Innate lymphoid cells sustain colon cancer through production of interleukin-22 in a mouse model

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Patients with inflammatory bowel disease (IBD) have an increased risk of colon cancer. However, the immune cells and cytokines that mediate the transition from intestinal inflammation to cancer are poorly understood. We show that bacteria-induced colon cancer is accompanied by differential accumulation of IL-17+IL-22+ colonic innate lymphoid cells (cILCs), which are phenotypically distinct from LTi and NK-22 cells, and that their depletion in mice with dysplastic inflammation blocks the development of invasive colon cancer. Analysis of the functional role of distinct Type 17 cytokines shows that although blockade of IL-17 inhibits some parameters of intestinal inflammation, reduction in dysplasia and colorectal cancer (CRC) requires neutralization of IL-22 indicating a unique role for IL-22 in the maintenance of cancer in this model. Mechanistic analyses showed that IL-22 selectively acts on epithelial cells to induce Stat3 phosphorylation and proliferation. Importantly, we could detect IL-22+CD3+ and IL-22+CD3− cells in human CRC. Our results describe a new activity of IL-22 in the colon as a nonredundant mediator of the inflammatory cascade required for perpetuation of CRC, highlighting the IL-22 axis as a novel therapeutic target in colon cancer.

Microbe-induced inflammatory pathways are important drivers of carcinogenesis in malignancies like Helicobacter pylori–induced gastric cancer and colitis-associated cancer (CAC; Trinchieri, 2012). In inflammatory bowel disease (IBD) patients, deranged inflammatory responses toward the intestinal microbiota increase the risk of colon cancer twofold (Eaden et al., 2001). However, the immune cells and cytokines that mediate the transition from colitis to colon cancer are poorly understood.

Cancer and chronic inflammation share common features and often use shared molecular pathways. The IL-23 axis is a key pathway for intestinal inflammation in many mouse models (Hue et al., 2006; Izcue et al., 2008), and its components IL-23R and Stat3 are not only associated with IBD susceptibility (Duerr et al., 2006; Barrett et al., 2008) but also involved in inflammation-associated cancer in the skin and colon (Langowski et al., 2006; Bollrath et al., 2009; Kortylewski et al., 2009; Grivennikov et al., 2012).

IL-23 was found to be increased in human colon adenocarcinoma and through induction of proinflammatory responses such as up-regulation of MMP9 and evasion of tumor immunosurveillance by exclusion of cytotoxic T cells promotes tumor growth and incidence (Langowski et al., 2006). One of the key functions of IL-23 is the ability to promote Th17-type responses characterized by the induction of the cytokines IL-17 and IL-22 (Liang et al., 2006). Although there is a good understanding of the role of other T cell subsets like Th1 cells and cytotoxic...
IL-22 is an important component of Type 17 responses and functions as a double-edged sword in the intestinal inflammatory response. On the one hand, it has been shown to promote colitis in some settings (Kamanaka et al., 2011) and IBD patients express increased levels of IL-22 in the colon (Brand et al., 2006; Geremia et al., 2011). The heterodimeric receptor for IL-22, IL-22R, and IL-10R2 is exclusively expressed on nonhematopoietic cells like intestinal epithelial cells (IECs) and signals through Stat3, a molecule involved in several models of CAC (Bollrath et al., 2009; Grivennikov et al., 2009). A procarcinogenic role for IL-22 via Stat3 activation was previously shown in several extracolonic cancers like hepatocellular carcinoma and non-small cell lung cancer (Zhang et al., 2008; Jiang et al., 2011). Surprisingly little is known about the role of IL-22 in CRC although polymorphisms in IL-22 were shown to be associated with a 1.46-fold increased risk for development of CRC (Thompson et al., 2010). Furthermore, IL-22 in conjunction with IFN-γ can induce iNOS production and procarcinogenic nitric oxygen species in human colon carcinoma cell lines (Ziesché et al., 2007).

In this paper, we have assessed the role of ILC and Type 17 cytokines in the development and maintenance of bacteria-driven CRC. Our results reveal a hitherto unappreciated role for ILC in the progression from colitis to colon cancer and, furthermore, show a specific role for IL-22 but not IL-6 or IL-17 in this process.

**RESULTS**

**Colonic innate lymphoid cells (cILCs) accumulate during cancer development**

To investigate whether ILCs are involved in bacteria-induced CAC, we used a new innate model in which genetically susceptible 129SvEv.RAG<sup>−/−</sup> mice are infected with *H. hepaticus* and treated with the carcinogen AOM. In this setting, mice progress from having chronic inflammation of the cecum and colon to invasive CRC within 3–5 mo (model based on Nagamine et al., 2008; Boulard et al., 2012; Fig. 1 A, left). CRCs are mainly located at the sites of highest inflammation like the cecum and distal colon (Boulard et al., 2012). Approximately 75% of mice develop between 1 and 4 (average per mouse, 1.2) colorectal carcinomas. In contrast to other CRC models, for example, DSS+AOM, the tumor number is low. However, the pathological features closely resemble human colitis-associated carcinoma and contained atypical glands breaching the muscularis mucosa accompanied by the induction of a desmoplastic stromal response (Fig. 1 A, right).

In *Hh+AOM*-treated mice, we found increased levels of IL-23, IL-17, and IL-22 mRNA in precarcinogenic aberrant crypt foci (ACF) compared with adjacent control tissue (Fig. 1 B). This was accompanied by increased levels of molecules involved in intestinal epithelial neoplasia, such as the EGFR ligand amphiregulin (*Areg*), a mediator of proliferation, and the matrix metalloproteinase *Mmp9* (Langowski et al., 2006), involved in invasion and angiogenesis, and *Hif1a* and *Vegfa*, but not *Pgk2* (Cow3) and *Mmp2* (Fig. 1 B). To characterize the populations of ILC present, we stained colonic lamina propria leukocytes (cLPs) for different ILC markers. The majority of...
CD45⁺, IL-7Rα⁺, lineage⁻ cells expressed Thy1 in uninfected as well as in Hh+AOM-treated mice (Fig. 2 A). The composition of ILC changed upon inflammation and cancer development as the percentage of Nkp46⁺ cells decreased from ~20 to 2% and the percentage of CD4⁻ Nkp46⁻ Sca1hi cILC increased from 70 to 95%, whereas CD4⁻ LTi cells remained stable (Fig. 2, A and B). Although the overall frequency of all three populations in the colonic lamina propria decreased upon Hh+AOM treatment as a result of the recruitment of other inflammatory cells, cILC numbers increased two- to three-fold, whereas Nkp46⁺ ILC decreased and LTi cells were unchanged (Fig. 2 C).

Further characterization of cILC surface markers showed that they expressed CD44, the IL-2 receptor α chain CD25, ICOS, and CCR6 (Fig. 2 D). They were negative for c-Kit and the IL-33R subunit ST2, markers which characterize Th2-type mucocytes and natural helper cells. ILCs, therefore, are phenotypically similar to ILCs previously identified by us during chronic Hh-induced inflammation. ILCs (IL-7Rα⁺ ROR-γ⁻) were localized in clusters (Fig. 2 E), here shown close to an E-cadherin⁺ invasive crypt, but also distributed within the lamina propria (not depicted).

ILCs promote bacteria-driven colon cancer

It was previously shown that depletion of ILC with a Thy1 antibody could block the development of colitis and also impair protective immunity to Citrobacter rodentium infection (Buonocore et al., 2010; Sonnenberg et al., 2011). To assess if ILCs were players in the transition from established chronic inflammation to colon cancer, we injected ILC-depleting anti-Thy1 into mice subjected 3 mo earlier to Hh infection and AOM treatment (Fig. 3 A). This therapeutic intervention led to a significant reduction of spleen mass, as well as cecal and colonic inflammation (Fig. 3 B), accompanied by reduced levels of IL-17, IL-22, and IFN-γ, cytokines previously shown to be produced by cILC (Buonocore et al., 2010; Fig. 3 C). We also detected reduced granulocyte recruitment to the colon (Fig. 3 D). To determine the impact of the treatment on carcinogenesis, we first analyzed methylene blue-stained colons for ACF. The area of ACF was strongly reduced upon treatment with anti-Thy1 (Fig. 3 E). Histopathological analysis of sequential H&E-stained sections of the colon revealed no invasive carcinomas but only dysplasia (9%) in the ILC depleted group, whereas in the isotype-treated group 62% of mice progressed to invasive adenocarcinoma (Fig. 3 G). Invasive adenocarcinomas were characterized by architecturally abnormal crypts penetrating the muscularis mucosa (Fig. 3 F, left). In contrast, in the colon of ILC-depleted mice, epithelial hyperplasia and inflammation was abrogated (Fig. 3 F, right).

IL-22 in tumor maintenance

Selective accumulation of cILC in the colon was accompanied by increased levels of IL-17, IL-22, and proinflammatory factors like TNF-α and IL-1β, but not IL-6, produced by cLPs (Fig. 4 A). We investigated which of those cytokines were produced by the ILC populations predominantly present in uninfected and Hh+AOM-treated mice. Whereas in uninfected mice the two dominant populations, cILC and Nkp46⁺ cells, expressed none of these cytokines in detectable amounts when cultivated ex vivo in the presence of monensin, cILC from Hh+AOM-treated mice expressed IL-17 and...
IL-22 (Fig. 4 B). We could also detect low levels of intracellular IL-6 and TNF in cILC.

CD4\(^+\) LTi and Nkp46\(^+\) NK-22, as well as CD11c\(^+\) cells, were previously shown to be major producers of IL-22 in response to IL-23 in different models of colitis, especially in a tissue-protective response to C. rodentium infection and DSS colitis (Cella et al., 2009; Pickert et al., 2009; Sonnenberg et al., 2011). To further investigate which cells were the
Next, we investigated if IL-22 or any of the other known cILC-associated cytokines could sustain bacteria-induced cancer. 

**Figure 3. Colitis to cancer transition is dependent on ILC.** (A) Treatment scheme. (B) Spleen mass (left), colitis (middle), and typhlitis (right) in isotype (n = 13) or anti-Thy1 (n = 11)–treated Hh+AOM mice. (C) Cytokines in supernatants of pooled cLP cells cultured overnight from one representative experiment of three independently performed experiments. (D) Flow cytometry of colonic CD11b+ Gr1+ granulocytes. (E) Aberrant crypt area on methylene blue–stained colons. (F) Representative photomicrographs (bars, 200 µm) of invasive CRC in an isotype treated mouse (left) and normal colon in an anti-Thy1–treated mouse (right). (G) Highest tumor grade per mouse. Data are shown as means. Results show a pool of two independent experiments (B, E, and G). **, P < 0.01; ***, P < 0.001, Mann-Whitney non-parametric t test (B and E), Fisher’s square test (G).
Figure 4. Nkp46−CD4−lin−Thy1hi ILCs are the major source of IL-22 in Hh+AOM CRC. (A) Secretion of cytokines into the supernatant of overnight cultured cLP cells from uninfected or Hh+AOM-treated mice. (B) Intracellular cytokine stain gated on different ILC populations from uninfected or Hh+AOM-treated mice after 3 h culture of cLPs in the presence of monensin. (C–E) Characterization of IL-22 producers in cLPs of Hh+AOM-treated mice stimulated overnight in the presence of IL-23 and for 3 h with PMA, Ionomycin, and monensin (n = 5). (C) Intracellular stain with isotype (left) or anti–IL-22 (right). (D) Anti–IL-22 stains mostly Thy1hi (left) Nkp46−CD4− (right) cells. (E) IL-17 and IL-22 expression in Thy1hi cells. Data are shown as means and SEM. Results are representative of at least two independent experiments. *, P < 0.05, Mann-Whitney nonparametric test.
Figure 5. IL-22 drives cancer perpetuation. (A) Treatment scheme. (B) Spleen mass (left), colitis (middle), and typhlitis (right). Means (n = 9–14). Results are a pool of two independent experiments. (C) Representative photomicrographs of colons (bars, 200 µm). (D) Granulocyte frequencies (left) and numbers (right) in cLPs as analyzed by flow cytometry for Gr-1<sup>hi</sup>, CD11b<sup>hi</sup>, and F4-80<sup>–</sup>, Ly6C<sup>int</sup> cells (n = 3–5). Data shown represent one out of two independent experiments. (E) CRC in antibody-treated Hh+aOM mice. Highest tumor grade per mouse. (F) Colonization with Hh. Results are a pool of two independent experiments (B, E, and F). Bars represent the mean value (B, D, and F). Significance refers to isotype control (B and E). *, P < 0.05; **, P < 0.01; ***, P < 0.001; ****, P < 0.0001, one-way ANOVA with Bonferroni’s post test (B) or Fisher’s square test (E).

Blockade of IL-17 or IL-22 significantly reduced systemic inflammation, as measured by spleen mass as well as cecal inflammation (Fig. 5 B). In both anti–IL-17 and anti–IL-22–treated groups there was a trend toward reductions in the number of infiltrating colonic granulocytes (Gr1<sup>hi</sup>, CD11b<sup>hi</sup>, Ly6C<sup>int</sup>, F4-80<sup>–</sup> cells) but this did not reach significance (Fig. 5 D).
Similarly, accumulation of Ly6C^hi, F4-80^hi inflammatory monocytes was not reduced (unpublished data). Strikingly, IL-17 and IL-22 appear to have distinct functional roles in inflammation and tumor development, as only IL-22 blockade ameliorated established colitis and reduced tumor burden significantly (Fig. 5, B, C, and E). The differential effects of anti–cytokine treatment cannot be explained by effects on Hh infection, as we could not detect differences in cecal Hh colonization levels upon antibody treatment (Fig. 5 F).

Remarkably, in both the anti–IL-22 and anti–Thy1 group the numbers of mice that showed high-grade dysplasia and invasive CRC were significantly lower (30%) than in the isotype-treated group (78%). As all mice sacrificed before therapeutic treatment with anti–IL-22 had developed CRC, these results suggest that IL-22 is a pivotal player not only in the transition from inflammation to CRC but also in the perpetuation of the cancerous state.

IL-22 regulates epithelial functions via Stat3
To investigate the mechanisms of how IL-22 sustains inflammation and cancer, we hypothesized that IL-22 acting on epithelial cells could lead to an inflammatory response and tissue repair program (Brand et al., 2006; Cella et al., 2009; Pickert et al., 2009) which becomes deregulated and promotes CRC. The heterodimeric receptor for IL-22 consisting of IL-22R and IL-10R2 is supposedly only expressed on nonhematopoietic cells (Witte et al., 2010). Indeed, detectable IL-22R expression was found only in colonic epithelial cells but not in the stromal cell fraction containing intestinal fibroblasts, myofibroblasts, and endothelial cells or cLPs (Fig. 6 A). Expression of IL-22R was also found in dysplastic epithelial cells in the colon of Hh^+AOM–treated mice (Fig. 6 B). We further dissected the therapeutic effect of blocking IL-22 on epithelial cells in our cancer model by treating mice in vivo for a short time period with anti–IL-22.

We found that phosphorylated Stat3 accumulates in the nuclei of colonic epithelial cells in isotype–treated Hh^+AOM mice (Fig. 7 A, top left). However, upon anti–IL-22 treatment for 1 wk the phosphorylation was almost undetectable in the epithelium (Fig. 7, A [top right] and B) but was still found at reduced levels in cells of the lamina propria (Fig. 7, A and B). Interestingly short-term treatment with anti–IL-6R or anti–IL-17 either failed or only minimally reduced Stat3 phosphorylation, indicating that epithelial Stat3 activation in this model is not regulated by these pathways (Fig. 7, A [bottom left and right] and B). We could detect an overall reduction of proliferating epithelial cells in colons of Hh^+AOM mice treated with anti–IL-22 using flow cytometry for Ki-67 (Fig. 7 C). Furthermore, Cyclin D1 levels were reduced in dysplastic areas of anti–IL-22–treated mice (Fig. 7 D, top), as well as Cyclin D1 message levels in colonic epithelial cell extracts (Fig. 7 D, bottom).

Blockade of IL-22 also affected other target genes of the IL-22–Stat3 axis. We found that RNA levels of the antimicrobial peptides RegIIg, RegIIb, and S100a8, the suppressor of cytotoxic signaling 3 (Socs3), and the neutrophil-attracting chemokine Cxcl1, which were up-regulated in Hh^+AOM–treated animals, were strongly decreased upon IL-22 blockade and, to a lower extent, by anti-Thy1 treatment but not upon IL-6R or IL-17 blockade (Fig. 7 E). The effect of anti–IL-22 treatment on Cxcl2 expression was less pronounced, indicating the possible involvement of other factors in granulocyte recruitment. The expression of the antiapoptotic molecules Bcl2 and Bcl-xl was also not inhibited by IL-22 blockade. Together, these results indicate a strong and almost exclusive role for IL-22 in the epithelial–Stat3 axis in this model, regulating epithelial proliferation, myeloid cell recruitment, and antimicrobial defense.

IL-22 in human CRC
To investigate the presence of IL-22 in human CRC, we obtained fresh and paraffin–embedded CRC tissue from patients who underwent surgery. Tumors were from different colonic locations and patients did not have IBD. As IBD patients undergo a rigorous screening procedure to avoid cancer development, IBD–associated CRC samples are rare and sporadic CRC samples provided the best available alternative for our study.
Figure 7. IL-22 regulates epithelial functions through Stat3 phosphorylation. (A–E) Hh+AOM-treated mice were injected with isotype control, anti–IL-22, anti–IL6R, anti–IL-17, or anti-Thy1 (B and E) for 1 wk. (A) Colonic phospho-Stat3-Y705 expression (bars, 25 µm). Arrows indicate pStat3+ cells in the cLP. (B) Immunoblot for phospho-Stat3 and total Stat3 in colonic epithelial and lamina propria cells. (C and D) Blocking of IL-22 reduces epithelial proliferation. (C) Ki-67 expression in freshly isolated EC from Hh+AOM mice (n = 4) and anti–IL-22 treated mice (n = 4). Gated on live, Epcam+, CD45− cells. (D) Cyclin D1 expression in dysplastic epithelial cells of Hh+AOM mice after 1 wk isotype or anti–IL-22 treatment (top). Cyclin D1 mRNA expression in isolated EC (bottom). (E) mRNA expression in EC. Data are shown as means and SEM of n = 2–5 per group. Results are representative of at least two independent experiments. *, P < 0.05; ***, P < 0.001; ****, P < 0.0001, one-way ANOVA with Bonferroni’s post test (E) or unpaired Student’s t test (C and D).
In tissue sections of human CRC specimens, cancerous areas showed an increased and aberrant expression of Ki-67 (Fig. 8 A, left). We could identify IL-22–positive cells close to dysplastic areas (Fig. 8 A, right) but also in normal tissue areas of the same patients. Immunofluorescence revealed that IL-22 was expressed by CD3+ and CD3− cells within tumors (Fig. 8 B). Although the majority of IL-22-positive cells within tumors are T cells, a lower percentage of lineage− cells produce IL-22.
We analyzed cytokine expression levels in 12 matched pairs of human CRC and adjacent tissue by quantitative real-time PCR (Fig. 8 E). In 58% of pairs, IL-22 mRNA expression was more than twofold higher in tumor versus normal tissue. Similarly, IL-17 was overexpressed in 75% of tumors and IFN-γ was more associated with CRC. In contrast, IL-10 was predominantly expressed in normal tissue. The presence of IL-22–producing cells within tumors could point toward a possible therapeutic application of IL-22 blockade in some CRC patients.

**DISCUSSION**

It is increasingly appreciated that inflammatory responses can be both protective and deleterious in cancer. In human CRC, Type 17 responses are associated with a poor prognosis, and in animal models of CRC, IL-23 and Th17 responses can provide an inflammatory environment that supports tumorigenesis (Wu et al., 2009; Liu et al., 2011; Tosolini et al., 2011). However, to date it has not been possible to separate pathways that are required for the inflammatory response from those that promote and sustain cancer. In this paper, using a bacteria-driven CAC model we identify Type 17 ILC and IL-22 in particular as pivotal for the development and maintenance of invasive CAC.

IL-22–producing ILCs are present in the intestine and contribute to the early innate response to enteric pathogens such as *C. rodentium* through induction of antimicrobial peptides and mucus production by IEC (Sugimoto et al., 2008; Zheng et al., 2008). IL-22–producing host-protective responses are contained within CD4+ LTi cells (Sonnenberg et al., 2011) and Nkp46+ NK-22 populations (Satoh-Takayama et al., 2008). A subpopulation of IL-12–responsive Nkp46+ LTi cells was also shown to be involved in antitumor immunity (Eisenring et al., 2010). In contrast with early protective responses, we observed accumulation of pathogenic double-negative Nkp46− CD4− ILC in chronic bacteria-induced inflammation (Buonocore et al., 2010). This population also differentially expands in bacteria-driven CAC at the expense of Nkp46+ and CD4+ ILC populations and produces IL-17 and IL-22. Thus, microbial insult in a genetically susceptible host leads to a shift from tissue-protective and potentially anti-tumorigenic ILC populations to chronic inflammatory pathogenic cILC. Although we describe ILC as the major source of IL-22 in this model, in lymphocyte-replete models of CRC, functionally similar IL-17– and IL-22–producing T cells may contribute to pathogenesis. Indeed, in human CRC we observed both T cell and non–T cell sources of IL-22. Further studies are required to determine the contribution of IL-22–producing T cells and innate cells to the natural history of human cancer development using normal and premalignant tissue samples.

IL-23 is a key driver of ILC-mediated colitis in the *Hh* model (Hue et al., 2006). IL-23 also drives colon cancer in an APC model, where it is induced in tumor-associated myeloid cells by microbial products, which are thought to specifically penetrate tumors through local down-regulation of mucins and tight junction proteins (Griennikov et al., 2012). However, in that model and in the enterotoxigenic bacterium *Bacteroides fragilis*–driven colonic Apc<sup>Min</sup> model, IL-17 is considered to be the key downstream mediator (Wu et al., 2009). Although the exact mechanism of IL-17 function in tumorigenesis is unclear, it was suggested to induce epithelial hyperproliferation and production of reactive oxygen and nitrogen species from granulocytes. However, it should be noted that both of these studies analyzed cancer development and not signals required for maintenance of the cancerous state. Indeed, we found that although anti–IL-17 blocks bacteria-driven innate colitis (Buonocore et al., 2010) and granulocyte recruitment, blockade of this pathway is not sufficient to reverse dysplasia and invasive adenocarcinoma. Instead, we observed a nonredundant role for IL-22 in this process.

Although IL-22 is host protective in acute intestinal infection and inflammation (Satoh-Takayama et al., 2008; Sugimoto et al., 2008), it is also implicated in the pathogenesis of chronic colitis through epithelial hyperproliferation (Kamanaka et al., 2011). We have also found a pathogenic role for IL-22 in development of *Hh*–induced colitis in immune-deficient and lymphocyte-replete mice in the presence of IL-10 deficiency (unpublished data). The reasons for the dual properties of IL-22 in terms of intestinal homeostasis are not known but may relate to the nature of microbial stimulation and that sustained production in response to chronic bacterial stimulation, such as that provided by the pathobiont *Hh* eventually becomes pathological. A recent study on IL-22 and its regulation by IL-22BP further supports this dual function of IL-22 and its role in inflammation and colorectal tumorigenesis but does not study tumor maintenance (Huber et al., 2012).

The presence of other cytokines also influences IL-22 functions. Thus, dual production of IL-17 and IL-22 by ILC led to immune pathology in a model of lung inflammation through recruitment of neutrophils (Sonnenberg et al., 2010). Granulocytes may contribute to carcinogenesis through production of reactive oxygen species; however, blockade of IL-22 or IL-17 did not significantly reduce granulocyte accumulation in the colon despite the former having anti-cancer properties, suggesting that other functions of IL-22, particularly its direct action on IEC, could be more relevant for tumor maintenance.

The main pathway of IL-22 action is via ligation of IL-22R on epithelial cells, leading to Stat3 activation. The role of Stat3 in colon cancer is well known but this has been attributed to activation by IL-6 and IL-11 (Ernst et al., 2008; Griennikov et al., 2009). Surprisingly, neutralization of IL-22, but not IL-6 or IL-17, almost completely abrogated Stat3 phosphorylation in the epithelium, as well as epithelial proliferation and cyclin D1 expression. This indicates that IL-22 is the main activator of the Stat3 pathway in this model and could facilitate cancer perpetuation via proliferative effects on the epithelium.
IL-22 may also promote cancer through direct effects on intestinal epithelial stem cells, previously shown to be key precursors for intestinal cancer (Barker et al., 2009). IL-22R expression was recently demonstrated in intestinal stem cells and IL-22 had protective effects on these cells in a graft-versus-host disease model (Hanash et al., 2012). A persistent IL-22 signal during chronic inflammation could therefore directly act on the “master regulator” of epithelial regeneration, leading to hyperproliferation and later to tumor maintenance. Such a mechanism could also explain the therapeutic effect of anti–IL-22 treatment on established tumors.

It is also possible that through the induction of antimicrobial peptides, IL-22 could induce a shift in the intestinal microbiota, favoring the outgrowth of “inflammatory allies” that support inflammation and cancer in conjunction with Hh. The composition of the gut microbiota influences cancer susceptibility as germ-free mice do not develop cancer (Uronis et al., 2009). Furthermore, certain species like the anaerobic Fusobacterium nucleatum were found to be associated with human CRC tissue (Castellarin et al., 2012; Kostic et al., 2012). Further studies are required to assess the role of microbiota changes in IL–22–driven cancers.

Although we found that IL-22 blockade very efficiently reduced tumor maintenance, a role for other cILC-produced factors in tumorigenesis is not excluded. Besides IL-22, cILCs can produce other molecules involved in epithelial repair. It was recently shown that lung ILC, phenotypically similar to nuocytes and natural helper cells, produced the EGFR ligand Areg in response to influenza virus-induced tissue damage in the lung (Monticelli et al., 2011). Administration of Areg in ILC-depleted mice was able to restore lung function upon influenza virus infection. In our cancer model, Areg is increased in aberrant crypt areas of Hh+AOM-treated mice. Furthermore, we found that cILCs express Areg message (unpublished data) and this could contribute to deregulated epithelial proliferation and tumor development. Further transcriptional profiling of cILCs could lead to the identification of other factors involved in tissue repair and cancer.

In summary, our results show that innate immune responses intended to protect the host from invading pathogens and to initiate tissue repair can become deregulated and chronic in genetically susceptible hosts, paving the way for cancer development. Together, our results identify a novel role for IL–22 in maintenance of colon carcinogenesis in chronic inflammation. The finding that established colon tumors in this model system are not cell autonomous but require the immune mediator IL–22 for their perpetuation raises the possibility that targeting the IL–22 axis may have therapeutic utility in human CRC.

**MATERIALS AND METHODS**

**Human study subjects.** CRC patients were recruited from the colorectal surgery department at the Churchill Hospital in Oxford. Colon specimens were collected from patients undergoing surgery for severe disease. Diagnosis of CRC was confirmed by a pathologist. Matched tumor and normal samples were selected from the same patients by a pathologist. Ethical approval was obtained from the Oxfordshire Research Ethics Committee (Reference numbers: 11/YH/0020 and 09/H0606/5) and informed written consent was given by all study patients.

**Mice.** 129SvEv.RAG<−/−> (129SvEvS6. RAG2<−/−>) mice were maintained in an accredited specific pathogen-free facility and experiments were conducted in accordance with the UK Scientific Procedures Act (1986). Mice were routinely screened for the absence of pathogens and *Helicobacter spp.*

**Hh CAC.** Hh NCI-Frederick isolate 1A (strain 51449) was grown as described previously (Maloy et al., 2003). Mice were fed 3% on consecutive days with Hh 1A (1.0 × 10^8 CFU) by oral gavage and injected with AOM (Sigma-Aldrich) i.p. once a week at a dose of 10 mg/kg for 5 wk, starting 6 wk after Hh infection (Nagamne et al., 2008). For experiments where animals were treated with blocking or depleting antibodies, treatment groups were mixed in cages to minimize cage effects and animals were treated at weeks 12–19 with the respective antibodies and analyzed at week 20. In some experiments, a preinjection group was analyzed at week 12.

**Hh colonization of mice** was quantified in fecal contents, which were collected upon sacrifice. DNA was isolated using DNA Stool kit (Qiagen) and qPCR with Hh-specific primers against the 16S rRNA gene was performed as previously described (Maloy et al., 2003).

For analysis of precarcinogenic aberrant crypts, crypts were fixed in 10% formalin, stained in 2% methylene blue, and analyzed using a dissection microscope and sliding caliper. For histological analysis, 200–300 sections of paraffin-embedded Swiss-rolled colons and cecums were prepared, and every 10th section was H&E-stained and blindly graded by a pathologist using criteria for grading dysplasia in human IBD as previously described (Boulard et al., 2012). Tumors were staged as per TNM classification of human CRC.

**In vivo antibody treatment.** Therapeutic administration of different antibodies was started at week 12 of Hh+AOM treatment in 129SvEv.RAG<−/−> mice, when inflammation was already fully developed, and was continued until the end of the experiment at week 20. To deplete Thy1<+> cells, 1 mg of depleting rat anti-Thy1 (YTS154.7.7.10) or isotype control (YKIX 337.217.1) was administered i.p. once a week. To specifically block different cytokines in vivo, 0.375 mg of IL-17 blocking Ab (UCB Celltech) or 1 mg of IFN-γ blocking Ab (clone AN18) was administered i.p. twice a week. A blocking mAb against IL–22 (Genentech, clone 8E11) or isotype control (GP120 10E7.1D2) was administered i.p. three times per week at 0.15 mg. For short-term treatment, anti–IL-12, anti–IL-17, anti-IL6R, or anti–Thy1 were administered to 5–mo Hh+AOM mice three times within 1 wk.

**Histological assessment of colitis and typhlitis.** Colonic and cecal inflammation in Hh+AOM mice was assessed upon sacrifice at 3 or 5 mo. H&E-stained samples, prepared as described above, were analyzed for inflammation in the proximal, mid, and distal part of the colon as well as the cecum. Scoring of inflammation was performed using previously published criteria (Izcue et al., 2008). In brief, each sample was graded semiquantitatively from 0 to 3 for the four following criteria: degree of epithelial hyperplasia and goblet cell depletion, leukocyte infiltration in the lamina propria, area of tissue affected, and presence of markers of severe inflammation (such as crypt abscesses, submucosal inflammation, and ulcers). Scores for each criterion were added to give a final quantification for intestinal inflammation ranging from 0 to 12. For colitis scores, a mean value of the three sections (proximal, mid, and distal colon) was calculated.

**Cell isolation and FACS staining.** For CLP cell isolation, mouse colon pieces were incubated (3×) in RPMI 1640 with 10% FCS and 5 mM EDTA at 37°C to remove epithelial cells. Tissue was then digested with 300 U/ml of type VIII collagenase for 45 min at 37°C. The isolated cells were layered on a 30/40/75% Percoll gradient (GE Healthcare), which was centrifuged for 20 min at 600 g, and the 40/75% interface, containing mostly leukocytes, was recovered. cILPs were stained with a fixable viability dye (eBioscience) and antibodies against CD45.2 and CD4 (RM4-5; both BD), a lineage mix (CD11b [M1/70], Ly6G [Gr1, RB6-8C5], and B220 [RA3-6B2]), Thy1 (53–2.1), Sca1 (D7), Nkp46 (29A1.4), and IL-7Rα (A7R34; all from eBioscience).
For intracellular cytokine staining, cLPS were either directly cultivated for 3 h in the presence of Golgi-stop-Monensin (BD) or stimulated overnight with 10 ng/ml of recombinant IL-23 (R&D Systems), and then for the last 3 h PMA and Ionomycin were added. Cells were stained intracellularly with antibodies against IL-22 (1H8PWSR) and IL-17 (17B7; both eBioscience) or isotype control antibodies using the Cytofix/Cytoperm kit (BD). cLPS were analyzed by flow cytometry using an LSRII flow cytometer (BD).

For the epitope of cellular colons were incubated for 5 min in 5 mM EDTA at 37°C. Cells in suspension were collected, immediately washed with cold PBS, and centrifuged. Flow cytometry was performed using a fixable viability dye and antibodies against EpCam (G8.8; BioLegend), CD45.2, and Ki-67 (both BD). Colonic stromal cells were collected from the 30%/40% Percoll interface during cLP isolation.

Human cLPS from tumors and normal tissue were isolated as previously described using enzymatic and mechanical disruption by GentleMACS (Miltenyi Biotec; Geremia et al., 2011). Cells were stained for the following lineage markers: CD11c (AbD Serotec), CD14 (BD), and CD16 (SG8; BioLegend); CD56 (HCD56), CD45 (OKT3), and CD4 (RPA-T4; both eBioscience); and intracellular using Cytofix/Cytoperm for IL-22 (R&D Systems) and IL-17A (BL168; BioLegend).

Cytokine detection. For detection of cytokines by the Flowcytomix Multiplex system (eBioscience), 10^6 isolated cLPS were cultivated overnight in 100 µl RPMI-1640 with 10% FCS at 37°C. Cytokines in the supernatant were detected using Simplex kits for mouse IL-17, IL-22, IFN-γ, IL-1β, IL-6, and TNF. Samples were analyzed on a FACSCalibur (BD).

Quantification of mRNA levels by real-time PCR. cLPS, stromal cells, or epithelial cells were lysed in RLT buffer (Qiagen) with β-mercaptoethanol. RNA isolation was performed using the RNeasy kit (Qiagen) including a DNase I digest step. Content and purity of RNA was controlled with a Nanodrop spectrophotometer (Thermo Fisher Scientific). cDNA synthesis was performed using the Superscript III reverse transcription kit (Life Technologies). Quantitative real-time PCR for the candidate genes was done using SYBR Green Mastermix (Applied Biosystems) and values were normalized to Hprt. Analysis was performed according to the ΔΔ Ct method. The following primer pairs were used for the quantification of gene expression in mouse tissue with SYBR green. Quantitect Primer Assay for Hprt (Qiagen): Mmp2, 5'-CACCA-CGCCAGACTATGACC-3' and 5'-TCTCTTGCTGACAGAAGG-3'; Mmp9, 5'-TTGCACACTGCAAGAAGTG-3' and 5'-CCACGACC-ATACAGATACGG-3'; Arg2, 5'-GGATCCAGCGGAGGATGAC-3' and 5'-GGCTTGCAAATGTCAACT-3'; Fgfa, 5'-TATCGCTGTGGACCTTCCAC-3' and 5'-ACAGGACGGCTGATTAGTG-3'; Hf1a, 5'-CGA-CACCATCATCCTCTGCG-3' and 5'-TGATTCAGTGACGATGACG-3'; and Ptp2 (Cox2), 5'-CAGTACAGGACTCTGCTACG-3' and 5'-TTGAGCATGGATGACG-3'. Primer pairs and probes for TaqMan Gene Expression Assays (Life Technologies) for mouse IL-22, IL-17, IL-23, Ccl2, Bcl2, Bcl3, Cox11, Cox12, Reg3g, Reg3h, S100A8, and S100A9: HPRT, 5'-GACCGGTCCCTGTACGCG-3'; 5'-CTACACCTGTGGATCTCCATCTCGC-3'; and VIC/TAMRA probe 5'-ACCACCGATCCCCAGC-GTCGGTCGT-3'. Primer pairs for the quantification of gene expression in mouse tissue with SYBR green. Quantitect Primer Assay for Hprt (Qiagen): Mmp2, 5'-CACCA-CGCCAGACTATGACC-3' and 5'-TCTCTTGCTGACAGAAGG-3'; Mmp9, 5'-TTGCACACTGCAAGAAGTG-3' and 5'-CCACGACC-ATACAGATACGG-3'; Arg2, 5'-GGATCCAGCGGAGGATGAC-3' and 5'-GGCTTGCAAATGTCAACT-3'; Fgfa, 5'-TATCGCTGTGGACCTTCCAC-3' and 5'-ACAGGACGGCTGATTAGTG-3'; Hf1a, 5'-CGA-CACCATCATCCTCTGCG-3' and 5'-TGATTCAGTGACGATGACG-3'; and Ptp2 (Cox2), 5'-CAGTACAGGACTCTGCTACG-3' and 5'-TTGAGCATGGATGACG-3'. Primer pairs and probes for TaqMan Gene Expression Assays (Life Technologies) for mouse IL-22, IL-17, IFN-γ, IL-10, and IκBα.

Immunohistochemistry, immunofluorescence, and immunoblot. For immunohistochemistry, subsequent 5-µm tissue paraffin sections were deparaffinized with Histo-Clear (National Diagnostics) and rehydrated. Anti-gen retrieval was performed using 10 mM citrate buffer, pH 6, and then intrinsic peroxidase activity was blocked (peroxidase block from Dako En Vision kit) as well as unspecific interactions of the secondary antibody using PBS with 2% BSA and 2% goat serum. Mouse tissue slides were stained with pS6-371 (Cell Signaling Technology), Cyclin D1 (NeoMarkers), IL-22R (R&D Systems), or rabbit isotype control (Zymed, Life Technologies). Human tissue slides were stained for Ki-67 (Abcam) or IL-22 (eBioscience). Staining was visualized using the appropriate secondary HRP-coupled antibodies and the EnVision System and slides were counterstained with hematoxylin.

For immunofluorescence in mouse, cryosections were fixed with 2% formaldehyde and for the nuclear ROR-γ stain additionally with Fix/Perm (eBioscience). Endogenous peroxidase activity was blocked with 3% H2O2 and 2% NaNO3, and unspecific binding with 10% donkey serum. Sections were either incubated with anti-IL-17R (A7R34; eBioscience) in 10% donkey serum in PBS or with ROR-γ in Perm buffer (eBioscience), followed by donkey anti-rat-HRP (Jackson ImmunoResearch Laboratories). Tyramide signal amplification was performed (Perkin Elmer). Then sections were stained with E-cadherin–FITC (612130; BD) and with DAPI.

For immunofluorescence of human tumor and normal sections, tissue was frozen in OCT, cut into 7 µm sections using a cryostat, and fixed with 1% paraformaldehyde for 5 min. Sections were stained for CD3 (Abcam) with Cy3-Perkin Elmer tyramide amplification, reblocked, stained for IL-22 (eBioscience) with FITC Perkin Elmer tyramide amplification, and mounted with Vectashield containing DAPI. Images were collected using a 710 microscope (Carl Zeiss).

Western blots were performed using Nupage gels (Life Technologies) and standard immunoblot technique. Inrivotrol (Life Technologies) membranes were treated with pStat3-Y705 (9145) and Stat3 (both from Cell Signaling Technology) and developed using ECL (Thermo Fisher Scientific).

Statistical analysis. Statistical analysis and graphical representations were done using Prism5 software (GraphPad Software). For comparison of Q-PCR results and histological analysis, the nonparametric Mann-Whitney test and unpaired Student's t test were used, and for multiple samples, one-way ANOVA with Bonferroni’s post-tests was applied. For analysis of dysplasia and CRC versus normal, the Fisher square test was used.

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S. Kirchberger and O. Boulard planned and performed experiments. D.J. Roxton graded and scored cancer pathology. R.L. Szabady, E. Thornton, O. Harrison, and F. Franchini were involved in experiments. F. Powrie directed the research. S. Kirchberger and F. Powrie wrote the manuscript.

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