Impact of the gut microbiome on mucosal inflammation

Warren Strober
National Institute of Allergy and Infectious Diseases (NIAID), NIH, Bethesda, MD, USA

In the past 10 years it has become increasingly apparent that the gut microbiome has profound effects on the immune system to which it is juxtaposed, the mucosal immune system. Here, I explore recent studies in which the effects of the microbiota expand or facilitate anti-inflammatory or regulatory immunological machinery or which favor development of proinflammatory immunological machinery in this system. I then focus on how these opposing processes play out in inflammatory bowel disease (IBD); a disease in which normal immune homeostasis is disturbed and inflammation takes hold.

Gut microbiota drives IBD
In recent years, studies probing the composition and function of the endogenous microbiota in the normal gastrointestinal (GI) tract have greatly expanded our appreciation for and understanding of how the microbiota shape mucosal immune responses, as well as more global GI tract activities. To some extent, these studies have been driven by the desire to understand better IBD.

Crohn’s disease (CD) and ulcerative colitis (UC) are thought to result from a breakdown in mucosal responsiveness to gut commensal organisms [1–3] (Figure 1). This concept is based first on the fact that in the many existing mouse models of colonic inflammation, either those induced by various external agents or those occurring spontaneously in genetically altered mice, one does not see inflammation in the absence of colonic microbiota [1]. In addition, there is now solid evidence that the most prominent genetic polymorphisms associated with IBD cause disease (or prevent disease) by affecting responsiveness of the mucosal immune system. For example, NOD2 deletion in mice or CD-associated NOD2 polymorphisms in humans lead to increased Toll-like receptor (TLR) responses because such responses are regulated by prolonged or repeated stimulation of NOD2 [4]. Similarly, deletion of the ATG16L1 gene in mice, another gene with CD-associated polymorphisms, results in hyperactivity of the Nod-like receptor family, pyrin domain-containing 3 (NLRP3) inflammasome and a polymorphism in the IL-23R gene in humans that is associated with decreased risk for developing IBD leads to decreased T cell interleukin (IL)-17 responses [5,6]. Finally, as discussed below, although colitogenic microbiota can be demonstrated in certain mouse models of colonic inflammation, there is as yet little evidence that such microbiota can cause persistent disease in the normal host.

In the review below I first summarize current information on how the microbiota of the GI tract either controls or prevents gut inflammation by the induction of regulatory T cells (Tregs), and then discuss data suggesting that changes in the microbiota can also result in the opposite, namely the induction or aggravation colitis. I then review the now extensive information on microbiota changes in IBD patients and its possible relation to the causation of this disease.

Mucosal homeostasis and Tregs induced by gut microbiota
In recent years, considerable evidence has accumulated supporting the notion that the gut microbiota induces mucosal Tregs that then play a vital role in maintaining gut homeostasis under normal conditions or in controlling inflammatory responses that would lead to disease. Evidence of this type was first obtained in studies in which the gut epithelial barrier was transiently perturbed by intrarectal administration of ethanol or Vibrio cholerae zonula occludens toxic hexapeptide; agents that cause increased epithelial permeability and increased exposure of lamina propria cells to luminal commensal microbiota [7]. Such treatment was shown to result in barely perceptible and transient inflammation accompanied by IL-10-dependent induction of forkhead box (Fox)p3-negative, cell-surface transforming growth factor (TGF)-β-positive CD4+ Tregs that could be shown to protect mice from induction of trinitrobenzene sulfonic acid (TNBS) colitis. Importantly, the development of these Tregs requires the presence of the gut microbiota and the presence of TLR2; it was thus established that innate TLR2 responses initiated by the microbiota are necessary for Treg development.

Further work confirming and expanding on these results utilized germ-free mice reconstituted with an ‘altered Schaedler’s flora’; a nonpathogenic mixture of commensal organisms [8]. Here, one observes induction of CD4+ Tregs that in this case are Foxp3-positive and IL-10-independent. The development of these Tregs is dependent on both innate and adaptive immune responses, because reconstituted myeloid differentiation primary response gene 88/TIR-containing adaptor molecule-1 [MyD88/Ticam-1 (TRIF)]
double-deficient mice whose T cells bear a T cell receptor (TCR) transgene specific for lymphocytic choriomeningitis virus (SMARTA mice) that do not respond to commensal organisms, exhibit greatly impaired Treg responses. In addition, in the absence of Treg development or IL-10, the mice manifest robust IL-17 and interferon (IFN)-γ responses in the colonic lamina propria, albeit in the absence of tangible inflammatory changes. Finally, in studies parallel to those involving mice treated with ethanol, recolonized mice pre-exposed to a low dose of dextran sulfate to cause injury to the intestinal barrier, developed heightened Treg responses and minimal cytokine responses, whereas recolonized SMARTA or MyD88/Ticam1-deficient mice failed to develop heightened Treg responses and exhibited vigorous cytokine responses; the latter associated with high mortality.

These studies, taken together, lead to the view that intestinal homeostasis, that is, the noninflamed state of the normal intestinal, is dependent on Tregs induced by commensal microbiota that gain entry into the lamina propria. Thus, although unfettered entry of commensals into the lamina propria due to gross epithelial damage [9] may cause severe inflammation, low level and transient entry has the opposite effect of girding the lamina propria from inflammatory influences. These concepts condition our understanding of IBDs because they make it likely that the inflammation of the GI tract that defines these diseases must initially overcome two anti-inflammatory barriers: the barrier imposed by Tregs induced by commensal microbiota and that created by Tregs that are generated by the inflammation itself (Figure 2).

**Induction of Tregs and prevention of colitis by specific commensal organisms**

So far, I have focused on the ability of the intestinal commensal microbiota as a whole to induce regulatory effects rather than on the ability of individual members of the commensal microbial community with a special propensity to induce such effects. However, there are, in fact, studies that show that certain bacteria are more effective than others in inducing Treg activity in the GI tract. Perhaps the most complete of these studies relate to the regulatory function of nonenterotoxigenic *Bacteroides fragilis*, a commensal organism existing within the colonic microbiota of both mice and humans. Early studies showed that mice mono-colonized with *B. fragilis* give rise to IL-10-producing Tregs that can protect recombination activating gene 2 (RAG2)-deficient mice from *Helicobacter-hepaticus*-induced colitis; furthermore, these cells are induced by a polysaccharide produced by *B. fragilis* known as polysaccharide A (PSA) [10]. Later studies expanded on these findings by showing that colonization of germ-free mice with PSA-producing *B. fragilis* but not PSA non-producing *B. fragilis* elicits induction of Foxp3+ T cells that produce IL-10 and exhibit a PSA-specific profile as they produce TGF-β2 and not TGF-β1, and do not manifest increased expression of cytotoxic T-lymphocyte antigen-4 (CTLA) or glucocorticoid-induced TNFR-related protein (GITR) when stimulated by PSA [11]. The regulatory function of the PSA-induced Foxp3+ T cells has been revealed in mice with TNBS colitis treated with PSA. PSA elicits Foxp3 Tregs that suppress effector cell responses and ameliorates colitis when administered before and after TNBS administration. On this basis, PSA has been proposed as a possible treatment of human IBD, but this possibility will have to
await studies showing that it can be given in a way that does not also induce T helper (Th1) effector cell responses, as shown in some studies [12]. In addition, PSA may be antigenic and lose effectiveness when repeatedly administered.

An interesting aspect of *B. fragilis* PSA regulatory activity is that it does not increase the number of Foxp3+ regulatory cells in TLR2-deficient mice and is thus TLR2 dependent [11]. This initially posed a problem in explaining its mechanism of action inasmuch as other bacteria also expressing TLR2 ligands did not have a similar capacity to induce regulatory cell activity. This problem was resolved in additional studies that showed that PSA, but not conventional TLR2 ligands, interacts directly with TLR2 on Foxp3+ T cells rather than on dendritic cells to induce production of IL-10, and to establish the profile of PSA-induced Foxp3+ cells mentioned above [12]. The possible biological significance of PSA stimulation of regulatory T cells via TLR2 was revealed in studies that showed that colonization of the gut with *B. fragilis* reduces IL-17 production via PSA induction of Tregs, and that *B. fragilis* expressing PSA (but not those not expressing PSA) are capable of colonizing colonic crypts in close proximity to the mucosal immune system, presumably because of their capacity to reduce local IL-17 production [12]. This leads to the concept that PSA is representative of a new class of TLR ligands that induce regulatory responses rather than inflammatory responses.

Although the ability of PSA to induce Treg activity is now well established by the studies discussed above, its overall function in mucosal homeostasis remains unclear. It may be that under normal homeostatic conditions it has a limited role in enabling *B. fragilis* to occupy a particular niche in the colonic crypts rather than a more global role in inducing suppressor T cells. This conclusion is suggested by compelling recent evidence that *Clostridium* rather than *Bacteroides* are the primary drivers of Foxp3+ T cell development in the colon.

Turning now to this *Clostridium*-related regulatory activity, it was shown initially that germ-free mice provided a specific-pathogen-free (SPF) flora that exhibited an increase in the numbers of Foxp3+ T cells in the colon, and that this change depends on the Gram-positive, spore-forming fraction of the SPF (i.e., a fraction that excludes *B. fragilis*) [13]. In subsequent studies designed to identify which bacterial species induce Tregs, the approach taken was to reconstitute germ-free mice with various cocktails of specific organisms and determine the effect of each cocktail on Foxp3+ cell levels in the colon. Although segmented filamentous bacteria (SFBs), and large collections of *Bacteroides* or *Lactobacillus*, had no capacity to increase the number of Foxp3+ cells, clostridial groups, particularly those belonging to clusters IV and XIVa, had a striking capacity to increase the number of Foxp3+ T cells. In addition, colonization with *Clostridium* enhanced colonic TGF-β concentration as well as other Treg-inducing molecules. These effects could also be observed in MyD88-, receptor interacting protein kinase-2 (RIP2)- and CARD9-deficient mice and were thus independent of TLR, NOD or Dectin receptor signaling. Finally, it was shown that *Clostridium* but not other bacteria (including *Bacteroides*) induced IL-10-producing cells in the colon. One caveat to these various findings is that they apply to colonic rather than small intestinal Foxp3+ T cells as reconstitution with clostridial organisms affected neither Foxp3+ T cells nor IL-10-producing T cells in the latter location.

Finally, in studies of the clinical significance of the above findings concerning clostridial organisms, it was shown that mice colonized with *Clostridium* at an early age (*Clostridium*-abundant mice) developed less severe dextran sodium sulfate (DSS) colitis than control mice, and exhibited a reduced tendency to mount Th2 cytokine and IgE antibody responses [13]. These findings mesh with the fact that *Clostridium* clusters IV and XIVa are proportionally reduced in patients with IBD, as discussed in greater detail below. Thus, overall, clostridial organisms emerge as a major inducer of Tregs, albeit by mechanisms that are as yet undefined (Figure 2).

**Colitogenic bacterial microbiota**

The mirror image (or rather, negative image) of bacterial microbiota that induce regulatory cells and thus protect organisms from inflammation are the so-called colitogenic organisms mentioned at the outset of this review, which cause *de novo* intestinal inflammation in normal mice. The first convincing evidence that such colitogenic organisms exist came from studies of recombination activating gene 2 (RAG2)-deficient, T-bet-deficient mice that develop spontaneous colitis and have been termed TRUC mice, reflecting both the immunodeficiency (TR) and the resemblance of the colitis to human IBD (UC) [14]. The origin of the colitis in these mice can be traced to the fact that T-bet is a transcriptional repressor of tumor necrosis factor (TNF-α) in dendritic cells and its absence leads to excessive TNF-α production by these cells, which then synergizes with IL-23 to drive IL-17 production by innate intestinal cells; in addition, the mice lack Tregs due to the RAG2 deficiency [14,15]. The TRUC model of colitis would not have been especially remarkable were it not for the fact that studies of this model showed for the first time that mice with experimental colitis could develop a colitogenic flora that transmitted colitis vertically to wild type (WT) pups nursed by TRUC mice and horizontally to co-housed WT mice that then exhibited some level of colitis for considerable lengths of time, even when separated from TRUC mice (Figure 3).

Subsequently, molecular characterization of fecal flora (based on 16s rRNA analyses) was conducted to define the organisms that were colitogenic in TRUC mice [16]. The most important findings to emerge from this analysis were evidence that two bacterial species, *Klebsiella pneumoniae* and *Proteus mirabilis*, were likely to be components of the colitogenic flora. This evidence consisted of the fact that these organisms are present in TRUC fecal flora as well as in the flora of WT mice fostered by TRUC mice, and antibiotic treatment of the TRUC mice that ameliorates the colitis reduces the numbers of these bacteria in the feces to levels below the limit of detection. However, these bacteria appear to require interaction with other bacteria in the normal flora to cause colitis because they do not cause colitis in germ-free TRUC mice free of colitis, but do cause colitis in wild type or RAG2/- mice with a normal
organisms. Bacteria activate cytokines. This is equivalent to findings in a more recent study of bacteria in TRUC mice in which it was found that a substrate of TRUC that does not develop colitis differs from mice that develop colitis by the fact that their microbiota lacked *Helicobacter typhlonius* and develop colitis if they are administered the latter organism [15]. Thus, it is possible that *H. typhlonius* is the most proximal colitogenic organism in TRUC mice.

In a final set of studies TRUC mice were administered either anti-TNF-α neutralizing antibody or Tregs to determine if restoration of immune function influences colitis in mice [15]. Both therapies ameliorate colitis and, although administration of anti-TNF-α diminishes *K. pneumoniae* or *P. mirabilis* levels, administration of Tregs does not. The latter fact suggests that these organisms and/or other organisms in the colitogenic microbiota require the presence of an appropriately abnormal mucosal immunological microenvironment to cause disease and are not intrinsically colitogenic, despite their ability to cause disease in WT recipients for some period of time. It is possible, for instance, that the colitogenic potential of these organisms is enhanced by interaction with proinflammatory cytokines.

Another set of studies revealing the presence of organisms that are associated with colitis involves studies of mice with deficiencies of NLRP6. The latter is a member of an NLR family of intracellular microbial recognition molecules that activate inflammasomes, that is, molecular complexes that result in the activation of caspase 1 and thus the proteolytic cleavage and then secretion of active IL-1β and IL-18. NLRP6 is a unique member of the NLR inflammasome family because its expression is localized to epithelial cells and thus it is particularly likely to be activated by microbial flora inhabiting the microbial–epithelial interface.

Initial studies of NLRP6 showed that mice deficient in this molecule manifest more severe induced colitis (DSS colitis) or even mild spontaneous colitis [17,18]. Later studies have provided evidence that NLRP3 deficiency leads to a change in the bacterial microbiota that mediates these susceptibilities [19]. This consists of the fact that WT mice co-housed for prolonged periods develop DSS colitis of equal severity to their NLRP6-deficient cohorts. Similarly, WT mice co-housed with mice deficient in apoptosis-associated speck-like protein containing a CARD (ASC), a key inflammasome component necessary for caspase I cleavage or co-housed with mice deficient in IL-18 also develop colitis, establishing that the NLRP6 deficiency is in fact an inflammasome-related defect. It is important to point out, however, that the increased susceptibility to DSS colitis in WT mice is a transient abnormality and WT mice separated from NLRP6-deficient mice eventually lose the increased susceptibility.

Subsequent studies of the microbiota in NLRP6–ASC- and IL-18-deficient mice revealed that these mice or indeed WT mice co-housed with these mice exhibited a microbiota enriched for the anaerobic taxa, *Prevotellaceae* and candidate bacterial phylum TM7 and that treatment of these mice with antibiotics abolished the transferability of DSS colitis susceptibility to co-housed WT mice [19]. Of interest, the *Prevotella* organisms in deficient mice were located adjacent to epithelial cells in the intestinal crypts, indicating that lack of epithelial inflammasome activity led to a defect in the ability of the deficient mice to control the proliferation of certain potentially pathogenic intestinal organisms normally occupying this micro-niche. The mechanism of this defect is, however, not yet known.

The ‘dysbiosis’ occurring in NLRP6-deficient mice may apply to mice with other inflammasome defects affecting hematopoietic cells rather than epithelial cells, and indeed to mice with other defects in innate immune responses; this possibility is in fact suggested by the finding that NLRP3-deficient and NOD2-deficient mice also exhibit increased severity of induced colitis and develop an altered microbiota [20,21]. In addition, there is evidence in the case of NOD2 deficiency that this altered microbiota causes increased susceptibility to induced colitis in WT mice. In any case, it is important to point out that the organisms occurring in mice with NLRP6 inflammasome defects (and possibly other defects) differ from those associated with TRUC mice discussed above in that they predispose normal (or deficient) mice to induced colitis rather than causing spontaneous colitis; thus, these organisms are (pathogen-free) flora. Thus, whether these organisms are true colitogenic bacteria that require help from other bacteria or are instead necessary helpers of true colitogenic bacteria awaits additional study. This question is relevant to findings in a more recent study of bacteria in TRUC mice in which it was found that a substrate of TRUC that does not develop colitis differs from mice that develop colitis by the fact that their microbiota lacked *Helicobacter typhlonius* and develop colitis if they are administered the latter organism [15]. Thus, it is possible that *H. typhlonius* is the most proximal colitogenic organism in TRUC mice.

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more properly called colitis-predisposing organisms rather than colitogenic organisms. The significance of this lies in the fact that these organisms, even more than the TRUC organisms, probably do not cause intestinal inflammation in themselves and thus cannot initiate colitis in humans in the absence of an intrinsic immune defect (Figure 3). A final point relating to the above findings in NLRP6-deficient mice concerns a recent comprehensive study of the intestinal microbiota of mice deficient in various TLRs or MyD88 that disclosed that, although the microbiota in the various mice differed markedly from each other, they generally did not differ greatly from WT littermates [22]. This suggests that the differences originated from an ‘extended husbandry in isolation’ rather from differences in innate immune responsivity. It can be argued that these findings call into question the idea that NLRP6 deficiency actually causes changes in the microbiota associated with the epithelium as implied above; however, this is not likely because the appearance of colitis-predisposing organisms in the intestine occurs in mice with ASC and IL-18 deficiency as well as NLRP6, indicating that a particular inflammasome-mediated mechanism in a variety of mouse colonies leads to the proliferation of a colitis-predisposing organism even in mice of differing origins.

Gastrointestinal microbiome in IBD

In the light of the above studies detailing either the anti-or proinflammatory effects of gut microbiota, it becomes of great interest to define the microbiota of patients with IBD and thus to determine if the organisms contained within this microbiota contributes to the occurrence of such disease.

A considerable number of surveys of the microbiota in patients with IBD and control individuals using metagenomic analyses of 16S rRNA in extracted gut specimens or fecal material have now been conducted. Although these molecular techniques are generally superior to culture-based techniques, because of inherent difficulties in culturing many members of the microbial community, they still have some limitations. Chief among these is that they rely on PCR-based techniques that may not detect bacterial species present in low abundance and they quantify copy number of 16S rRNA species rather than true bacterial numbers. Other difficulties in such surveys that adhere to culture-based surveys as well is that patient populations are both environmentally and genetically heterogeneous so that results obtained with studies (especially with small numbers of patients) may not reflect general abnormalities or may pertain to only a subclass of patients.

One landmark study conducted by Frank et al. in 2007 in large groups of patients provides a general framework for microbiota abnormalities observed in IBD patients, and serves as a basis for evaluation of changes found in studies of smaller patient groups [23]. These investigators analyzed surgically obtained gut-wall biopsy specimens and found that in colons of patients with IBD (both CD and UC) Bacteroidetes and Firmicutes (Lachnospiraceae family) were depleted whereas Proteobacteria (which contain Escherichia coli species) and the Bacillus group of Firmicutes were increased. In small intestines of patients, the Bacillus group of Firmicutes were decreased and Proteobacteria were increased relative to controls, whereas Bacteroidetes were unchanged. Upon principle component analysis, however, these overall differences were due to microbiota changes in only one-third of CD patients and one-quarter of UC patients, and the remainder exhibiting a normal microbiota pattern. In these subgroups, the decrease in Firmicutes (Lachnospiraceae family) and Bacteroidetes were particularly apparent in both colon and small intestine.

A notable feature of this analysis is that it did not disclose a particular bacterium present at levels expected of a pathological agent; in particular, the analysis detected few if any copies of Mycobacterium avium ssp. paratuberculosis rRNA; an agent that has been linked to CD in some studies [24]. In addition, the subset of IBD patients with abnormal intestinal microbiota was younger and more likely to have disease associated with abscess formation. These associations suggest that the abnormal microbiota are a feature of more severe disease and could thus be a factor that aggravates disease. This view fits with the fact that an abnormal microbiota was not found in the majority of patients and is thus unlikely to be a primary etiological factor. Finally, this analysis revealed that IBD small intestinal microbiota was characterized by reduced diversity in the Bacteroidetes and Firmicutes phyla, meaning that the number of distinctly different bacterial clones in these phyla was decreased.

Other groups of investigators, also using molecular methodology to assess bacterial populations associated with gut specimens, obtained results that were similar in some ways and different in others (reviewed in [25]). In summary, these studies of smaller and mostly CD patient populations provide data that add to those obtained in the study by Frank et al., in that they emphasize that decreases in the population of Firmicutes include Faecalibacterium prausnitzii decreases and increases in the population of Proteobacteria include E. coli increases; in addition, they suggest that these Firmicutes / Proteobacteria abnormalities may be limited to CD patients with small bowel disease [26–31]. The picture with respect to Bacteroidetes is somewhat less clear in that several studies found decreased number of these bacteria whereas in other studies, most notably those examining the ilea, increased concentrations of Bacteroidetes (B. fragilis) were found [29,32,33].

The molecular analysis of the gut microbiome in patients with IBD summarized above, as well as previous culture-based studies of the microbiota not discussed here, offers several insights into the role of commensal bacteria in the etiology of this disease. They provide strong evidence that a single pathogenic bacterial organism is not the root cause of IBD-related inflammation. This evidence begins with the fact that the molecular analyses did not reveal the presence of any known pathogen in sufficient numbers to cause inflammation [23], but also includes the fact that in several studies distortions in the microbiota were present in uninvolved tissue and/or were absent from involved tissue (such as colonic tissue), and that distortions tended to disappear when patients were administered agents that ameliorated immunological abnormalities, suggesting that they were secondary effects of underlying immunological defects [34].
Distortions in the gut microbiota are not found in all patients and tend to occur in only certain parts of the intestine and in patients with more severe disease [21,26]. In addition, such changes in microbiota are more evident in CD than in UC [31]. These facts argue against the idea that the distortions are a universal driver of the IBD disease process and for the idea that they are secondary and situational. However, these views need to be tempered by the realization that the methods so far used to analyze the gut microbiota may be grossly deficient in identifying the presence of more subtle bacterial abnormalities.

A consistent finding in CD is the selective loss of *Firmacutes* and *Bacteriodetes* organisms that conceivably could be members of the microbiota important in the induction of regulatory cells, as suggested by the murine studies of *B. fragilis* and *Clostridium*, as discussed above. The most striking of these findings relates to *Firmicutes prausnitzii*, which, as discussed above, is a clostridial organism that is consistently decreased in patients with CD; particularly those with small bowel inflammation [29]. In direct studies of the immunoregulatory function of this organism, Sokol *et al.* showed that human peripheral blood mononuclear cells (PBMCs) cultured in the presence of *F. prausnitzii* exhibited a higher ratio of IL-10 to IL-12 production than cells cultured in the presence of several other commensal organisms; moreover, supernatants from *F. prausnitzii* cultures suppressed IL-1β-induced IL-8 secretion and nuclear factor (NF)-κB reporter gene activity in CaCo-2 cells [35]. Perhaps more importantly, intragastric administration of *F. prausnitzii* or a supernatant obtained from its culture reduces the severity of TNBS colitis and lowers colonic IL-12 production; in addition, such treatment tends to correct the dysbiosis observed in mice with TNBS colitis. These studies support the idea that changes in *F. prausnitzii* in the microbiota are a significant factor in the severity of CD, but this requires further investigation because the cytokine changes observed in vivo, although statistically significant, are nevertheless rather marginal. Finally, with respect to *B. fragilis*, one can hardly claim that lack of this bacterium is playing an immunoregulatory role in IBD, because, as noted above, these organisms are greatly increased in the biofilm of IBD patients.

A final insight of the analyses of bacterial populations in IBD concerns the finding that *E. coli* is frequently increased in patients with CD [27]. This finding coincides with a series of studies showing that a subtype of *E. coli* with semipathogenic properties, called adherent–invasive *E. coli* (AIEC), is frequently found in patients with CD and play some role in the pathogenesis of this inflammation [36,37]. AIEC occurs in the inflamed ileum of ~22% of CD patients with chronic inflammation and a somewhat higher percentage in the terminal small bowel in postsurgical patients, but in only 6% of the ilea of control patients [36]. However, it is usually not found in affected colons of patients and is found in 22% of the ilea of patients without ileal inflammation; thus, AIEC is by no means a universal accompaniment of Crohn’s inflammation. As indicated by how they have been named, AIEC adheres to and colonizes small intestinal epithelial cells [37]. This important property of the organism is explained by the fact that AIEC expresses a unique type I pilus that has the capacity to bind to CEACAM6 on the surface of epithelial cells and thus facilitates AIEC attachment to such cells [38]. Not unexpectedly, such binding is dependent on the level of carcinoembryonic antigen-related cell adhesion molecule 6 (CEACAM6) expression. This is shown by the fact that mice bearing a CEACAM (CEABAC) transgene in epithelial cells and expressing large amount of epithelial CEACAM6 exhibit massive AIEC colonization [39]. This relation between CEACAM6 expression and colonization might explain the occurrence of AIEC colonization in CD inasmuch as the secretion of inflammatory cytokines such as TNF-α upregulates CEACAM6 expression.

Two important questions relating to the role of AIEC in CD concerns first its capacity to penetrate the epithelial barrier and thus invade the mucosa, and second, the inflammatory properties once such penetration is achieved.

Regarding their invasive properties, AIEC organisms are found in epithelial cells and in the lamina propria of inflamed CD patients, but it is not clear if such penetration is occurring through an already damaged (and possibly ulcerated) epithelium or through an intact epithelium. The fact that there are no reports showing that AIEC occurs within cells or tissues of uninflamed mucosa suggests that some damage to the epithelium is antecedent to such invasion. The latter conclusion is concordant with the fact that although the invasive (and proinflammatory) properties of AIEC are impressive, these organisms do not manifest the invasive properties of true pathogens such as *Shigella* and *Salmonella* because, if this were the case, they would cause inflammation throughout the bowel and not just in areas of Crohn’s inflammation. These caveats concerning the invasive capacity of AIEC are counterbalanced by the fact that mice expressing a transgene expressing human CEACAM6 mentioned above do exhibit AIEC translocation and passage into the lamina propria [39]. In addition, these mice express increased amounts of claudin in the plasma membrane (a pore-forming molecule) and manifest increased epithelial permeability [40]. Thus, when CEACAM6 is highly expressed, penetration of pre-existing normal epithelium may occur and this may be the case in inflamed IBD tissue exhibiting upregulated CEACAM6.

Related to the fact that AIEC can gain access to the mucosa via epithelial cells is recent evidence that AIEC, in common with several gut bacterial pathogens, express long polar fimbriae that allow them to bind to an M cell glycoprotein called GP2 and thus enter M cells [41]. Inasmuch as M cells are specialized epithelial cells that take up and translocate bacteria and soluble molecules into the Peyer’s patches, this property of AIEC is a means by which these bacteria can interact with the organized mucosal immune system.

AIEC taken up by both epithelial cells and macrophages tends to replicate and survive within these cells and induce the macrophages to produce inflammatory cytokines such as TNF-α and IL-6 [42,43]. Recently, it has been shown that such survival in macrophages is controlled, at least in part, by autophagic machinery, and cells with various autophagic defects due to impaired expression of genes associated with CD that adversely affect autophagy, such as *ATG16L1* and *NOD2*, support increased intracellular
AIEC survival and cytokine induction [44]. This raises the possibility that CD patients with these genetic abnormalities are more prone to AIEC colitogenic effects.

The adherent, invasive, and proinflammatory properties of AIEC discussed above have led to the suggestion that these organisms are primary causative factors in CD, at least in some patients. However, several observations mitigate against this possibility. First, as alluded to above, AIEC colonization is dependent on CEACAM6 expression and it is known that the latter can occur as a result of inflammatory cytokine secretion; thus, it is possible or even likely that AIEC colonization is secondary to an underlying inflammation rather than its cause. To counter this argument one would have to have evidence that AIEC can induce CEACAM6 expression in cells that cannot themselves produce cytokines that would have this effect. Second, as already mentioned, penetration of AIEC organisms in the Crohn’s mucosa lacks the uniformity that one might expect of a colitogenic organism, and the inflammation of the lamina propria in CEACAM6 transgenic mice, in which penetration is more, is a neutrophil-dominant inflammation that does not resemble the granulomatous inflammation of CD [39]. Third and finally, if AIEC does play a major role in CD, there should be evidence that those patients with AIEC colonization exhibit responsiveness to treatment with antibiotics to which AIEC are sensitive. To date, antibiotic treatment of CD patients with a variety of agents has had only marginal success and, perhaps more importantly, no patient subgroups have emerged (presumably those with AIEC colonization) who are particularly responsive to such treatment [45].

On the basis of the above, AIEC is most likely a secondary and aggravating factor in CD whose colitogenic potential is dependent on the presence of pre-existent and genetically determined proinflammatory immune abnormalities [46] (Figure 4). This view obtains additional support from a recent study of mice with TLR5 deficiency, that is, mice that do not bind flagellated organisms on the basolateral surface of epithelial cells due to absence of the TLR5 flagellin receptor [47]. It was found that such mice exhibit spontaneous colitis and that the latter is associated with the development of a microbiota that includes organisms that bind to the intestinal surface and contains increased numbers of proteobacteria and enterobacteria such as E. coli. That such organisms initiated the colitis was suggested by the fact that germ-free TLR5-deficient mice, but not WT mice, were subject to the development of chronic colitis when exposed to a reference strain of AIEC, and that such colitis persisted after this organism was no longer present. One explanation of these findings is that bacteria that occasionally penetrate the mucosal barrier interact with TLR5 and thus stimulate epithelial cell production of factors (such as chemokines causing neutrophil infiltration) that ordinarily limit proliferation of such penetrant bacteria. Thus, in the absence of such TLR5 function, local proliferation of bacteria in the lamina propria is more likely to occur and this causes general changes in epithelial function that cause colitis. This may include loss of barrier function that leads on the one hand to the excessive TLR responses that define the colitic state, and on the other, to changes in the microbiota such as the appearance of AIEC-like organisms that aggravate the proinflammatory process.

Overall then, these studies of TLR5-deficient mice again suggest that organisms with the properties of AIEC are colitogenic but only in the sense that they trigger inflammation in the presence of an underlying genetic defect (in this case involving epithelial cell function). Thus, we come back to the conclusion that whereas colitogenic organisms capable of initiating de novo inflammation in a normal host have been identified in experimental (murine) models of GI inflammation, such as TRUC mice, they have not yet been identified in human IBD.

Concluding remarks

The study of the relation of the gut microbiota to the development and maintenance of the mucosal immune system is a dynamic and rapidly expanding area of research. We know that the organisms comprising the gut microbiome act severally and singly to shape both the anti-inflammatory (i.e., regulatory) as well as the proinflammatory aspects of mucosal function. In addition, we know that the gut microbiome can be altered by disease and that such alterations may be primary or secondary factors in disease pathogenesis. Finally, we know that genetic factors underlying the development of disease in many or all instances affect the latter because of defective responses to organisms in the gut microbiome. Thus, one of the major research challenges in the area of gut inflammation is to define the precise mechanisms that underlie these defective responses.

References
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