Diet and specific microbial exposure trigger features of environmental enteropathy in a novel murine model

Eric M. Brown1,2, Marta Wlodarska1,2, Benjamin P. Willing3, Pascale Vonaesch4, Jun Han5, Lisa A. Reynolds2, Marie-Claire Arrieta2, Marco Uhrig1, Roland Scholz2, Oswaldo Partida2, Christoph H. Borchers5,6, Philippe J. Sansonetti4 & B. Brett Finlay1,2,7

Environmental enteropathy (EE) is a subclinical chronic inflammatory disease of the small intestine and has a profound impact on the persistence of childhood malnutrition worldwide. However, the aetiology of the disease remains unknown and no animal model exists to date, the creation of which would aid in understanding this complex disease. Here we demonstrate that early-life consumption of a moderately malnourished diet, in combination with iterative oral exposure to commensal Bacteroidales species and Escherichia coli, remodells the murine small intestine to resemble features of EE observed in humans. We further report the profound changes that malnutrition imparts on the small intestinal microbiota, metabolite and intraepithelial lymphocyte composition, along with the susceptibility to enteric infection. Our findings provide evidence indicating that both diet and microbes combine to contribute to the aetiology of EE, and describe a novel murine model that can be used to elucidate the mechanisms behind this understudied disease.
Malnutrition is a significant worldwide health issue, causing one-fifth of all deaths in children <5 years of age. Early-life malnutrition results in childhood growth stunting and decreased cognitive development, along with significant changes in intestinal microbiota composition. During the first 2 years of life in malnourished children, there is an immature development of the intestinal microbiota compared with healthy, case-controlled children, and a comprehensive study across multiple geographical regions concluded that children consistently harbour a significantly more diverse early-life microbiota composition in low socioeconomic regions of the world. Clinical trials have shown that therapeutic food interventions are not sufficient to aid in the development of a healthy early-life intestinal microbiota and fail to fully restore health in malnourished children. However, antibiotic treatment preceding the nutritional intervention leads to an improvement of the health defects cause by malnutrition, indicating the importance of understanding the microbial-driven disease pathologies and underlying defects in intestinal function contributing to malnutrition.

A poorly understood disorder of small intestine called environmental enteropathy (EE), also referred to as tropical enteropathy or environmental enteric dysfunction, is now widely recognized to be a major contributor to childhood malnutrition, and is thought to be the reason why current therapeutic interventions in malnourished children are efficacious less than one-third of the time. This disease impacts the function of the small intestine, and features include chronic intestinal inflammation, villous blunting and increased intestinal permeability, ultimately resulting in diminished nutrient uptake, adversely affecting growth and development. As a subclinical sequelae of malnutrition happens independently of diarrhoea. Malnutrition induces an altered small intestinal microbiota with significant changes in intestinal microbiota composition in low socioeconomic regions of the world. This hypothesis would also explain the benefit of antibiotic treatment before therapeutic food interventions for malnourished children. Unfortunately, EE is largely understudied owing to the difficulty in obtaining samples directly from the small intestine in humans, which requires invasive biopsies deemed unethical because of the subclinical nature of EE. Thus, insight into the connection between metabolism, microbiota and immunity in the malnourished intestine is limited. Furthermore, many human studies take only faecal samples to assess the microbiota composition in malnutrition and EE cases, yet this does not capture the small intestinal bacterial community, an important consideration, since the faecal microbiota composition is not representative of those microbes colonizing the small intestine where pathology of this disease occurs. Thus, there is a major unmet need for a preclinical animal model of EE, and these models will be crucial to understand the pathogenesis of EE, its impact on malnutrition and to develop therapies.

Here we describe the first animal model of EE and we demonstrate that both diet and oral microbial exposure to specific bacteria are required for the induction of EE in a mammalian host. Utilizing this model, we were able to achieve for the first time, an in-depth characterization of the impact of malnutrition on the microbiota, metabolism and immune system in the mammalian small intestine, along with assessing the impact of enteric infection. These studies provide a preclinical model to test therapeutics and will enable a greater understanding of the pathophysiological nature of EE.

**Results**

**Impact of malnutrition on growth and gut barrier function.** We first sought to characterize the impact of a moderately malnourished diet on the growth rate, intestinal architecture and barrier function of the small intestine in young mice. Post weaning, mice were randomized into receiving either a malnourished diet (7% protein and 5% fat) or isocaloric control diet (20% protein and 15% fat; Fig. 1a and Supplementary Table 1). Both groups consumed equivalent weights per day of the chow (Supplementary Fig. 1a). Mice fed the malnourished diet gained significantly less weight over time in comparison with mice fed the isocaloric control diet (Fig. 1b). Growth stunting of the malnourished mice was evident as they gained an average of 30% less weight after 21 days (Fig. 1c). At this time point, malnourished mice also had considerably shorter tails, a surrogate marker for length of the mouse (Fig. 1d). There was a decreased expression of insulin-like growth factor 1, which encodes a growth factor hormone, and angiotensin-converting enzyme 2, which is necessary for protein uptake, in the jejunum of malnourished mice (Supplementary Fig. 1b,c). Next, the barrier function of the jejunum was assessed by analysing mRNA expression levels of tight-junction protein-1 (TJP1), claudin-2 (CLDN2), claudin-4 (CLDN4) and claudin-15 (CLDN15). TJP1, the product of which is ZO-1, a major component of tight-junction assembly and function, was expressed at lower levels in malnourished mice (Fig. 1e). In contrast, the expression of CLDN2 was upregulated in the jejunum of malnourished mice, which is also indicative of increased permeability based on previous reports. However, the expression of other claudin transcripts (CLDN4/15) was similar between the groups of mice (Fig. 1e). To measure whether these gene expression changes also had a functional impact on barrier function, mice were fed fluorescein isothiocyanate (FITC)-labelled dextran and subsequent FITC levels were measured in the serum. There was a 3.5-fold increase in the serum concentration of FITC in malnourished mice compared with controls, indicating an increase in intestinal permeability (Fig. 1f). Barrier dysfunction did not significantly impact total secretory immunoglobulin A (IgA) levels in the jejunum, an important antibody for host epithelial defence (Supplementary Fig. 1d). In addition, sections of the small intestine (duodenum, jejunum and ileum) were visualized microscopically and no significant signs of inflammation, goblet cell depletion, histopathology or villous blunting were observed in the mice fed the malnourished diet (Fig. 1g and Supplementary Fig. 1e). These data suggest that the malnourished diet led to a moderate growth stunting and increased intestinal permeability relative to the isocaloric control diet independent of small intestinal histopathology.

**Malnutrition induces an altered small intestinal microbiota.** To test whether malnutrition alters the small intestinal microbiota in mice, we performed 454 pyrosequencing of the 16S rRNA genes from the microbiota in the duodenum and ileum of the mice 3 weeks after being fed the malnourished or isocaloric control diet (Supplementary Data 1). We found marked shifts in the bacterial species composition of both the ileum and duodenum in the malnourished mice, compared with those on the control diet (Fig. 2a). The ileum of malnourished mice showed both an...
expansion of species from the Bacteroidetes and Proteobacteria phyla, and decrease in the Lachnospiraceae and Clostridiales families compared with control mice (Fig. 2a and Supplementary Fig. 2a). The duodenum of malnourished mice also had an abnormal and significant expansion in the proportion of bacteria belonging to the phyla Bacteroidetes and Proteobacteria (largely the Enterobacteriaceae family; Fig. 2a,b and Supplementary Fig. 2a,b), which are normally found at high abundance in the

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Figure 1 | Assessment of the growth rate and intestinal barrier function in C57BL/6 mice fed a malnourished or isocaloric control diet. (a) A schematic of the components of each diet, expressed as a per cent of total calories. (b) Mice were given each diet post weaning, and daily weight change was assessed daily over a period of 21 days. Data are representative of the three independent experiments, n = 8 mice per group. Data points represent the mean and error bars indicate the s.d. (**P < 0.01, repeated measures analysis of variance). After 3 weeks of being fed each diet, the total amount of (c) weight gained and (d) final tail lengths were calculated. In the box and whisker plots shown, the middle bar represents the mean, the bottom and top of the box are the first and third quartiles, respectively, and whiskers indicate the range of the data. Data are representative of three independent experiments, n = 8 mice per group (Student’s t-test **P < 0.01). (e) The jejunal mRNA expression of TJP1, CLDN2, CLDN4 and CLDN15 are expressed as fold change relative to the control-fed mice. Bars indicate the mean with s.e.m., and are representative of two independent experiments, eight mice per group (*P < 0.05, Student's t-test). (f) Concentration of FITC in the serum was assessed 4 h post administration orally, after mice were fed each diet for 3 weeks. Bars indicate the mean with s.e.m., and are representative of two independent experiments, n = 8 per group (*P < 0.05, Student's t-test). (g) The average measured villous height in the duodenum (DUO), jejunum (JEJ) and ileum (ILE) with representative H&E-stained images of jejunum sections from mice exposed to each diet. Scale bar, 100 μm (in length). Bar graph indicates the mean with s.e.m., and is representative of three independent experiments, eight mice per group. CON, control; MAL, malnourished.
lower intestinal tract nearer the colon in C57BL/6 mice. Normal residents of the duodenum, the Gram-positive bacteria from Lactobacillaceae and Erysipelotrichaceae families were decreased in abundance in the malnourished duodenum with a striking 45% reduction in the abundance of Lactobacillaceae (Fig. 2b). This decrease coincided with an increase in abundance of a more diverse set of operational taxonomic units (OTUs) from the Bacteroidales, Prevotella, Lachnospiraceae, Clostridiales, Peptostreptococcaceae, and Ruminococcaceae groups (Supplementary Table 2). As visualized on the principal component analysis (PCA) plot, the dietary changes cause the microbial community of the duodenum to have greater similarity with the ileum microbial community, and this can be quantified by assessing the UniFrac distance, a measure of community similarity (Fig. 2c). Using this measure, we found that the resident microbiota in the malnourished duodenum was significantly more similar (P < 0.01, Student’s t-test) to the ileal community than the community in the control duodenum samples (Fig. 2c). There was also a marked increase in bacterial diversity of the duodenum in malnourished mice compared with those in the control diet, as seen in the phylogenetic distribution at the genus level and quantified using the inverse Simpson’s diversity index (Supplementary Fig. 2b,c). Furthermore, quantitative PCR analysis of 16S copies in the duodenal microbiota verified that the family-level microbial differences observed in the pyrosequencing data were consistent across multiple experiments (Fig. 2d–h). Overall, these data suggest that the malnourished diet remodelled the bacterial
community in the duodenum to more closely resemble the ileum, where Gram-negative species (Bacteroidetes and Proteobacteria) were more abundant, displacing the indigenous Gram-positive Firmicutes, such as Lactobacillaceae (Supplementary Fig. 2d).

Impact of malnutrition on small intestinal metabolome. As malnutrition leads to drastic shifts in the intestinal microbiota, we hypothesized that this would also impact the metabolic environment of the host small intestine. After 3 weeks of treatment with either diet and concurrent with our microbial analysis, contents of the small intestine in malnourished and control diet mice were assessed for abundance changes of the metabolites using untargeted metabolomic analysis by ultrahigh-performance liquid chromatography–Fourier transform (FT) mass spectrometry (UPLC–FTMS). The method detected and relatively quantified over 3,500 unique metabolite features, and ~420 showed differential abundances between malnourished and control mice (P < 0.05; >2-fold change). Overall, the metabolite profile of the malnourished small intestine substantially shifted from the profile of control mice, as shown by unsupervised PCA (Fig. 3a). When all metabolites detected in the positive ion mode were visualized on a heat map, malnourished and control groups clustered together in the dendogram indicating that the diet shifted the metabolome to an alternate state (Fig. 3b). Each of these observations held true for metabolites detected on the negative ion channel (Supplementary Fig. 3a,b).

Using the metabolite matches based on mass to charge ratio (m/z) values found with the Kyoto Encyclopedia of Genes and Genomes database, we determined specific pathways that were enriched by the malnourished or control diet. Metabolites in the bile acid biosynthesis pathway (37% of pathway impacted; Fisher’s exact test P < 0.001), linoleic acid metabolism and amino-acid biosynthesis were all significantly over-represented in the control diet (Fig. 3c). The changes in the amino-acid biosynthesis pathways were expected in the control mice, given the higher protein content of their diet, supporting the accuracy of our method and analysis. The most striking over-representation in malnourished mice was the greater abundance of metabolites in the steroid biosynthesis pathway (28% of pathway impacted; P < 0.001) (Fig. 3d). More specifically, many of these steroid metabolites represented those from the vitamin D biosynthesis pathway. Random forest analysis was implemented to identify discriminatory features between the two diets, and it ranked a vitamin D metabolite as the most significant feature between the samples from the malnourished mice and controls (Supplementary Table 3). Of note, the vitamin D pathway is known to impact many aspects of intestinal homeostasis, including gut inflammation25 and bile acid production.26 We confirmed these metabolome changes with a targeted metabolomics approach to quantify the concentrations of bile acids, vitamins and microbial-produced short-chain fatty acids (SCFAs) in the small intestine. Bile acid-targeted metabolomics revealed a number of significant changes in the bile acid pool between the malnourished and control small intestine, supporting the pathway analysis data from the untargeted metabolomics screen (Fig. 3e,f). In the malnourished small intestine, there was a shift towards lower concentrations of tauro-conjugated bile acids, and, conversely, higher levels of unconjugated bile acids, with the ratio between the two being significantly altered (Supplementary Fig. 4a–c). Each tauro-conjugated bile acid detected in this analysis was lower in the malnourished small intestine, with the concentrations of tauro-α-muricholic acid, tauro-α-muricholic acid, tauroursodeoxycholic acid, taurohodeoxycholic acid and taurochenodeoxycholic acid reaching statistical significance (Mann–Whitney U-test; Fig. 3e). In the unconjugated pool, the concentrations of muricholic acid, ursodeoxycholic acid and 7-ketodeoxycholic acid were all significantly higher in the malnourished small intestine (Mann–Whitney U-test; Fig. 3f).

Results from the vitamin-targeted metabolomics analysis showed that the malnourished diet induced large shifts in the luminal concentrations of many vitamins in the small intestine (Supplementary Fig. 5), which are not due to altered concentrations of vitamins in the diets as they are identical (Supplementary Table 1). Malnutrition also induced significant reductions in vitamin A metabolites (retinol and retinal), vitamin B1, vitamin B6, vitamin B12 and vitamin E (α-tocopherol), and elevated concentrations of vitamin D3 and vitamin B7 (Supplementary Fig. 5). No significant changes in SCFAs were observed, although there was a general trend towards greater total concentrations of each in the malnourished small intestine (Supplementary Fig. 6). These data clearly show that a malnourished diet exerts a strong, measurable impact on both the microbial and metabolite content in the small intestine, mainly in the biosynthesis or bioavailability of steroids, vitamins and bile acids, and may be contributing to or a result of changes in bacterial composition.

A Bacteroidales and E. coli cocktail replicates features of EE. Given the malnourished-induced shifts in the small intestinal microbiota and metabolome, we postulated that oral exposure to only specific species of bacteria would be necessary to induce EE features. We utilized the small intestinal microbial data to rationally select taxonomically grouped mixtures of species with a propensity to colonize the small intestine more readily in malnourished mice, and, subsequently, we orally exposed mice on both diets to these cocktails of cultured bacterial species (Supplementary Fig. 7 and Supplementary Table 4). In this model, young mice were randomized onto either the control or malnourished diet for 2 weeks, then gavaged every 2 days with each bacterial cocktail for an additional 6 days and finally the characteristic pathological features of EE were assessed 1 week after the last gavage (Fig. 4a). Alternatively, to measure growth rates, malnourished mice were gavaged every 3 days post weaning for a total of 2 weeks, and EE features were assessed 1 week later (Supplementary Fig. 8). Remarkably, we found that in combination with the malnourished diet, the oral exposure to defined mixture of Bacteroidales and E. coli (BG) was able to trigger villous blunting and exacerbate the effect of the malnourished diet on growth stunting, intestinal permeability and inflammation resulting in intestinal disease, which is strikingly similar to features of human EE (Fig. 4b–h and Supplementary Fig. 8a–e). Importantly, mice given a standard diet and exposed to BG did not show any pathological consequences and maintained normal growth compared with untreated, control mice (Supplementary Fig. 8a,b).

Malnourished mice exposed to BG showed increased growth stunting (35% less weight) as compared with unexposed malnourished mice (Supplementary Fig. 8a,b). The increased weight loss in the BG-exposed malnourished mice was accompanied with increased intestinal permeability (Fig. 4b–d). When comparing jejunal gene expression of tight-junction proteins of the malnourished mice with the BG-exposed malnourished mice, we found that there is a further increase in CLDN2 expression, both at the gene and protein level (Fig. 4b,c), and a further reduction of CLDN4 expression in the BG-exposed malnourished mice (Fig. 4b), suggesting an exacerbation of intestinal permeability. The heightened intestinal permeability of the BG-exposed malnourished mice was functionally relevant as increased levels of FITC in the serum were measured after oral administration of FITC–dextran compared with unexposed malnourished mice and control mice (Fig. 4d). Zonulin levels in the serum have been
shown previously to correlate with increased intestinal permeability in many intestinal diseases, by regulating tight-junction proteins and is a biomarker for human EE. We observed that BG-exposed malnourished mice have a fourfold increase of zonulin in the serum, compared with unexposed malnourished mice and control mice (Fig. 4e).

Figure 3 | Untargeted and targeted metabolomics of the small intestinal metabolome. (a) A PCA plot showing separation of metabolomic data as detected by the positive ion channel from malnourished mice (green) and control mice (red). (b) A heat map of the relative abundance of all metabolites identified from the small intestinal metabolome as detected by the positive ion channel from malnourished and control mice (n = 4). The malnourished (MAL) and control (CON) samples clustered together in the dendogram based on cluster analysis by the Ward method, with a Pearson distance measure. The heat map scale is a log-base, from the range of −3 (blue) to +3 (red). (c,d) Plots showing over-represented pathways using Metaboanalyst 2.5 software in (c) control mice and (d) malnourished mice, as identified by the Kyoto Encyclopedia of Genes and Genomes database. The plot is graphed based on the log(P value; Welch’s t-test, y-axis) and per cent of pathway impacted (x axis). The size of the circles represents the number of metabolites identified to be part of the pathway. (e) Tauro-conjugated bile acid and (f) unconjugated bile acid concentrations in the small intestine of malnourished (n = 3) and control (n = 4) as determined by targeted metabolomics. Bars represent the mean ± s.e.m., (*P < 0.05, Mann–Whitney U-test). CA, cholic acid; CDCA, chenodeoxycholic acid; HDCA, hyodeoxycholic acid; KDCA, ketodeoxycholic acid; MCA, muricholic acid; T, tauro conjugation; UDCA, ursodeoxycholic acid.
Villous blunting is a histological hallmark of human EE, and we show that BG exposure induced villous blunting, as BG-exposed malnourished mice had significantly shorter villi in the jejunum (250 μm on average) compared with the villi of unexposed malnourished mice (350 μm on average; Fig. 4f and Supplementary Fig. 9c,d). An alternative measure of villous blunting is the ratio of the villous length to crypt depth, which decreased from 4:1 in the malnourished mice to 2.5:1 in the BG-exposed malnourished mice (Fig. 4f). Given the histological differences, we assessed whether BG exposure induced small intestinal inflammation in malnourished mice. We measured *ex vivo* jejunal cytokine secretion from cultured jejunal sections and found significantly elevated interleukin (IL)-6 and monocyte chemotactic protein-1 (MCP-1) production in BG-exposed malnourished mice only (Fig. 4g). To further confirm the presence of chronic inflammation in this model, we tested the impact of BG exposure on another EE biomarker10,12, faecal calprotectin, whose levels correlate positively with intestinal...
inflammation. The concentration of calprotectin in the faeces of BG-exposed malnourished mice was 2.5 greater than controls or unexposed mice, confirming the presence of significant intestinal inflammation (Fig. 4h). Notably, the BG exposure had no effect on TJ expression and protein levels, intestinal permeability, serum zonulin or faecal calprotectin levels in mice fed the control diet (Fig. 4b–d). Given that the BG cocktail contains Gram-negative bacteria, we sought to understand the role of TLR4 signalling in the induction of villous blunting and inflammation. Five of the seven Bacteroidales and E. coli strains used were able to sufficiently activate TLR4 (Supplementary Fig. 9a). However, TLR4-deficient mice exposure to BG still resulted in significant blunting of the jejunal villi, along with upregulated IL-6 secretion, where MCP-1 secretion remained unchanged (Supplementary Fig. 9b–e). Furthermore, the metabolic activity of the BG bacteria are required to induce any measurable level of villous blunting or small intestinal inflammation in malnourished mice, as mice given a heat-inactivated BG did not develop these characteristic EE features (Supplementary Fig. 9c–e).

Unexpectedly, neither the administration of the Bacteroidales species alone nor the E. coli species alone were sufficient to trigger the growth stunting, inflammation or villous blunting characteristic of EE as seen with the combined BG (Supplementary Fig. 8a–e). This implies that the Bacteroidales species and E. coli isolates work synergistically to impart their effect on the small intestine. We also assessed five additional bacterial mixtures including a Ruminococcus mix, Clostridium mix, Prevotella mix and Peptostreptococcaceae mix, along with VSL#3, a commercial probiotic, and found that only exposure to BG was able to exacerbate the effects of malnourishment and result in the development of EE features (Supplementary Fig. 8a–d and Supplementary Table 4).

The BG challenge alters microbial colonization patterns. We next analysed the bacterial colonization levels in the small intestine of mice on each diet, with or without exposure to the BG. We found malnourished mice and BG-exposed malnourished mice had a significant increase in adherent, cultivable anaerobic bacteria in the jejunum compared with control mice (Fig. 5a). Quantification of 16S rRNA levels in jejunal tissue showed a similar trend to the culture-based analysis, as mice fed a malnourished diet had more bacteria associated with the small intestinal tissue (Fig. 5b). Surprisingly, BG-exposed malnourished mice did not have appreciably more bacteria adhering to the small intestine in comparison with the malnourished, unexposed group, despite the extensive villous blunting and increase in intestinal inflammation (Fig. 5b). Bacteroidetes 16S rRNA expression analysis showed similar levels of Bacteroidetes (MIB) in BG-exposed and unexposed malnourished mice, which was greater than the control groups (Fig. 5c). Enterobacteriaceae 16S rRNA expression analysis showed significantly more adherent Enterobacteriaceae in the small intestine of BG-exposed malnourished mice compared with unexposed malnourished mice, and importantly this was not seen in BG-exposed control mice (Fig. 5d). These data suggest that mice given a malnourished diet are more susceptible to colonization of bacteria in the small intestine, especially those from the Enterobacteriaceae family.

To distinguish the localization of bacteria in the small intestine and visually confirm the increase in tissue-associated bacteria in the small intestine, histological slices of the tissues were subjected to analysis using fluorescence in situ hybridization (FISH). By using a universal bacterial DNA probe, a larger number of stained bacteria could be visualized between the villi and associated with the epithelium of the jejunum of malnourished mice (Fig. 5e). Even more prominent was the increase of tissue-associated bacteria after the administration of BG to malnourished mice, where large numbers of bacteria were found to reside along the epithelium and deep within the villi (Fig. 5e). Using γ-Proteobacteria-specific probes, the increase in tissue-associated bacteria in this phylum was most striking in the BG-exposed malnourished mice (Fig. 5e). Complementing the pyrosequencing and qPCR data, FISH analysis also revealed an expansion of Bacteroidetes in the jejunal samples, a decrease in the amount of Firmicutes and more adherent bacteria along the epithelium in all groups (Supplementary Figs 10 and 11). Overall, of the groups analysed by FISH, γ-Proteobacteria penetrated deep into the crypts more than species from the Firmicutes or Bacteroidetes phyla (Fig. 5e and Supplementary Figs 10 and 11). Of note, this increase in adherent bacteria coincides with an increased expression of antimicrobial peptides or defence molecules by the host epithelium in BG-exposed malnourished mice, as matrix metalloproteinase 7 is significantly elevated, along with resistin-like molecule-β, and trends to higher expression of REG3-γ and cryptdin (Supplementary Fig. 12). These data indicate that the observed increased bacterial load adhering to the intestinal epithelium is not because of decreased antimicrobial peptide production in the jejunum.

Influx of IELs contribute to inflammation after BG exposure. Owing to the presence of intestinal inflammation and villous blunting observed after BG exposure, we investigated the impact of diet and BG on the intestinal immune cell populations. We

Figure 4 | Characterizing the impact of Bacteroidales-E. coli oral exposure on small intestinal histopathology, inflammation and intestinal permeability in malnourished and control mice. (a) A schematic of the experimental design used to administer the Bacteroidales-E. coli cocktail. (b) The jejunal mRNA levels of TJP1, CLDN2, CLDN4 and CLDN5 were determined on unexposed mice on each diet (white and black bars), and BG-exposed mice on each diet (blue and red bars). The values are expressed as fold change relative to the control, unexposed mice. Representative of three independent experiments, n = 5 per group. (c) Levels of CLDN2 protein blotted from extracted protein from jejunal IECs relative to the actin control. Blot is representative of three samples and two independent experiments (Supplementary Fig. 14). (d) Four hours after FITC-dextran administration, concentration of FITC in the serum in malnourished and control mice (n = 8), and BG-exposed malnourished and control mice (n = 8). Representative of two independent experiments. (e) Box and whisker plot of the concentration of zonulin in the sera of malnourished and control mice (n = 8) and BG-exposed malnourished and control mice (n = 8). Data are pooled from two independent experiments. (f) Histological assessment of H&E-stained jejunal tissues from BG-exposed and unexposed malnourished and control mice. Scale bars, 100 μm (in length). Images and graphs are representative of three independent experiments, five mice per group. (g) Concentrations of IL-6 and MCP-1 released in cultured jejunal tissue sections from BG-exposed and unexposed malnourished and control mice as measured by ELISA. Data are representative of two independent experiments, five mice per group. (h) Box and whisker plot of the concentration of faecal calprotectin in BG-exposed or unexposed malnourished mice (n = 4 per group). All data in standard bar graphs are presented as the means with s.e.m. Box and whisker plots have a middle band that represents the mean, the bottom and top of box are the first and third quartiles, respectively, and the whiskers indicate the range of the data. Statistical analysis was performed using the one-way analysis of variance with post hoc Tukey’s test (**P<0.05, ***P<0.01, ****P<0.001). CON, control; MAL, malnourished; Wks, weeks.
Hypothesized that shifts in lymphocytes in the small intestine could be driving inflammation, and, consequently, the barrier defect observed in the malnourished mice. We first isolated lymphocytes from the small intestine and discovered that the malnourished diet led to an increase in numbers of intraepithelial lymphocytes (IELs) in the duodenum compared with control mice, and that this difference was greater in BG-exposed malnourished mice, although not statistically significant (one-way ANOVA, P = 0.051; Fig. 6a). Within the duodenum IEL compartment, there was a fivefold increase in the numbers of γδ T cells in the malnourished mice and a similar fivefold increase in BG-exposed malnourished mice, compared with controls (Fig. 6b). The majority of the influx of γδ T cells consisted of CD8+ γδ T cells in both BG-exposed and unexposed malnourished mice (Fig. 6c and Supplementary Fig. 8a). CD4+ and CD8+ αβ T-cell numbers in the IEL compartment were not significantly different between groups; however, abundance of double-positive CD4+CD8+ γδ T cells, counterintuitively, decreased in abundance after BG was administered (Supplementary Fig. 13b–d). Much of this influx of IELs in the duodenum was driven by the malnourished diet alone; however, NK-T-cells were significantly more abundant in BG-exposed malnourished mice compared with malnourished mice (Fig. 6d).

Cultured IELs from the duodenum of BG-exposed malnourished mice secreted significantly greater quantities of tumour necrosis factor alpha (TNF-α) and interferon gamma (IFN-γ), as well as a trend for elevated IL-17A compared with unexposed malnourished and control mice, after T-cell specific stimulation (Fig. 6e). These findings suggest that the malnourished diet is sufficient to induce the IEL influx; however, heightened cytokine production by these cells is only triggered with exposure to the BG, and this may contribute to the inflammation and villous blunting observed specifically in the BG-exposed malnourished mice.

Impact of malnutrition and BG on enteric infection. To understand the impact of malnutrition and BG exposure on enteric infection, we utilized the well-studied enteric pathogen Salmonella enterica subsp. typhimurium (S. Typhimurium) as its colonization dynamics in the small intestine are well known30. Furthermore, its greatest burden is in regions of the world where children are also affected by EE31. In mice fed a standard diet and orally infected with S. Typhimurium, very few S. Typhimurium colony-forming units (c.f.u.’s) were detected in the small intestine (Fig. 7a). In contrast, mice infected orally after being fed a
malnourished diet for 3 weeks post weaning had a 3 log fold increase in the burden of *S. Typhimurium* adherent to the jejunum 3 days post infection in BG-exposed and unexposed mice (Fig. 7a). There was a trend towards increased *S. Typhimurium* colonization in the caecum of malnourished mice, but *S. Typhimurium* colonization significantly increased in the caecum of BG-exposed malnourished mice compared with controls (Fig. 7b). Malnourishment led to increased systemic burdens of *S. Typhimurium* with significantly increased colonization in the liver and spleen (Fig. 7c,d). The liver of the infected malnourished mice exhibited striking gross and histological changes. Histological examination showed greater levels of cell necrosis and the presence of fatty lipid droplets (Fig. 7e). Higher levels of inflammatory cytokines, including IL-6, IFN-γ, TNF-α, and MCP-1, were released in the livers of BG-exposed and unexposed malnourished mice 3 days post-infection compared with control, infected mice (Fig. 7f). Overall, malnourished mice were more susceptible to *S. Typhimurium* infection, and this increased susceptibility was maintained on exposure to BG, exhibiting higher bacterial burdens in the intestine, while also invading systemically to greater levels, and inducing a prominent pathological change in the liver.

**Discussion**

These results indicate that a malnourished diet profoundly alters the small intestinal ecosystem, permitting a greater probability of colonization from environmentally acquired microbes. We have identified a set of commensal Bacteroidales and *E. coli* strains that, only in combination with a malnourished diet, are able to reproduce the features of human EE in young mice. To date, it has been unclear which species of bacteria, if any, contribute to EE phenotypes. This observed impact of the Bacteroidales-*E. coli* challenge (BG) is consistent with our hypothesis and previous literature suggesting a microbial aetiology for EE, where environmental ingestion of particular microbes can trigger EE and worsen the impact of childhood malnutrition.

A number of epidemiological and microbiological studies indicate that increased oral bacterial exposure early in life (due to poor environmental sanitation) is behind the aetiology of EE, and in a number of these cases, children exhibit small intestinal bacterial overgrowth. Notably, contamination of the household environment with *E. coli* and Bacteroidales species was common and correlated with increased intestinal permeability, as well as growth rate deficiencies in rural areas of Zimbabwe and Bangladesh. However, this observation remains a correlative one and other microbes could be involved. In the limited studies assessing the small intestine of malnourished children with EE features, breath test and culture-based analysis indicated the presence of small intestinal bacterial overgrowth and Gram-negative bacteria colonizing in the duodenum, consistent with our observations in malnourished mice. To date, no studies have performed high-throughput sequencing analysis on the small intestine of children with EE, thus we do not have information on the prevalence of Bacteroidales and *E. coli* in these patients, or whether other Gram-negative bacteria may be involved.

By utilizing FISH, we were able to visualize an increased abundance of tissue-adherent bacteria triggered by the diet and exacerbated with exposure to Bacteroidales-*E. coli* mixture, this
Unexpectedly, when administered the malnourished diet promotes the abundance of Bacteroidales and Enterobacteriaceae after BG exposure, and also an increased proliferation in the upper small intestine 49, and can impact the growth potential of these microbes in the small intestine. The inflammatory-driving properties of Enterobacteriaceae species have been well documented, and human commensal E. coli species are linked to the development of intestinal inflammation in susceptible hosts 44–46; however, in our model, they are not pathogenic unless combined with commensal E. coli.

The metabolomic analysis of the small intestine revealed for the first time that malnutrition leads to drastic shifts in small intestinal bile acid pool. Specifically, a reduction in tauro-conjugated bile acids was observed in malnourished mice, which are important for fat and nutrient uptake in the small intestine 47, and based on previous reports in patients with tauro-conjugated bile acid deficiency, also could be linked to the observed alterations the luminal bioavailability of the fat-soluble vitamins A, D and E 48. Bile acid changes could also be a factor in the observed resemblance of the microbiota in the malnourished duodenum to that of the ileum. Many studies report that bile acids are one of the major mechanisms to restrict bacteria proliferation in the upper small intestine 49, and can impact host–microbial interactions during intestinal inflammation 50. Thus, the shift in bile acids could be the mechanism that permits for the increased colonization of commensal Bacteroidales and E. coli strains in the small intestine, and may also be behind the ability of S. Typhimurium to proliferate in the malnourished small intestine to higher abundance. The infection with S. Typhimurium highlighted the observations from the BG exposure that malnutrition induces an increased proliferative capacity and colonization potential of Gram-negative bacteria in the small intestine. The increased ability for S. Typhimurium to invade systemically also gives biological relevance to the increase in intestinal permeability observed pre-infection in malnourished mice 51.

The expansion of duodenal IELs observed in malnourished mice seems contradictory to the literature, since protein
malnutrition is known to lead to depressed immune function52. However, in the limited studies where biopsies of the small intestine were taken from children with EE features, there is evidence of a marked infiltrate of lymphocytes to the epithelium, notably γδ T cells and activated CD8+ T cells53, an observation consistent with our results in this model. While BG exposure did not alter total numbers of CD8+ γδ T cells in malnourished mice, IELs isolated from BG-exposed malnourished mice secreted more inflammatory cytokines after stimulation, and this increase in inflammatory potential after stimulation may be driving the histopathology observed after BG exposure. γδ T cells also proliferate to high numbers in coeliac disease, which shares histopathology observed after BG exposure. Mice, IELs isolated from BG-exposed malnourished mice secreted consistent with our results in this model. Furthermore, this mouse model showed increased inflammatory cytokines after stimulation, and this increase in how malnutrition in mice alters the small intestinal microenvironment and utilized these data to develop this model. Furthermore, this mouse model showed increased concentrations of zonulin in serum and in faecal calprotectin, both proposed biomarkers of EE in humans10,12, showcasing the relevance of this model to human disease.

In summary, we show that an orally administered cocktail of specific bacteria are able to reproducibly replicate features of EE only in mice fed a malnourished diet. Utilizing this model, we were able to achieve for the first time an in-depth characterization of the impact of malnutrition on the microbiota, metabolism and immune system in the mammalian small intestine. Future studies will utilize this murine model to gain a deeper understanding of the pathophysiological nature of EE. Furthermore, this model will allow further studies testing the impact of early-life interventions to reverse EE induction.

**Methods**

**Animal studies.** All animal work was done according to the Canadian Council on Animal Care guidelines by utilizing protocols that were approved for use by the Animal Care Committee at the University of British Columbia. Three-week-old female C57BL/6 mice or TLR4−/− mice were ordered for each experiment (Jackson Laboratory, Bar Harbor, ME) and housed in a barrier animal facility at the University of British Columbia with a 12-h light–dark cycle. Upon arrival, mice were randomized and housed into separate groups (4–5 per cage), which were either fed a control diet with 20% protein and 15% fat, similar to one used in previous studies56–58. However, these studies did not characterize the impact of the diet on the microbiota, metabolism and intestinal immune cells, or assess the impact of oral challenge of defined set of bacteria on EE features. In doing these analyses, we utilized a holistic approach to demonstrate novel observations in how malnutrition in mice alters the small intestinal microenvironment and utilized these data to develop this model. Furthermore, this mouse model showed increased concentrations of zonulin in serum and in faecal calprotectin, both proposed biomarkers of EE in humans10,12, showcasing the relevance of this model to human disease.

In summary, we show that an orally administered cocktail of specific bacteria are able to reproducibly replicate features of EE only in mice fed a malnourished diet. Utilizing this model, we were able to achieve for the first time an in-depth characterization of the impact of malnutrition on the microbiota, metabolism and immune system in the mammalian small intestine. Future studies will utilize this murine model to gain a deeper understanding of the pathophysiological nature of EE. Furthermore, this model will allow further studies testing the impact of early-life interventions to reverse EE induction.

**Real-time qPCR analysis for host gene expression.** Real-time qPCR analysis for host gene expression was performed utilizing Quantitite SYBR-Green Mastermix (Qiagen) and group-specific primers for 16S rRNA (Supplementary Table S). PCR was performed on an Applied Biosystems 7500 machine. The abundance of 16S rDNA in the small intestinal sample was determined by comparing the Ct values to the values generated by standard curves. The standard curves were developed from applying the group-specific primers to DNA purified from cultured American Type Culture Collection (ATCC) (Manassas, VA, USA) strains of bacteria with a known value of 165 copies per nanogram of DNA. Results were expressed in total 16S copies per gram of tissue extracted.

**Real-time qPCR analysis for bacterial abundance.** The assessment of bacterial abundance was performed using Quantitate SYBR-Green Mastermix (Qiagen) and group-specific primers for 16S rRNA (Supplementary Table S). PCR was performed on an Applied Biosystems 7500 machine. The abundance of 16S rRNA in the small intestinal sample was determined by comparing the Ct values to the values generated by standard curves. The standard curves were developed from applying the group-specific primers to DNA purified from cultured American Type Culture Collection (ATCC) (Manassas, VA, USA) strains of bacteria with a known value of 165 copies per nanogram of DNA. Results were expressed in total 16S copies per gram of tissue extracted.

**DNA extraction and microbiota analysis.** To assess the composition of the microbiota, sections from the small intestine of malnourished or control-fed mice were homogenized using a bead-beating method (FastPrep instrument, MP Biomedicals, Solon, OH), and total DNA was extracted using a Stool DNA Extraction Kit (Qiagen). 16S rRNA gene fragments were PCR amplified with nucleotide bar-coded primer pairs 27F: 5′-AGAGTTTGATCMTGGCTCAG-3′ and 510R: 5′-GWATACCGCGGCKGCTG-3′. PCR products were gel-purified (QIAquick gel extraction kit, Qiagen). Each amplicon (100 ng) was pooled and sequenced using a 454 Titanium platform (Roche, Branford, CT).

**Bioinformatics.** All sequences were processed using MOTHUR according to the standard operating as previously described59, accessed on 10 July 2013. Quality sequences were obtained by removing sequences with ambiguous bases, a quality read length <200 bases and/or chimeras identified using chimera.uchime. Quality sequences were aligned to the silva bacterial reference alignment and OTUs were generated using a dissimilarity cutoff of 0.03. Sequences were classified using the classify.seqs command with ribosomal database project (RDP) as reference. Inverse Simpson’s diversity index was used to calculate diversity. Differences in microbial communities between groups and intestinal sites were investigated using the phylogenies-based weighted UniFrac distance metric. The Bray Curtis index was used as a measure of similarity in microbial composition. Diversity, similarity and abundance of bacterial OTUs and families were compared using the Mann–Whitney U-test or Student’s t-test, and the Bonferroni correction was applied in cases of multiple comparisons.

**Metabolite extraction.** Mouse small intestine digesta samples (luminal contents) were collected, weighed and homogenized in water followed by addition of acetonitrile for metabolite extraction. After vortex mixing, 30s sonication and centrifugation, the clear supernatant, the collection was dissolved under gentle nitrogen gas flow. The dried metabolite residues were dissolved in 40% acetonitrile containing 0.01% formic acid, 4 μl g⁻¹ of the raw material. After 15 s vortex and 30 s sonication, each sample was further diluted 1:2 with water. Twelve-microlitre aliquots were injected for untargeted metabolic fingerprinting by UPLC–FTMS.

**UPLC–FTMS.** A Waters Acuity UPLC system coupled to a Thermo LTQ-Orbi-trap Velos Pro mass spectrometer was equipped for metabolic analysis. The mass spectrometer was equipped with a heated electrospray ionization (ESI) source and was operated in the FT mass spectrometry (MS) scan mode with mass resolution of 60,000 full-width at half-maximum at m/z 400. The m/z detection range was 80–1,200. A Waters BEH C18 UPLC column (2 mm × 50 mm, 1.7 μm) was used for chromatographic separation with the mobile phases being water–formic acid (50:50) and acetonitrile (0.01% formic acid in water) at 0.2 ml min⁻¹. The flow rate of the binary solvent elution gradient was 5–40% B in 6 min and 40–100% B in 15 min. The mobile phase was kept at 100% B for 2 min before column equilibration with 5% B
for 4 min between injections. The column temperature was 30 °C and the flow rate was 0.3 ml min⁻¹. Two liquid chromatography (LC–MS) runs per sample were performed, except for the negative ion mode, which was duplicated. Retention time shift correction was applied to ensure mass accuracy throughout LC–MS runs.

Data processing. The positive and negative ion UPLC–FTMS data sets were converted to the mzXML files using the MSConvert tool (ProteoWizard) and processed using the XCMS package (http://metlin.scripps.edu/xcms/) in the R platform (http://www.r-project.org/). For each data set, peak detection and integration were performed using the centWave algorithm⁶⁰. Retention time shift correction was achieved considering at least 200 peak groups. After two iterations of peak grouping, imputation of missing data was performed by returning to the raw spectral data and integrating the areas of the missing peaks using the ‘fillPeaks’ algorithm. Finally, a data matrix was generated from each UPLC–MS data set and imported into Microsoft Excel. After peak de-isotoping and removal of the significant background noise signals observed in each UPLC–MS blank run, the data were saved as two-dimensional (m/z retention time versus peak area) matrices amenable to subsequent statistics and further data analysis.

Metabolic pathway analysis. The resulting m/z values after data processing were searched against the METLIN database (http://metlin.scripps.edu/metabolometrix.php?return=yes) for metabolic identification. The metabolic pathways that the identified metabolites were involved in were matched using the Kyoto Encyclopedia of Genes and Genomes pathway tool, permitting the over-representation analysis to determine over-represented pathways in each treatment group. Multi-variate statistical analysis (random forest, PCA) and over-representation analysis were carried out using the Metaboanalyst version 2.5 software (http://www.metaboanalyst.ca/MetaboAnalyst/) based on recommendations from previous published protocols⁶¹. The data from the positive and negative ion mode detection were treated separately in the statistical analysis. In the analysis, missing values were assumed to have no deviation. By default, the level of detection in the parameters. Each data set was normalized to the median of each of the observed peaks, following a log transformation of the data. The Welch’s two-sample t-test was used to determine significant changes between groups (P < 0.05; fold change > 2).

Vitamin-targeted metabolomics. The samples were analysed by UPLC–MRM (multiple-reaction-monitoring)/MS on a Dionex UltiMate 3400 RSLC system coupled to an AB Sciex 4000 QTRAP triple-quadrupole mass spectrometer equipped with an electrospray ionization source. The standard substances of vitamin A (retinol, retinal and retinoic acid), B₁ (thiamine), B₂ (riboflavin), B₃ (niacinamide), B₆ (pyridoxamine, pyridoine, pyridoxal and pyridoxal-monophosphate), B₇ (biotin), B₁₂ (folic acid), D₁, D₃, E (α-tocopherol, β-tocotrienol), K₁ and K₂ were purchased either from Sigma-Aldrich or from Cayman Chemicals Inc. The MRM transitions of individual analytes were optimized using the centWave algorithm⁶⁰. Retention time shift correction was achieved considering at least 200 peak groups. After two iterations of peak grouping, imputation of missing data was performed by returning to the raw spectral data and integrating the areas of the missing peaks using the ‘fillPeaks’ algorithm. Finally, a data matrix was generated from each UPLC–MS data set and imported into Microsoft Excel. After peak de-isotoping and removal of the significant background noise signals observed in each UPLC–MS blank run, the data were saved as two-dimensional (m/z retention time versus peak area) matrices amenable to subsequent statistics and further data analysis.

Bile acid-targeted metabolomics. Each sample was homogenized in LC–MS grade water at a ratio of 150 μl per 10 mg raw material and with the aid of two 3-mm stainless steel metal balls. Bile acids were extracted by addition of acetonitrile at a ratio of 350 μl per 10 mg raw material followed by vortexing and sonication (1 min) in an ice–water ultrasonic bath. The samples were centrifuged. Twenty microtiter of the supernatants were precisely taken out and mixed with a predefined mix of 14 deuterium-labelled bile acids as the internal standards. The mixtures were subjected to phase-phase extraction with 1 ml methanol, 1 ml water, and 0.2 ml was injected into the Bruker Scion 456 gas chromatograph, using a Stabilwax-DA 30 m × 0.53 mm × 0.5 μm column (Restek, Bellefonte, PA). A standard solution containing acetic acid, propionic acid, isobutyric acid, butyric acid, isovaleric acid, valeric acid and capric acid, combined with internal standard, was injected in every run. The programmed temperature vaporizing (PTV) injector and flame ionization detector (FID) detector temperatures were held at 250 °C for the entire run. The oven was started at 80 °C and immediately ramped to 210 °C at 45 °C per min, where it was held for 5.11 min. Total run time was 8.00 min. Helium was used at a constant flow of 20.00 ml min⁻¹.

SCFA analysis. Small intestinal samples were weighed and combined with 25% phosphoric acid, vortexed and centrifuged until a clear supernatant was obtained. Supernatants were submitted for gas chromatography (GC) analysis to the Department of Agricultural, Food and Nutritional Science of the University of Alberta. Samples were analysed as previously described with modifications. Briefly, aliquots were combined with 4% methyl-vanillin and 1 ml of 2% methanol. Ten microlitres of residues were precisely taken out and mixed with a predefined mix of 14 deuterium-labelled bile acids as the internal standards. The mixtures were subjected to phase-phase extraction with 1 ml methanol, 1 ml water, and 0.2 ml was injected into the Bruker Scion 456 gas chromatograph, using a Stabilwax-DA 30 m × 0.53 mm × 0.5 μm column (Restek, Bellefonte, PA). A standard solution containing acetic acid, propionic acid, isobutyric acid, butyric acid, isovaleric acid, valeric acid and capric acid, combined with internal standard, was injected in every run. The programmed temperature vaporizing (PTV) injector and flame ionization detector (FID) detector temperatures were held at 250 °C for the entire run. The oven was started at 80 °C and immediately ramped to 210 °C at 45 °C per min, where it was held for 5.11 min. Total run time was 8.00 min. Helium was used at a constant flow of 20.00 ml min⁻¹.

Bacterial strains. The S. Typhimurium strain SL1344 was used, as this strain is virulent to mice and resistant to streptomycin⁶⁰. Colonies of SL1344 were grown on Luria-Bertani (LB) agar with 100 μg ml⁻¹ streptomycin. Colonies were selected and incubated for growth in LB broth at 37 °C and plated on Luria-Bertani (LB) agar. All commensal bacterial strains that are used in our study (Supplementary Table 4) were of human origin, and were provided to us by Emma Allen-Vercoe (University of Guelph) or ordered from the ATCC and DSMZ culture collections (Leibniz Institute, Germany). Commensal bacteria strains were all grown on fastidious anaerobe agar (FAA; LaLB) in anaerobic conditions.

Bacterial cocktail preparation and inoculation. In an anaerobic chamber, bacterial cultures from frozen stock were first plated on FAA, and subsequently pure cultures were selected and mixed together at a 1:1 ratio in sterile, reduced PBS. Bacterial mixtures in PBS were removed from the anaerobic chamber and were immediately transported to the animal facility to carry out gavage experiments. The volume of the mixture received per mouse was 100 μl at a concentration of 10⁵ cells per ml. The concentration of the mixture in cell per ml was carried out using a ultraviolet spectrometer, and gavage doses were confirmed after by back-titrating the inocula on FAA. In experiments using heat-inactivated bacteria, the bacterial mixture was incubated for 2 h at 80 °C and plated on FAA to confirm the absence of growth.

Intestinal epithelial cell (IEC) and IEL isolation. The upper 5 cm of the small intestine of mice was excised, attached fat and Peyer’s patches were removed, and tissues were cut longitudinally to further remove luminal contents by washing with ice-cold PBS. Epithelial cells were isolated using a PBS buffer containing 1 mM EDTA, 1 mM dithiothreitol and 5% fetal bovine serum (FBS), shaking at 37 °C for 10 min. This supernatant was filtered and combined with RPMI 1640 and centrifuged at 1,500 r.p.m. to isolate the IECs. The intact intestinal tissue was resuspended in 2 ml of Hank’s balanced salt solution with 5% FBS and further purified using a 40% Percoll gradient and were resuspended in RPMI 1640 with 5% FBS.

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Flow cytometry. Purified lymphocytes were counted using an automatic cell counter (Countess, BD Biosciences). Cells were stained with a 1:200 dilution of fluorochrome-conjugated antibodies against CD45 (clone 30F11-11, CD3ε, eBilo500-A2), CD4 (RM4–5), CD8 (53-6.7), NK1.1 (PKI36) and γδ TCR (eBiloGL3) (eBioscence), and their populations were analysed by an LSR II flow cytometer (BD Biosciences) using software packages from CellQuest and FlowJo version 8.7.

Western blotting. Total IECs were lysed with MP-40 and protease inhibitor cocktail (Roche Diagnostics, Basel, Switzerland) for 10 min on ice. The resulting lysis solution was centrifuged at 13,000 r.p.m. (16,360g) for 10 min, and the supernatant was collected. The protein concentration of each sample supernatant was quantified using a bichinchoninic acid assay (Sigma-Aldrich). Equal concentrations of the samples were electrophoresed through a 12% SDS–polyacrylamide gel and transferred onto nitrocellulose membrane (Millipore, Darmstadt, Germany). Membranes were probed with 1/500 dilutions of anti-CLDN2 and anti-actin primary antibodies (Invitrogen, Carlsbad, CA) and then a 1/1,000 dilution of a secondary anti-rabbit/mouse horseradish peroxidase antibody.

Fecal calprotectin determination. Fecal content of mice was collected, weighed, homogenized in PBS, centrifuged at 15,000 r.p.m. (16,360g) and the resulting supernatant was stored at −80 °C. The concentration of calprotectin in feces was performed on the fecal supernatants using a S0008A/S0010A calprotectin enzyme-linked immunosorbent assay (ELISA) kit as per manufacturer’s instructions (Hycult Biotechnology). The resulting concentration of calprotectin was determined by assaying the OD 450 nm using a plate reader (Tecan), comparing with a standard curve of known concentrations of calprotectin.

Total IgA determination. Total IgA concentrations were measured in the feces and small intestinal luminal content using a commercially available ELISA kit (eBioscience, cat. no. 88-50450-22). Fecal and small intestinal homogenates were weighed and diluted to 1:1,000 for the assay. Sample concentrations were determined by assaying the OD 450 nm using a plate reader (Tecan), compared with a standard curve, and normalized to weight of the fecal content.

Infection models of mice. A frozen stock of S. Typhimurium (SL1344) was first plated on LB agar with 100 μg/mL streptomycin and a single colony was subsequently inoculated into LB broth and grown, with shaking, for 18 h at 37 °C. Using this culture, mice were infected by oral gavage with 100 μl of complete growth medium without selection antibiotics at an OD600 of 0.5. Twenty-four hours after injection, the supernatant was cleared by centrifugation at 740 × g for 10 min at room temperature and assayed for NFκB-responsive, secreted alkaline phosphatase activity using QUANTI-Blue reagent (InvivoGen). A650 was assessed for four hours post stimulation, the supernatant was cleared by centrifugation at 740 × g for 10 min at room temperature and assayed for NFκB-responsive, secreted alkaline phosphatase activity using QUANTI-Blue reagent (InvivoGen). A650 was assessed at 37 °C. The averages of the cell-free medium and the Null2 cell controls were subtracted to obtain TL4 activation of HEK-Blue hTLR4 cells.

IEC and IEL cytokine quantification analysis. For ex vivo quantification of cytokine secretion in the small intestinal tissues, jejunal slices of tissue were washed with complete tissue culture media (RPMI, 10% FBS, 1% glutamine and 1% of 1 penicillin/streptomycin) and cultured in 1 ml of the same media in 24-well plates for 24 h at 37 °C, 5% carbon dioxide. The supernatants were serially diluted in PBS, plated on LB agar with 100 μg/mL streptomycin and incubated overnight at 37 °C. Bacterial colonies were enumerated the following day to determine the c.f.u. per gram of tissue. To determine the cytokine concentrations, tissue homogenates were centrifuged three times at 15,000 × g for 21,660 rpm for 20 min at 4 °C to remove cell debris and the resulting supernatants were aliquoted and stored at −80 °C. Cytokine levels in the homogenates were measured using a Cytometric Bead Array Mouse Inflammation Kit (BD Biosciences), according the manufacturer’s recommendations.

Fluorescence in situ hybridization. A 1-cm section of the jejunum was excised from mice, immediately submerged into methanol–Carney’s fixative for 2 h at 4 °C. Following fixation, fixed tissues were paraffin embedded, cut with a microtome to a thickness of 6–10 μm and transferred to glass slides. The slides were let dry overnight and the tissues were then fixed with 4% paraformaldehyde for 10 min at room temperature before permeabilization with a solution of 1 mg/mL lysozyme from Eggwhite at 37 °C for 1 h. The tissues were then pre-incubated for 1 h at 42 °C in hybridization buffer (20 mM Tris-HCl, pH 8.0, 0.9 M NaCl, 0.01% SDS and formamide, depending on the probe; Sigma-Aldrich), and then the slides were let dry at 37 °C and incubated overnight at 42 °C with the respective probe linked to Alexa Fluor 647 (Invitrogen) in the corresponding hybridization buffer at a probe concentration of 50 nM. The following probes were used: all-Bacterial probe (Eub338), Firmicutes (equimolar mixture of LGC354A, B and C), Bacteroidetes (Bac203) and Gamma-Proteobacteria (Gam42a). Details on the probes can be found at ProbeBase and the sequences of the probes used are listed in Supplementary Table 6. The tissues were then incubated for 20 min with Alexafluor 488-Phallolidin (130 nM) and 4,6-diamidino-2-phenylindole (1 μg/mL) before acquiring fluorescent images. The slides were imaged in a Zeiss Axioscop using a Zeiss AxiosCam MRm camera and a Zeiss Inverted Observer Z1 microscope (Carl Zeiss Microimaging GmbH, Germany). Images were captured in a Zeiss AxiosCam MRm camera and a Zeiss Inverted Observer Z1 microscope (Carl Zeiss Microimaging GmbH, Germany). Images were captured using AxioVision Rel. 4.7.1. Images were evaluated using Image J software. The area of the villi was evaluated using the “region of interest” tool in Image J software. The area was evaluated in the middle to the lower part of the villi. The percentage of invasion was calculated based on the evaluation of each villus. The statistical analysis of the data was performed using GraphPad Prism Software Version 6.00 (GraphPad Software, San Diego California USA; www.graphpad.com). Statistical significance was given as **P value <0.001, *P value <0.01 and P value <0.05; NS (not significant) P value >0.05. The results are expressed as the mean value with s.e.m., unless otherwise indicated.

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Author contributions
E.M.B conceived the project, wrote the manuscript and performed, analysed or designed all experiments in this manuscript. M.W., B.P.W., P.V., J.H., L.A.R., M.-C.A., M.U., R.S. and O.P. performed the experiments and analysed the data. C.H.B. and P.J.S. contributed analytical tools and supervised participants. B.B.F. helped conceive the project, write the manuscript mentored and supervised its participants.

Additional information
Accession codes. The microbiome 454 pyrosequencing data have been deposited in the NCBI SRA database under the Project Accession ID SRP058646. The metabolomics data obtained in this study have been deposited in the National Institutes of Health Common Fund’s Data Repository and Coordinating Center website (http://www.metabolomicsworkbench.org) under accession code PR000182.

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