Loss of intestinal core 1–derived O-glycans causes spontaneous colitis in mice

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Mucin-type O-linked oligosaccharides (O-glycans) are primary components of the intestinal mucins that form the mucus gel layer overlying the gut epithelium. Impaired expression of intestinal O-glycans has been observed in patients with ulcerative colitis (UC), but its role in the etiology of this disease is unknown. Here, we report that mice with intestinal epithelial cell–specific deficiency of core 1–derived O-glycans, the predominant form of O-glycans, developed spontaneous colitis that resembled human UC, including massive myeloid infiltrates and crypt abscesses. The colitis manifested in these mice was also characterized by TNF-producing myeloid infiltrates in colon mucosa in the absence of lymphocytes, supporting an essential role for myeloid cells in colitis initiation. Furthermore, induced deletion of intestinal core 1–derived O-glycans caused spontaneous colitis in adult mice. These data indicate a causal role for the loss of core 1–derived O-glycans in colitis. Finally, we detected a biosynthetic intermediate typically exposed in the absence of core 1 O-glycan, Tn antigen, in the colon epithelium of a subset of UC patients. Somatic mutations in the X-linked gene that encodes core 1 β1,3-galactosyltransferase–specific chaperone 1 (C1GALT1C1, also known as Cosmc), which is essential for core 1 O-glycosylation, were found in Tn-positive epithelia. These data suggest what we believe to be a new molecular mechanism for the pathogenesis of UC.

Introduction

Ulcerative colitis (UC) is an immune-mediated disorder that results from an abnormal interaction between colonic bacteria and mucosal immune cells in a genetically susceptible host (1, 2). The mechanisms underlying this interaction remain to be defined (1, 2).

Intestinal bacterial antigens are normally shielded from encountering mucosal immune cells by the intestinal barrier, which comprises an epithelial layer and a mucus layer (3, 4). The function of the intestinal epithelial cells, limiting antigens from entering the tissues while simultaneously allowing the immune system to sample antigens in the gut, has been extensively studied (2, 5–7). The colon mucus has two layers: an outer layer, where the bacteria sample antigens in the gut, has been extensively studied (2, 5–7). The colon mucus layer comprises the polymerized mucins, primarily Muc2, that are produced by goblet cells (4, 9). Mucins are glycoproteins that carry large numbers of O-linked oligosaccharides (O-glycans), which account for up to 80% of the mass of the mucin molecules and are responsible for many of the properties of mucins. O-glycans are synthesized post-translationally in the Golgi apparatus (10–12). All O-glycans are initiated with a primary structure referred to as Tn antigen (GalNAcα-O-Ser/Thr; Figure 1A), which is normally masked by additional glycosylation to form the main type of O-glycans, core 1–derived structures (12). The biosynthesis of core 1 is controlled by core 1 β1,3-galactosyltransferase (C1gal1, also known as T-synthase) (12), for which expression requires the specific molecular chaperone C1GALT1C1 in the ER (13).

Most UC patients have colitis only in the distal colon (14). Although the reasons for this regional variation are unknown, UC patients have deterioration of the mucus layer and abnormal mucin expression in the distal colon (1, 15–17). Altered intestinal O-glycosylation also occurs in patients with UC. However, the nature of the impaired O-glycosylation in UC patients is unclear (18). Whether normal O-glycosylation impairs the mucus inner layer and causes spontaneous colitis is unknown.

Here, we report that mice with intestinal epithelial cell–specific deficiency of core 1–derived O-glycans exhibited a breached inner mucus layer and developed spontaneous colitis primarily in the distal colon, which is similar to human UC. These results indicate an etiological role for impaired O-glycosylation in the pathogenesis of colitis. Notably, we discovered that Tn antigen was expressed in colon biopsy samples from a subset of UC patients and some of the Tn-expressing epithelia contained C1GALT1C1 missense mutations. These findings support a molecular mechanism for colitis development in a subset of UC patients whereby core 1 O-glycosylation is altered through somatic mutations in C1GALT1C1.

Results

Mice with targeted deletion of C1gal1 in intestinal epithelial cells do not express O-glycans and develop spontaneous colitis. Core 1–derived O-glycans are expressed in many tissues, including the intestine. Global

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deficiency of mouse C1galt1 causes embryonic lethality (12). To
determine the role of O-glycans in intestinal epithelium, we gener-
amized mice lacking C1galt1 specifically in the intestinal epithelium
by crossing mice with loxP sites flanking C1galt1 (C1galt1fl/fl) mice
(11) with an intestinal epithelium–specific Cre-expressing trans-
genic line (VillinCre mice) (19). The resultant mice are referred to
as intestinal epithelial cell–specific C1galt1−/− (IEC C1galt1−/−) mice
(Figure 1B). Antibody to Tn antigen (anti-Tn) detected Tn antigen
in the IEC C1galt1−/− and small and large intestinal epithelium but
not in other cell types or in WT intestinal epithelium (Figure 1C),
confirming the specificity of the deletion. Desialylation did not appre-
ciably affect the intensity of anti-Tn staining (data not shown),
suggesting that most exposed Tn antigens are not sialylated.

O-glycan structural analysis of the mucins purified from colon
mucus demonstrated that WT mucins contained core 1–derived
O-glycans such as fucosylated core 1, core 2 (Galβ1-3[GalNAcβ1-6]
GalNAc), and sialylated core 2 at m/z of 530, 587, and 766, respec-
tively, but these structures were absent in IEC C1galt1−/− mucins
(Supplemental Figure 1; supplemental material available online
with this article; doi:10.1172/JCI45538DS1), thus verifying that
deletion of C1galt1 abolishes core 1–derived O-glycans. Core 3–
derived O-glycans were detected in IEC C1galt1−/− and to a lesser
extent in WT colon samples (Supplemental Figure 1), which may
suggest a relative compensatory increase in core 3 O-glycosylation
in the absence of core 1 O-glycosylation.

As early as 2.5 weeks after birth, IEC C1galt1−/− mice in the C57BL/6j
and 129/SvImJ mixed or C57BL/6j congenic background that
were raised in a specific pathogen–free facility (no detectable Helicobacter) began showing signs of colitis, including loose stools and
fecal occult blood (Figure 1D). The disease progressed for approximately 2 weeks and then improved, suggesting a potential role for regulatory T cell function at this stage. At 8 weeks, approximately
20% of IEC C1galt1−/− mice had diarrhea, and by 12 weeks, all IEC
C1galt1−/− mice had diarrhea. Weight loss was evident by 8 weeks in
IEC C1galt1−/− mice. Gross and microscopic examination revealed
no significant abnormalities in the liver, stomach, spleen, and thy-
mus; however, nearly 15% of IEC C1galt1−/− mice had rectal prolapse
progressing over time (Figure 1E), a feature associated with murine
colitis. The colons of IEC C1galt1−/− mice, especially their distal
colons and rectums, had dilated and thickened walls (Figure 1F). In
fact, in all mice this region was diseased and was the most severely
affected; it was characterized by epithelial ulceration, inflamma-
tory cell infiltration, goblet cell loss, epithelial hyperplasia, and,
frequently, crypt microabscesses (Figure 1G). Although VillinCre
deletes C1galt1 in both small and large intestine, no significant
inflammation was found in the ileum (Supplemental Figure 2).
Colon has a thick, firmly attached inner and an outer loose mucus
layer, whereas the small intestine only has the loose mucus layer (4).
These results suggest the importance of mucus layer in maintain-
ing the homeostasis between mucosal cells and commensal bacte-
ria, which are at their greatest concentrations in the colon.

Adaptive immunity is not required for the initiation of colitis in IEC
C1galt1−/− mice. To identify the cell type that initiates colon inflam-
mation, we first compared various immune cell subsets in different
immune compartments between WT and IEC C1galt1−/− mice. The
profiles of lymphocytes and NK cells of 2.5-week-old IEC C1galt1−/−
mice were similar to those of WT littermates (Supplemental Fig-
ure 3, A and B). The number of F4/80-positive macrophages was,
Adaptive immunity is not required for the initiation of colitis in IEC C1galt1−/− mice. (A) Representative H&E-stained colonic sections indicate similar colonic inflammation in IEC C1galt1−/− mice with and without Rag1 deficiency at 2 weeks. (B) Clinical DAI (mean ± SD, n = 10 mice/group) of IEC C1galt1−/− mice based on diarrhea, fecal occult blood, and rectal prolapse. *P < 0.05. (C) Cryosections of IEC C1galt1−/−Rag1−/− colon stained with mAbs to myeloid cells (Gr1) and TNF at 3 weeks. Arrows mark infiltrates in epithelium. Inset (original magnification, ×400) highlights TNF-positive myeloid cells. Scale bars: 100 μm in A, 50 μm in C. Data are representative of at least 3 experiments.

However, significantly higher in IEC C1galt1−/− colon tissues (Supplemental Figure 3B). These results suggest that lymphocytes and NK cells are not important in colitis development in this model (2, 20). To provide definitive evidence that lymphocytes are not required for the initiation of colitis in IEC C1galt1−/− mice, we bred IEC C1galt1−/− mice with Rag1-deficient mice (Rag1−/− mice) (21), which lack lymphocytes. IEC C1galt1−/−Rag1−/− mice developed colitis similar to that of their IEC C1galt1−/− littermates (Figure 2, A and B), indicating a dispensable role for lymphocytes for the colitis initiation in our model. The transient alleviation of disease severity was not observed in IEC C1galt1−/−Rag1−/− mice around 6 weeks old (Figure 2B), suggesting a lack of a potential Rag-dependent T regulatory cell function in this model. At an older age (>16 weeks), the disease was less severe in IEC C1galt1−/−Rag1−/− mice than in IEC C1galt1−/− littermates (Figure 1D and Figure 2B). This result suggests that lymphocytes are important for the progression of colitis at a later stage in IEC C1galt1−/− mice.

To confirm the role of myeloid cells in colitis development, we stained 3-week-old WT and IEC C1galt1−/−Rag1−/− colons with antibodies to Gr1, an established marker for myeloid cells, and to TNF, a known inflammatory mediator in the pathogenesis of colitis. The 3-week-old IEC C1galt1−/−Rag1−/− colon exhibited large numbers of TNF/Gr1+ myeloid cells (Figure 2C). Consistent with this, 2.5-week-old IEC C1galt1−/−Rag1−/− sera had significantly increased levels of cytokines such as TNF, IL-6, and MCP-1 compared with WT sera (Supplemental Figure 3C), suggesting an innate immunity–driven inflammatory response.

Myeloid cells are essential for the initiation of colitis in IEC C1galt1−/− mice. To establish the role of myeloid cells in incipient disease, we first analyzed peripheral granulocytes and monocytes in IEC C1galt1−/− mice by flow cytometry. Notably, peripheral blood from IEC C1galt1−/− mice had elevated numbers of peripheral granulocytes (Ly6G+CD11b+) at 2 weeks and had a higher level of the inflammatory subset of monocytes (Ly6G−CD11b−Ly6C−) at 3 weeks (Figure 3A and Supplemental Figure 4). Blockage of P- and E-selectins, which inhibit myeloid cell transmigration into inflammatory sites (22), or of TNF significantly improved colitis in IEC C1galt1−/− mice as compared with IEC C1galt1−/− mice treated with control agents (Figure 3B), indicating that myeloid cells and TNF are key inflammatory initiators in our mouse model.

Further immunostaining revealed substantial infiltration of TNF-producing granulocytes and monocytes/macrophages in IEC C1galt1−/− colon tissues (2.5-week-old; Figure 3C) but not before the onset of colitis (1 week of age), suggesting that these are major cell types that sense the early microbial intrusion and initiate inflammation. We bred IEC C1galt1−/− with mice lacking Myd88, a universal adaptor protein used by most TLRs, or TLR4, which recognizes bacterial antigens, and found that neither Myd88 nor TLR4 deficiency protected IEC C1galt1−/− mice from developing colitis (Supplemental Figure 5), suggesting that TLRs are not essential for recognition of bacterial components in our mouse models.

Loss of intestinal O-glycans results in a breached inner mucus gel layer and defective mucosal integrity. Staining with Alcian blue, which recognizes acidic carbohydrates, revealed that 2.5-week-old IEC C1galt1−/− colon sections had a loss of the inner mucus gel layer (Figure 4A). Consistent with this finding, anti–Muc2–stained inner mucus layer was substantially impaired in the IEC C1galt1−/− mice as compared with WT mice (Figure 4B and ref. 23). Consequently, a significant increase in 16S bacterial ribosomal DNA (rDNA) was found in IEC C1galt1−/− as compared with WT colon (Figure 4C and ref. 23), indicating that impaired mucosal integrity increases in mucosa-associated bacteria in IEC C1galt1−/− mice.

Resident intestinal bacteria are assumed to play an important role in IBD pathogenesis (3). We found that broad-spectrum antibiotics (neomycin sulfate, metronidazole, vancomycin, and ampicillin [NMVA]) depleted gut aerobic and anaerobic bacteria (Supplemental Figure 6) and reduced colitis in IEC C1galt1−/− mice (Figure 4D), supporting a requirement for intestinal bacteria in the development of colitis in our models.

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Inducible deficiency of O-glycans in the intestinal epithelium causes spontaneous colitis in adult mice. In IEC C1galt1/− mice, the inactivation of C1galt1 occurs during embryonic development, eliciting secondary host compensation, such as changes in maternal milk immunoglobulins and intestinal commensal microbiota (3). Accordingly, the colitis phenotypes of IEC C1galt1/− neonates could reflect these secondary changes. To circumvent these potential problems and to determine whether loss of intestinal epithelial core 1–derived O-glycans in adult mice causes colitis, we generated mice with an inducible deficiency of intestinal epithelial O-glycans by crossing C1galt1+/− mice with VillinCre-ER² transgenic mice (ref. 24 and Figure 5A), in which an inert form of Cre can be activated in intestinal epithelium by administration of tamoxifen (TM). Specific and efficient inducible deletion of O-glycans in intestinal epithelium was achieved 2 days after 5 consecutive daily intraperitoneal injections of TM in 7-week-old C1galt1+/−/VillinCre-ER² (TM-IEC C1galt1/−) mice, as indicated by anti-Tn (Figure 5B). The TM-IEC C1galt1/− colon started to lose Alcian blue–stained glycans 5 days after TM induction, with a complete loss of the staining in the mucus layer and goblet cells by 10 days (Figure 5B). In TM-IEC C1galt1/− colon, the thickness of the mucus gel layer was significantly reduced by 5 days and was lost by 10 days (Figure 5, B and C). Significantly, the TM-IEC C1galt1/− colon began to show signs of colitis, such as inflammatory infiltrates, mucosal proliferation, and loss of goblet cells by 5 days after TM, and developed full-fledged colitis 10 days after TM induction, with progressive deterioration thereafter (Figure 5, B, D, and E). Like disease in IEC C1galt1/− mice, colitis in TM-IEC C1galt1/− mice was dependent on intestinal bacteria (Figure 5D). Thus, the loss of O-glycans directly caused spontaneous colitis.

Loss of intestinal O-glycans results in a defective mucus barrier. TM-IEC C1galt1/− mice represent an ideal model to address how the lack of core 1–derived O-glycans causes colitis. Loss of O-glycans did not cause significant changes in the miRNA levels of Muc1, Muc2, Muc3 (17), Muc13, and MucSat in WT and mutant colons (Supplemental Figure 7). However, the staining of Muc2, the major secreted mucin, was reduced in IEC C1galt1/− colons 5 days after TM induction (Figure 6A). Compared with WT colons, TM-IEC C1galt1/− colons had significantly increased intestinal permeability and mucosa-associated bacteria as determined by FITC-dextran feeding and 16S-rDNA PCR, respectively, 5 days after TM induction (Figure 6B and C). 16S rDNA in feces remained unaltered at different time points after TM, suggesting that the lack of O-glycans did not cause appreciable changes in the density of commensal bacteria based on PCR. FISH revealed that WT colon epithelium was covered by an intact inner layer of mucus that was devoid of bacteria. In contrast, TM-IEC C1galt1/− epithelium had no mucus layer and directly contacted commensal bacteria (Figure 6D). These results suggest that loss of O-glycans causes an increased association of intestinal bacteria with TM-IEC C1galt1/− colonic epithelia before...
full-fledged colitis occurs. Future studies of bacterial variety and density in colonic microbiota before and after colitis onset in this model are necessary to determine whether the composition of the resident bacterial population is altered.

Similar to the infiltration found in IEC C1galt1−/− mice (Figure 3C), TNF-producing myeloid cells were observed in the TM-IEC C1galt1−/− colon at the onset of colitis (5 days after TM) and became more prominent over time (Figure 6E). TNF induces tight junction molecule endocytosis and the loss of tight junction barriers (25). We observed increased internalization of the tight junction molecule occludin in TM-IEC C1galt1−/− but not in the WT colonoocytes 5 days after TM (Supplemental Figure 8), suggesting that the observed barrier loss at this time point is secondary to this cellular effect of TNF on epithelial tight junctions in the TM-IEC C1galt1−/− colon.

The unfolded protein response (UPR) has been genetically implicated in human IBD (26). Unlike N-linked glycosylation, which regulates protein folding in the ER, however, O-glycosylation occurs exclusively in the Golgi apparatus and is not known to regulate protein trafficking in the ER (27, 28). Nevertheless, we examined whether UPR contributed to the colitis in O-glycan-deficient mouse model. Our results demonstrated that TM-IEC C1galt1−/− colon epithelium did not have significant accumulation of the non–O-glycosylated Muc2 precursor in the ER (Supplemental Figure 9A). RT-PCR did not detect a spliced form of XBP-1, which is the activated form of the ER stress response transcription factor (29), in the inflamed TM-IEC C1galt1−/− colon 15 days after TM (Supplemental Figure 9B). These results indicate that an abnormal O-glycosylation–related UPR response does not occur, or at least is of such small scale that it does not induce a detectable biologic response in our model.

Colon epithelial cells from patients with UC express Tn antigen, and a subset has mutated C1GALT1C1. To investigate the clinical relevance of our O-glycan–deficient colitis mouse models, we stained archived paraffin-embedded colon biopsy samples from UC patients with anti-Tn. Epithelia in 8 of the 46 UC patients were Tn positive (Figure 7, A and B), whereas all 16 non-IBD samples were negative. Approximately 30% of the UC biopsy colon tissue paraffin blocks for the Tn antigen staining contained little epithelium as a result of severe inflammation/ulceration; therefore, this rate of Tn detection may be an underestimate. Tn-positive and -negative crypts often occurred within a single colon section, suggesting a clonal abnormality of O-glycosylation. Defective core 1 or core 3 O-glycosylation could lead to the exposure of Tn (12, 23). Immunohistochemical staining demonstrated that the expression of core 3 β1,3-N-acetylgalactosaminyltransferase in UC colons was comparable to that in normal control colons (Supplemental Figure 10), suggesting defects in pathways controlling the core 1 O-glycosylation. Recently, somatic mutations in X-linked CIGALT1C1, which is essential for core 1 O-glycosylation (Figure 1A), were reported (30, 31). We identified 4 missense mutations in the coding sequence of CIGALT1C1 from genomic DNA extracted from Tn-positive epithelia (Supplemental Figure 11, A–C). Sequencing of PCR products amplified from independently collected Tn-positive crypts from the same patient produced the same result, indicating that the mutations were not artifacts. Site-directed mutagenesis analysis indicated that these mutations resulted in impaired function of C1GALT1C1 (Supplemental Fig-
Although large-scale high-throughput genetic analyses with sufficient power are necessary to make definitive conclusions, the observations suggest that loss-of-function mutations in C1GALT1 contribute to aberrant O-glycosylation in some UC patients (Figure 7 and Supplemental Figure 11).

**Discussion**

Our results demonstrate that loss of intestinal O-glycans causes reduced levels of mucins; a breached colon mucus inner barrier, leading to increased interaction of intestinal bacteria with epithelia; and subsequently spontaneous colitis in mice (Figure 7C). This study highlights the important contribution of O-glycans, a primary component of mouse colon mucus, to intestinal barrier function and to the pathogenesis of UC.

Mucin synthesis and secretion decrease in active UC (15, 32, 33). The relevance of Muc2 and its O-glycan decoration is validated by colitis susceptibility of mice bearing genetic deficiencies in these traits (23, 34, 35). Mechanistically, their contribution to colitis sus-
ceptibility is due at least in part to the crucial role of these structures as physical barriers to microbial intrusion, and by providing selective glycans for cognate receptors specific to restricted microbial phylotype (36–38). Few changes in mucin gene expression or polymorphisms have, however, been detected in IBD, except for the changes found in Muc3A (2, 39). Therefore, the cause of the abnormal mucin expression in UC remains to be defined. The IBD-associated variants of XBP1 and ATG16L1, which are involved in either ER stress response or autophagocytosis, may cause secondary mucin dysfunction (26, 40). However, the causes of abnormal mucin expression and whether these alterations are causative in human UC are largely unknown. Our data indicate that O-glycans are essential for the expression of both secreted and membrane-associated mucins and for the integrity of the mucus layers, thus underscoring the importance of altered O-glycosylation as a causal factor in the pathogenesis of colitis. Loss of O-glycans may impair the expression and function of many intestinal mucins, thereby causing more profound defects in intestinal barrier function than an abnormality caused by the deficiency of an individual mucin. Indeed, compared with Muc2-deficient mice, which develop colitis only in a 129/Sv background (34), O-glycan–deficient mice develop more severe spontaneous colitis on either C57BL/6 and 129SV mixed or C57BL/6 congenic background.

The nature of the impaired O-glycosylation in patients with UC is unclear (18). Our studies provide the first demonstration to our knowledge that some UC colons express Tn antigen, underscoring the clinical relevance of the O-glycan–deficient colitis models. Significantly, our pilot study identified missense somatic mutations in C1GALT1C1 in Tn-positive epithelial cells from UC patients. Mutations in C1GALT1C1 are reported in patients with several diseases, including colorectal tumor cells that express Tn antigens (30, 31). These promising results suggest that altered O-glycosylation causes impaired mucus barrier function in a subset of UC patients and may serve as an etiological factor for UC. Notably, only one of the genes regulating glycosylation has been identified as relevant to known IBD loci (2, 41). Glycosylation pathways are, however, complex (42), such as the indirect regulation of C1gal1 activity by C1GALT1C1. More-focused genetic studies will be needed to determine whether mutations in genes controlling O-glycosylation are commonly associated with UC.
Activation and recruitment of myeloid cells in colon mucosa are hallmarks of active UC and are associated with epithelial injury and clinical disease activity (1). However, the significance of myeloid cells in the initiation of colitis and in disease progression remains unclear. Our data indicate that myeloid cells are among the major early responders to bacterial intrusion when the mucosal barrier is breached and contribute significantly to tissue damage in the C1galt1−/− colon. In general, microbial recognition is achieved through pattern receptors, such as TLR and Nod-like receptors (NLRs) (1, 2, 43). The activation of myeloid cells in our models is Myd88 independent, which is similar to that in T-bet−/− mice that exhibit colitis in the background of Rag1 deficiency (44), although, unlike in T-bet−/− mice, spontaneous colitis in C1galt1−/− mice is not dependent on the Rag1 deficiency. The underlying mechanism of colitis susceptibility in T-bet−/− mice includes an enteric dysbiosis resulting from their host genetic deficiency in mucosal defense, with resultant recruitment and activation of proinflammatory myeloid cells. This is different from the mechanism in C1galt1−/− mice. In C1galt1−/− mice, increased association of intestinal bacteria with colonic epithelia may result from impaired ability of the O-glycan–dependent mucus barrier to impede nonspecific bacterial invasion. Recent work has demonstrated that some symbiotic bacteria have developed O-glycan sensing and degradation pathways (45). Based on this, it can be anticipated that the changes in O-glycans will alter the composition of the gut microbiota. The ability of bacteria to penetrate the mucus barrier may also contribute to the pathogenesis of colitis in our models. Nevertheless, in both models, Myd88 independence is an surprising feature, suggesting that the relevant microbial sensing in this setting of innate inflammation predominates with Myd88-independent TLR, NLR, or other families of pathogen-associated molecular products (1, 43).

Methods

Details regarding O-glycan structural analysis, detection of expression of mucin genes and XBPI by RT-PCR, detection of non–O-glycosylated Muc2 precursor in the ER, anti-C3GalT1 staining, and C1GALT1C1 mutation analysis are provided in Supplemental Methods.

Mice. IEC C1galt1−/− mice and TM-IEC C1galt1−/− mice were generated by breeding C1galt1+/− mice with VillinCre transgenic mice (Tg[Vil-cre]997Gum; Jackson Laboratory) and VillinCre-ER(Tg) transgenic mice (provided by Sylvie Robine, CNRS-Institut Curie, Paris, France, and Robert Coffey, Vanderbilt University, Nashville, Tennessee, USA) (11, 19, 24, 47), respectively. To induce a postnatal deficiency in intestinal O-glycans, 6- to 8-week-old TM-IEC C1galt1−/− mice were injected intraperitoneally with 1 mg TM (MP Biomedicals) in an ethanol/sunflower seed oil mixture (1:9 [v/v]) for 5 consecutive days. Rag1−/− mice (Jackson Laboratory), Tlr4−/− mice, and Myd88−/− mice (originally developed by Shizuo Akira, Osaka University, Osaka, Japan) (48) were crossed to IEC C1galt1−/− mice to establish mice with combined gene deficiencies. The animal studies were conducted with protocols that had been approved by the Institutional Animal Care and Use Committee of the Oklahoma Medical Research Foundation and UCLA. Mice were raised in a specific pathogen–free barrier facility and genotyped routinely by PCR assay on genomic DNA isolated from tail clips. The primers used for genotyping include Flox1: 5′-TGACAGCCAGGAATGGAACTTG-3′ and Flox2: 5′-GCCCTTTCCTGCCAACAAATATCCTC-3′; CRE1: 5′-AGGTGATAGAGAAGCTATAG-3′, and CRE2: 5′-CTAATCGCCATCTTCCAGCAGG-3′. Sex- and age-matched littermates were used as controls in all experiments. Unless specified, all mice were on the C57BL/6J congenic background.

Histology. For histology, tissues were freshly harvested from mice and fixed with 10% neutral buffered formalin or in Carnoy’s fixative (60% meth-

Figure 7

Colon epithelium of a subset of UC patients expresses Tn antigen. Representative immunohistochemical staining of colon biopsy samples from a normal control (A) and UC patient (B). Arrows indicate Tn-positive crypts. Arrowhead marks a Tn-negative crypt. Scale bars: 50 μm. (C) Model illustrating that loss of intestinal core 1–derived O-glycans results in an impaired mucus barrier, allowing intestinal microflora to interact abnormally with the epithelium and mucosal immune cells, which causes myeloid cell–initiated colitis that is eventually amplified and/or diversified by other innate cells and lymphocytes.
anol, 30% chloroform, and 10% acetic acid) prior to paraffin embedding. Paraffin-embedded sections (5 μm) were stained with H&E. The pictures were obtained with an Eclipse E1000 microscope (Nikon). Histologic severity of colitis was assessed in a blinded fashion (23).

Evaluation of clinical activity of colitis. Mice were weighed biweekly and scored for colitis-associated symptoms, including stool consistency (0, normal; 2, loose stools; 4, diarrhea), presence of fecal blood (occult fecal blood: 0, negative; 2, positive), and rectal prolapse (0, normal; 2, present sometimes; 4, present always) (23).

Immunohistochemical and immunofluorescence staining. For Tn antigen staining, deparaffinized sections were incubated with or without 0.5 U/ml sialidase from Arthrobacter ureafaciens (Roche) at 37 °C for 3 hours. Sections were incubated for 30 minutes with biotinylated anti-Tn (mouse IgM, provided by Tongzhong Ju and Richard Cummings, Emory University, Atlanta, Georgia, USA) or with isotype-matched control mouse IgM. Bound antibodies were detected with horseradish peroxidase–conjugated streptavidin (Vector Laboratories) (11, 23).

For immunohistochemical detection of Muc2 and Muc3, deparaffinized sections were boiled for 20 minutes in 0.01 M citrate buffer at pH 6.0 for epitope retrieval. The sections were then blocked for 20 minutes using a protein blocking kit (Dako). Endogenous avidin-binding proteins were blocked using avidin/biotin blocking reagents (Vector Laboratories). Sections were incubated overnight at 4 °C with rabbit anti-Muc2 (1:200 dilution; Santa Cruz Biotechnology Inc.; epitope corresponding to amino acids 4,880–5,179, which maps to the C-terminus of mucin 2 of human origin), rabbit anti–muc3(17)-S2, which is glycosylation independent (raised against the peptide KYTPGFENTLDTVVKNLETKIKNAT), or control IgG. The sections were subsequently incubated with biotinylated anti-rabbit (Vector Laboratories) for 45 minutes, followed by a 20-minute incubation with 0.6% H2O2 in methanol to inhibit endogenous peroxidase activity. The sections were finally incubated with horseradish peroxidase–streptavidin (Vector Laboratories) for 30 minutes and developed with a diaminobenzidine substrate and counterstained with hematoxylin.

For the detection of infiltrated inflammatory cells, distal colons were dissected from mice and fixed in 4% paraformaldehyde at 4°C overnight, followed by cryoprotection in 20% sucrose, and then embedded in a mixture of OCT compound (Sakura Finetek) and tissue freezing medium (Electron Microscopy Sciences). Cryosections (20 μm) were stained with H&E. The pictures were obtained with a Nikon C1 confocal laser-scanning unit equipped with a 3-laser launcher (Nikon). Histologic severity of colitis was assessed in a blinded fashion (23).

To deplete the commensal microflora in C1galt1−/− mice, and their controls were fed 200 μl of FITC-dextran at 600 mg/kg body weight (molecular weight, 4 kDa; Sigma-Aldrich) by gavage. Blood was collected 4 hours later by retro-orbital bleeding. The serum concentration of the FITC-dextran (excitation, 490 nm; emission, 530 nm) was determined using an Infinicon M200 fluorometer (Tecan). Serial diluted FITC-dextran was used to generate a standard curve. Animals were sacrificed immediately after bleeding, and intestinal cryosections were prepared for fluorescence microscopy.

Bacterial translocation. Bacterial translocation in IEC C1galt1−/− mice was evaluated with real-time PCR (23). Two sets of universal primers specific for the conserved regions of the bacterial 16S rDNA gene were used. These included 16S1 (forward, 5′-CCATGAATCGGAATCTGCTAG-3′; reverse, 5′-ACTCCTCATGTTGTGACGG-3′) and 16S2 (forward, 5′-TCCTAGGGAGGCACGAGT-3′; reverse, 5′-GGACTACGGGATTCTAATCTCGTTT-3′). DNA extracted from colonic tissues and colonic fecal materials was used as a template. Colonic tissues were washed carefully before extraction of DNA to remove residual feces. For the quantification of 16S rDNA in colonic tissues, a pair of primers specific to the mouse Selp gene (forward, 5′-ATGATTAGAAATTTCACTGCTGTGT7-3′; reverse, 5′-TAGGTCTCTTAGATCTCCCTTTCAAT-3′) was used in a separate reaction as the endogenous control to normalize the DNA loading between samples. For the quantification of fecal bacterial 16S rDNA, DNA from each mouse fecal sample was quantified, and 25 ng DNA was used as a template in each PCR reaction. Real-time PCR was performed on an ABI Prism 7000 spectrophuorometric thermal cycler (Applied Biosystems) using SYBR green as a double-stranded DNA–specific binding dye. The relative amount of 16S rDNA in each sample was estimated using the ΔΔCt method (23). Each sample was assayed in duplicate.

**FISH.** FISH was performed as described previously (8). Briefly, colon sections fixed with Carnoy’s fixative were incubated with 250 μg Alexa Fluor 555–conjugated EUB (5′-GCTGCCCTCCGCTAGGATG-3′; bp 337–354 in bacteria EU822773) in 50 μl hybridization buffer (20 mM Tris-HCl, pH 7.4; 0.9 M NaCl, 0.1% SDS) at 50 °C overnight. The sections were rinsed in wash buffer (20 mM Tris-HCl, pH 7.4; 0.9 M NaCl) and washed at 50 °C for 15 minutes. Co-immunosstaining with glycosylation-independent anti-MUC2 (C2) (8) was performed at 4 °C without antigen retrieval to preserve the in situ hybridization, and sections were mounted in ProLong Gold antifade (Invitrogen). Pictures were obtained with a Radiance 2000 (Bio-Rad) confocal microscope.

Marine P- and E-selectin and TNF blockade. mAbs against murine P-selectin (SH1, rat IgG1; 50 μg/mouse) and E-selectin (9A9, rat IgG2b; 50 μg/mouse), etanercept (TNF blocker, 5 mg/kg body weight; Amgen), or irrelevant control mAbs were injected intraperitoneally twice a week for 2 weeks, starting 1 week before the start of the UC patient sample study was approved by the Institutional Review Boards of the Oklahoma Medical Research Foundation and the University of Oklahoma Health Science Center. De-identified, archived paraffin sections of human UC colon biopsy samples were first stained for expression of Tn antigens using anti-Tn antibody. Deparaffinized sections were incubated with biotinylated mAb against Tn (mouse IgM), after streptavidin/biotin and nonspecific protein blocking. The sections were then incubated with HRP-streptavidin (Vector Laboratories) for 30 minutes, and then developed with a DAB kit (Vector Laboratories) and counterstained with hematoxylin.

**Statistics.** All data are presented as the mean ± SEM, unless otherwise indicated. Unpaired 2-tailed Student’s t test was used to analyze the difference between 2 groups. A P value less than 0.05 was considered to be significant.

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