Intestinal barrier regulates immune responses in the liver via IL-10–producing macrophages

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Introduction

Under physiological conditions, the liver is continuously exposed to gut-derived antigens through the portal vein, and bacterial degradation of metabolites fermented by commensal bacteria reach the liver and influence its innate and adaptive immune responses (1–3). Intestinal inflammation can lead to a leaky gut, and live bacteria and bacterial products migrate from the gastrointestinal tract to portal or systemic circulation following intestinal barrier disruption (4), after which they can activate immune cells in the liver. Liver tolerance is a well-known organ-specific phenomenon possibly caused by its unique anatomical location. That immune responses in the liver trend toward immune permissiveness was first recognized in a porcine orthotopic liver transplantation model (5). The liver is indispensable to the maintenance of oral tolerance against nutrients, gut-derived bacterial metabolites, LPS from the cell walls of gram-negative bacteria, and cellular debris that enters the liver via the portal vein from the intestine. In general, the presentation of antigens by liver antigen-presenting cells promotes initial proliferation, followed by clonal exhaustion or apoptosis of T cells (6), resulting in the maintenance of immunological balance under homeostatic conditions. Recently, gut microbiota has been reported to both help and overcome liver tolerance and contribute to the pathogenesis of liver diseases (7–10). Clinically, primary sclerosing cholangitis (PSC) is a well-documented example of chronic hepatobiliary disease highly associated with inflammatory bowel disease (IBD) (11), indicating that the gut-liver axis plays a role on the pathogenesis (7, 12). These phenomena suggest that increased intestinal permeability with continuous exposure to gut-microbial products might influence immunological responses in the liver. However, the underlying mechanisms by which subsets of immune cells contribute to immunolog-
ical activation and tolerance in the liver remain unclear.

In this study, we demonstrated that preexisting intestinal barrier damage induced by dextran sulfate sodium (DSS) unexpectedly ameliorated the severity of subsequent immune-mediated acute liver injury models induced by concanavalin A (Con A). Using a tandem model of DSS colitis and Con A hepatitis, we explored the roles of the gut-liver axis in hepatic immune regulation, demonstrating that colitis prior to hepatitis preconditioned hepatic CD11b+ F4/80+ innate immune cells as IL-10–producing regulatory cells in vivo. Furthermore, the amelioration of Con A hepatitis was canceled by gut sterilization, suggesting that immune balance in the liver is potentially regulated by gut microbiota and metabolites.

**Results**

Severe mucosal barrier degradation leads to significantly milder liver injury following a sublethal injection of Con A. To clarify whether excessive gut-derived antigen stimulation affects immune responses in the liver, mice were orally administered DSS to induce mucosal barrier degradation, followed by Con A administration to induce acute liver injury. Mice were either untreated or orally administered 2.0% DSS for 7 days, followed by i.v. PBS or Con A (20 mg/kg) exposure, in distilled water–PBS (DW–PBS), DSS–PBS, DW–Con A, or DSS–Con A experimental groups. DSS–treated mice developed substantial colitis, as demonstrated by body weight loss and disease activity index (DAI) (Figure 1A). Interestingly, DSS–treated mice exhibited significantly milder liver injury, as determined by serology and histology of H&E staining and TUNEL assay to detect apoptosis, than did untreated mice following a sublethal dose of Con A (Figure 1B–D). As shown previously, Con A administration upregulated the mRNA expression of proinflammatory cytokines — such as TNF-α and IL-6 — in the liver, whereas their expression was significantly decreased in DSS–Con A mice (Figure 1E).

To examine the relationship between the severity of colitis and subsequent liver injury, mice were pretreated with DSS for 3–7 days. We observed that the severity of DSS-induced colitis was correlated with the duration of DSS administration (Figure 1F). Although mice pretreated with DSS for 3 days exhibited more severe liver injury, pretreatment with DSS for 7 days significantly attenuated subsequent liver injury (Figure 1G). Furthermore, mice that recovered from DSS colitis and that had received DW for 3 weeks after DSS treatment developed more severe hepatitis following Con A administration (DSS–DW–Con A) than did DSS–Con A mice (Supplemental Figure 1A, A and B; supplemental material available online with this article; https://doi.org/10.1172/jci.insight.91980DS1), suggesting that the severity of colitis directly affected susceptibility to subsequent liver injury in this model. Similarly, 2,4,6-trinitro-benzene sulfonic acid–treated (TNBS-treated) mice, another murine model of colitis, exhibited less susceptibility to Con A–induced acute liver injury. Additionally, DSS–treated mice exhibited less susceptibility to α-galactosylceramide–induced (α-Galcer–induced) acute liver injury, another murine model of immune-mediated acute liver injury (Supplemental Figure 1, C–F). These results suggested that preexisting colitis with intestinal barrier degradation directly contributed to the attenuation of subsequent liver injury. Additionally, to confirm whether coexisting inflammations compensated for one another, we performed the experiment in the reverse order (i.e., Con A–injected mice were subsequently administered DSS [Supplemental Figure 2A]). We observed that Con A pretreatment did not ablate DSS-induced colitis (Supplemental Figure 2, B and C), suggesting that the unidirectional gut-liver axis contributed to immune tolerance in the liver in this model.

Monocyte-derived macrophages accumulating in the liver following DSS–Con A treatment display decreased TNF-α production and increased IL-10 production. We then examined the cell types responsible for attenuating liver injury in this model. We found that DSS administration induced an increase in CD11b+ macrophages in the liver, as reported previously (13), and Con A administration also induced a robust increase in CD11b+ macrophages in the liver (Figure 2, A and B). Notably, DSS–Con A–treated mice exhibited the greatest increase in CD11b+ macrophage number in the liver among the 4 groups (Figure 2, A and B). We then assessed whether DSS–Con A–induced hepatic CD11b+ macrophages were ontogenically distinct. As shown in Supplemental Figure 3, A and B, hepatic CD11b+ cells from DSS–Con A mice showed a higher expression of conventional macrophage markers F4/80 and lymphocyte antigen 6 complex, locus C1 (Ly6C), and CSF1 receptor (CSF1R), a marker of BM-derived monocytes (14). The expression of C-type lectin–domain family 4 member F (CLEC4F), a marker of resident liver macrophages (Kupffer cells) (15), was unchanged between the 4 groups, and NKp46 (a marker of NK cells) and Ly6G (a marker of neutrophils) were not increased in the DSS–Con A group. Furthermore, DSS–Con A macrophages displayed marked downregulation of major histocompatibility complex class II-IAb expression as compared with that observed in...
DW–Con A macrophages. We previously reported that BM-derived, TNF-α–producing C-C chemokine receptor9+CD11b+ (CCR9+CD11b+) macrophages play a critical role in Con A–induced acute liver injury by inducing Th1 responses (16, 17). Consistently, CCR9 expression in DW–Con A macrophages was significantly elevated, whereas DSS–Con A macrophages lacked CCR9 expression. Unlike endotoxin-tolerant mice (18), upregulation of IL-receptor–associated kinase 3 (IRAK3) was not observed in macrophages derived from the livers of DSS–Con A mice (Supplemental Figure 3C). To examine potential inflammatory cytokine production, liver mononuclear cells (MCs) and colon lamina propria MCs (LPMCs) were stimulated with LPS for 4 hours in vitro, followed by intracellular TNF-α staining. As shown in Figure 2, C and D, hepatic CD11b+ cells derived from DSS–Con A mice produced less TNF-α than did DW–Con A mice.

Figure 1. Severe mucosal barrier degradation leads to significantly milder liver injury following a sublethal injection of concanavalin A (Con A). (A) Mice were either untreated or orally administrated 2.0% dextran sulfate sodium (DSS) followed by intravenous PBS or Con A (20 mg/kg) administration. Mean body weight and disease activity index (DAI) on days 0–7 of each experimental group (n = 5–7/group). (B) Serum alanine aminotransferase (ALT) level for each experimental group (n = 5–7/group). (C) Histopathology of H&E-stained sections of liver. Data are representative of each experimental group. Scale bars: 500 μm (n = 5/group). (D) Representative TUNEL assay images of mouse liver sections from each experimental group (n = 5/group). Scale bars: 50 μm. (E) Expression of TNF-α and IL-6 mRNA in whole liver cells (n = 4/group). (F) DAI for each duration of DSS administration. (G) Serum ALT levels after Con A injection for each duration of DSS pretreatment. Data represent means ± SEM. *P < 0.05, **P < 0.01, ***P < 0.001 according to 1-way ANOVA with Tukey’s multiple-comparison correction.
whereas colon lamina propria CD11b+ cells from DSS–Con A mice exhibited robust TNF-α production (Figure 2, E and F), clearly demonstrating the distinct phenotypes of the liver and colon cells. Moreover, we assessed antiinflammatory cytokine production by DSS–Con A macrophages upon LPS stimulation in vitro. Notably, DSS–Con A macrophages in the liver produced larger amounts of IL-10 following LPS stimulation in vitro (Figure 2, G and H). The numbers of other immune-regulatory cell subsets, such as

Figure 2. Liver DSS–Con A monocyte–derived macrophages display decreased TNF-α production and potentially increased IL-10 production upon LPS stimulation in vitro, and antigen-presenting potential in hepatic macrophages is reduced following DSS–Con A treatment. (A) Representative CD11b and CD11c staining of whole liver mononuclear cells (MCs) isolated from each experimental group. Each box and number indicates the population and the percentage of CD11b+ cells in whole liver MCs. (B) Percentage (left) and number (right) of CD11b+ cells in whole liver MCs of each experimental group (n = 4/group). (C) Representative surface CD11b and intracellular TNF-α staining of whole liver MCs isolated from each experimental group after 4 hours of LPS stimulation. Each number indicates the percentage resident in whole liver MCs. (D) Percentage and number of TNF-α+CD11b+ cells in the livers of mice from each experimental group (n = 4/group). (E) Representative surface CD11b and intracellular TNF-α staining of whole lamina propria MCs isolated from each experimental group after 4 hours of LPS stimulation. Each number indicates the percentage resident in whole lamina propria MCs. (F) Percentage and number of TNF-α+CD11b+ cells in the colon lamina propria of each experimental group (n = 4/group). (G) Representative surface CD11b and intracellular IL-10 staining of whole liver MCs isolated from each experimental group after 4 hours of LPS stimulation. Each number indicates the percentage in whole liver MCs. (H) Percentage and number of IL-10+CD11b+ cells in the livers of mice from each experimental group (n = 4/group). Data represent the mean ± SEM. *P < 0.05 according to 1-way ANOVA with Tukey’s multiple-comparison correction.
CD11c⁺Siglec H⁺ plasmacytoid DCs (pDCs) (19) and CD4⁺CD25⁺Foxp3⁺ Tregs (20), were decreased in the livers of DSS–Con A mice (Supplemental Figure 4, A–D).

Liver DSS–Con A macrophages show impaired antigen-presenting function to T cells in vitro and exert protective functions against excessive liver injury induced by Con A. Next, we assessed the antigen-presenting function of hepatic CD11b⁺F4/80⁺ cells isolated from DSS–Con A–treated mice and compared it with the same function of CD11b⁺F4/80⁺ cells from DW–Con A–treated livers. DSS–Con A– or DW–Con A–derived CD11b⁺F4/80⁺ cells were cocultured with naive carboxyfluorescein succinimidyl ester–labeled (CFSE-labeled) CD4⁺CD62L⁺T cells from the spleens of OT-II mice in the presence of ovalbumin peptides (Figure 3A). After a 72-hour culture, naive CD4⁺ T cells extensively proliferated in the presence of DW–Con A–treated macrophages, whereas they exhibited less proliferation in the presence of DSS–Con A–derived macrophages (Figure 3, B and C). These data suggested that hepatic CD11b⁺ macrophages derived from DSS–Con A–treated mice displayed an immune-regulatory phenotype that might contribute to the attenuation of acute liver injury. We then investigated whether DSS–Con A–derived CD11b⁺ cells directly suppressed liver injury. CD11b⁺ cells obtained from either DW–Con A–treated or DSS–Con A–treated Ly5.1⁺ WT mice were transferred to clodronate-pretreated Ly5.2⁺ mice to evaluate the role of adoptively transferred macrophages. We confirmed that CD45.1⁺CD11b⁺ cells substantially accumulated in the inflamed liver of the mice (Figure 3D). As shown in Figure 3E, transfer of DSS–Con A–derived CD11b⁺ cells significantly protected mice from subsequent Con A–induced liver injury. These
results clearly demonstrated that CD11b+ macrophages derived from DSS–Con A–treated mice were capable of attenuating Con A–induced acute liver injury. Activated Th1 cells play a critical role on the pathogenesis of Con A–induced acute liver injury, given that Rag2−/−, Ifng−/−, Stat1−/−, and Tbx21−/− mice develop less severe liver injury (21–23). As expected, DSS–Con A–treated mice exhibited diminished accumulation of both CD3+CD4+ and CD3+CD8+ T cells in the liver (Supplemental Figure 5, A–C). To examine potential inflammatory cytokine production, liver MCs were stimulated with PMA/ionomycin for 4 hours in vitro, followed by intracellular IFN-γ staining. Accordingly, as shown in Supplemental Figure 5, D–G, both CD3+CD4+ and CD3+CD8+ T cells derived from DSS–Con A–treated livers produced less IFN-γ than did DW–Con A mice. These data indicated that preexisting colitis with intestinal barrier degradation affected Con A–induced T cell activation in the liver.

IL-10 and TGF-β are involved in the attenuation of liver injury by tandem DSS–Con A administration. Given that the number of IL-10–producing macrophages increased in DSS–Con A–treated livers (Figure 2), we hypothesized that enhanced expression of regulatory cytokines promoted the attenuation of acute liver injury. Indeed, IL-10 and TGF-β mRNA expression were significantly upregulated in whole livers derived from DSS–Con A–treated mice as compared with levels observed in DW–Con A–treated livers (Figure 4, A and B). To investigate the direct role of IL-10 and TGF-β in attenuating liver injury in this model, both DW- and DSS-administered mice were treated with an anti–IL-10 or anti–TGF-β antibody 30 minutes prior to Con A administration. Neutralization of IL-10 and TGF-β levels had little effect on Con A–induced liver injury, whereas neutralization of IL-10 and TGF-β partially ablated the protective effect of DSS–Con A treatment on liver injury (Figure 4, C and D). These data suggested that IL-10 and TGF-β were directly involved in the attenuation of liver injury by tandem DSS–Con A administration.

Continuous exposure to gut-derived products directly affect the susceptibility to subsequent liver injury. We then examined whether gut sterilization affects the severity of DSS–Con A–induced liver injury to clarify the involvement of gut microbiota in these mechanisms. Mice were orally administered a broad-spectrum antibiotic cocktail (ampicillin [Amp], vancomycin [VCM], neomycin [Neo], and metronidazole [MNZ]) 3 times weekly for 2 weeks, followed by DSS–Con A treatment (Figure 5A). We confirmed that both the severity of colitis and body weight loss were comparable between the control and gut-sterilized groups (Figure 5B). Notably, gut sterilization by the antibiotic cocktail followed by DSS–Con A treatment ablated the protective effect against liver injury, as measured by serum alanine aminotransferase (ALT) levels and compared with control mice without gut sterilization (Figure 5C), suggesting that gut microbiota play a substantial role in the protective mechanism. Importantly, pretreatment with Neo, which mainly depletes gram-negative bacteria, followed by DSS–Con A treatment worsened colitis symptoms but did not ablate the protective effect against Con A–induced liver injury, as compared with that observed in control mice without gut sterilization (Figure 5, B and C). These results suggested that increased exposure to microbial products (especially gram-positive microbiota) accompanied by intestinal barrier disruption played a substantial role in the induction of immune tolerance in the liver. We further determined whether the TLR/
myeloid differentiation primary response gene 88 (MyD88) pathways are responsible for hepatic immune tolerance in DSS–Con A–treated mice (Supplemental Figure 6A). MyD88-deficient mice administered DSS developed substantial colitis (Supplemental Figure 6B), and we observed a similarly suppressed liver inflammation in MyD88-deficient mice administered DSS–Con A (Supplemental Figure 6C), suggesting that LPS/TLR-independent pathways play a role in this model.

Nicotinamide N-methyltransferase (NNMT) and concomitant increase in 1-methylnicotinamide (1-MNA) levels play integral roles in the induction of immune tolerance via the gut-liver axis. We then investigated whether a damaged intestinal barrier (leaky gut) and concurrent leakage of gut-derived products through the portal vein directly affect the severity of subsequent liver injury. As shown in Figure 6A, intestinal permeability evaluated by serum FITC-dextran concentration was significantly aggravated in DSS–Con A–treated mice as compared with other groups. The LPS level in the serum derived from each portal vein was also increased in DSS-PBS and DSS–Con A mice (Figure 6B). We finally investigated the molecules exhibiting protective functions in the DSS–Con A model. We initially demonstrated that pretreatment with plasma derived from DSS–Con A mice significantly protected mice against subsequent Con A–induced liver injury (Figure 6C). Moreover, liver MCs derived from DSS-PBS–treated mice were cultured in vitro for 6 hours with plasma from DSS–Con A–treated mice or DW-PBS–treat-
ed mice (Supplemental Figure 7A). Consistently, DSS–Con A–derived plasma induced higher IL-10 production from liver CD11b+ cells as compared with DW-PBS–derived plasma (Supplemental Figure 7B). These data suggested that molecules in plasma from DSS–Con A–treated mice exhibited a protective effect against Con A–induced liver injury; therefore, we determined the metabolites present in plasma derived from DSS–Con A–treated mice by capillary electrophoresis TOF mass spectrometry–based (CE-TOFMS–based) metabolome analysis (Figure 6D). Several metabolites in plasma derived from DSS–Con A–treated mice showed higher levels relative to those in plasma derived from DW–Con A–treated mice, including 1-MNA, a terminal product of nicotinamide (NAM) clearance. We then confirmed that 1-MNA administration protected mice against Con A–induced liver injury, along with increased mRNA expression of TGF-β and IL-10, and especially induction of IL-10–producing macrophages in the liver (Figure 6, E–G). Moreover, the mRNA expression of NNMT, which catalyz-
es NAM methylation to produce 1-MNA, was significantly higher in DSS–Con A–treated livers than in control mice. Finally, gut sterilization via the antibiotic cocktail followed by DSS–Con A treatment ablated NNMT upregulation in the liver (Figure 6H). These results collectively suggest that NNMT upregulation by exposure to gut-derived products and concomitant increases in 1-MNA play an integral role in the induction of immune tolerance via the gut-liver axis.

**Discussion**

The gut-liver axis is of clinical importance as a potential therapeutic target in a wide range of liver diseases, including nonalcoholic steatohepatitis (NASH) (24), alcoholic liver diseases (25), and PSC (7). Recent reports have suggested that altered gut microbiota and increased intestinal permeability contribute to liver disease initiation and progression (26–28). The present study revealed an unexpected role of the gut-liver axis in maintaining immunological balance in the liver. Herein, we demonstrated that a distinctive state with intestinal barrier disruption and continuous exposure to microbial products protected mice from further liver injury, along with the emergence of immunosuppressive macrophages. As a proposed mechanism, induction of immunological tolerance in the liver was IL-10 and TGF-β dependent, as the protective effect against DSS–Con A administration was partially canceled by treatment with anti–IL-10 or anti–TGF-β antibodies. Furthermore, certain gut microbiota or microbial products played a substantial role via the gut-liver axis based on the attenuated protective effect against DSS–Con A administration by gut sterilization. We identified 1-MNA as a candidate metabolite capable of directly suppressing liver injury by inducing IL-10–producing macrophages in the liver (Figure 7).

We previously reported that an anti-Fas–induced fulminant-hepatitis model exhibited a deteriorated response to DSS-induced colitis (13). Similar liver inflammation linked with increased intestinal permeability and dysbiosis has been observed both in human diseases and murine models, including those related to jejunal self-filling blind loops (29) and alcoholic and nonalcoholic liver diseases. Con-
trary to these previous findings, we demonstrated that DSS-colitis pretreatment protected mice from immune-mediated hepatitis initiated by Con A. It is worth mentioning that the severity of intestinal inflammation is conversely correlated with subsequent liver injury. Specific subsets of innate immune cells in the liver adequately respond to gut-derived antigens, whereas excessive antigen stimuli induce immune permissiveness to avoid further liver injury (3, 30). On the other hand, pretreatment with an extremely low dose of endotoxin has an opposite effect in that it potentiates a proinflammatory response to subsequent endotoxin challenge, referred to as the Shwartzman-like reaction (31). It is generally believed that the initial magnitude of the immune reaction determines the immunological fate, namely immune priming or immune tolerance (32). This 2-pronged phenomenon might account for the discrepancy between our findings and those from previous reports illustrating that continuous mild intestinal inflammation induced by chronic DSS administration enhanced hepatic inflammation and fibrogenesis in an experimental NASH model (33).

Whether the mechanism behind the liver tolerance demonstrated in the present study is distinct from LPS tolerance is an important issue. First, the expression of IRAK3, a molecule contributing to LPS tolerance (34), was not induced but was decreased in the livers of DSS–Con A–treated mice. Second, we observed equally suppressed liver inflammation in MyD88-deficient mice administered DSS–Con A (Supplemental Figure 6C). Third, pretreatment with Neo to examine the role of gram-negative bacteria failed to cancel the induction of immune tolerance in DSS–Con A–treated mice. These findings collectively suggest that immune tolerance in this model was independent of LPS tolerance.

We recently reported the critical role of TNF-α–producing CCR9⁺CD11b⁺ macrophages in the pathogenesis of Con A–induced hepatitis (16, 17). These macrophages are characterized as classically activated macrophages that contribute to host defenses against a variety of bacteria and viruses and play a substantial role in antitumor immunity. However, in the present study, we found that continuous exposure to gut-derived antigens with diminished intestinal integrity induced immunological tolerance in the liver, along with the emergence of IL-10–producing macrophages, and directly protected mice from Con A–induced acute liver injury. It is reasonable that recruited hepatic macrophages shifted from inflammatory to regulatory functions in the hepatic microenvironment continuously exposed to microbial products through the portal vein, rather than being differentially recruited directly to the liver as regulatory macrophages. Accordingly, Con A–pretreated mice (a state of severe hepatitis) were not protected from subsequent DSS-induced colitis, suggesting that the emergence of immune-suppressive macrophages via the gut-liver axis was unique to the liver. As candidate products potentially contributing to the induction of immune tolerance in the liver, we confirmed that 1-MNA directly contributed to the attenuation of Con A–induced liver injury by inducing IL-10–producing macrophages. Recent studies have suggested that 1-MNA possesses biological activity and is not merely a terminal product of NAM clearance. 1-MNA is methylated from NAM by NNMT, which is specifically expressed in hepatocytes (35), and its expression in the liver is associated with serum cholesterol, triglycerides, and other metabolic parameters (36). Mechanistically, increasing NNMT expression or 1-MNA levels stabilizes the sirtuin 1 protein, an effect that is required for its metabolic benefits (36). Increased NNMT expression and 1-MNA release have been reported in humans and rodent models of inflammation, including those involving Con A–induced acute liver injury and pulmonary hypertension (37–39), indicating a protective role for NNMT in compensating for inflammation. We demonstrated that increased NNMT expression in the liver following DSS–Con A administration was abrogated by gut sterilization, suggesting potential regulation of NNMT expression in the liver by gut microbes.

In summary, our findings showed that IL-10–producing CD11b⁺ macrophages contribute to immune tolerance in the inflamed liver under intestinal inflammation conditions accompanied by diminished intestinal barrier function. Our results provided a potentially novel mechanism of immune tolerance in the liver via the gut-liver axis, as well as insight into a therapeutic strategy for treating gut-related liver diseases.

Methods

Animals. WT C57BL/6J mice were purchased from Japan CLEA. Mice were maintained under specific pathogen–free (SPF) conditions in the Animal Care Facility of Keio University School of Medicine. Experiments were performed with age- and sex-matched mice at 8–12 weeks of age.
Experimental protocols of Con A–induced hepatitis. Con A (type IV) was purchased from MilliporeSigma. Con A (15 mg/kg) were i.v. administered into the tail vein of mice 12 hours before liver resection.

Experimental protocols of DSS-induced colitis as model of mucosal barrier degradation. DSS (molecular weight 36–50 kDa) was purchased from MP Biomedicals. To induce mucosal barrier degradation by acute experimental colitis, mice received 1.5%–2.0% DSS dissolved in sterile DW ad libitum for 4–7 days.

Assessment of clinical activity in DSS-treated mice. Assessment of clinical activity in DSS-treated mice were evaluated as described previously (40). Mice were weighed daily and visually inspected for diarrhea and rectal bleeding. DAI was assessed in a blinded fashion. No weight loss was registered as 0, weight loss of 1%–5% from baseline was assigned 1 point, with 6%–10% as 2 points, 11%–20% as 3 points, and >20% as 4 points. For stool consistency, 0 points were assigned for well-formed pellets, 2 points for pasty and semiformal stools that did not adhere to the anus, and 4 points for liquid stools that adhered to the anus. For bleeding, 0 was assigned for no blood, 2 points for positive bleeding, and 4 points for gross bleeding.

Preparation of liver MCs and intestinal LPMCs. Liver MCs were separated from the liver as described previously (16). Single-cell suspensions of intestinal LPMCs were prepared as previously described (41), with slight modifications. Details are described in the Supplemental Methods.

Flow cytometry. After blocking with anti-FcR (CD16/32, 2.4G2, BD Pharmingen) for 20 minutes, the cells were incubated with the specific fluorescence–labeled mAbs used for surface staining at 4°C for 30 minutes, followed by permeabilization and intracellular staining in some cases. Antimouse mAbs used are listed in the Supplemental Methods.

Quantitative PCR (qPCR). All qPCR assays were performed as described previously (16). Details and the predesigned primers are described in the Supplemental Methods.

Macrophage depletion. To deplete macrophages, mice were i.v. injected Clodronate Liposome (200 μg/body; FormuMax) 24 hours before Con A administration.

Adoptive transfer. CD11b+ cells were isolated using FACS Aria (Becton Dickinson). Isolated cells (5 × 10^6 cells) were transferred immediately into Con A.

Neutralizing IL-10 or TGF-β. To neutralize IL-10 or TGF-β, mice were treated with neutralizing antibodies to IL-10 (JES5-2A5; 200 μg/body, eBioscience) or TGF-β (1D11; 100 μg/body, Gene Tex) or isotype-matched antibodies 30 minutes before Con A administration.

Evaluation of intestinal permeability. Mice were orally provided with 4,000 Da FITC dextran (MilliporeSigma) via nasogastric tube after being fasted for 6 hours. Blood samples were collected from the tail vein at 4 hours. The concentration of FITC from obtained serum was measured by spectrophotofluorometry according to the instructions. Serum from mice not administered with FITC-dextran is used to determine the background.

Gut sterilization with broad-spectrum antibiotics. Mice were treated with broad-spectrum antibiotics via nasogastric tube (500 μl each) 3 times a week for 2 weeks. The following amount of antibiotics were used; Amp (6.7 g/l), Neo (6.7 g/l), VCM (3.3 g/l), and MNZ (6.7 g/l).

Pretreatment with plasma derived from portal vein injection in vivo. Mice were i.v. administered with plasma derived from portal vein (200 μl/body) derived from DW-PBS or DSS–Con A mice at 2 hours before Con A injection.

CE-TOFMS–based metabolome analysis. A quantitative analysis of charged metabolites by CE-TOFMS was performed as described previously (42). Details are described in the Supplemental Methods.

Statistics. Results are expressed as mean ± SEM. Data groups were analyzed with GraphPad Prism software using 1-way ANOVA with Tukey’s multiple-comparison correction or 2-tailed Student’s t test. A P value less than 0.05 was considered statistically significant.

Study approval. All animal protocols described in this study were performed in accordance with Institutional Guidelines on Animal Experimentation at Keio University and were approved by The Keio University IACUC.

Author contributions
NT helped design the study, performed experiments, and analyzed the data; NN conceived and designed the study, analyzed the data, and wrote the paper; PSC, TA, T. Teratani, TS, AY, SS, RM, and T. Katayama performed experiments; T. Tsukimi and SF performed metabolome analysis; YM and HE helped design the study; and T. Kanai helped conceive and supervise the study.
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