ORIGINAL ARTICLE

Bacteria penetrate the normally impenetrable inner colon mucus layer in both murine colitis models and patients with ulcerative colitis

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ABSTRACT

Objective The inner mucus layer in mouse colon normally separates bacteria from the epithelium. Do humans have a similar inner mucus layer and are defects in this mucus layer a common denominator for spontaneous colitis in mice models and ulcerative colitis (UC)?

Methods and results The colon mucus layer from mice deficient in Muc2 mucin, Core 1 O-glycans, Tlr5, interleukin 10 (Il10) and Sla9a3 (Nhe3) together with that from dextran sodium sulfate-treated mice was immunostained for Muc2, and bacterial localisation in the mucus was analysed. All murine colitis models revealed bacteria in contact with the epithelium. Additional analysis of the less inflamed Il10−/− mice revealed a thicker mucus layer than wild-type, but the properties were different, as the inner mucus layer could be penetrated both by bacteria in vivo and by fluorescent beads the size of bacteria ex vivo. Clear separation between bacteria or fluorescent beads and the epithelium mediated by the inner mucus layer was also evident in normal human sigmoid colon biopsy samples. In contrast, mucus on colon biopsy specimens from patients with UC with acute inflammation was highly penetrable. Most patients with UC in remission had an impenetrable mucus layer similar to that of controls.

Conclusions Normal human sigmoid colon has an inner mucus layer that is impenetrable to bacteria. The colon mucus in animal models that spontaneously develop colitis and in patients with active UC allows bacteria to penetrate and reach the epithelium. Thus colon mucus properties can be modulated, and this suggests a novel model of UC pathophysiology.

INTRODUCTION

Inflammatory bowel disease, which is on the rise in the Western world, can be divided into Crohn’s disease (CD) and ulcerative colitis (UC). Both these diseases are dependent on the presence of intestinal microbiota, although specific bacteria have not been linked to disease.1,2 Classical CD has been genetically and functionally linked to proteins such as NOD2 and ATG16L1.3 UC is not as well understood as CD, is restricted to the colon and rectum, and is characterised by superficial mucosal inflammation.4 The disease is likely to occur in genetically predisposed individuals in combination with environmental and microbial exposure.5 A morphological characteristic used to assess disease activity is the typical goblet cell depletion reflecting emptied goblet cells, something that may suggest a role for the mucus system in UC.

The gut epithelium manages enteric bacterial flora in a way that allows a symbiotic relation between host and microbes. The functions, and thus the needs, of the epithelium are different along the intestine, and protection of these surfaces are accordingly handled in different ways.6 The
small-intestinal epithelium mediates nutritional uptake, and the mucus covering these surfaces allows this transport. The bacteri­al load of the intestine, however, increases dramatically in the distal direction with massive colonisation in the colon. These numerous bacteria are normally handled without causing inflammation, but sometimes the immune system is strongly activated and severe inflammation is observed. How this balance is controlled is still not fully understood, but the recently discovered inner mucus layer in the colon, which is impervious to bacteria and physically separates bacteria from the epithelium, has provided a novel insight into this function. It has been suggested that there is also a separation between bacteria and epithelium in humans, but it is still not clear if humans have a similar protective mucus layer formed by MUC2 mucin, although a stratified mucus layer has been suggested.

Colonic mucus is produced by the goblet cell, a cell type devoted to this task. The main mucus component in the intestine is MUC2 mucin, a large and heavily O-glycosylated gel-forming mucin that forms enormous polymeric nets by C-terminal dimerisation and N-terminal trimerisation. Upon secretion from the goblet cells, the mucus expands and builds a stratified dense layer that is attached to the epithelium. This mucus contains additional proteins, of which the FCGBP protein is important as it can stabilise the mucus by additional cross-links. At a distance, far from the epithelial surface, the inner mucus is transformed into a soluble and less organised outer mucus layer which, by proteolytic expansion, generates the preferred habitat for the commensal microbes. Development of spontaneous colitis in Muc2-deficient mice confirms the protective properties of the inner colonic mucus layer. In these mice, bacteria are found in direct contact with the epithelial cells, far down in the colonic crypts and inside enterocytes, locations never observed in wild-type (WT) animals. These observations and earlier studies on humans with UC showing bacteria in contact with the epithelium may suggest that defects in the inner mucus layer that allow bacteria to reach the epithelium in larger quantities than normal can activate the immune system. This prompted us to address, both in animal models and human UC, whether there are defects of the inner colon mucus layer that can be linked to colitis. We now show that colitic mouse models and patients with UC have dysfunctional colonic mucus that can be penetrated by bacteria.

**MATERIALS AND METHODS**

**Animals**

Experiments were performed using WT C57/Bl6, IL-10−/−, Slc9a3−/−, C1galt−/− and Tlr5−/− mice all on a C57/Bl6 background (male 8–12 weeks) according to local ethics committee guidelines. All experiments included controls from the same animal facility as the knock-out strains. Dextran sodium sulfate (DSS) experiments were performed as described.

**Human subjects and biopsies**

Subjects were recruited from among patients referred for colonoscopy at Sahlgrens’ University Hospital, Gothenburg, Sweden. Biopsy specimens from the sigmoid colon of 28 patients with UC were assessed. Disease activity was determined by the Mayo endoscopic score. Clinical information on the patients with UC is presented in table 1. Biopsy samples were also obtained from 12 patients with macro- and micro-scopically normal mucosa (control group). Clinical information on these patients is presented in online supplementary table S1. Written and informed consent was obtained from all study subjects, and approval for this study was granted by the Human Research Ethics Committee, University of Gothenburg. Biopsy samples were collected one at a time using single-use large-capacity forceps (Olympus) and instantly put into ice-cold oxygenated Krebs solution or fixed in Carnoy’s fixative.

**Fluorescence in situ hybridisation (FISH) and immunostaining**

Paraffin-embedded Carnoy-fixed sections were dewaxed and stained with H&E or Alcian blue/periodate acid Schiff (PAS) or hybridised with 10 ng/μl of a general bacterial 16S rRNA probe (EUB338) and immunostained for Muc2 using the MUC2C3 antiseraum or for DNA by 4,6-diamidino-2-phenylindole (DAPI). Images were obtained with an Axio Examiner Z1 LSM 700 confocal microscope and ZEN 2010 software (Zeiss).

**Mucus penetration score, histology score and goblet cell measurements**

Mucus penetration by bacteria was blindly evaluated on DNA-stained sections (n=5 for IL-10−/− and WT) by two independent observers. The scores (0–4) were based on observations from a whole intestinal cross-section per sample, and increased scores correspond to increased contact between bacteria and epithelium. Histological scoring of inflammation was performed on H&E- and Alcian blue/PAS-stained sections (n=3 per genotype) in a blinded fashion by two independent observers evaluating one whole tissue section per sample. The sum of scores for inflammatory cell infiltration (score, 0–4), goblet cell depletion or decreased mucous accumulation (score, 0–4), mucosal thickening (score, 0–4), destruction of architecture (score, 0 or 3–4) and loss of crypts (score, 0 or 3–4) was calculated (maximum 20). All data are presented as mean±SEM.

Number of goblet cells was counted as described in the online supplement. Data are presented as mean±SEM with 34 human samples (10 controls, 14 patients with UC in remission, 10 patients with active disease) and 10 mouse samples (five WT, five IL-10−/−). Mucus-filled theca area of goblet cells was measured on pictures of anti-MUC2C3-stained sections as described in the online supplement. Data are presented as mean area ±SEM with 34 human samples (10 controls, 14 patients with UC in remission, 10 patients with active disease) and 10 mouse samples (five WT, five IL-10−/−).

**Preparation of mouse tissue for in vivo and ex vivo experiments**

For the in vivo experiments, mice were anaesthetised by continuous administration of isoflurane (Isoba vet; Schering Plough). The abdomen was opened and a cup was placed on the opened intestinal segment and filled with warm saline. For the ex vivo experiments, mice were anaesthetised with isoflurane and killed by cervical dislocation. The distal colon was dissected, flushed, and the longitudinal muscle layer removed. The tissue explant was mounted in a horizontal perfusion chamber.

**Mucus thickness**

The thickness of the colonic mucus was measured as described previously. Briefly, the upper surface of the colonic mucus was visualised by addition of charcoal particles. The mucus thickness was determined by measuring the distance between the epithelial surface and the mucus surface by a micropipette viewed through a stereomicroscope (Leica MZ12).

**Mucus penetrability**

Mucus penetrability was measured as described previously. Briefly, colonic explants were mounted in a perfusion chamber
The inner mucus layer of the distal colon is formed from sheets of Muc2 mucin organised into stacked layers in a stratified way that does not allow bacteria to penetrate. This physically separates bacteria from the epithelium (figure 1, WT). Animals deficient in the Muc2 gene are devoid of secreted mucus layers, resulting in constant contact between bacteria and the epithelium (figure 1, Muc2−/−). The presence of bacteria in contact with the epithelium was estimated (figure 2A and online supplementary figure S1A), and the level of inflammation was estimated on H&E- and Alcian blue/PAS-stained tissue sections evaluating immune cell infiltration, crypt architecture and goblet cell depletion (‘depletion’) (figure 2B,C and online supplementary figure S1B). As observed previously, the Muc2−/− animals showed severe inflammation marked by infiltration of leukocytes and increased cell proliferation as evidenced by crypt elongation (figure 2C, Muc2−/−). A more subtle way to modify Muc2 mucin is to alter its glycosylation by shortening the O-glycans. Blocking the core 1 extensions by deletion of the C1galt1 glycosyltransferase results in reduced mucin glycosylation, which causes spontaneous colitis. These animals show a defective inner mucus layer that allows bacteria to penetrate and come in contact with the epithelium (figure 1 and online supplementary figure S1A, C1galt1−/−). Sodium–hydrogen exchanger Scl9a3 (Nhe3)-deleted mice have been shown to develop spontaneous colitis and colon tissue sections from these mice were analysed, essentially normal mucus was observed with regard to thickness and

Table 1 Clinical information on the analysed patients with UC

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*For patients with recent disease onset, the number of relapses cannot be given. Instead year of debut is stated in parentheses.
**PSC, primary sclerosing cholangitis. x, means present.
†Medication with TNFa inhibitor for rheumatoid arthritis.
‡This patient had inflammation in more proximal part.
5-ASA, 5-aminosalicylic acid; A, alterations in crypt architecture; C, cryptitis; CA, crypt abscesses; D, denuded epithelium; E, eosinophils; F, female; L, lymphocytes; M, male; N, neutrophils; NAD, nothing abnormal detected; P, plasma cells; PSC, primary sclerosing cholangitis; TNF, tumour necrosis factor; U, ulceration; UC, ulcerative colitis.
stratified appearance, but it was penetrated by bacteria (figure 1 and online supplementary figure S1A, Slc9a3−/−). As previously described, the tissue sections also showed signs of inflammation with infiltration of immune cells (figure 2C and online supplementary figure S1B, Slc9a3−/−). The reason for this mucus phenotype is not understood, but an optimal local ion milieu is believed to be necessary for normal mucus expansion and organisation.

The most widely used colitis model is the DSS model where rodents are given 2–5% DSS in the drinking water. Inflammation typically appears after 3–5 days. When distal colon tissue sections were examined after 12 h exposure, the
inner mucus layer looked relatively normal with regard to thickness and with a stratified pattern. However, bacteria had penetrated the mucus and were found close to the non-inflamed epithelium (figure 1 and online supplementary figure S1A, DSS). After 5 days of DSS, the inner mucus layer was lost and the tissue severely inflamed.

Bacteria recognition is part of the innate response, and a subgroup of Tlr5-deficient mice spontaneously develop colitis. When such animals were studied, an inner mucus layer was observed, but it was penetrated by bacteria, and massive infiltration of immune cells was observed (figures 1 and 2C, Tlr5−/− Infl.). The non-inflamed Tlr5-deficient mice revealed a more intact and bacteria-free inner mucus (figure 1, Tlr5−/− Non-Infl.).

One of the earliest knock-out animals generated was the one lacking IL-10, leading to inflammation that is dependent on the presence of bacteria. When these animals were analysed, they had very mild inflammation and normal thickness and stratified pattern of the inner mucus layer. Interestingly, their inner mucus was also penetrated by bacteria (figures 1 and 2A, IL-10−/−). In general terms, more severe inflammation often correlated with a high number of bacteria in contact with the epithelium. The major exception to the rule was the animals exposed to DSS for only 12 h where the bacteria had penetrated the inner mucus layer before development of colitis. This indicates that bacterial contact with the epithelium precedes inflammation and that bacterial contact could trigger subsequent inflammation. The IL-10−/− mice showed a low histology score and also a low mucus-penetration score, suggesting a relation between bacterial penetration and inflammation. Thus all tested mouse models that developed colitis showed bacteria in direct contact with the epithelium in the distal colon.

The secreted mucus layers are not thinner in IL-10-deficient mice with colitis

Because the IL-10−/− mice have become a prototype of experimental colitis and do not have a primary defect in any mucus component, we analysed this model in more detail. As the penetration of the inner mucus layer by bacteria appears to be a hallmark of murine colitis, one could speculate that this is due to a thinner mucus layer. To address this, we measured mucus thickness in the distal colon both in vivo and ex vivo in young IL-10−/− mice with low-grade inflammation and WT mice (figure 3A,B). In vivo the thickness of the total (inner and outer) mucus layer was measured directly; the loose outer mucus was then aspirated, and the thickness of the inner firmly adherent mucus was measured. The total thickness did not differ between IL-10−/− and WT mice, but the adherent mucus was significantly thicker in the IL-10−/− mice (figure 3A). Distal colon tissue was also mounted in a horizontal perfusion chamber, and mucus growth was followed ex vivo for 1 h. Mucus thickness and growth during this time did not differ between IL-10−/− and WT mice (figure 3B). Thus, a thinner mucus layer cannot explain why bacteria are found close to the epithelium in the IL-10−/− mice.

The mucus stored in the goblet cells is the material used to build up and renew the secreted mucus layer. The number of goblet cells in the upper 100 μm of the crypt were counted in IL-10−/− and WT mouse colon sections. The amount of goblet cells did not differ between the two (figure 3C). The amount of stored mucus was assessed by measuring the area

Figure 2  Bacteria penetration and inflammation in different colitis models. (A) Bacteria penetration of the inner mucus was scored for IL-10−/− (n=5) and wild-type (WT; n=5) mice, with Muc2+/−, which have high numbers of bacteria in contact with the epithelium, as comparison (n=3). Data are presented as mean±SEM. Penetration scores for all the genotypes are presented in online supplementary figure S1. nd, not detected. (B) Inflammation was monitored as histology scores for IL-10−/− and WT (n=5) mice, with Muc2+/− mice as a severely inflamed comparison (n=3). Data are presented as mean ±SEM. Histology scores for all the genotypes are presented in online supplementary figure S1. (C) H&E-stained tissue sections corresponding to samples in figure 1. Scale bars in all panels are 100 μm. DSS, 3% dextran sodium sulphate; Infl. inflamed.
Inflammatory bowel disease

Figure 3  Mucus thickness in wild-type (WT) and interleukin 10-deficient (IL-10−/−) mice. (A) In vivo measurements of the initial mucus thickness in WT (n=7) and IL-10−/− (n=5) mice. The total mucus thickness (Total) was measured followed by aspiration of the mucus and measurement of the remaining mucus thickness (Adherent). (B) Ex vivo measurements of the increase in total mucus thickness over time in WT (n=5) and IL-10−/− (n=6) mice. (C) Number of goblet cells per upper crypt in WT (n=5) and IL-10−/− (n=5) mice. (D) Area of the goblet cell theca in anti-MUC2C3-stained colon section of WT (n=5) and IL-10−/− (n=5) mice. Data are presented as mean±SEM, and the two-tailed Mann–Whitney U test was used to compare the mucus thickness in WT and IL-10−/− mice.

of the mucus-filled theca in goblet cells of the upper part of the crypt. There was no significant difference in the area of the mucus-filled theca of the goblet cells between WT and IL-10−/− mice (figure 3D). The stored mucus available for secretion is thus not altered in IL-10−/− mice.

The mucus quality is defective in IL-10-deficient mice

As the inner mucus was not found to be thinner in the IL-10−/− mice, we instead analysed the penetrability of the mucus to beads the size of bacteria. Colonic explants were mounted in a horizontal perfusion chamber and allowed to secrete mucus for 20 min; fluorescent beads (0.5 and 2 μm) were then placed on top of the mucus formed. The beads were allowed to sediment for 40 min, and the position of the beads in relation to the epithelium was determined by confocal microscopy. WT explant mucus did not allow the beads to penetrate and kept both 0.5 and 2 μm beads separated from the epithelium (figure 4A). On the other hand, mucus produced by the IL-10−/− mice was unable to create such a separation (figure 4A). To quantify mucus penetrability, we estimated the bead intensity in close proximity to the epithelium (figure 4B). In the IL-10−/− mice, almost 50% bead intensity was found close to the epithelium compared with 1% in the WT mice. Mucus from IL-10−/− mice was completely penetrable, while the controls had ~200 μm thick non-penetrable inner mucus (figure 4C). Thus, the quality of the inner mucus is compromised in IL-10−/− mice. The thickness is based on the mucus containing beads, and as the mucus in IL-10−/− mouse colon is very penetrable, this results in a smaller value than in figure 3B where the mucus thickness is based on charcoal binding to the mucus surface.

Bacteria in normal human sigmoid colon are well separated from the epithelium

As previously described, mouse colon has a two-layer mucus system where the inner layer is devoid of bacteria, so we first asked if the same organisation was present in humans. Patients referred for colonoscopy were included after they had given informed consent. Clinical information on the control patients is compiled in online supplementary table S1. The penetrability of the secreted mucus from sigmoid biopsy samples was measured, and representative Z-stack projections and corresponding normalised intensity plots for the different groups were collected (figure 5). Control patients showed an impenetrable (IP) mucus layer, on average 400 μm thick, separating beads from the epithelium (figure 5A, D). To confirm that the human colonic mucus forms an inner mucus layer similar to that observed in mouse colon,7 we analysed control sigmoid samples not pretreated with laxative (figure 6C,D). Biopsy samples were fixed to preserve the mucus, immunostained for MUC2, and counterstained for DNA to visualise cell nuclei and bacteria (FISH probes do not detect all bacteria associated with the biopsy samples). Similar to the mice, a stratified mucus layer was observed with bacteria only present at the luminal surface of the mucus.

Patients with active UC and a subgroup of patients with UC in remission have penetrable mucus

Considering that increased bead penetrability correlated with bacteria close to the epithelium in mouse colitis models, we asked whether this also translated to human patients with UC. Patients with UC were divided into two groups: those in remission (Mayo endoscopic score 0) and those with active disease (Mayo endoscopic score 1–3). Clinical information on the patients with UC is compiled in table 1. The patients with active disease displayed many more inflammatory-associated histological changes than patients in remission. The penetrability of the secreted mucus from sigmoid biopsy samples was measured, and representative Z-stack projections and corresponding normalised intensity plots for the different groups were collected (figure 5). All patients in remission (Mayo score 0) except three had a mucus thickness similar to the control patients (figure 5D). In contrast, most patients with active UC had a thinner mucus layer that was penetrable by beads. Mucus penetrability, quantified as the number of beads close to the epithelium (20 μm into the tissue and 120 μm into the lumen), was ~40% in patients with active disease, 10% in patients in remission, and close to zero in the controls (figure 5B). The individual values for penetrability versus mucus thickness (defined as bead-containing material) show heterogeneity especially in the patients in remission (Mayo score 0) (figure 5C). Interestingly, three of the patients in remission (numbers 1, 2 and 12) produced mucus that was penetrable to the beads. When these patients with penetrable mucus (P) were plotted separately from the remission group, the IP group looked like the control patients and the P group like those with active UC (Mayo score 1–3) (figure 5D). Thus, mucus secreted from
human sigmoid colon was normally dense and thick, whereas that in patients with active inflammation and in a subgroup of patients in remission was thinner and penetrable. The mucus quality did not correlate with azathioprine treatment. The number of goblet cells in the upper 150 μm crypt was determined, and no differences were observed in any of the groups, indicating that patients who are either in remission or have active inflammation do not have fewer goblet cells (figure 6A).

Mucus stored in goblet cells of the upper crypts measured as theca area did not differ between controls and patients in remission (figure 6B). The patients in remission who were identified to have penetrable mucus (figure 5B,C) had a similar goblet cell theca area, arguing for normal amounts of stored mucus (figure 6B). The patients with active inflammation were, however, observed to have fewer filled goblet cells, seen as smaller theca areas (figure 6B), which would correspond to the thinner mucus layer observed.

We finally asked if the difference in mucus quality of the patients with active UC, which allowed beads to sediment down to the epithelium, reflected bacteria penetrating the inner mucus. Biopsy specimens taken from the same individuals as used for penetrability measurements were fixed to preserve the mucus, immunostained for MUC2, and counterstained for DNA to visualise cell nuclei and bacteria. These patients were all pre-treated with oral laxative to clear the intestine for routine colonoscopy. This treatment affects the mucus and its turnover, and the fixed material shows a more voluminous and less well structured mucus in control patients also (figure 6C-b,D-b). The
amount of mucus that remained attached to the epithelial surface varied between the biopsy specimens, which excluded evaluation of all patients. In comparison with the laxative-treated control patients, the patients with inflammation had less attached mucus, and bacteria penetrated this mucus and reached the epithelium (figure 6C-c, D-c).

**DISCUSSION**

As demonstrated previously, the inner mucus layer of normal healthy colon of mice forms a barrier that separates bacteria from the epithelium. This was shown by Carnoy fixation of intact tissue without previous washing of the luminal content. We have now studied a number of different colitis models and confirmed mucus defects in all of them. It is well established that intestinal bacteria are necessary for all these models and that the composition of the microbiota influences the severity and outcome of the inflammation. Muc2-null mice totally lack mucus and spontaneously develop severe inflammation, while other models all have a more or less intact inner mucus layer which is, however, penetrated by bacteria. The glycan-deficient Core 1-deficient mice have less well glycosylated Muc2 mucin, which is probably easier to degrade by the bacteria. DSS induces inflammation initiated by a direct toxic effect on the mucus. Tlr5−/− mice have a thinner and defective inner mucus layer. Nhe3-deficient and IL-10−/− mice display a morphologically stratified inner mucus layer that is still penetrable to bacteria. These data argue for an impaired mucus layer that allows bacteria to penetrate as a potential mechanism involved in the development of colon inflammation.

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Figure 5  Mucus penetrability in human colonic biopsy samples from controls (n=12) and patients with ulcerative colitis (Mayo 0, n=17, Mayo 1–3, n=11). (A) Representative Z-stack projections with the respective normalised intensity plots. Scale bars 100 μm. (B) Percentage of total bead intensity in close proximity to the epithelial surface (<120 μm). Differences between groups were analysed using the Kruskal–Wallis test with Dunns’ correction for multiple comparisons (primary analysis p=0.0009 for the 2 μm beads and p=0.0004 for the 0.5 μm beads). (C) Percentage of beads in relation to total mucus thickness of individual patients. Numbers refer to patient numbers given in table 1. (D) Relation between penetrable and impenetrable mucus of the 2 μm green beads. C, control; M0 IP, patients with Mayo endoscopic score 0 and impenetrable mucus; M0 P, Mayo score 0 and penetrable mucus (patients number 1, 2 and 12); M1–3, Mayo score 1–3.

Figure 6  Mucus-filled goblet cells and bacteria in fixed human sigmoid biopsy samples. Human colon biopsy samples were Carnoy fixed to preserve the mucus and immunostained for MUC2 in combination with 4',6-diamidino-2-phenylindole (DAPI) for DNA in nuclei and bacteria. (A) The number of goblet cells per upper crypt was determined in fixed and MUC2-stained sections from patients with ulcerative colitis (UC) and controls. (B) The goblet cell theca area was measured in fixed and MUC2-stained sections from patients with UC and controls. Data are presented as mean±SEM, and the Kruskal Wallis test with Dunn’s correction for multiple comparisons was used to compare the UC patients with the control group. Ctrl, control; M0 IP, patients with endoscopic Mayo score 0 and impenetrable mucus; M0 P, Mayo score 0 and penetrable mucus (patients number 1, 2 and 12); M1–3, Mayo score 1–3. (C) Sections from human sigmoid biopsy samples stained for MUC2 (green) and DAPI (blue). (a) A biopsy specimen collected and directly fixed from sigmoid colon of a control patient without preceding laxative treatment. (b) A biopsy specimen from a control patient included in the penetrability study who was pretreated with laxatives before colonoscopy. (c) A biopsy specimen from patient with UC pretreated with laxative and with a Mayo endoscopic score of 2 at colonoscopy. Pictures to the right only show the DAPI staining. Bacteria (arrows) are found on the outer surface of the mucus in control patients (a and b). Bacteria are found inside the inner mucus and close to the epithelium in the patient with active UC (c). Some detached cells can be observed (arrowhead). (D) H&E-stained tissue each corresponding to parts a–c in (C). Scale bars are 10 μm (C) and 100 μm (D).

Assuming that the inner mucus layer of the colon is vitally important for colon barrier function, a thinner mucus layer could be associated with inflammation. However, mucus thickness alone was not a useful indicator of mucus barrier function since, for example, the IL-10-deficient mice had a thicker inner attached mucus layer. Instead, these animals showed increased mucus secretion, maybe in an attempt to overcome the mucus barrier defect.

We can now show that the sigmoid colon of healthy humans also has an inner stratified MUC2-containing mucus layer that separates bacteria from the epithelium. Bacterial contact with the epithelium has previously been observed in inflamed intestinal samples, but it was only analysed on fixed biopsy samples. The mucus is then often lost or degraded, problems that can hamper the evaluation. We have now studied the protective quality of the mucus by assessing its ability to restrict small spherical particles from sedimenting down to the epithelial surface to illustrate its property of excluding bacteria close to the epithelium. This was performed on live explant under physiological conditions to gain information about its normal in vivo function. The distance between beads and cells was ∼200 μm in the mouse, something that should be compared with ∼50 μm thick inner mucus layer as measured after aspiration of the outer loose mucus layer. That the beads did not pass all the way down to 50 μm suggests that the outer mucus layer is not homogeneous and that it is denser close to the border between the inner and outer mucus layer. This is in line with the prediction of proteolytic activity being responsible for the conversion from inner dense to outer loose mucus layer.

The mucus separating the beads was ∼400 μm thick in humans, and thus the inner mucus layer can be predicted to be about twice as thick in humans as in mice.

The IL-10<sup>−/−</sup> model is one of the oldest colitis models studied, in which the inflammation can be explained by the loss of the anti-inflammatory effects of IL-10. The proinflammatory initiating effect is mediated by colonic mononuclear phagocytes of the innate defence system. The IL-10<sup>−/−</sup> mice kept in our animal house display only minor histological inflammatory signs, but still have a mucus layer that is penetrable to both beads and bacteria. This argues for a link between mucus properties and the immune system and cytokines produced. The mechanism behind the IL-10 effect on the mucus-producing goblet cells is not understood, but the number of goblet cells and size of the theca with the mucus granulae in the goblet cells do not differ from WT mice. This implies changes not related to mucus amount and may suggest other mechanisms. Mucus organisation and expansion at secretion depend on the external milieu, and alternations could have devastating effects.

Mucus organisation is also a result of additional cross-linking properties and the immune system and cytokines produced. The property of excluding bacteria close to the epithelium has previously been observed in in vitro experiments. The surface epithelial cell layer can probably withstand and handle some bacteria, but will most probably have difficulty withstanding substantial direct bacterial contact for a long period. Extensive bacterial contact will cause bacterial leakage into the tissue, something that could trigger the subepithelial adaptive immune system.

Most of the patients with UC in remission (Mayo score 0) had, like the control patients, a thick inner mucus layer that was...
not penetrated by fluorescent beads. However, three of these patients had mucus that was more or less totally penetrable by the beads, but did not have smaller goblet cell theca area in analogy with the IL-10−/− mice. Two of these patients had primary sclerosing cholangitis. As all the spontaneous mouse colitis models had mucus that was penetrable to bacteria, it can be expected that some patients with UC might also have mucus defects making their mucus less protective. The reason for the observed mucus alterations in patients with active colitis may be higher mucus secretion as shown by the same number of, but less well filled, goblet cells—observed as smaller theca area or more empty goblet cells—described after PAS staining as goblet cell depletion. This characteristic is mainly attributable to more severely inflamed tissue and not observed in inactive disease or the less inflamed IL-10−/− animals. As generation of the enormously large MUC2 polymeric mucus network is time consuming and difficult, a high demand and fast turnover, as in inflammation, are likely to generate mucus of poorer quality.17 This was recently illustrated by studies of colon ischaemia where reperfusion emptied the crypt goblet cells with a concomitant clearance of bacteria.18 After this, the goblet cells were not refilled for many hours, and another challenge would be less efficiently handled. An inner mucus gel that is less dense with MUC2 mucin is likely to be more penetrable, but other more sophisticated mechanisms are likely to be involved. The genetic and environmental factors causing and maintaining UC are probably numerous. Our observations suggest that the inner mucus and its barrier function is an important factor in limiting bacterial contact with the epithelium and that defects in this system might trigger inflammation. The properties of colon mucus are also shown to be dynamic and probably modulated by both bacteria and host factors using only partly known mechanisms. Understanding these mechanisms may provide novel approaches for prolonging remission periods for patients with UC.

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Contributors

MEVI and GCH conceived the original idea. MEVI, JKG, JH-L, HS, KSJ and GCH designed the study, analysed the data and wrote the manuscript. MEVI, JKG and JH-L conducted the experiments, and analysed the data. LX, HX, FKG, FAC and ATG provided genetically modified mouse tissues. KSJ and HS analysed the patients.

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Competing interests

None.

Ethics approval

All animal studies were approved by the animal ethics committee of the respective universities: Gothenburg, Emmo, Arizona and Oklahoma. The human studies were approved by the Human Research Ethics Committee, University of Gothenburg.

Patient consent

Obtained.

Provenance and peer review

Not commissioned; externally peer reviewed.

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Supplemental material to:
Bacteria penetrate inner colon mucus layer in both murine colitis models and in patients with ulcerative colitis

MATERIALS AND METHODS

Animals
Experiments were performed using wild type C57/Bl6, IL10<sup>−/−</sup>, Slc9a3<sup>−/−</sup>, C1galt<sup>−/−</sup>, Muc2<sup>−/−</sup> and Tlr5<sup>−/−</sup> mice all on a C57/Bl6 background (male 8-12 weeks) according to local ethics committee guidelines. All experiments included controls from the same animal facility as the knock-out strains. DSS experiments were performed as described (1;2). The animals were housed in standardized conditions of temperature (21-22°C) and illumination (12h light and 12 dark) with food and water provided ad libitum.

Human subjects and biopsies
Subjects were recruited among patients referred for colonoscopy at Sahlgrens’ University Hospital, Gothenburg, Sweden. Biopsies from the sigmoid colon were assessed from 28 patients with UC. Disease activity was determined by the endoscopic Mayo Score (3) and histological evaluation was performed by clinical pathologist. Clinical information regarding the UC patients is presented in Table 1. Biopsies were also obtained from 12 patients referred for colonoscopy for reasons such as anemia, bleeding and altered bowel habits. These subjects all had macroscopically normal mucosa at colonoscopy, and were used as a control group. Clinical information on these patients is presented in Table S1. Written and informed consent was obtained from all study subjects and approval for this study was granted by the Human Research Ethical Committee, University of Gothenburg. Biopsies were collected one at the time using single-use large capacity forceps (Olympus) and instantly put into ice-cold oxygenated Krebs buffer for studies of mucus penetrability or fixed in Carnoy’s fixative (4) for immunohistological analysis.

FISH and immunostaining
Animals were euthanized by cervical dislocation and the colon dissected. A piece of the distal colon containing a fecal pellet was cut and fixed in Carnoy’s fixative (60% dry methanol, 30% chloroform, 10% glacial acetic acid). Paraffin embedded sections were dewaxed and stained with H&E, Alcian blue/PAS or hybridized with 10 ng/µl of a general bacterial 16S rRNA probe (EUB 338) and immunostained for Muc2 using the MUC2C3 antisera or DNA was stained by DAPI (4).

Images were obtained with an Axio Examiner Z1 LSM 700 confocal microscope using a plan apochromat 40x/1.3 oil DIC objective and the ZEN 2010 software (Zeiss, Germany). In the immunohistological analysis of mouse colon the pictures were taken with a resolution of 2048x2048 pixels, 129 µm pinhole, averaging 4 pictures in two tracks (excitation: 555 nm and 488 nm). Immunohistochemistry of human biopsies were examined using the same instrument with 1024x1024 resolution, 92 µm pinhole, averaging 8 pictures in two tracks (excitation: 405 nm and 488 nm). Brightness and contrast adjustments were made for the whole image and all samples were equally processed.
Mucus penetration score and histology score. Mucus penetration by bacteria was blinded evaluated on DNA stained sections by two independent observers. The IL10^−/+ and WT groups (n=5) were used for statistical analysis. Mucus penetration by bacteria was in addition scored in the other genotype groups (n=3). The scores (0-3) were based on observations from 3 whole intestinal cross section per sample. No bacteria in contact with the epithelium and no bacteria in the inner mucus layer was assigned a value of 0. Bacteria found in the inner mucus layer with up to 20% of the surface epithelial cells in contact with bacteria was assigned a value of 1. Bacteria in the inner mucus layer with about 50% of the surface epithelial cells in contact with bacteria was assigned a value of 2, and bacteria in the inner mucus layer with more than 80% of the cells in contact with bacteria or bacteria located in the crypts was assigned a value of 3. Data is presented as mean±SEM.

Histological scoring of inflammation was performed on H&E and Alcian blue/PAS sections in a blinded fashion by two independent observers evaluating a whole tissue section per sample. The IL10^−/+ and WT groups (n=5) were used for statistical analysis. Inflammatory histology scores were also evaluated in the other genotype groups (n=3). The sum of scores for inflammatory cell infiltration (score, 0–4), goblet cell depletion or decreased mucus accumulation (score, 0–4), mucosa thickening (score, 0-4), destruction of architecture (score, 0 or 3-4) and loss of crypts (score, 0 or 3-4) was calculated (maximum 20). Inflammatory cell infiltration (0-4): The presence of no or occasional inflammatory cells in the lamina propria was assigned a value of 0, increased number of inflammatory cells in the lamina propria was assigned a value of 1, high numbers of number of inflammatory cells in the lamina propria was assigned a value of 2, confluent inflammatory cells in the lamina propria was assigned a value of 3 and confluent inflammatory cells in the lamina propria, extending into the submucosa was assigned a value of 4. Goblet cell depletion or decreased mucus accumulation (0-4): Normal goblet cell distribution with densely filled goblet cells was assigned a value of 0. 5-20% decrease in number of densely filled goblet cells was assigned a value of 1. 20-40% decrease was assigned a value of 2, 40-50% decrease was assigned a value of 3, and more than 50% reduction was assigned a value of 4. Mucosa thickening and crypt elongation (0-4): Normal thickness was assigned a value of 0, 20-40% increased thickness was assigned a value of 1, 40-80% increased thickness was assigned a value of 2, 80-100% increased thickness was assigned a value of 3, and more than 100% increased thickness was assigned a value of 4. Destruction of architecture (0, 3-4): Normal epithelial structure was assigned a value of 0, changed architecture of the epithelium with altered crypt structure and irregular luminal surface was assigned a value of 3 and gross structural changes including ulcers was assigned a value of 4. Crypt loss (0, 3-4): Normal crypt frequency was assigned a value of 0, 5-20% decreased number of crypts was assigned a value of 3 and more than 20% loss of crypts was assigned a value of 4. Data is presented as mean±SEM.

Goblet cell number and theca area measurement.

Number of goblet cells were counted for a defined distance (100 μm of mouse samples and 150 μm for human samples) from the surface epithelium of longitudinally cut crypts. A total of 12 crypts (4 crypts per section) were analyzed for each mouse and 3 crypts (from 3 different sections) were analyzed for every human subject. Data is presented as area±SEM.
with 34 human samples (10 controls, 14 UC patients in remission, 10 UC patients with active disease) and 10 mouse samples (5 WT, 5 IL10⁻/⁻).

Mucus filled theca area of goblet cells were measured on pictures of Anti-MUC2C3 stained sections using the Volocity software (version 6.1, Perkin Elmer). The pictures were acquired with the same settings, and the intensity threshold was set to 15% and a size limit exclusion of 30 µm² was used. Region of interests were chosen to only include goblet cells of the upper third of the crypt and 25 goblet cells (in 4 sections of 1 biopsy) were measured per human sample and 60 goblet cells (20 goblet cells per section in 3 sections) per mouse sample. Data is presented as area±SEM (µm²) with 34 human samples (10 controls, 14 UC patients in remission, 10 UC patients with active disease) and 10 mouse samples (5 WT, 5 IL10⁻/⁻).

Preparation of mouse tissue for in vivo and ex vivo experiments
For the in vivo experiments mice were anesthetized by spontaneous inhalation of 3.5% isoflurane (Isoba vet, Schering Plough) and moved to a breathing mask (AgnThos) for continuous isoflurane administration. Body temperature was kept at 37°C by a heating pad and controlled by a rectal thermometer. The abdomen was opened through a midline incision, and the distal colon positioned on a holder (in house) at the level of the abdomen. The intestinal segment was cut open (1 cm). Fecal pellets were removed and the mucosal surface was rinsed with warm saline. A cup exposing 0.07 cm² of the tissue was gently positioned on the mucosa and filled with warm saline. The isoflurane dosage was lowered to 2% prior to mucus thickness measurement.

For the ex vivo experiments mice were anaesthetized with isoflurane and euthanized by cervical dislocation. The distal colon (approximately 2 cm) was dissected, flushed with ice-cold oxygenized Krebs’ buffer and opened along the mesenteric border. The longitudinal muscle layer was removed by blunt dissection and the tissue was mounted in horizontal perfusion chambers for measurements of mucus thickness or mucus penetrability.

Perfusion chamber characteristics and buffer compositions
Horizontal perfusion chambers with an open apical chamber and a closed basolateral chamber were used in both the mucus thickness measurements and mucus penetrability measurements. The chamber characteristics were identical except for two features. The apical chamber volume was 150 µl for the mucus thickness measurements and 1.5 ml for the mucus penetrability measurements. The diameter of the circular opening was 2.5 mm in the thickness measurements and 1.5 mm in the penetrability measurements. The chambers were mounted in a heating block and kept at 37°C during the whole experiment. The apical Krebs’ buffer was kept unstirred during the experiment to limit disturbance to the mucus layer while the serosal chamber (volume 165 µl) was perfused at a rate of 5ml/h. The buffer compositions were the same in both chambers; Krebs’ buffer (mM): NaCl 116, CaCl₂ 1.3, KCl 3.6, KH₂PO₄ 1.4, NaHCO₃ 23 and MgSO₄ 1.2 (Merck, Germany). The apical Kreb-Mannitol buffer also contained Na-Pyruvate (5.7 mM) (Sigma-Aldrich, Germany), Na-L-Glutamate (5.1 mM) (Merck, Germany) and D-Mannitol (10 mM) (Sigma-Aldrich, Germany) whereas the serosal Kreb-Glucose buffer contained D-Glucose (10mM) (Sigma-Aldrich, Germany) instead of D-
Mannitol. All solutions were oxygenized (95% O₂, 5% CO₂) for 20 min prior to the experiment and pH was set to 7.4 using 37% HCl.

**Mucus thickness**
The thickness of the colonic mucus was measured as described previously (2). Briefly, the upper surface of the colonic mucus was visualized by addition of charcoal particles to the apical surface of the tissue. The thickness of the mucus layer was then determined by measuring the distance between the epithelial surface and the mucus surface by a micropipette (tip diameter ~10 µm) connected to a digimatic indicator (Mitutoyo, Japan) and viewed through a stereomicroscope (Leica MZ12). For the *in vivo* experiments, the initial mucus thickness was measured followed by aspiration of mucus and measurement of the remaining layer thickness. For the *ex vivo* experiments, the initial mucus thickness was measured followed by repeated measurements in 15 min intervals for 60 min.

**Mucus penetrability**
Mucus penetrability was measured as described previously (2). Briefly, human colonic biopsies or mouse colonic explants were mounted in the perfusion chamber as for mucus thickness measurements, and incubated for 20 min followed by addition of a suspension of 2 µm green beads and 0.5 µm red beads to the apical surface (Fluospheres, Invitrogen). The beads were left to sediment through the mucus for 40 min after which the position of the beads in relation to the epithelium was analyzed by taking confocal images in XY stacks (320x320 µm).

An LSM 700 Axio Examiner Z.1 confocal imaging system with a Plan-Apochromat 20x/1.0DIC water objective (Zeiss, Germany) was used in the penetrability studies. XY sections were obtained with an optical thickness of 2.8 µm in 10 µm intervals. The pictures were obtained at a 512x512 pixels resolution using bidirectional scan, 12 bit depth, 45 µm pinhole, 6.3 µs pixel dwell time and an average of 2 pictures. In track 1 the excitation was at 405 nm and emission at SP 490 nm, DBS 493 nm and master gain 730-820. In track 2 the excitation was at 488 nm and emission at SP 555 nm, DBS 604 nm and master gain 329. In track 3 the excitation was at 555 nm the emission at SP 640 nm, DBS 626 nm and master gain was set to 350. Data was analyzed using the Volocity 5.5.1 software (Perkin-Elmer). The density parameter was set to 70% in track 1 and (track 1) and 50% in track 2 and 3 and the brightness was set to 2x and 3x respectively. Data was obtained as total intensity measurements in ROI of 20 µm sections of the Z-stack consisting of the raw data from the ZEN 2010 acquisition software. The level of the tissue surface was set manually from the XY sections and values above the tissue surface in track 1 were considered background.

The mucus penetrability was analyzed in two ways. We first determined how large proportion of the total bead intensity that was in close proximity to the epithelial surface. In mouse colon the analysis window was set -20 to +40 µm above the epithelial surface. In human colon the analysis window was set -20 to +120 µm. The total bead intensity in the region was compared to the total bead intensity in the entire Z-stack. In the second analysis we determined how large part of the mucus that was penetrated by the beads. The distance between the outer border of the beads and the epithelial surface was compared to the distance between the inner most beads and the epithelial surface. The lowest point of beads was
defined as the section where the bead intensity was <5% of the maximum bead intensity, representing background fluorescence.

**Statistical analysis**
Non-parametric tests were chosen as normal distribution could not be verified due to the small sample sizes. Mouse data was analyzed using a two-tailed Mann-Whitney U test. For the human data the Kruskal-Wallis test with Dunns’ correction for multiple comparisons was used. A p-value <0.05 was regarded as statistically significant. Mean of each analysis per individual was used as value for the statistical analysis.

**REFERENCES**


**Table S1:** Clinical information about the control patients.

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*M: male, F: Female*
Figure S1. Bacteria penetration and inflammation in the different colitis models. (A) Bacteria penetration of the inner mucus was scored for all the genotypes (n=5 for WT and IL10−/− and n=3 for all other groups). Data is presented as mean±SEM. The WT data is representative for controls from different animal facilities. (B) Inflammation was monitored as histology scores for the different genotypes (n=5 for WT and IL10−/− and n=3 for all other groups). Data is presented as mean±SEM.