Iron Sequestration in Microbiota Biofilms As A Novel Strategy for Treating Inflammatory Bowel Disease

Jean-Paul Motta, PhD, Thibault Allain, PhD, Luke E Green-Harrison, BSc, Ryan A Groves, MSc, Troy Feener, MSc, Hena Ramay, PhD, Paul L Beck, MD/PhD, Ian A Lewis, PhD, John L Wallace, PhD, Andre G Buret, PhD
Iron Sequestration in Microbiota Biofilms As A Novel Strategy for Treating Inflammatory Bowel Disease

Jean-Paul Motta, PhD,*,† Thibault Allain, PhD,*, Luke E. Green-Harrison, BSc,*, Ryan A. Groves, MSc,*, Troy Feener, MSc,† Hena Ramay, PhD,‡ Paul L. Beck, MD/PhD,§ Ian A. Lewis, PhD,*, John L. Wallace, PhD,*,‡ and Andre G. Buret, PhD,*,#

Significant alterations of intestinal microbiota and anemia are hallmarks of inflammatory bowel disease (IBD). It is widely accepted that iron is a key nutrient for pathogenic bacteria, but little is known about its impact on microbiota associated with IBD. We used a model device to grow human mucosa-associated microbiota in its physiological anaerobic biofilm phenotype. Compared to microbiota from healthy donors, microbiota from IBD patients generate biofilms ex vivo that were larger in size and cell numbers, contained higher intracellular iron concentrations, and exhibited heightened virulence in a model of human intestinal epithelia in vitro and in the nematode Caenorhabditis elegans. We also describe an unexpected iron-scavenging property for an experimental hydrogen sulfide-releasing derivative of mesalamine. The findings demonstrate that this new drug reduces the virulence of IBD microbiota biofilms through a direct reduction of microbial iron intake and without affecting bacteria survival or species composition within the microbiota. Metabolomic analyses indicate that this drug reduces the intake of purine nucleosides (guanosine), increases the secretion of metabolite markers of purine catabolism (urate and hypoxanthine), and reduces the secretion of uracil (a pyrimidine nucleobase) in complex multispecies human biofilms. These findings demonstrate a new pathogenic mechanism for dysbiotic microbiota in IBD and characterize a novel mode of action for a class of mesalamine derivatives. Together, these observations pave the way towards a new therapeutic strategy for treatment of patients with IBD.

Key Words: microbiota biofilms, colitis, iron, hydrogen sulfide, mesalamine, inflammatory bowel disease

INTRODUCTION

Therapeutic limitations, high costs, and adverse effects of currently available drugs for treatment of inflammatory bowel disease (IBD) underscore the need for medications that are more effective and safe. Although anemia is one of the most common extraintestinal complications of IBD, dietary iron supplementation leads to disease exacerbation and a higher risk of infection, possibly through alterations of commensal microbiota and increased abundance of Enterobacteriaceae. Beyond the taxonomic microbiota abnormalities associated with IBD (decreased diversity, increased representation of Proteobacteria), the presence of pathobionts (for example adherent-invasive strains of Escherichia coli) or abnormal host responses toward commensals are suspected. A better understanding of functional alterations of gut microbiota during IBD is sorely needed and may pave the way toward more precise therapies.

We hypothesized that microbiota with increased ability to access iron, and subsequently increased virulence, contribute significantly to intestinal inflammation. We thus aimed to evaluate if novel derivatives of a widely used drug for treating inflammation in patients with IBD (ie, hydrogen-sulfide releasing mesalamine) could have a direct and beneficial effect toward dysbiotic microbiota associated with IBD in a biologically relevant model. Microbiota indeed naturally exist as biofilms in the large intestine, both in health and during colitis. Biofilms are

Supported by: Jean-Paul Motta was funded by postdoctoral fellowships from the Alberta-Innovate Health Services (AIHS), University of Calgary Eye's High Fellowship, Izaak Walton Killam Fellowship, and AgreenSkills Fellowship from the EU. This research was funded by grants to Andre G. Buret and John L. Wallace from the Natural Sciences and Engineering Research Council of Canada (NSERC Discovery and CREATE grants to Andre G. Buret), from Crohn's and Colitis Canada (Andre G. Buret and John L. Wallace), and the Canadian Institutes of Health Research (John L. Wallace and Andre G. Buret). Metabolomics data were collected by Ryan A. Groves at the Calgary Metabolomics Research Facility, supported by the International Microbiome Centre and the Canada Foundation for Innovation (CFI-JELF 34986); Ian A. Lewis is supported by a Alberta Innovates Translational Health Chair.

Address correspondence to: Andre G. Buret, PhD, Department of Biological Sciences, Faculty of Science, University of Calgary, 2500 University Drive NW, Calgary, Alberta, T2N 4N1, Canada. E-mail: aburet@ucalgary.ca

© 2018 Crohn’s & Colitis Foundation. Published by Oxford University Press. All rights reserved. For permissions, please e-mail: journals.permissions@oup.com

doi: 10.1093/ibd/izy116
Published online 17 May 2018
defined as communities of bacteria encased in a complex extracellular matrix composed of polysaccharides, proteins, nucleic acids, and glycolipids.10 We used a well-established model to study multispecies microbiota biofilms obtained from human colonic biopsy tissues ex vivo under anaerobic and static conditions.6, 11–13 This model reproduces some of the complexity of microbial-microbial interactions in nature and thus served as an attractive platform to perform preclinical studies of a drug designed to correct dysbiotic behavior associated with IBD.

MATERIALS AND METHODS

Patients
Descending colon biopsies of human donors who had not taken antibiotics in the previous 16 weeks were provided by the Inflammation Tissue Bank of the University of Calgary. Ethics approvals were delivered by the Research Ethics Board and the Calgary Health Region (#REB14-2430 REN1). Clinical characteristics of the donors are presented in Table 1. Biopsies were transported in BBL Port-a-cul (anaerobic transport, BD Bioscience, Mississauga, Canada) from the hospital to the lab and mucosa-associated bacteria were cultured from these biopsies immediately under an anaerobic hood.

Drugs
ATB-428 and ATB-429 are compounds consisting of mesalamine covalently linked to an H2S-donor: 4-hydroxythiobenzamide (TBZ) in the case of ATB-428, and (5-(4-hydroxyphenyl)-3H-1,2-dithiole-3-thione, ADT-OH) in the case of ATB-429. These drugs were provided by Antibe Holdings Inc. (Calgary, Canada).

Growth of Biofilms
All steps described herein were performed in anaerobic conditions (Bactron II, Sheldon Manufacturing, Oregon, USA) as previously described,11–13. Colon biopsies were homogenized in a microtube pestle and mucosa-associated microbiota was cultured in tryptic soy broth supplemented with yeast extract (5 g/L, BD Bioscience), L-cysteine (5%, Sigma-Aldrich, Oakville, ON, Canada), hemin (100 mg/L), and menadione (2 mg/L, Sigma-Aldrich). The Calgary Biofilm Device (Innovotech, Edmonton, Canada) features a lid with 96 polystyrene extensions on which biofilms grow from the shear stress generated by agitation.11, 47 Biofilms were generated from identical optical density of bacteria (OD600nm = 0.1) and matured for 3 days under agitation (125 g). Mature biofilms were transferred onto a plate with wells containing fresh media or containing working concentrations of drugs (in broth media as vehicle), and after 24 hours biofilm-dispersed bacteria were collected in the bottom plate. Viability was assessed by colony-forming unit counting in agar plates.

Human Intestinal Epithelial Cells
Human intestinal epithelial cells (Caco-2, ATCC, Manassas, USA) were grown in Dulbecco’s modified Eagle medium (Gibco, Invitrogen) on flat-bottom 6- or 12-well plates for 7 days (Costar, Thermo Fischer) or on polycarbonate 12- or 24-transwells plates for 21 days (Costar). Cells were co-cultured with bacteria (8 × 105 CFU/ml) in Hanks Balanced Salt solution (HBSS, Gibco), in 5% CO2 incubator at 37°C. During co-culture, bacterial growth rate was equivalent in all groups. Alternatively, 1 ml of biofilm spent media (OD600nm = 1) was filter-sterilized through 0.22 µm filters, and cells were stimulated with 10% dilution of spent media in HBSS. Epithelial necrosis was quantified by lactate dehydrogenase activity (LDH) assay (Roche, Mississauga, Canada) and absorbance was read (SpectraMax, Molecular Devices Corporation, Menlo Park, USA). Bacteria adhesion (90 minutes) and invasion (3 hours) protocols were based on standardized antibiotic assay and agar plate counting, using a broad-spectrum cocktail of antibiotics: gentamicin (200 µg/ml), ampicillin (100 µg/ml), and vancomycin (100 µg/ml). For translocation assays, Caco-2 were differentiated on 8 µm transwells and stimulated apically with live bacteria for 5 h. Translocation was assessed by agar plating of the basolateral media and was expressed as a translocation percentage of total bacteria. Total RNA was extracted in Trizol after 6 hours co-culture (Qiagen, Montreal, Canada).

<table>
<thead>
<tr>
<th>TABLE 1: Characteristics of Patients in the Study</th>
</tr>
</thead>
<tbody>
<tr>
<td>No.</td>
</tr>
<tr>
<td>-----------------------------------------------</td>
</tr>
<tr>
<td>Healthy Controls</td>
</tr>
<tr>
<td>Crohn’s Disease</td>
</tr>
<tr>
<td>Steroids or mesalamine (3) Others (1)</td>
</tr>
<tr>
<td>Ulcerative Colitis</td>
</tr>
</tbody>
</table>

Values are total number of patients (n). F, female; M, male; SEM, standard error of mean.
Quantitative PCR for CXCL8 mRNA (RT-qPCR Primer, Qiagen) was performed on IQ5 PCR machine (SYBRgreen, BioRad, Mississauga, ON, Canada). Data were analyzed by 2-ΔΔCt method using human B2M (b2-microglobulin, Qiagen) and GAPDH (Qiagen) as control genes and expressed as a fold-change compared to control cells with no bacteria.

**Confocal Staining of Biofilms and Epithelial Cells**

Biofilms were stained, without fixation, with acridine orange (unspecific biomass stain, BD Bioscience), TO-PRO-3 (DNA-RNA, Invitrogen) and visualized on a confocal microscope (Leica DM IRE2 Microsystems, Richmond Hill, ON, Canada). Total biofilm biomass (acridine orange + TOP-PRO-3 stains) was measured using surface staining measurement in Imaris Bitplane (v7, Concord, MA, USA).

**Microbiota 16S Sequencing and Analysis**

DNA from biofilm-dispersed bacteria were extracted by a silica-membrane-based purification method according to the manufacturer’s instructions (QI Amp DNA Mini Kit, Qiagen), with an additional step of incubation in lysozyme solution (20 mg/ml, 1 hour). Amplicon sequencing libraries were obtained from the V4 region of the 16S SSU rRNA using 515P-806R primers. Paired-end amplicons (250 bases) were sequenced on Illumina Mi-Seq (Génome Québec Innovation Centre, Montreal, Canada). Bioinformatics analysis was performed by first removing the primers and quality trimming the reads. UPARSE pipeline was used to generate the OTU (Operational Taxonomic Unit) table and taxonomy was assigned using the RPD database and classifier. Downstream analysis was done in R using phyloseq 1.16.2 package.

**Liquid Chromatography-Mass Spectrometry (LC-MS) Analysis**

Mature biofilms were incubated in fresh media. After 16 hours, spent media was collected and diluted 1:1 in pure methanol. Methanolic extracts were separated by Ultra High-Performance Liquid Chromatography (UHPLC) mass spectrometry (MS), using a hydrophilic interaction liquid chromatography column (Synchronis HILIC, Thermo Fisher, stationary phase). Solvent A (20 mM ammonium formate pH 3 in H2O) and B (MS Grade Acetonitrile with 0.1% formic acid) were used for LC Gradient. High-resolution full-scan MS data were acquired on a Thermo Fisher Scientific Q-Exactive Mass spectrometer using positive-mode electrospray ionization. Metabolites were identified using retention times of standards using MAVEN freeware.

**Caenorhabditis elegans Model**

The *Caenorhabditis elegans* toxicity model was based on a recently published protocol. Briefly, Bristol N2 wild-type *C. elegans* (Caenorhabditis Genetics Center, University of Minnesota, St. Paul, MN, USA) synchronized in L4 stage were fed in 96-well microtiter plates with biofilm-dispersed bacteria in PBS/S-Basal complete media (OD600nm = 0.1). Plates were incubated with bacteria at 20°C for 72 hours. Worm mortality (no active movement, stiff body structure, and no response to mechanical stimuli) was assessed by an experimented investigator blinded to treatments using a binocular microscope (Leica MZ75 microscope).

**Dinitrobenzene Sulfonic Acid (DNBS) Colitis Model**

All experiments using animals were approved by the University of Calgary’s Animal Care Committee (Certificate #AC13-0006). Colitis was induced by intracolonic instillation of 5 mg of DNBS dissolved in 50% ethanol. Control groups were treated similarly with PBS. Mice were given 50 mg/kg of ATB-429 per os in a vehicle of 1% carboxymethylcellulose, twice daily, for 5 days. The severity of colitis was blindly evaluated using previously described endpoints. Liver biopsies were collected aseptically, weighed, homogenized, and plated on blood agar for 24 hours. Carnoy’s-fixed mice colon tissues were paraffin-embedded. Visualization of microbiota biofilm was performed by FISH staining (EUB338-Cy3, Images were acquired using a Leica DM IRE2 confocal microscope and analyzed on Fiji freeware (v.1.51).

**Iron Quantification**

Thirty microliters of detection buffer (5 mM ferrozine, 6.5 mM neocuproine, and 2.5 M ammonium acetate; Sigma-Aldrich) were added to 170 µl of samples, and specific ferrozine-iron absorbance (562 nm) was measured after 30 minutes on a spectrophotometer. Iron concentrations were extrapolated from a standard curve generated with FeCl2. Changes in absorbance of ferrozine-iron complexes after 24 hours were monitored to evaluate the binding ability of chelators. Biofilms were incubated 24 hours to ATB-428 (1 mM), ATB-429 (1 mM), or 2,2-bipyridil (1 mM, Sigma Aldrich), after which biofilm-dispersed bacteria were collected, normalized to OD600nm = 1, and lysed by sonication for measurement of intracellular iron. Mouse feces were mechanically homogenized in 1 ml of 0.1 mM HCl, and iron concentrations were first measured in the fecal spent media (Fig. S7C). Cellulose debris were discarded after centrifugation. Bacteria left were centrifuged and sonicated for measurement of intracellular iron.

**Statistical Analysis**

Graphic representation and statistical analysis were performed using GraphPad Prism (v6, La Jolla, USA). One-way ANOVA with Dunnett’s or Fisher’s test and Kruskal–Wallis with Dunn’s test were used accordingly after D’Agostino–Pearson normality test. For multiple variables, we used 2-way ANOVA with Dunn’s test. An associated P value less than 5% was considered significant. All center values are means for histograms and dot plots, median for box plots. Error bars...
RESULTS

Microbiota from IBD Patients Generate Larger BioFilms and Exhibit Greater Iron Intake than Microbiota from Healthy Controls

Microbiota biofilms abnormally adhere to the epithelial surface of intestinal tissues in patients with IBD. In an attempt to more precisely characterize these biofilms, mucosa-associated microbiota in colonic biopsies from either healthy donors (healthy; n = 7) or IBD patients (Crohn's disease, CD, n = 13; or ulcerative colitis, UC, n = 11) were first homogenized in modified tryptic soy broth, normalized to cell numbers, and plated onto the Calgary Biofilm Device to allow the growth of bacteria under their natural adherent biofilm phenotype (Fig. S1). All procedures were performed under anaerobic conditions and experimental conditions were kept identical among the different cohorts of patients. Taxonomic identification confirmed that biofilms generated ex vivo from the biopsy samples reproduced a complex community of microbiota containing all major bacterial phyla reported in the human gut (Fig. S2). Confocal laser scanning microscopy allowed 3-dimensional representations of biofilms (Fig. 1A). Total biomass quantifications revealed that IBD biofilms had a significantly greater biomass (bacteria and their extracellular matrix) compared to healthy control biofilms (Fig. 1B). IBD-biofilms contained more viable bacteria compared to healthy control biofilms (Fig. 1C). Pathogenic strains of bacteria (eg, pathogens from Enterobacteriaceae family) are known to have high iron uptake capacity, but data regarding human microbiota living as a complex biofilm community are lacking. Bacteria from

![Figure 1](https://academic.oup.com/ibdjournal/advance-article-abstract/doi/10.1093/ibd/izy116/4997020)

**FIGURE 1.** Microbiota from IBD patients form larger biofilms than microbiota from healthy controls and contains higher intracellular iron. 

A. Human microbiota biofilms (healthy; Crohn's Disease, CD; ulcerative colitis UC) were grown on the Calgary Biofilm Device. Confocal laser scanning microscopy of biofilms was performed, the extracellular matrix of biofilms was stained by acridine orange (green) and bacteria DNA with TO-PRO-3 (red). Scale bars correspond to 70 μm. B. Biofilms from IBD patients had a higher relative biomass compared to healthy biofilms (Crystal violet staining, n = 145 to 313 biofilms per group). C. Biofilms from IBD patients contained higher numbers of colony forming units (CFU per biofilm) compared to biofilms from healthy biofilms (n = 21 to 24 biofilms per group). D. Equivalent numbers of bacteria from IBD biofilms contained higher concentration of intracellular iron compared to healthy biofilms (n = 33 biofilms per group). B, 1-way ANOVA followed by Dunnett’s tests. C, D, Kruskal-Wallis followed by Dunn’s tests versus healthy group, **P < 0.01, ***P < 0.001
IBD-biofilms contained significantly higher concentrations of intracellular iron (1.5-fold higher in CD and 1.4-fold in UC) compared to concentrations measured in bacteria from healthy control biofilms (Fig. 1D). Overall, these data demonstrate for the first time that mucosa-associated microbiota from IBD patients have an increased affinity for iron and a greater capacity to grow under a biofilm phenotype compared to microbiota from healthy controls.

**Bacteria from IBD-biofilms Have a More Virulent Phenotype**

To determine adhesion, invasion and virulence characteristics of IBD microbiota biofilms, we used dispersed bacteria from these biofilms (Fig. S1). Indeed, bacteria released by adherent biofilms have strikingly different characteristics than their sessile counterparts, including elevated antimicrobial resistance and virulence.17 Human intestinal epithelial cells (Caco-2) were co-cultured with identical numbers of these biofilm bacteria (8 × 10^6 CFU/ml). After 3 hours of co-culture, bacteria from IBD biofilms were 5.6-fold (for CD) and 14-fold (for UC) more invasive into Caco-2 cells compared to healthy controls (1 ± 0.2 × 10^3 CFU/ml, Figs. 2A, S3A). Finally, bacteria from CD (88-fold) and UC biofilms (13-fold) translocated significantly more across epithelia than bacteria from healthy control biofilms (5 × 10^5 CFU/ml, Fig. 2B). Confocal imaging revealed that translocation followed both the paracellular (Fig. 2C, upper panel) and transcellular (Fig. 2C, lower panel) pathways. These results suggest that mucosa-associated microbiota from IBD patients, grown under a biofilm phenotype, have increased virulent properties compared to microbiota from healthy controls.

**IBD Biofilms Induce Elevated CXCL-8 and Pathogenic Responses Compared to Healthy Biofilms**

To evaluate and compare the proinflammatory host response induced by microbiota from IBD patients or healthy controls, we co-cultured Caco-2 cells with bacteria dispersed from their biofilm and measured CXCL8 mRNA transcription. At 6 hours, CXCL8 mRNA was increased versus vehicle (media only) by 130-fold (for CD biofilms) and 174-fold (for UC biofilms), hence significantly more than the 37-fold increase induced by healthy biofilms (Fig. 2D). IBD microbiota also caused significantly more epithelial necrosis after 5 hours than those from healthy controls (Fig. S4A). Filter-sterilized spent
media from IBD biofilms significantly increased paracellular permeability to dextran-FITC (Fig. S4B). Bacteria from IBD biofilms induced significantly greater lethality in the nematode Caenorhabditis elegans than bacteria from healthy biofilms (Fig. S3B). Together, these data demonstrate that microbiota biofilms from IBD triggered a more severe inflammatory host response than that caused by microbiota from healthy controls, and induced negative effects in epithelia in vitro and are lethal to C. elegans in vivo.

**Hydrogen Sulfide-releasing Mesalamine Derivatives Have Potent Iron-chelating Properties**

ATB-428 and ATB-429 are novel antiinflammatory drugs consisting of mesalamine (5-aminosalicylate; 5-ASA) covalently linked to 2 different H₂S-donors (TBZ or ADT-OH, respectively). Considering the independent interactions of H₂S and mesalamine with iron, we tested the ability of these drugs to chelate iron. Both ATB-428 and ATB-429 (Fig. 3A) effectively chelated iron in a concentration-dependent manner. ATB-429 was more potent than ATB-428 (maximum chelation of 93% for ATB-429 versus 55% for ATB-428), possibly attributable to the more potent iron-scavenging property of ADT-OH (maximum chelation of 39%) versus TBZ (maximum chelation 14%, Fig. 3A). At equimolar concentrations, the 2 drugs had markedly enhanced iron-scavenging properties compared to mesalamine (maximum chelation of 9%). These results demonstrate that ATB-429, in particular, has significant iron-scavenging properties.

**ATB-429 Limits Iron Intake in Colitis-associated Biofilms and Modifies their Metabolome, But Not Their Composition**

Considering the newly revealed iron-chelating properties of ATB-429, and in a lesser way of ATB-428, we exposed human microbiota biofilms to 1 mM of these drugs dissolved in media for 24 hours. ATB-429, but not ATB-428 (Fig. S5A), reduced intracellular iron content in bacteria from biofilms, and this effect was reversed by the addition of iron to the medium (Fig. 3B). At 1 mM, H₂S-releasing mesalamine derivatives did not modify the pH of bacterial media (Figs. S5B, S6A), did not affect bacteria survival (Figs. S5C, S6B), and did not significantly alter biofilms overall species composition (Fig. S2C). Biofilms were incubated for 16 hours in fresh medium containing ATB-429 (1 mM), then metabolomic analyses were performed. Purine and pyrimidine metabolites were measured in the spent media because their metabolism relies on intracellular iron intake and is important for host colonization. Upon exposure to ATB-429, biofilm intake of purine nucleosides (guanosine, Fig. S6C) was reduced, whereas release of urate (Fig. S6D) and hypoxanthine (Fig. S6E), 2 metabolites of purine catabolism, were increased. Biofilm secretion of uracil (Fig. S6F), a pyrimidine nucleobase, was significantly reduced with the addition of ATB-429. The ATB-429 alone (media + ATB-429) had no effect on relative concentrations of metabolites in bacterial media. Together, these data demonstrated that ATB-429 reduces iron intake by colitis-associated microbiota. ATB-429 had a profound effect on microbiota biofilm metabolism, without significantly changing the microbiota species composition.

**FIGURE 3.** Hydrogen sulfide-releasing mesalamine chelates iron and reduces iron intake in microbiota associated with colitis. A, Drugs were dissolved in bacterial media containing 100 µM of FeCl₂. After 24 hours, the remaining concentration of free iron was markedly reduced in a positive control iron chelator (2,2-bipyridil) and in ATB-429, ATB-428, and ADT-OH, but not in TBZ and mesalamine (5-ASA). n = 3 to 9 duplicates per concentrations of drugs. B, Biofilms (healthy controls; Crohn’s disease, CD; and ulcerative colitis, UC) were incubated for 24 hours in media containing 2,2-bipyridil (200 µM; iron chelator positive control), ATB-429 (1 mM), or ATB-429 + FeCl₂ (1 mM and 100µM, respectively). Concentrations of intracellular iron were reduced in microbiota exposed to positive control (2,2-bipyridil, 2,2bp) or to ATB-429; addition of iron abolished the effect of ATB-429 (n = 12 biofilms per group for 2,2-bipyridil, n = 32 biofilms per group for vehicle and ATB-429). C, Intracellular concentrations of iron in fecal microbiota (normalized to 1 g of feces) were significantly higher in microbiota from the colitis group (DNBS) compared to microbiota from control animals (vehicle, veh) and groups treated with ATB-429 (50 mg/kg, twice daily, per os). n = 8 mice per group. B, C, Kruskal-Wallis followed by Dunn’s tests. *P < 0.05, **P < 0.01, ***P < 0.001.
ATB-429 Reduces the Virulent Properties and Associated Proinflammatory Effects of IBD Biofilms In Vitro and Reduced the Severity of Experimental Colitis in Mice

Using in vitro (Caco-2 cells) and in vivo (experimental colitis in mice) models, additional experiments sought to assess whether the reduction of iron intake in microbiota biofilms by ATB-429 was therapeutically meaningful.

When biofilms were preexposed to ATB-429, bacteria translocated significantly less (60% reduction from healthy and CD, 50% reduction for UC) through epithelial monolayers compared to bacteria from biofilms exposed to vehicle alone (Fig. 4A). Preexposure to ATB-429 also reduced by 51% (for CD) and 27% (for UC) the induction of epithelial CXCL8 mRNA by biofilm bacteria (Fig. 4B). Supplementation of iron (10 μM of FeCl₃) partially reversed the effect of ATB-429 (1 mM) by increasing levels of bacterial adherence to Caco-2 cells (Fig. S4C). These data suggest that ATB-429 is able to directly reduce virulence of gut microbiota grown in their natural state of complex multispecies biofilms through an iron-dependent mechanism.

Concentrations of iron in fecal bacteria from mice with experimental colitis (dinitrobenzene sulfonic acid, DNBS) were greater than in fecal bacteria from healthy mice (Fig. 3C), consistent with the effects observed using human IBD microbiota biofilms ex vivo (Fig. 1D). In healthy mice, ATB-429 (50 mg/kg twice daily per os) was well tolerated with no signs of disease (Figs. 4C, S7A). In the DNBS colitis group, coinciding with twice daily per os) was well tolerated with no signs of disease biofilms ex vivo (Fig. 1D). In healthy mice, ATB-429 (50 mg/kg twice daily per os) was well tolerated with no signs of disease (Fig. S7A). In the DNBS colitis group, coinciding with the decreased concentrations of intracellular iron in fecal bacteria (Fig. 3C), administration of ATB-429 reduced the severity of colitis (Figs. 4C, S7A) and inhibited the translocation of commensal bacteria into the liver (Fig. S7B). Staining of microbiota biofilms in situ revealed that during colitis, bacteria invaded the mucus layer and adhered to the intestinal epithelium (arrow, Fig. 5). Treatment with ATB-429 abolished these abnormalities (Fig. 5). Together, these results indicate that ATB-429 reduced iron intake in colitis-associated microbiota and restored the normal localization of the commensal microbiota biofilm, thus efficiently reducing the severity of experimental colitis.

DISCUSSION

Mesalamine is a commonly prescribed drug for treating IBD patients. In addition to its well-known effects on the host, a potential action on microbiota has been scarcely investigated. Indeed, mesalamine concentrations in the lumen of patients can reach 100 mM, thus exceeding by 100-fold concentrations in the host mucosa or in the circulation. It can modify bacterial gene expression and appears to cause a transient increase in the abundance of Firmicutes in humans. It can also suppress polyphosphate accumulation in bacteria, which in turn may serve as an intracellular iron chelator. Independently, antiinflammatory properties of H₂S have been well established in the gastrointestinal tract and H₂S-releasing drugs have become part of extensive research and development programs. In an effort to combine beneficial effects of H₂S and mesalamine, H₂S-releasing mesalamine derivatives have been synthesized and their efficacy in protecting against colitis was demonstrated, but exact mechanisms of action of these novel classes of drugs remain incompletely understood. Little is known about the effects of H₂S on microbiota. Recent reports demonstrated that dietary-derived H₂S donors stimulates the growth of anaerobic microbiota biofilms obtained from experimental animals with colitis.

FIGURE 4. ATB-429 reduces the virulence of IBD microbiota biofilms and protects from DNBS colitis in mice. Human microbiota biofilms (healthy; Crohn’s disease, CD; and ulcerative colitis, UC) were first incubated with ATB-429 (1 mM) for 24 hours. Caco-2 cells were then co-cultured with equal numbers of biofilm-dispersed bacteria. A, Bacteria from IBD biofilms exposed to ATB-429 had a reduced capacity to translocate across Caco-2 cells monolayers compared to bacteria exposed to vehicle (n = 21 to 28 transwells per group, *versus corresponding vehicle group). B, Induction of CXCL8 mRNA transcription in Caco-2 cells was significantly lower when induced by bacteria dispersed from IBD biofilms exposed to ATB-429 versus bacteria from biofilms exposed to vehicle (n = 8 to 15 wells per group, *versus corresponding vehicle group). C, Macrosopic damage scores in the colon of mice from animals with colitis were significantly greater than the scores measured in controls (Healthy); treatment with ATB-429 (DNBS/ATB-429) maintained the damage scores at control levels (DNBS/vehicle; n = 8 mice per group. A, B, Kruskal-Wallis followed by Dunn’s tests. C, 1-way ANOVA followed by Fisher’s tests. *P < 0.05, **P < 0.01, ***P < 0.001
from healthy human donors. These findings motivated our present work to investigate the precise effects of H\(_2\)S-releasing antiinflammatory drugs on the microbiota living in their natural biofilm phenotype.

Anemia is the most common extraintestinal complication of IBD. However, dietary iron supplementation leads to disease exacerbation and a higher risk of infection, possibly through alterations of commensal microbiota. Whether chronic blood loss associated with IBD could elevate iron concentrations in the intestinal lumen, and thus favor populations of pathogenic bacteria, is unknown. In this study, biofilms grown ex vivo from either CD or UC patients were larger in size than healthy control biofilms (Fig. 1) and exhibited elevated adhesion, invasion and virulence compared to healthy control biofilms. Although iron concentrations in the fecal spent media were identical between healthy and colitis group (Fig. S7C), we observed an increase of intracellular iron in fecal microbiota (mice, Fig. 3C) and in human IBD biofilms (Fig. 1D). It was demonstrated in vitro in animal models and in humans that Proteobacteria can thrive at the expense of other gut bacteria in an iron-rich environment. Proteobacteria abundance was indeed higher in UC biofilms versus healthy biofilms, but also versus CD biofilms, with no differences between healthy and CD (Fig. S2). Because iron is an essential nutrient for several other microorganisms, including enterococci, these findings suggest that a unique taxon could not totally explain increased affinity for iron in IBD-associated microbiota. Elevated biofilm growth and virulence were observed for both CD and UC biofilms, although no major alterations in relative abundance were revealed by 16S taxonomic sequencing. Further whole genome sequencing and transcriptomic analyses thus are warranted to determine whether there is a specific strain-advantage or an iron-favored genotype/phenotype in IBD microbiota biofilms, which will help identify new genetic alterations of these pathogenic IBD biofilms.
The present findings show that ATB-429 does not affect bacterial survival, and, importantly, that it does not modify overall bacterial composition of complex multispecies biofilms from human patients (Fig. S2). Nevertheless, exposure of microbial communities to exogenous H$_2$S might lead to a community shift, as this molecule has been demonstrated to be lethal for mouse intestine or to proliferate in the human bloodstream. 20–22 Their environment to efficiently colonize and persist in the Vibrio species (members of Firmicute) Bacillus as well as deprivation by ATB-429 in gut microbiota biofilms warrant further investigation for translational applications. It is well known that pathogenic bacteria require iron during infection. Local iron deprivation may induce some bacteria to increase their expression of iron-chelating siderophores and/or hijack host iron-specific receptors and transport proteins (eg, heme, transferrin). 42 Surrounding low iron concentrations can have either negative (Escherichia coli and Pseudomonas) 43,44 or positive (Staphyloccocus) 45 influence on the formation of monospecies biofilm, but the influence of iron in the context of polymicrobial gut biofilms needs to be further elucidated. Furthermore, knowing that iron can generate powerful radicals, and that iron chelators are strong antioxidants, future research will need to study the potential antioxidant properties of ATB-429, which could have significant implications for its application in other clinical contexts.

The culture of gut mucosal microbiota in their complex, multispecies, anaerobic biofilm phenotype has been a missing link in studies trying to understand the role of microbiota in IBD. Ex vivo models, like the one we used in the present report, help reproduce key components of the microbial-microbial and microbial-host interactions occurring in the gut. Several biases are inevitably associated with these ex vivo models, and some of these include the lack of host components, which are likely to be essential components of the microbial biofilm in vivo (eg, mucus and host DNA). Future research in vivo on molecular characteristics of microbiota biofilms in healthy and disease states, on geolocalization of various bacterial species within the commensal biofilm, and metagenomics will help improve understanding of how gut microbiota may influence health in the gut, and beyond.

In summary, this study offers new evidence that gut microbiota biofilms grown ex vivo from patients with IBD are characterized by increased iron uptake and increased virulent properties compared to biofilms from healthy subjects. The findings also provide proof-of-principle evidence that a new H$_2$S-releasing antiinflammatory drug can suppress iron intake in complex multispecies gut microbiota grown in their natural biofilm phenotype, hence reducing their virulence without modifying overall composition and survival. These are promising observations in our attempts to generate advances towards the development of a novel therapeutic for IBD that will help target both host inflammation and microbiota dysbiosis.

SUPPLEMENTARY DATA
Supplementary data are available at Inflammatory Bowel Diseases online.

ACKNOWLEDGMENTS
The authors are grateful to Dr. R. Leong-Quong (BioCore Facility), Dr. D. Hansen for providing C. elegans, Dr. D.W. Morck for access to the anaerobic hood, Dr. J.J. Harrison for access to the confocal microscope and software, and the Animal Care facility staff. G. Kasendra (translational research technician) helped to provide human biopsies. H$_2$S-mesalamine and H$_2$S-donors were provided by Antibe Holdings Inc., and Innovotech donated the Calgary Biofilm Device.
REFERENCES


29.把你提供的内容转换为自然语言。