediaries. Although only present in small numbers, cells with the phenotypic and morphological features of Ly6C\textsuperscript{hi} monocytes can be found in the steady state mucosa. However, major transcriptional differences exist between these and Ly6C\textsuperscript{hi} monocytes found in blood, suggesting that the differentiation process occurs immediately after monocytes enter the colonic mucosa (6, 10). Importantly, as monocytes mature they progressively acquire anti-
inflammatory properties, such as constitutive production of IL-10 and hyporesponsiveness to e.g. TLR ligands (6, 11, 12). However, this physiological process is disrupted during inflammation (6, 12, 13), and understanding of how this change in mØ behaviour occurs would be an important advance in our knowledge of how to prevent and treat IBD. Studies of the colitis induced by feeding dextran sodium sulphate (DSS) suggest that the tissue pathology is paralleled by the accumulation of highly pro-inflammatory monocytes, whereas the fully differentiated mØs that remain do not alter their behaviour and retain their anti-inflammatory characteristics (6, 12). However, it is not clear if this pattern extends to other forms of intestinal inflammation, particularly under conditions where there are intrinsic defects in the mechanisms that normally control resident mØ function.

The IL-10 – IL-10R axis is an important brake on mØ activation, particularly in the intestine, where deletion of either the cytokine or its receptor leads to spontaneous onset of inflammation in association with hyper-responsiveness of mØ (13-19). Furthermore, humans with non-functional mutations in IL10, IL10RA, or IL10RB develop severe enterocolitis within the first months of life (reviewed in (20)). Here we have examined the relationship between IL-10 and the differentiation of intestinal mØ in inflammation in more depth by exploring mØ behaviour during colitis induced by inoculation of mice with Helicobacter hepaticus (Hh) in the absence of IL-10 signalling (21, 22). We report that Hh-induced colitis is characterized by the accumulation of pro-inflammatory CD11b⁺ myeloid cells that are hyper-responsive to TLR stimulation. Most of these cells are monocytes and their immediate progeny that express intermediate levels of CX3CR1 and that have not differentiated fully into anti-inflammatory, resident-type CX3CR1ʰ⁻ mØ. We further show that the expansion of cells in the monocyte-mØ compartment is maintained for several months before returning to normal levels as the inflammation resolves.
MATERIALS AND METHODS

Infection and antibody treatment of mice

Female C57BL/6 (B6) Il10−/−, B6 WT, B6 CD45.1+ Cx3cr1<sup>gfp/gfp</sup> mice (obtained from Jackson Laboratory), and B6 CD45.1+ Cx3cr1<sup>+/gfp</sup> (generated by crossing Cx3cr1<sup>gfp/gfp</sup> mice with B6 CD45.1+ WT animals) were bred and maintained in an accredited SPF facility, and the experiments were conducted in accordance with the UK Animals (Scientific Procedures) Act 1986 under a Project License authorized by the UK Home Office and approved by the University of York Animal Welfare and Ethical Review Body. The mice tested negative for antibodies to specific murine viruses, were free of Helicobacter spp. as assessed by PCR, and were >6-weeks old when used.

Mice were allocated to treatment groups and inoculated i.g. with 1.5 x 10<sup>7</sup> H<sub>h</sub> NCI-Frederick isolate 1A (23), derived originally from the same mouse colony as isolate Hh-1 (24) (American Type Culture Collection strain 51449). IL-10-sufficient mice were also treated i.p. with 1 mg of anti-IL-10R (clone 1B1.3a) on days 0, 7, 14, 21, 28, 35, 42, 49, 56, 63, and 70 of H<sub>h</sub> infection. Age- and sex-matched uninfected animals were included as controls. One week after the last mAb injection, mice were sacrificed and intestines collected for analysis.

Lamina propria (LP) cell isolation

For the experiments shown in Figures 1 and 2, colon and caecum were pooled from individual mice and digested with Liberase Cl (0.42 mg/ml, Roche, Burgess Hill, UK) and DNase I (125 U/ml; Sigma-Aldrich, Gillingham, UK), followed by enrichment of LP cells on 40/80% Percoll gradients as described previously (25). For the experiments shown in Figures 3-5 and in Supplementary Figures, colonic LP cells were isolated as described previously without the use of Percoll gradients (4, 26).
Flow cytometric analysis and FACS

After blocking Fc receptors with anti-FcγRII/III mAb (2.4G2; BD Biosciences), LP cells were stained at 4°C with fluorochrome-conjugated antibodies (see Table S1) for 20-30 min in the dark before being washed in FACS buffer (2% FCS and 1mM EDTA in PBS) and then analyzed on a CyAn ADP (Beckman Coulter, High Wycombe, UK) or LSRII/FACSAria I (BD Bioscience) flow cytometer. Data were analyzed using FlowJo software (Tree Star Inc. OR, USA). Myeloid cell populations (as defined in figure legends) were sorted using a MoFlo Astrios or a FACSAria I sorter, with purities of >96%. For flow cytometric analyses on BD instruments, automatic compensation was performed in FACSDiva using UltraComp or OneComp beads together with fluorescence minus one controls.

Intracellular cytokine staining (ICS)

Large intestinal LP cells (2 x 10^6/ml) from uninfected and 2-week Hh-infected Il10−/− mice were cultured in medium alone or in the presence of 10 μg/ml Pam3CSK4 (Invivogen, Toulouse, France) for 4 hr at 37°C with 10 μg/ml brefeldin A during the last 3 hr. Thereafter, cells were stained for surface markers (CD45 and CD11b) and intracellular TNF-α as described previously (25).

Analysis of cytokine protein and mRNA expression

For cytokine protein analysis, FACS-purified CD45+CD11b+ cells were cultured in 96-well round-bottomed plates (5x10^4/ml; 0.2ml/well) at 37°C and 5% CO2 in medium alone or with 10 μg/ml ultrapure E. coli LPS, 10 μg/ml Pam3CSK4 (both from Invivogen, Toulouse, France) or 10 μg/ml soluble Hh antigen (SHELAg) prepared as described (21, 27). After 24 hr,
supernatants were collected and analyzed by ELISA for IL-12p40 (Mabtech, Nacka Strand, Sweden) and IL-6 (R&D Systems), and by FlowCytomix for TNF-α (Bender MedSystems, Vienna, Austria).

For cytokine RT-qPCR analysis in Fig. 2, FACS-purified CD45+CD11b+ cells were homogenized in TRIzol, and total RNA isolated by chloroform extraction and reverse transcribed using SuperScript II and random hexamers. cDNA was amplified using SYBR green reagents and an ABI Prism RT-PCR system (Applied Biosystems). Cytokine expression levels for each individual sample (run in duplicates) were normalized to HPRT using ΔCt calculations and the 7000 system SDS software (Applied Biosystems). For myeloid cell populations in Fig. 5, FACS-purified cells were lysed in RLT buffer (Qiagen) and homogenised using Qiashredders (Qiagen). RNA was then isolated and purified using an RNeasy micro kit (Qiagen) and reverse transcribed using a High Capacity RNA-to-cDNA kit (Life Technologies) and random hexamers. cDNA was amplified using SYBR green reagents and an ABI 7900HT Prism RT-PCR system (Applied Biosystems). Cytokine expression levels for each individual sample (run in triplicates) were normalized to TBP using ΔCt calculations. Specific primers pairs are detailed in Table S2.

Statistical analysis
Multiple group comparisons were performed by one-way ANOVA, while Student’s t-test and Mann Whitney test was used to compare two groups. Differences were considered statistically significant with P<0.05.
RESULTS

Expansion of myeloid cells in the large intestine of Hh-infected colitic Il10–/– mice.

To begin to characterize the innate immune response in the large intestine following Hh inoculation, WT and Il10–/– mice were inoculated i.g. with the bacterium, and the cellular composition of LP cells from pooled ceca and colons was examined 2 weeks later, the time point at which pathology peaks in Il10–/– hosts (25). As expected, Hh-infected Il10–/– mice displayed enhanced cellularity of the large intestinal LP compared with uninfected controls (Fig. 1A, left panel), with greatly expanded proportions and numbers of CD45+ hematopoietic cells (Fig. 1A, middle and right panels), consisting of B cells, CD4+ T cells, and CD11b+ myeloid cells (Fig. 1B). Only minor increases were observed in the proportion and number of LP CD45+ cells in Hh-infected WT mice (Fig. 1A), and the size of CD11b+ myeloid compartment was unaffected in these hosts (Fig. 1B, right panel). More detailed analysis of the early cellular kinetics following Hh inoculation of Il10–/– animals revealed that the numbers of total CD45+ cells and of CD11b+ myeloid cells were significantly increased in Hh-infected Il10–/– mice on day 5 pi, and expanded steadily until day 11 pi (Fig. 1C). Thus, Hh-driven typhlocolitis in Il10–/– mice is associated with massive infiltration by both lymphoid and myeloid cells.

Myeloid cells from Hh-infected Il10–/– mice display a pro-inflammatory phenotype

To understand how the expansion of the myeloid compartment might contribute to the colitis in Il10–/– hosts, we examined the cytokine secretion profile of CD11b+ cells from uninfected and Hh-infected Il10–/– mice. To this end, large intestinal LP cells were stimulated with the TLR ligand Pam3CSK4, and TNFα production examined by intracellular cytokine staining (ICS). TLR ligation of LP cells from Hh-infected Il10–/– animals resulted in ~3-fold increase in
the proportion of CD11b+ cells producing TNFα compared with cells cultured in medium alone (Fig. 2A, bottom panels), and this translated to a highly significant increase in the absolute number of TNFα+CD11b+ cells in the LP of infected Il10−/− mice compared with uninfected controls (Fig. 2B). In contrast, the proportion of TNFα+ CD11b+ cells from uninfected Il10−/− animals was unaffected by Pam3CSK4 stimulation and remained at levels similar to those seen in naïve WT colon (Fig. 2A, upper panels and Fig. 2B).

To extend these analyses, we next FACS-purified CD11b+ cells from the large intestinal LP of uninfected and 2-week Hh-infected Il10−/− animals and cultured them overnight with LPS, Pam3CSK4, or soluble Hh antigen (SHelAg), before assessing the levels of a wider range of cytokines using ELISA and FlowCytoMix. CD11b+ cells from Hh-infected Il10−/− mice secreted higher amounts of IL-12p40, IL-6, and TNFα after all forms of stimulation compared with CD11b+ cells from uninfected Il10−/− hosts (Fig. 2C). When analysed directly ex vivo, FACS-purified CD11b+ cells from Hh-infected Il10−/− mice also contained higher levels of Il12a and Il23a transcripts compared with CD11b+ cells from uninfected controls, whereas the levels of Il12b accumulation were similar in the two populations (Fig. 2D). Thus, the expanded myeloid cell compartment in the large intestine of Hh-infected colitic Il10−/− mice displays pro-inflammatory characteristics.

**Composition of the myeloid compartment in Hh-infected Il10−/− mice**

We next set out to explore what cell types accounted for the change within the CD11b+ compartment during Hh colitis. To do this, we exploited multiparameter flow cytometry and rigorous gating strategies we have developed recently to characterise the myeloid compartment of the intestinal LP, allowing precise identification of monocytes, mØ, eosinophils, neutrophils and dendritic cells (DC) (6; Fig. S1). We also omitted the Percoll...
gradient step during the purification to exclude the possibility of selective loss of individual cell types. These approaches confirmed marked changes in the composition of the myeloid cell compartment in the colon 2 weeks after Hh inoculation of Il10–/− mice, compared with uninfected Il10–/− mice or Hh-infected WT mice (Fig. 3A). Ly6G+ neutrophils and SSChi (MHCII−) eosinophils (Figs. 3A, B and C) accounted for a substantial part of the infiltration found in Hh-infected Il10–/− mice and no significant differences were seen in these granulocyte populations in Hh-infected WT mice, or in uninfected Il10–/− mice compared with uninfected WT mice (Figs. 3A, B, and C).

Hh-infected Il10–/− mice also showed marked expansion in the numbers of non-granulocytic myeloid cells (Figs. 3A, D and E), and we therefore explored the contribution of monocytes and mφ in more detail. To do this, we focused on cells expressing the pan-mφ markers F4/80 and/or CD64 (Fig. S1) and examined the monocyte-mφ differentiation continuum that we and others have defined in the colonic LP (6, 28, 29). This so-called ‘monocyte waterfall’ consists of newly arrived Ly6C+MHCII− monocytes, differentiating Ly6C−MHCII+ monocytes, and more mature Ly6C−MHCII+ cells that include tissue-resident mφ (Fig. 3D). Hh-infected Il10–/− mice showed marked increases in the proportions and absolute numbers of Ly6C+MHCII− and Ly6C−MHCII+ cells, which were increased by ~125-fold and 155-fold respectively compared with their numbers in uninfected Il10–/− mice (Figs. 3D and E). There was also a modest, but significant increase in the number of more mature Ly6C−MHCII+ cells in Hh-infected Il10–/− mice (Fig. 3E, right panel). Hh-infected WT mice, which do not develop colitis, showed some evidence of increased infiltration by Ly6C+MHCII−, Ly6C+MHCII+ and Ly6C− cells, while uninfected Il10–/− mice had slightly higher numbers of Ly6C+MHCII+ cells compared with their naïve WT counterparts; however, these differences were modest and did not attain statistical significance (Fig. 3E).
DCs were identified as CD11c^+MHCII^+CD64^- cells amongst LP leukocytes (30) and their numbers increased in *H. hepaticus*-infected compared to control *Il10^-/-* mice (Fig. S2A). The frequency of DCs among LP cells of *H. hepaticus*-infected *Il10^-/-* mice decreased compared to uninfected controls (data not shown), most likely reflecting the increase in other leukocytes. Finally, there were minor changes in the proportion of DC subsets identified on the basis of CD11b and CD103 expression, although none of these changes reached statistical significance (Fig. S2B).

**Altered monocyte-mø differentiation in *H. hepaticus*-infected colitic mice**

The accumulation of monocytes we found in *H. hepaticus* colitis is reminiscent of our own and other results from DSS and T-cell-mediated colitis, where there appeared to be an arrest in the local differentiation continuum that normally generates anti-inflammatory, resident mø (6, 12, 28). To examine whether a similar block was present during *H. hepaticus*-induced colitis, we used *Cx3cr1^+/gfp* reporter mice in which one allele of the *Cx3cr1* gene has been replaced with the gene encoding green fluorescent protein (GFP) (31). This allows fully differentiated resident CX3CR1^hi* mø to be distinguished from cells in the earlier CX3CR1^int* stages in the developmental continuum, some of which would have been included amongst the Ly6C^- MHCII^+ population we defined earlier. In this way, we could examine the relative roles of resident and recently recruited mø in *H. hepaticus*-induced inflammation, as well as explore how these cells behave during the resolution of pathology that occurs at later stages after bacterial inoculation (25). Because the *Cx3cr1^+/gfp* reporter mice are IL-10 sufficient, we had to induce colitis by *H. hepaticus* inoculation plus weekly administration of anti-IL-10R mAb (22). Consistent with our studies in *Il10^-/-* mice, this resulted in massive accumulation of total leukocytes and CD11b^+ myeloid cells in the colonic mucosa by day 14 pi (Figs. 4A and B). However, by day
41 pi, both compartments had contracted significantly, and by day 77 pi they were both reduced almost to baseline levels (Figs. 4A and B).

As in colitic II10−/− mice, the composition of the myeloid compartment was markedly altered in anti-IL-10R-treated Hh-infected Cx3cr1+/+ mice, with a large expansion of granulocytes and monocyte-derived cells at day 14 pi, followed by resolution at later times (Fig. 4C). Moreover, the non-granulocyte component of the myeloid infiltrate contained large numbers of Ly6C−MHCII− and Ly6C+MHCII+ cells at day 14 pi (Figs. 4C-E). The Ly6C−MHCII− population also expanded in number early in colitis, but most of this expansion was accounted for by CX3CR1int cells (Fig. 4F), a population that we have shown previously to be a further intermediary stage in the local differentiation of monocytes (6). In parallel, there was a substantial decrease in the proportion of fully-mature CX3CR1hi mø amongst the Ly6C− MHCII+ cells at this time (Fig. 4F), although their absolute numbers were increased and remained so throughout the experiment (Fig. 4G). By day 41 pi, the proportions and numbers of CX3CR1int cells amongst Ly6C−MHCII+ cells had contracted significantly, and by day 77 pi, CX3CR1hi cells had again come to dominate the Ly6C− mature mø compartment (Fig. 4F & G). At this time the numbers of Ly6C−MHCII− and Ly6C+MHCII+ cells had also returned to baseline levels (Fig. 4E).

The numbers of Ly6G+ neutrophils and CD11c+CD11b+F4/80− DC were greatly increased by day 14 pi in anti-IL-10R-treated Hh-infected Cx3cr1+/+ mice, before falling at later times in parallel with the reduction in monocytes and mø (Fig. S3A and B). Interestingly however, the numbers of colonic eosinophils remained high in these mice until the end of the experiment on day 77 pi (Fig. S3C). All these Hh-induced changes in myeloid cells were seen only when Hh-infected Cx3cr1+/+ mice were also co-administered with anti-IL-10R (Fig. 4 and data not shown). Anti-IL-10R treatment alone of uninfected Cx3cr1+/+ mice did not result in
leukocyte accumulation in the colonic mucosa (Fig. S4), findings that are in agreement with those we previously reported for uninfected WT mice (22).

Together our results demonstrate that the acute phase of Hh-induced colitis is associated with accumulation of CX3CR1\textsuperscript{int} monocytes and early-stage m\ø, consistent with the idea that there may be a block in the normal differentiation process.

**Colonic CX3CR1\textsuperscript{int} monocytes/m\øs are the major producers of pro-inflammatory cytokines during Hh colitis**

Finally we explored whether elicited CX3CR1\textsuperscript{int} cells were responsible for the production of pro-inflammatory mediators by myeloid cells during colitis, or if this reflected altered behaviour of the more mature CX3CR1\textsuperscript{hi} m\ø. RT-qPCR analysis showed marked upregulation of mRNA transcripts for \textit{Il1b}, Nos2 (iNOS), \textit{Il23p19} and \textit{Il12p35} in CX3CR1\textsuperscript{int} cells from anti-IL-10R- treated Hh-infected mice at the peak of inflammation on day 14 compared with CX3CR1\textsuperscript{int} cells from control mice (Fig. 5A-D). In contrast, colonic CX3CR1\textsuperscript{hi} m\ø from anti-IL-10R-treated Hh-infected mice showed only minor changes in mRNA levels of pro-inflammatory mediators (Fig. 5). Thus, taking into account their numerical dominance in the myeloid compartment, newly recruited CX3CR1\textsuperscript{int} monocytes/m\øs form the predominant pro-inflammatory population in this model of colitis.

**DISCUSSION**

The results presented here underline the importance of the myeloid compartment in the inflammatory colitis that occurs in mice infected with \textit{Helicobacter hepaticus} when IL-10-mediated signalling is absent. Using a variety of approaches to identify myeloid lineages,
including Cx3cr1^+/gfp mice, we show that an intense infiltrate of CD11b^+ cells appears early during infection and that this is made up of neutrophils, eosinophils, Ly6C^hiMHC^–/+ monocytes and CX3CR1^int^ mφ at the time of peak disease at 2 weeks after Hh inoculation. While this confirms other work (32), we also show here that this expansion of the monocyte-mφ compartment is sustained for up to 11 weeks after infection, by which time other parameters of inflammation, such as changes in myeloid cell subset composition, have returned to steady-state levels. Importantly, the inflammatory changes required both infection with Hh and neutralisation of IL-10R signalling, with very few changes being seen in mice with Hh infection alone or loss of IL-10 alone. Similarly, the greatly heightened production of pro-inflammatory mediators in response to activation in vitro by intestinal myeloid cells was fully dependent on these factors operating together. These results confirm previous conclusions that Hh plays a crucial role in provoking intestinal inflammation in the absence of IL-10-mediated immunoregulation and that pro-inflammatory responses to this organism are normally restrained by IL-10 (21, 22, 33).

In previous work, we have shown that resident mφ in normal intestinal mucosa are replenished continuously by Ly6C^hi monocytes that differentiate via a number of intermediary stages into mature, resident mφ (6, 11, 28). At early times after inoculation with Hh, this so-called monocyte-mφ “waterfall” expanded dramatically, and consisted mostly of Ly6C^hiMHCII^- monocytes and CX3CR1^int^ mφ, with less expansion of fully mature CX3CR1^hi^ mφ. Although these findings are consistent with previous work, e.g. in chemically-induced colitis (6), and suggest that the normal process of monocyte-mφ differentiation is arrested, it is important to note that the absolute numbers of mature mφ were actually increased in our mice with Hh-induced colitis and remained so for the duration of the 11-week experiment. Thus, although the majority of infiltrating monocytes may be short-lived in the colitic mucosa, a proportion of
these cells may still develop into mature CX3CR1<sup>hi</sup> mϕ, even in the face of inflammation. How the mϕ recruited under these conditions might contribute to the inflammation or its resolution and if they eventually acquire all the properties of the normal, resident population remain to be determined, as do the relative contributions of newly recruited and pre-existing cells to the “resident” mϕ compartment.

Although methods were not available to examine and track infiltrating monocytes directly following their arrival in the gut, we were able to explore whether the total mature mϕ population was altered in function during H<sup>h</sup> colitis. In steady-state intestine, resident mϕ exhibit an anti-inflammatory phenotype characterised by constitutive production of IL-10 and low levels of TNFα, together with an inability to respond to stimuli such as TLR ligands, but it has been unclear whether these properties can change during inflammation (34). In contrast to their CX3CR1<sup>hi</sup> counterparts, as we show here, cells within the CX3CR1<sup>int</sup> compartment are actively pro-inflammatory, expressing much higher levels of mRNA for pro-inflammatory mediators such as IL1β, iNOS, IL-23, and IL-12 during H<sup>h</sup> colitis than their control counterparts. In parallel, CX3CR1<sup>hi</sup> mϕ taken at the peak of inflammation showed no increased expression of IL-1β or IL-12 mRNA, and only a modest increase in iNOS and IL-23 mRNA, suggesting some, but limited ability to adopt a pro-inflammatory phenotype under these circumstances. In DSS colitis, we and others have shown that resident mϕ do not acquire pro-inflammatory characteristics (6, 12), but this has been reported in models of colitis where IL-10-mediated signalling is absent, with wider alterations in gene expression than we tested here (18). While this contrasting behaviour in the presence and absence of IL-10 is yet to be confirmed in other models of inflammation, the results are consistent with recent evidence that IL-10 drives permanent, epigenetic silencing of pro-inflammatory genes (35). As a result, the absence of IL-10R-mediated signalling profoundly alters the genetic landscape of
mature intestinal μφ, allowing them to respond to stimuli to which they are normally resistant (18). Whether IL-10 also controls other aspects of intestinal μφ differentiation is unclear, although it has been reported that their expression of the scavenger receptor CD163 is dependent on IL-10 (14). Arnold et al. showed reduced levels of CD206 on CX3CR1^hi^ μφ in Hh colitis (32), but it is not clear whether this reflects an intrinsic effect of IL-10, or is secondary to the effects of inflammation. That resident μφ retain a substantial proportion of their homeostatic properties in the absence of IL-10 is suggested by our finding that sustained expansion of this population is not accompanied by maintenance of Hh-dependent immunopathology.

The colitis induced by Hh is dependent on IL-23 (22) and upregulation of this cytokine was one of the major changes we and others observed in CX3CR1^int^CD11b^+^ myeloid cells in this model (32). It has often been difficult to identify precisely the cell responsible for producing IL-23 in infection or inflammation, in great part because μφ and DC share many phenotypic features such as CD11c, MHCII, CX3CR1 and CD11b. Therefore it is important to note that by using rigorous gating strategies based on CD64 as a μφ marker, both we and Arnold et al. (32) show that IL-23 was derived from the μφ lineage, which is consistent with work in other models (7, 36, 37). Notably, however, we detected increased IL-12p35 and IL-23p19 transcript levels in elicited CX3CR1^int^ monocyte/μφs, suggesting that these cells produce both IL-12 and IL-23 during Hh-induced intestinal inflammation. Thus we propose that the Th17 response and subsequent phenotype switching of Th17 cells into IFN-γ^+^IL-17A^+^ lymphocytes in Hh colitis (24) is driven by IL-23 derived from μφ, presumably activated directly by products of the organism. Nevertheless it should be noted that both CD103^+^CD11b^+^ and CD103^−^CD11b^+^ DC from the intestine have been shown to be capable of driving Th17 responses in vivo and in vitro respectively, via production of IL-23 and/or IL-6.
(30, 38-40). Thus there may be flexibility between the mΦ and DC lineages in their ability to stimulate T_{h}17 activity depending on the stimulus. Alternatively, these cell types may cooperate in driving such T-cell responses, perhaps with migratory DC being responsible for priming T_{h}17 cells in the draining lymph node, while tissue resident mΦ may sustain the survival of these cells once they migrate back to the mucosa (41).

Together our results show that elicited monocyte-derived CX3CR1^{int} macrophages form the predominant pro-inflammatory macrophage population in Hh-induced colitis. Given that these cells appear to derive from the same Ly6C^{hi} blood monocytes as their homeostatic counterparts, identifying the factors that govern monocyte fate in the colon during homeostasis versus inflammation could provide new therapeutic targets for the treatment for IBD.
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AUTHOR CONTRIBUTIONS

CCB, CJO, CAT, and MCK performed the experiments, analysed the data, and assisted with the manuscript. CCB, MCK, and AMM designed and coordinated the research, interpreted the data, and wrote the manuscript.
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FIGURE LEGENDS

Figure 1. Expansion of myeloid cells in the large intestine of *Hh*-infected colitic *Il10−/−* mice. WT and *Il10−/−* mice were inoculated with *Hh*, and LP cells were isolated from pooled caecum and colon 2 weeks (A-B) or 1, 3, 5, 8, and 11 days (C) later and examined by flow cytometry. Uninfected mice were included as controls. (A) Total numbers of LP cells (left panel), percentage (middle panel) and total numbers (right panel) of CD45+ hematopoietic cells from uninfected (white bars) and 2-week *Hh*-infected (black bars) WT and *Il10−/−* mice. (B) Numbers of B220+CD19+ B cells (left panel), CD3+CD4+ T cells (middle panel), and CD11b+ myeloid cells (right panel) from uninfected (white bars) and 2-week *Hh*-infected (black bars) WT and *Il10−/−* mice. Data in A-B are from one representative experiment out of at least three performed, and bars represent mean + SD of 3 mice per group. (C) Total numbers of LP cells (left panel), CD45+ hematopoietic cells (middle panel) and CD11b+ cells (right panel) from pooled caecum and colon of 1-, 3-, 5-, 8-, and 11-day *Hh*-infected *Il10−/−* mice. Data are pooled from two individual experiments, and are the mean + SD of 5-6 LP cell preparations (each pooled from 1-2 animals) from *Hh*-infected mice and 8 cell preparations (each pooled from 2-3 animals) from uninfected mice. One-way ANOVA followed by Tukey’s multiple comparison test. ***P<0.001, ****P<0.0001 when comparing *Hh*-infected and uninfected mice (A-B) or when compared to uninfected mice (C).

Figure 2. Large intestinal CD11b+ myeloid cells from *Hh*-infected *Il10−/−* mice secrete elevated amounts of pro-inflammatory cytokines following TLR stimulation. *Il10−/−* mice were inoculated with *Hh*, and LP cells were isolated from pooled caecum and colon 2 weeks later. Uninfected mice were included as controls. (A-B) Large intestinal LP cells were cultured
in medium alone or stimulated with 10 μg/ml Pam3CSK4 for 4 hours with 10 μg/ml brefeldin A during the last 3 hours, and then stained for CD45, CD11b, and TNFα or appropriate isotype control. Dot plots in A are gated on CD45⁺CD11b⁺ LP cells from uninfected (Uninf.; upper panels) and 2-week Hh-infected Il10⁻/⁻ mice (lower panels), and numbers (± SD) of TNFα⁺ CD11b⁺ cells per mouse in B were calculated from the percentages in A and the total number of cells isolated from each mouse. Data in A-B are representative of two independent experiments performed with 3 individual mice per group. One-way ANOVA followed by Tukey’s multiple comparison test. **P<0.01 and ****P<0.0001. (C) Large intestinal LP CD11b⁺ myeloid cells were FACS-purified from uninfected (white bars) and 2-week Hh-infected (black bars) Il10⁻/⁻ mice and stimulated in vitro with LPS (10 μg/ml), Pam3CSK4 (10 μg/ml), or SHelAg (10 μg/ml) or cultured in medium alone. After 24 hours, supernatants were collected and analyzed for the presence of IL-12p40, IL-6, and TNFα. Bars represent mean ± SD of quadruplicate (IL-12p40 and IL-6) or duplicate (TNFα) ELISA values (where each value represents a separate culture) combined from two independent experiments. (D) RT-qPCR analysis of IL-12p35 (Il12a), IL-12p40 (Il12b), and IL-23p19 (Il23a) transcripts relative to HPRT in sorted CD11b⁺ cells from uninfected (white bars) or 2-week Hh-infected (black bars) Il10⁻/⁻ mice. Bars represent means ± SD of three (for uninfected mice) and four (for Hh-infected mice) individual experiments, each consisting of cells pooled from 5-6 mice. Mann Whitney test. *P<0.05.

**Figure 3. Composition of the myeloid compartment in Hh-infected Il10⁻/⁻ mice.** WT and Il10⁻/⁻ mice were inoculated with Hh and colonic LP cells were isolated 2 weeks later and examined by flow cytometry. Uninfected mice were included as controls. (A) Relative frequencies amongst total colonic CD11b⁺ cells of Ly6G⁺ neutrophils, SSC⁺ MHCII⁻
eosinophils, CD64− CD11c+ MHCII+ CD11b+ DC and Ly6C/MHCII-defined cells of the CD64+ monocyte/macrophage compartment (Ly6ChiMHCII−, Ly6C*MHCII+, Ly6C−MHCII+) from uninfected or *Hh*-infected WT and *Il10−/−* mice. (B-C) Absolute numbers of Ly6G+ neutrophils (B) and SSChi MHCII− eosinophils (C) per colon of uninfected or *Hh*-infected WT and *Il10−/−* mice. (D) Representative expression of Ly6C and MHCII by CD11b+Ly6G−SSCloCD64+ cells from uninfected or *Hh*-infected WT and *Il10−/−* mice. Bar graph shows the frequency amongst CD11b+Ly6G−SSCloCD64+ cells of Ly6ChiMHCII−, Ly6C*MHCII+ and Ly6C−MHCII+ cells. (E) Absolute numbers of Ly6C*MHCII−, Ly6C*MHCII+ and Ly6C−MHCII+ cells per colon of uninfected or *Hh*-infected WT and *Il10−/−* mice. Data are from one of two independent experiments performed. Bars represent the mean + SD of 4 individual mice per group. One-way ANOVA followed by Tukey’s multiple comparison test. ****P<0.0001

**Figure 4.** Monocyte differentiation is dysregulated in anti-IL-10R-treated *Hh*-infected colitic *Cx3cr1*+gfp mice. *Cx3cr1*+gfp mice were inoculated with *Hh* and treated weekly with anti-IL-10R to induce colitis. The composition of the colonic myeloid compartment was then examined at 14, 41 and 77 days post infection and compared to uninfected mice or mice given *Hh* alone. (A-B) Absolute numbers of total CD45+ cells (A) and CD11b+ cells (B) per colon of *Hh*anti-IL-10R-treated mice 14, 41 and 77 days after infection, compared with control mice (uninfected and *Hh* alone; data pooled from day 14-77 for these groups). (C) Relative frequencies amongst total colonic CD11b+ cells of Ly6G+ neutrophils, SSChiMHCII− eosinophils, F4/80+CD11c+MHCII+CD11b+ DC and Ly6C/MHCII-defined cells of the F4/80+ monocyte/πφ compartment (Ly6C*MHCII−, Ly6C*MHCII+, Ly6C−MHCII+) of mice as in A. (D-E) Representative expression of Ly6C and MHCII by CD11b+Ly6G−SSCloF4/80+ cells (D) and absolute numbers of Ly6C*MHCII−, Ly6C*MHCII+ and Ly6C−MHCII+ cells per colon (E) of


mice as in A. (F) Representative expression of CX3CR1-GFP by Ly6C–MHCII+ cells from uninfected mice, mice inoculated with Hh alone, and mice receiving Hh plus anti-IL-10R when analysed at 14, 41 and 77 post inoculation. The histograms for the uninfected and Hh alone are taken from the day 14 experiment. (G) Absolute numbers of CX3CR1\textsuperscript{int} (top panel) and CX3CR1\textsuperscript{hi} (bottom panel) Ly6C\textsuperscript{−} MHCII\textsuperscript{+} cells per colon of mice as in A. Data are from one experiment and bars represent the mean ± SD of 4 individual mice per group. One-way ANOVA followed by Tukey’s multiple comparison test. *P<0.05, **P<0.01, ***P<0.001.

Figure 5. Colonic CX3CR1\textsuperscript{int} monocytes/macrophages produce pro-inflammatory mediators during Hh colitis. Cx3cr1\textsuperscript{+/gfp} mice were inoculated with Hh and treated weekly with anti-IL-10R to induce colitis. Two weeks later colonic LP CD64\textsuperscript{+}CX3CR1\textsuperscript{int} and CD64\textsuperscript{+}CX3CR1\textsuperscript{hi} cells (both pre-gated on CD45\textsuperscript{+}CD11b\textsuperscript{+}Ly6G\textsuperscript{−}SiglecF\textsuperscript{−}) were FACS-purified and processed for RT-qPCR analysis of pro-inflammatory mediators. Uninfected mice, mice given Hh alone, and mice given anti-IL-10R alone were included as controls. Transcript levels of IL-1\ β (\textit{Il1b}) (A), iNOS (\textit{Nos2}) (B), IL-23 (\textit{Il23a}) (C) and IL-12p35 (\textit{Il12a}) (D) relative to TATA-binding protein (TBP) in uninfected, Hh alone, anti-IL-10R alone, and Hh plus anti-IL-10R groups. Each symbol represents a pool of 3 mice (for uninfected, Hh alone, and anti-IL-10R alone) or individual mice (for Hh plus anti-IL-10R). Data are pooled from two individual experiments. Unpaired Student’s t test. *P<0.05, **P<0.01, ***P<0.001.
Figure 2

A

B

C

D

- Uninf. Il10–/–
- Hh-inf. Il10–/–

Medium
Pam3CSK4

TNFα
CD11b
Isotype

WT
Il10–/–

Uninf. + Hh
Uninf. + Hh

0
1
2
3

TNFα, CD11b cells (×10^6)

0
2
4
6
8
10

Medium
LPS
Pam3CSK4
SHelAg

0
100
200
300
400

IL-12p40 (ng/ml)

0
2
4
6

IL-6 (ng/ml)

0
2
4
6
8
10

mRNA expression relative to Hprt (×10^-3)

Il12a
Il12b
Il23a

<1
1
2
5

****
***
**

Medium
LPS
Pam3CSK4
SHelAg

- Medium
- LPS
- Pam3CSK4
- SHelAg

- TNFα (pg/ml)

- mRNA expression relative to Hprt (×10^-3)

Il12a
Il12b
Il23a

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****
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* * * *
Figure 3

A. 

Uninf.  

Hh  

WT  

Il10–/–

Ly6G+ neutrophils

SSC<hi> MHCII– 

eosinophils

Ly6C<hi> MHCII– 

Ly6C+ MHCII+ 

Ly6C<hi> MHCII– 

CD11b<hi> DC

B. 

Neutrophils per colon (×10^5)

Eosinophils per colon (×10^5)

C. 

Subset of CD64<hi> cells (%)

D. 

Gated: CD11b<hi> Ly6G– SSC<lo> CD64<hi>

Uninf.  

Hh  

WT  

Il10–/–

Ly6C<hi> MHCII– 

Ly6C+ MHCII+ 

E. 

Ly6C<hi> MHCII<hi> cells/colon (×10^5)

Ly6C<hi> MHCII<hi> cells/colon (×10^5)

Ly6C<hi> MHCII<hi> cells/colon (×10^5)
Figure 4

A

B

CD45+ cells per colon (x10^6)

CD11b+ cells per colon (x10^6)

C

D


E


F

G

Events (% of max)
Figure 5

A  Il1b

B  Nos2

C  Il23p19

D  Il12p35