Contributions of microbiome and mechanical deformation to intestinal bacterial overgrowth and inflammation in a human gut-on-a-chip

The MIT Faculty has made this article openly available. Please share how this access benefits you. Your story matters.

<table>
<thead>
<tr>
<th>Citation</th>
<th>Kim, Hyun Jung et al. “Contributions of Microbiome and Mechanical Deformation to Intestinal Bacterial Overgrowth and Inflammation in a Human Gut-on-a-Chip.” Proceedings of the National Academy of Sciences 113,1 (January 2016): E7–E15. © 2017 National Academy of Sciences</th>
</tr>
</thead>
<tbody>
<tr>
<td>As Published</td>
<td><a href="http://dx.doi.org/10.1073/pnas.1522193112">http://dx.doi.org/10.1073/pnas.1522193112</a></td>
</tr>
<tr>
<td>Publisher</td>
<td>National Academy of Sciences (U.S.)</td>
</tr>
<tr>
<td>Version</td>
<td>Final published version</td>
</tr>
<tr>
<td>Accessed</td>
<td>Sun Nov 05 15:37:49 EST 2017</td>
</tr>
<tr>
<td>Citable Link</td>
<td><a href="http://hdl.handle.net/1721.1/111591">http://hdl.handle.net/1721.1/111591</a></td>
</tr>
<tr>
<td>Terms of Use</td>
<td>Article is made available in accordance with the publisher's policy and may be subject to US copyright law. Please refer to the publisher's site for terms of use.</td>
</tr>
<tr>
<td>Detailed Terms</td>
<td></td>
</tr>
</tbody>
</table>
Contributions of microbiome and mechanical deformation to intestinal bacterial overgrowth and inflammation in a human gut-on-a-chip

Hyun Jung Kim,a,1, Hu Li,2, James J. Collins,a,b,c,d,e,f,3 and Donald E. Ingber,a,g,h,3

aWyss Institute for Biologically Inspired Engineering at Harvard University, Boston, MA 02115; bInstitute for Medical Engineering and Science, Massachusetts Institute of Technology, Cambridge, MA 02139; cDepartment of Biological Engineering, Massachusetts Institute of Technology, Cambridge, MA 02139; dSynthetic Biology Center, Massachusetts Institute of Technology, Cambridge, MA 02142; eHarvard-Massachusetts Institute of Technology Program in Health Sciences and Technology, Cambridge, MA 02139; fVascular Biology Program, Boston Children’s Hospital, Harvard Medical School, Boston, MA 02115; and gHarvard John A. Paulson School of Engineering and Applied Sciences, Harvard University, Cambridge, MA 02138

Contributed by James J. Collins, November 15, 2015 (sent for review August 18, 2015; reviewed by David J. Beebe and Rustem F. Ismagilov)

A human gut-on-a-chip microdevice was used to coculture multiple commensal microbes in contact with living human intestinal epithelial cells for more than a week in vitro and to analyze how gut microbiome, inflammatory cells, and peristalsis-associated mechanical deformations independently contribute to intestinal bacterial overgrowth and inflammation. This in vitro model replicated results from past animal and human studies, including demonstration that probiotic and antibiotic therapies can suppress villus injury induced by pathogenic bacteria. By cessation peristalsis-like motions while maintaining luminal flow, lack of epithelial deformation was shown to trigger bacterial overgrowth similar to that observed in patients with ileus and inflammatory bowel disease. Analysis of intestinal inflammation on-chip revealed that immune cells and lipopolysaccharide endotoxin together stimulate epithelial cells to produce four proinflammatory cytokines (IL-8, IL-6, IL-1β, and TNF-α) that are necessary and sufficient to induce villus injury and compromise intestinal barrier function. Thus, this human gut-on-a-chip can be used to analyze contributions of microbiome to intestinal pathophysiology and dissect disease mechanisms in a controlled manner that is not possible using existing in vitro systems or animal models.

Various types of inflammatory bowel disease (IBD), such as Crohn’s disease and ulcerative colitis, involve chronic inflammation of human intestine with mucosal injury and villus destruction (1), which is believed to be caused by complex interactions between gut microbiome (including commensal and pathogenic microbes) (2), intestinal mucosa, and immune components (3). Suppression of peristalsis also has been strongly associated with intestinal pathology, inflammation (4, 5), and small intestinal bacterial overgrowth (5, 6) in patients with Crohn’s disease (7) and ileus (8). However, it has not been possible to study the relative contributions of these different potential contributing factors to human intestinal inflammatory diseases, because it is not possible to independently control these parameters in animal studies or in vitro models. In particular, given the recent recognition of the central role of the intestinal microbiome in human health and disease, including intestinal disorders (2), it is critical to incorporate commensal microbes into experimental models; however, this has not been possible using conventional culture systems.

Most models of human intestinal inflammatory diseases rely either on culturing an intestinal epithelial cell monolayer in static Transwell culture (9) or maintaining intact explanted human intestinal mucosa ex vivo (10) and then adding live microbes and immune cells to the apical (luminal) or basolateral (mucosal) sides of the cultures, respectively. These static in vitro methods, however, do not effectively recapitulate the pathophysiology of human IBD. For example, intestinal epithelial cells cultured in Transwell plates completely fail to undergo villus differentiation, produce mucus, or form the various specialized cell types of normal intestine. Although higher levels of intestinal differentiation can be obtained using recently developed 3D organoid cultures (11), it is not possible to expose these cells to physiological peristalsis-like motions or living microbiome in long-term culture, because bacterial overgrowth occurs rapidly (∼1 d) compromising the epithelium (12). This is a major limitation because establishment of stable symbiosis between the epithelium and resident gut microbiome as observed in the normal intestine is crucial for studying inflammatory disease initiation and progression (13), and mechanical deformations driven by peristalsis are required to both maintain normal epithelial differentiation (14) and restrain microbial overgrowth in the intestine in vivo (15).

Significance

The main advance of this study is the development of a microengineered model of human intestinal inflammation and bacterial overgrowth that permits analysis of individual contributors to the pathophysiology of intestinal diseases, such as ileus and inflammatory bowel disease, over a period of weeks in vitro. By studying living human intestinal epithelium, with or without vascular and lymphatic endothelium, immune cells, and mechanical deformation, as well as living microbiome and pathogenic microbes, we identified previously unknown contributions of specific cytokines, mechanical motions, and microbiome to intestinal inflammation, bacterial overgrowth, and control of barrier function. We provide proof-of-principle to show that the microfluidic gut-on-a-chip device can be used to create human intestinal disease models and gain new insights into gut pathophysiology.

Author contributions: H.J.K., J.J.C., and D.E.I. designed research; H.J.K. performed research; H.J.K., H.L., J.J.C., and D.E.I. analyzed data; and H.J.K. and D.E.I. wrote the paper.

Conflict of interest statement: D.E.I. is a founder of Emulate Inc., holds equity in the company, and chairs its Scientific Advisory Board.

Data deposition: Human Gene 1.0 ST Array mRNA data of Caco-2 cells cultured in various conditions were deposited to the Gene Expression Omnibus (GEO) database, www.ncbi.nlm.nih.gov/geo (accession no. GSE65790).

1Present address: Department of Biomedical Engineering, The University of Texas at Austin, Austin, TX 78712.

2Present address: Center for Individualized Medicine, Department of Molecular Pharmacology and Experimental Therapeutics, Mayo Clinic, Rochester, MN 55905.

3To whom correspondence may be addressed. Email: jimjc@mit.edu or don.ingber@wyss.harvard.edu.

This article contains supporting information online at www.pnas.org/lookup/suppl/doi:10.1073/pnas.1522193112/-/DCSupplemental.

www.pnas.org/cgi/doi/10.1073/pnas.1522193112
Thus, we set out to develop an experimental model that would overcome these limitations. To do this, we adapted a recently described human gut-on-a-chip microfluidic device that enables human intestinal epithelial cells (Caco-2) to be cultured in the presence of physiologically relevant luminal flow and peristalsis-like mechanical deformations, which promotes formation of intestinal villi lined by all four epithelial cell lineages of the small intestine (absorptive, goblet, enteroendocrine, and Paneth) (12, 16). These villi also have enhanced barrier function, drug-metabolizing cytochrome P450 activity, and apical mucus secretion compared with the same cells grown in conventional Transwell cultures, which made it possible to coculture a probiotic gut microbe (Lactobacillus rhamnosus GG) in direct contact with the intestinal epithelium for more than 2 wk (12), in contrast to static Transwell cultures (17) or organoid cultures (11) that lose viability within hours under similar conditions. In the present study, we leveraged this human gut-on-a-chip to develop a disease model of small intestinal bacterial overgrowth (SIBO) and inflammation. We analyzed how probiotic and pathogenic bacteria, lipopolysaccharide (LPS), immune cells, inflammatory cytokines, vascular endothelial cells and mechanical forces contribute individually, and in combination, to intestinal inflammation, villus injury, and compromise of epithelial barrier function. We also explored whether we could replicate the protective effects of clinical probiotic and antibiotic therapies on-chip to demonstrate its potential use as an in vitro tool for drug development, as well as for dissecting fundamental disease mechanisms.

Results

Establishing a Complex Human Intestinal Model in Vitro. To study interactions between cultured microbiome and human intestinal epithelial cells in an organ-like context and mimic the human intestinal microenvironment undergoing injury and inflammation, we leveraged a gut-on-a-chip microfluidic device (12, 16) composed of optically clear, flexible polydimethylsiloxane (PDMS) polymer with three parallel hollow microchannels (Fig. 1A). The central channel is split into an upper (lumen) and lower (capillary) channel by a flexible, extracellular matrix (ECM)-coated PDMS membrane containing an array of pores (10 μm in diameter) lined by human Caco-2 intestinal epithelial cells. The cells are exposed to trickling flow of culture medium (30 μL/h, equivalent to 0.02 dyne/cm² shear stress) through the upper and lower microchannels and to cyclic peristalsis-like mechanical deformations (10% in cell strain, 0.15 Hz in frequency) (Movie S1) generated by applying cyclic suction to hollow side chambers that rhythmically deflect the central membrane with the attached cells (Fig. 1B). By mimicking the physical microenvironment of the intestine in this manner, the cultured human intestinal epithelial cells spontaneously form intestinal villi (Fig. 1C and Movie S1) lined by a highly polarized, differentiated columnar epithelium with a tight apical brush border (Fig. 1D) that appear similar in form to living intestinal villi in vivo (17).

To more clearly demonstrate the influence of culture conditions on the differentiation state of the Caco-2 cells, we performed genome-wide gene profiling analysis of the cells cultured as a static monolayer in a conventional Transwell insert, in the mechanically active gut-on-a-chip microdevice, or on-chip when cocultured with living commensal gut microbes. The commensal microbes used were contained in a therapeutic formulation of probiotic bacteria (VSL#3) that has been studied in humans (18), which contains eight strains of beneficial probiotic bacteria, including six that were originally isolated from human gut microbiome (Lactobacillus acidophilus, Lactobacillus plantarum, Lactobacillus paracasei, Bifidobacterium breve, Bifidobacterium longum, and Bifidobacterium infantis). Gene microarray analysis revealed that Caco-2 cells cultured in the gut-on-a-chip exhibit a significantly distinct gene expression profile ($P < 0.00001$) across 22,097 genes compared with the cells cultured in a static Transwell, and their phenotype again changed significantly ($P < 0.00001$) when

Fig. 1. The human gut-on-a-chip microfluidic device and changes in phenotype resulting from different culture conditions on-chip, as measured using genome-wide gene profiling. (A) A photograph of the device. Blue and red dyes fill the upper and lower microchannels, respectively. (B) A schematic of a 3D cross-section of the device showing how repeated suction to side channels (gray arrows) exerts peristalsis-like cyclic mechanical strain and fluid flow (white arrow) generates a shear stress in the perpendicular direction. (C) A DIC micrograph showing intestinal basal crypt (red arrow) and villi (white arrow) formed by human Caco-2 intestinal epithelial cells grown for ~100 h in the gut-on-a-chip under medium flow (30 μL/h) and cyclic mechanical stretching (10%, 0.15 Hz). (Scale bar, 50 μm.) (D) A confocal immunofluorescence image showing a horizontal cross-section of intestinal villi similar to those shown in Fig. 1C, stained for F-actin (green) that labels the apical brush border of these polarized intestinal epithelial cells (nuclei in blue). (Scale bar, 50 μm.) (E) Hierarchical clustering analysis of genome-wide transcriptome profiles (Top) of Caco-2 cells cultured in the static Transwell, the gut-on-a-chip (with fluid flow at 30 μL/h and mechanical deformations at 10%, 0.15 Hz) (Gut Chip), or the mechanically active gut-on-a-chip cocultured with the VSL#3 formulation containing eight probiotic gut microbes (Gut Chip + VSL#3) for 72 h compared with normal human small intestinal tissues (Duodenum, Jejunum, and Ileum; microarray data from the published GEO database). The dendrogram was generated based on the averages calculated across all replicates, and all branches in the cluster have the approximately unbiased (AU) $P$ value equal to 100. The y axis next to the dendrogram represents the metric for Euclidean distance between samples. Corresponding pseudocolored GEDI maps analyzing profiles of 650 metagenomes between samples described above (Bottom).
that the transcriptomes obtained from Caco-2 cells cocultured with multiple normal gut microbes (VSL#3) in the gut-on-a-chip are distinct from the profiles exhibited by the same cells cultured in the other conditions without microbes ($P < 0.00001$), and they are more similar to normal human ileum than to duodenum or jejunum ($P < 0.00001$) (Fig. 1E and Fig. S1B).

Reconstituting Human Intestinal Inflammation and Injury On-Chip.

We explored whether this system could be used to model human intestinal inflammation in vitro. Some of the key hallmarks of intestinal inflammatory diseases are the destruction of intestinal villi and associated compromise of the intestinal permeability barrier, which are believed to result from complex pathological interplay between the intestinal epithelium, gut microbes, and immune cells, as well as from changes in luminal flow resulting from altered peristalsis (1, 3).

To explore how noncommensal microbes contribute to intestinal inflammation and gut injury, we cocultured Gram-negative Escherichia coli bacteria with the intestinal villus epithelium on-chip. When a nonpathogenic laboratory strain of green fluorescent protein-labeled E. coli (GFP-EC) was allowed to adhere to the apical (luminal) surface of villi for ~1.5 h under static conditions, these bacteria subsequently colonized and spontaneously inhabited regions overlying the crypts localized between adjacent villi (Fig. S2). We next introduced a pathogenic strain (serotype O124:NM) of enteroinvasive E. coli (EIEC) that causes intestinal cell destruction and extreme diarrhea in humans (20) into the lumen of the epithelial channel. These EIEC bacteria initially distributed to similar locations between neighboring villi, but then they rapidly overgrew across the whole apical surface of villi within 24 h (Fig. 2A and Movie S2), and planktonic cells appeared in the culture medium within the channel lumen. Furthermore, when the gut-on-a-chip devices were maintained in coculture with GFP-EC for 4 additional days with cyclic mechanical deformation, the GFP-EC bacteria failed to alter normal intestinal barrier function as indicated by maintenance of a relatively constant transepithelial electrical resistance (TEER). When we added LPS endotoxin ($15 \mu g/mL$) isolated from pathogenic E. coli (serotype O111:B4) that elicits strong immune responses when administered in vivo (21), it similarly failed to disrupt the intestinal barrier on-chip (Fig. 2B), which mimics results previously observed in both other in vitro studies (22, 23) and animal models (24). There also was no detectable evidence of intestinal injury when analyzed using either phase contrast or fluorescence confocal microscopy (Fig. 2C). In contrast, when EIEC cells were added to the apical surface of epithelium, intestinal barrier function (Fig. 2B, Left) and normal villus architecture were completely lost within 24–36 h (Fig. 2C).

**Fig. 2.** Reconstitution of pathological intestinal injury induced by interplay between nonpathogenic or pathogenic enteroinvasive E. coli bacteria or LPS endotoxin with immune cells. (A) DIC images showing that the normal villus morphology of the intestinal epithelium cultured on-chip (Control) is lost within 24 h after EIEC (serotype O124:NM) are added to the apical channel of the chip (+EIEC; red arrows indicate bacterial colonies). (B) Effects of GFP-EC, LPS (15 \mu g/mL), EIEC, or no addition (Control) on intestinal barrier function (Left). Right shows the TEER profiles in the presence of human PBMCs (+PBMC). GFP-EC, LPS, and EIEC were added to the apical channel (intestinal lumen) at 4, 12, and 35 h, respectively, and PBMCs were subsequently introduced through the lower capillary channel at 44 h after the onset of experiment (0 h) ($n = 4$). (C) Morphological analysis of intestinal villus damage in response to addition of GFP-EC, LPS, and EIEC in the absence (−PBMC) or the presence of immune components (+PBMC). Schematics (experimental setup), phase contrast images (horizontal view, taken at 57 h after onset), and fluorescence confocal micrographs (vertical cross-sectional views at 83 h after onset) were sequentially displayed. F-actin and nuclei were coded with magenta and blue, respectively. (D) Quantification of intestinal injury evaluated by measuring changes in lesion area (Top; $n = 30$) and the height of the villi (Bottom; $n = 50$) in the absence (white) or the presence (gray) of PBMCs. Intestinal villi were grown in the gut-on-a-chip under trickling flow (30 \mu L/h) with cyclic deformations (10%, 0.15 Hz) during the preculture period for ~100 h before stimulation (0 h, onset). Asterisks indicate statistical significance compared with the control at the same time point ($^*P < 0.001$, $^{**}P < 0.05$). (Scale bars, 50 \mu m.)
To mimic the chronically inflamed microenvironment of patients with intestinal inflammatory diseases, such as IBD, in which increased numbers of immune cells are recruited from the lamina propria (13), we carried out similar studies where isolated human peripheral blood mononuclear cells (PBMCs) were introduced into the lower capillary channel of the device and allowed to interact with the lumen without flow for 2 h (SI Text, Inflammation Study). PBMCs contain a mixed population of innate (e.g., monocytes and granulocytes) and adaptive (e.g., lymphocytes) immune cells (Fig. S3) and have been used previously in various in vitro models of human intestinal inflammation (9, 23). Addition of PBMCs did not induce any detectable injury on their own; however, PBMCs synergized with either GFP-EC or LPS eliciting a significant decrease of intestinal barrier and major destruction and shortening of the intestinal villi (Fig. 2 B–D). Also, damage of the intestinal epithelium provoked by EIECs was accelerated and exacerbated by the presence of PBMCs (Fig. 2 B–D).

Another key hallmark of intestinal inflammatory diseases is the production and secretion of inflammatory cytokines, which have been previously implicated in development of IBD (3, 25). We first analyzed production of a panel of cytokines relevant to intestinal inflammation in the gut-on-a-chip exposed to either LPS, PBMCs (Fig. 3 A), or GFP-EC (Fig. S4 A). LPS or GFP-EC alone did not alter production of any of these cytokines, and the PBMCs merely increased abluminal secretion of interleukin (IL)-8 into the lower capillary channel (Fig. 3 A). In contrast, when PBMCs were combined with either LPS (Fig. 3 A) or GFP-EC (Fig. S4 A), there was a significant increase ($P < 0.01$) in polarized secretion of IL-1β, IL-6, and TNF-α into the basal channel, which is consistent with past observations (9), and there was strong activation of IL-8 production and toll-like receptor 4 (TLR4) expression in the epithelium stimulated with PBMCs with LPS (Fig. S4 B). Polarized basal secretion of cytokines should stimulate recruitment of additional circulating immune cells in vivo, which could further enhance cytokine production, thereby creating a positive feedback loop that might augment or exacerbate the inflammatory response. To investigate whether the detected cytokines were sufficient to induce damage to intestinal epithelial cells, we added purified IL-1β, IL-6, IL-8, and TNF-α alone (Fig. S4 C) or in different combinations (Fig. 3 B and Fig. S4 D) at the same concentrations they were produced on-chip (Fig. 3 A). None of these individual cytokines induced villus injury (Fig. S4 C), nor did addition of a mixture of IL-1β, IL-6, and TNF-α (Fig. S4 D), even though these cytokines have been reported to contribute to intestinal damage in Crohn’s disease and ulcerative colitis in vivo (25). However, when these three cytokines were combined with IL-8, we were able to fully recapitulate the villus injury response observed by combined administration of LPS and PBMCs (Fig. 3 A). In addition, coadministration of a blocking monoclonal IL-8 antibody completely prevented villus disruption in the presence of all four cytokines (Fig. 3 B), even though addition of IL-8 alone was not sufficient to induce injury (Fig. S4 C). Thus, our results suggest that although IL-1β, IL-6, and TNF-α were involved in induction of intestinal villus injury as suggested by in vivo studies (25), this combination is not sufficient to induce this pathological response unless IL-8 is also present at extremely high levels because low levels of production induced by PBMCs alone did not produce intestinal injury (Fig. 3 A). To confirm that the cytokines were...
specifically produced by the intestinal epithelium, we separated the circulating cells from the PBMCs and then carried out qPCR analysis for 92 known human inflammatory genes. These studies revealed that after being challenged simultaneously with LPS and PBMCs, the isolated intestinal epithelial cells significantly up-regulated (P < 0.05) the expression of 36 genes relative to untreated controls, whereas the same panel of inflammatory genes was undetectable in unstimulated PBMCs under these conditions (Fig. S5A). The intestinal epithelial cells increased their expression of genes encoding proteins that mediate early inflammatory signaling (e.g., PLCB3, PLCB4, PLCG1, PLCG2, TNFRSF1A, and PTGS2 also known as COX-2), as well as many that are important for leukocyte recruitment (e.g., LTB4R2, HRH1/2/3, NOS2, and ITGAM), by more than 25-fold (Fig. S5B). In addition, TLR activation is known to drive cytokine production of TLR4 when costimulated by PBMCs and LPS. During inflammation in the intestine, pathophysiological recruitment of circulating immune cells is regulated via activation of the underlying vascular endothelium. To analyze this organ-level inflammatory response in our in vitro model, a monolayer of human microvascular endothelial cells (Fig. 3 C and D and Fig. S6 A and B) or lymphatic endothelial cells (Fig. S6 B and C) was cultured on the opposite (abluminal) side of the porous ECM-coated membrane in the lower microchannel of the device to effectively create a vascular channel (Fig. 3C). To induce intestinal inflammatory responses, LPS (Fig. 3 C and D) or TNF-α (Fig. S6) was flowed through the upper epithelial channel for 24 h, and then PBMCs were added to the vascular channel for 1 h without flow (Fig. 3 C and D). Treatment with both LPS (or TNF-α) and PBMCs resulted in the activation of intercellular adhesion molecule-1 (ICAM-1) expression on the surface of the endothelium (Fig. 3 C and D, Left, and Fig. S6) and a significant increase (P < 0.0001) in the number of PBMCs that adhered to the surface of the capillary endothelium compared with controls (Fig. 3D). These results are consistent with our qPCR results, which also showed up-regulation of genes involved in immune cell trafficking (Fig. S5). Neither addition of LPS nor PBMCs alone was sufficient to induce ICAM-1 expression (Fig. S5D). These findings correlate the effects of LPS and PBMCs on epithelial production of inflammatory cytokines (Fig. 3A) as well as on villus injury (Fig. 2 B and D).

Recapitulating Organ-Level Intestinal Inflammatory Responses. During inflammation in the intestine, pathophysiological recruitment of circulating immune cells is regulated via activation of the underlying vascular endothelium. To analyze this organ-level inflammatory response in our in vitro model, a monolayer of human microvascular endothelial cells (Fig. 3 C and D and Fig. S6 A and C) or lymphatic endothelial cells (Fig. S6 B and C) was cultured on the opposite (abluminal) side of the porous ECM-coated membrane in the lower microchannel of the device to effectively create a vascular channel (Fig. 3C). To induce intestinal inflammatory responses, LPS (Fig. 3 C and D) or TNF-α (Fig. S6) was flowed through the upper epithelial channel for 24 h, and then PBMCs were added to the vascular channel for 1 h without flow (Fig. 3 C and D). Treatment with both LPS (or TNF-α) and PBMCs resulted in the activation of intercellular adhesion molecule-1 (ICAM-1) expression on the surface of the endothelium (Fig. 3 C and D, Left, and Fig. S6) and a significant increase (P < 0.0001) in the number of PBMCs that adhered to the surface of the capillary endothelium compared with controls (Fig. 3D). These results are consistent with our qPCR results, which also showed up-regulation of genes involved in immune cell trafficking (Fig. S5). Neither addition of LPS nor PBMCs alone was sufficient to induce ICAM-1 expression (Fig. S5D). These findings correlate the effects of LPS and PBMCs on epithelial production of inflammatory cytokines (Fig. 3A) as well as on villus injury (Fig. 2 B and D).

Evaluating Antiinflammatory Probiotic and Antibiotic Therapeutics On-Chip. To investigate how the gut microbiome modulates these inflammatory reactions, we cocultured the human intestinal villi with the eight strains of probiotic bacteria in the VSL. #3 formulation that significantly enhanced intestinal differentiation (Fig. 1E and Fig. S1B). To mimic the in vivo situation, we colonized our microengineered gut on a chip with the commensal microbes (VSL. #3) first and then subsequently added immune cells (PBMCs), pathogenic bacteria (EIEC), or both in combination. The VSL. #3 microbial cells inoculated into the germ-free lumen of the epithelial channel primarily grew as discrete microcolonies in the spaces between adjacent villi (Fig. 4A and Movie S3) for more than a week in culture (Fig. S7A), and no planktonic growth was detected. These microbes did not overgrow like the EIEC (Fig. 2A and Movie S2), although occasional microcolonies also appeared at different spatial locations in association with the tips of the villi (Fig. S7 B and C). The presence of these living components of the normal gut microbiome significantly enhanced (P < 0.001) intestinal barrier function, producing more than a 50% increase in TEER relative to control cultures (Fig. 4B) without altering villus morphology (Fig. 4C). This result is consistent with clinical studies suggesting that probiotics, including VSL. #3, can significantly enhance intestinal barrier function in vivo (18).

When the multispecies probiotic VSL. #3 microbes were cocultured in the lumen of the epithelial channel, the devastating injury induced by EIEC infection (Fig. 2) was significantly reduced (P < 0.01), as evidenced by maintenance of high TEER values (>1,500 Ω cm²) (Fig. 4B) and retention of villi with normal morphology (Fig. 4 C and D). Although the probiotic bacteria failed to completely prevent the intestinal damage induced by the EIEC infection in the presence of immune cells, the onset of intestinal injury was delayed by ~18 h (Fig. 4 B–D) compared with addition of EIEC in the absence of VSL. #3 microbes (Fig. 2 B–D). PBMCs also tolerated the presence of probiotic bacteria in the chip as intestinal barrier functions (Fig. 4 B and D) and tissue morphology (Fig. S8) remained unchanged when both were cocultured on-chip. However, the probiotic bacteria were not able to prevent intestinal injury when the epithelium was challenged simultaneously with both EIEC and PBMCs (Fig. 4 B–D). These findings are reminiscent of clinical findings with VSL. #3 therapy, which suggest that it is most helpful in treatment of patients with ulcerative colitis during the early phase of the disease (27, 28).

To mimic the effects of antibiotic therapies that are sometimes used clinically in patients with intestinal inflammatory disease (29), we identified a dose and combination of antibiotics (100 units per mL penicillin and 100 μg/mL streptomycin) that produced effective killing of both EIEC and VSL. #3 microbes in liquid cultures (Fig. S9) and then flowed this drug mixture into the epithelial channel of gut-on-a-chip devices infected with EIEC. When we added PBMCs to these devices 1 h later, intestinal barrier function (Fig. 4B) and villus morphology (Fig. 4C) were largely protected from injury, and there was a significant reduction in lesion area (Fig. 4D). Thus, the gut-on-a-chip was able to mimic suppression of injury responses previously observed clinically using other antibiotics that produce similar bacterialidal effects.

Analyzing Mechanical Contributions to Bacterial Overgrowth. Finally, we used the gut-on-a-chip to analyze whether physical changes in peristalsis or villus motility contribute to intestinal pathologies, such as the small intestinal bacterial overgrowth (SIBO) (5, 6) observed in patients with ileus (8) and IBD (7). When the GFP-EC bacteria were cultured on the villus epithelium under normal flow (50 μL/h), but in the absence of the physiological cyclic mechanical deformations, the number of colonized bacteria was significantly higher (P < 0.001) compared with gut chips that experienced mechanical deformations (Fig. 5A). Bacterial cell densities more than doubled within 21 h when cultured under conditions without cyclic stretching compared with gut chips that experienced physiological peristalsis-like mechanical motions, even though luminal flow was maintained constant (Fig. 5B). Thus, cessation of epithelial distortion appears to be sufficient to trigger bacterial overgrowth, and motility-induced luminal fluid flow is not the causative factor as assumed previously (7).

Discussion
One of the critical prerequisites for mimicking the living human intestine in vitro is to establish a stable ecosystem containing physiologically differentiated intestinal epithelium, gut bacteria, and immune cells that can be cultured for many days to weeks. Here we leveraged a mechanically active gut-on-a-chip microfluidic device to develop an in vitro model of human intestinal inflammation that permits stable long-term coculture of commensal microbes of the gut microbiome with intestinal epithelial cells. The synthetic model of the human living intestine we built recapitulated the minimal set of structures and functions necessary to mimic key features of human intestinal pathophysiology during chronic inflammation and bacterial overgrowth including epithelial and vascular inflammatory processes and destruction of intestinal villi.

Interest in the human gut microbiome is burgeoning because it has been implicated in control of intestinal homeostasis (30) and...
immune modulation (31), as well as in the pathogenesis of Crohn’s disease (32). The coculture method in the gut-on-a-chip described here enables stable host–microbe coexistence because of the presence of continuous fluid flow and peristalsis-like motions that enhance intestinal differentiation (33) and permit bacterial populations to reach a dynamic steady-state (12) that can be sustained over weeks in culture. In addition, the high level of mucus production on microengineered intestinal villi (16) also may provide a protective barrier to villus epithelium to maintain long-term stable coexistence with gut microbes.

One potential limitation of this model system is that the Caco-2 intestinal epithelial cells were originally isolated from a human colorectal tumor, although they were subsequently shown to exhibit features more similar to human small intestine (12, 16, 34). However, the key question in developing in vitro microphysiological systems is whether the method effectively recapitulates human organ physiology and whether it offers capabilities superior to other existing in vitro models. In our past studies, we showed that Caco-2 cells retain stem cell-like functional capabilities in that they faithfully recreate intestinal villi that exhibit small intestinal villus morphology and functions observed in vivo (17). In the present study, transcriptome analyses clearly revealed that Caco-2 cells that form small intestinal villi in the gut-on-a-chip exhibit an entirely different phenotype compared with the same cells cultured in static Transwells, and additional coculture with commensal gut microbes pushed their level differentiation on-chip even further so that it more closely resembled that of normal human ileum. This finding is relevant because the ileum is the last segment of the small intestine...
where commensal gut microbes are much more abundant than in the proximal regions. In addition, these data indicate that coculture with normal commensal gut microbes has a significant normalizing effect on the differentiation state of the Caco-2 intestinal epithelial cells in the gut-on-a-chip model, even beyond that demonstrated in past publications (12, 16). Thus, we believe that this new method represents a significant leap forward relative to existing in vitro models of human intestinal inflammation.

Using this microdevice, we also discovered that immune cells must be present along with LPS or nonpathogenic bacteria to induce production of a key set of proinflammatory cytokines (IL-8, IL-6, IL-1β, and TNF-α) that act together to produce villus injury and intestinal barrier disruption. However, although our results confirm that IL-6, IL-1β, and TNF-α may contribute to development of intestinal inflammatory disease as suggested in the past based on in vivo studies (35), our ability to manipulate these factors independently revealed that these cytokines must act in the presence of high levels of IL-8 to exert these disease-promoting effects. This novel result would be difficult to detect using animal models.

We also studied the effects of PBMCs as a way to mimic the effect of immune cell recruitment into the lamina propria, which is observed in chronic intestinal inflammation (13). PBMCs have been similarly used in many in vitro models of intestinal inflammation in the past because they can differentiate into intestinal tissue resident macrophages and dendritic cells (9, 23, 36). However, because the gut-on-a-chip is a synthetic system, it would be interesting to integrate other immune cells that contribute to maintenance of the immune milieu of the normal gut (e.g., gut-associated lymphoid tissue) in this model in the future.

The level of LPS found in the human intestine is generally considered harmless because the commensal Gram-negative gut microbiota routinely produce it, but it is at least partially inactivated by mammalian cell-derived acylacyl hydrolase in vivo (37). Although an excessive amount of LPS was shown to produce acute intestinal inflammation in one mouse model (38), another study in mice with defects in macrophage function revealed that they failed to produce inflammatory cytokines when challenged with LPS, suggesting that immune cell function is critical in LPS-induced inflammation (24). To examine how immune cells alter responses to commensal Gram-negative bacteria and their products (i.e., LPS) during chronic inflammation, we cocultured PBMCs with a nonpathogenic laboratory strain of *E. coli* that produces only the core LPS (i.e., not long-chain LPS) (39). We confirmed that this nonpathogenic *E. coli* strain does not produce gut damage on its own. However, it did induce villus injury and compromise epithelial barrier function when PBMCs were present, suggesting that the inflammatory microenvironment does indeed sensitize the epithelial barrier to injury. Immune cells also were shown to stimulate greater increases in expression of IL-8 and other inflammatory cytokines compared with LPS alone in cultured human intestinal cells (23). Thus, our finding that immune cells must be present for LPS to produce intestinal inflammation in the human gut-on-a-chip both confirms prior work and provides additional evidence to show that LPS is not inducing injury on its own.

In addition, we showed that both inflammatory activation of the intestinal endothelium and associated surface adhesion of immune cells can be studied in this model. Because this human intestine model included both isolated immune cells and living gut microbiome, it also was able to mimic the destructive effects of pathogenic bacteria on intestinal villi in vitro and successfully replicate the protective effects of clinical probiotic and antibiotic therapies previously observed in IBD patients in vivo. Furthermore, among the 12 potential cytokines involved in human intestinal inflammation (40, 41) that we tested blindly in this study, a specific combination of 4 cytokines (IL-8, IL-1β, IL-6, and TNF-α) was discovered to be necessary and sufficient to induce injury in our model. Moreover, exposure of the epithelium to combinations of immune cells with either LPS or GFP-EC produced a similar injury response, which was also accompanied by increased production of this same set of cytokines. The same cytokines have been reported to be up-regulated in clinical studies of human inflammatory diseases, including in patients with active ulcerative colitis patients (41), Crohn’s disease (40), and pouchitis (42). Because intestinal macrophages capable of producing IL-8 are activated in IBD patients (43) and intestinal epithelial cells exposed to Gram-negative bacteria also produce this cytokine (23, 43), IL-8 has been implicated as a potential contributing factor in IBD as well. However, although our in vitro studies using a blocking antibody show that inhibition of IL-8 action completely prevents villus injury in the presence of all four cytokines, addition of IL-8 alone was not capable of inducing an injury response. Thus, apparently, IL-8 is only able to manifest its pivotal ability to compromise the intestinal barrier if it acts in an environment where IL-1β, IL-6, and TNF-α are also inflammatory. These observations raise the possibility that this particular combination of four cytokines and their receptors could be regarded as potential immunotherapeutic targets for a combination therapy to suppress inflammatory responses in patients with IBD. These types of mechanistic insights would have been extremely difficult to obtain using animal models.

Gene expression profiling revealed the huge extent of inflammatory changes that are induced in the human intestinal epithelial cells when challenged by LPS and immune cells in combination on-chip, as both proinflammatory and anti-inflammatory processes became activated. For instance, there was a dramatic up-regulation of receptors for histamine (HRH1, HRH2, and HRH3) and leukotriene B4 (LTB4R and LTB4R2) that mediate immune cell recruitment (44, 45), which is consistent with our finding that these conditions stimulate ICAM-1 expression and immune cell recruitment, as well as with clinical reports that show high levels of both histamine and leukotriene B4 are routinely found in IBD patients (46). Early inflammatory signaling molecules, including prostaglandin-endoperoxide synthase 2 (47) (PTGS2/COX-2), TNF receptors (48) (TNFRSF1A and TNFRSF1B), and phospholipase C enzymes (49) (PLCB3/4, PLCε1, and PLCG1/2) were up-regulated as well. However, at the same time, these cells increased expression of adrenergic receptors (50) (ADRB1 and ADRB2), annexin (51) (ANXA1 and ANXA3), and melanocortin neuropeptide receptor family member MC2R (52), which can suppress inflammatory reactions.
These studies therefore suggest that the intestinal epithelial cell plays a central role in controlling both the dynamics and directionality of the inflammatory response.

Our experiments also showed that a therapeutic probiotic formulation (VSL#3) protects against the overgrowth of EIEC and subsequent gut injury. Although we did not analyze the population dynamics of individual species in this study, the maximum specific growth rates of Lactobacillus spp. (53), Bifidobacterium spp. (54), and Streptococcus sp. (55) are similar under high-nutrient conditions. Thus, it is unlikely that microbial species-level competition for nutrients significantly altered species diversity. One potential caveat, however, is that although the VSL#3 formulation contains multiple obligate or facultative anaerobic strains of commensal gut microbes, it is not fully representative of the highly complex commensal microbiome of the normal intestine. Although a positive effect of probiotic coculture on barrier function was demonstrated in past studies using static Transwells (56, 57), they were only able to perform the coculture for a short period (4–48 h). They also were not able to detect the important effect of mechanical deformation on bacterial overgrowth as we did in this study. We also found that the microbial cells predominantly colonized the intervillus spaces even though they were seeded uniformly. This may be because crevices between villi stimulate these bacteria from fluid shear stress inside the microchannel, whereas bacteria positioned in regions above the tips of thin villi will tend to be washed out from the device. In the confined intervillus space within a densely packed microchannel, facultative microbes may rapidly consume available oxygen, which would produce a local anoxic microenvironment that might support growth of nearby anaerobic microbes, even when the overall culture conditions are aerobic (5% CO2); this cannot be achieved in conventional 2D cell culture models. In contrast to VSL#3, the EIEC cells exhibited excessive overgrowth in the cultures. This might be because the pathogenic EIEC cells adhere more tightly to the epithelial surface or they create a more interconnected biofilm in this system. Overgrowth by the EIEC cells resulted in rapid injury of the epithelium, loss of normal villus morphology, and disruption of cell-cell junctions (Fig. 2A), likely from drastic nutrient depletion and waste accumulation. In contrast, the VSL#3 microcolonies remained limited to the spaces between villi that retained well-delineated epithelial cell borders when analyzed at up to 8 d of coculture. Taken together, these findings suggest that gut microbes may be particularly useful for modeling SIBO given that bacterial overgrowth is currently assessed clinically by analyzing microbes in samples of small bowel fluid aspirates obtained by endoscopy (58).

Clinical and experimental studies suggest that antibiotic therapy can selectively decrease the invasion of pathogenic bacteria and broadly reduce luminal and mucosal bacterial densities in IBD patients (59). Our results show that administration of a mixture of bactericidal antibiotics can suppress intestinal injury induced by pathogenic EIEC bacteria demonstrate that this form of adjuvant therapy can be studied in vitro using the gut-on-a-chip method. However, the antibiotics we used are different from those used most commonly in the clinic, and thus, the effects of different antibiotic therapies should be tested in future studies.

Finally, by leveraging the modular capability of gut-on-a-chip to vary mechanical parameters independently, we discovered that the loss of physiological mechanical deformation is itself sufficient to induce small intestinal bacterial overgrowth similar to that observed in some patients with ileus (8) as well as IBD (6, 7). In the case of ileus, which results from complete cessation of peristalsis (e.g., following surgery), the cause of bacterial overgrowth remains unknown. This is because although past in vivo studies suggested that it is caused by reduced movement of luminal contents due to compromised gut motility (8), inhibition of peristalsis also results in decreased mechanical deformation of the epithelium, which is known to have potent effects on cell function (60). Thus, our finding shifts the paradigm from current mechanistic models based on in vivo studies, which assume the cause of bacterial overgrowth to be slowing of luminal flow due to decreased motility that would concentrate bacterial populations. All of these discoveries were made possible by our ability to integrate and independently modulate different system parameters (e.g., peristalsis-like motions, presence of immune cells, and microbes) in the gut-on-a-chip, and it would be extremely difficult, if not impossible, to uncover these direct relationships using animal models or even human clinical studies.

Taken together, these results demonstrate that the human gut-on-a-chip offers a robust modular platform not only for analysis of intestine–microbe interactions and interrogating etiology and development of human intestinal inflammatory diseases but also for defining potential therapeutic targets and testing drug candidates. Because all of the cellular components of the model can be varied independently, this microphysiologic system could be improved further by incorporating primary or iPSC cell-derived human intestinal epithelial cells, connective tissues, and various subsets of immune cells, as well as more complex commensal gut microbial communities (e.g., isolated from human IBD patients to model their specific contributions to intestinal inflammation in vitro). This potentially could be done in a patient-specific manner to advance personalized medicine in the future.

Materials and Methods

Device Microfabrication. The gut-on-a-chip microdevice used in this study was fabricated from PDMS as reported previously (12, 16, 33). The dimensions of cell culture microchannel (1 mm wide × 10 mm long × 0.15 mm high) and vacuum chambers (1.68 mm × 3.3 mm × 0.15 mm) were the same in the upper and lower layers; the porous PDMS membrane contained an array of circular holes (10 μm diameter × 20 μm high with 25 μm spacing). Gas-permeable silicone tubing (Tygon 3350, ID 1/32″, OD 3/32″) with a connector (hub-free stainless steel blunt needle, 18G; Kimble Chase) was connected into each microchannel to supply cell culture medium and vacuum suction.

Microfluidic Culture. Caco-2BBe human colorectal carcinoma-derived (Caco-2) intestinal epithelial cells (obtained from Harvard Digestive Disease Center) were cultured on the upper surface of the ECM-coated, flexible, porous membrane in the gut-on-a-chip device, with culture medium (DMEM, 20% FBS, antibiotics) being perfused constantly (30 μL/h; 0.02 dyn/cm2) and exposed to cyclic mechanical deformation (10% strain, 0.15 Hz); the Caco-2 cells spontaneously formed villi after ~100 h of culture. To model inflammatory activation, human capillary endothelial cells or human lymphatic microvascular endothelial cells were cocultured on the opposite side of the porous ECM-coated membrane in the presence of flowing (30 μL/h) coculture medium with cyclic mechanical deformations (10%, 0.15 Hz). To study inflammatory activation of the endothelium, LPS (15 μg/mL) and PBMCs (3.3 × 106 cells per mL) were introduced into the upper and lower microchannels, respectively. For host-microbe coculture, villi grown in a gut-on-a-chip device were preconditioned in antibiotic-free culture medium for 12 h before microbial cells (VSL#3, GMP-EC, or EIEC) suspended in an antibiotic-free cell culture medium were introduced into the upper microchannel. The microbes and cells were then incubated without flow for 1.5 h before flow (40 μL/h) and mechanical deformations (10%, 0.15 Hz) were restored to establish a stable host-microbe ecosystem. PBMCs were isolated from human blood drawn from healthy volunteers using a ficoll gradient. Blood collection procedures were approved by the Institutional Review Board of the Harvard University Faculty of Medicine.

Transcriptome Analysis. Gene expression profiling of Caco-2 cells cultured in static or microfluidic conditions was carried out using Human Genome 1.0 ST Affymetrix GeneChip arrays. Gene expression data for normal human small intestine (duodenum, jejunum, and ileum) were obtained from the National Center for Biotechnology Information (NCBI) Gene Expression Omnibus (GEO) database. Transcriptome profiles of Caco-2 cells performed in this study were deposited to the NCBI GEO database (accession no. GSE65790). Global gene expression profiles were also visually represented using self-organizing maps generated using the Gene Expression Dynamics Inspector (GEDI) program (19).

Morphological Analysis. Images of the intestinal villus morphology were recorded by using phase contrast or differential interference contrast (DIC) microscopy (Zeiss). The villus microarchitecture was also visualized using immunofluorescence microscopy with a laser scanning confocal microscope (Leica). Acquired images were processed using MetaMorph (Molecular Devices),

ACKNOWLEDGMENTS. We thank Sri Kosuri (Wyss Institute) for providing GFP-labeled E. coli strain; Jaeyuk Lee and Carolina Lucchesi (Wyss Institute) for help with the qPCR analysis; and Eunyoung Chun (Harvard T. H. Chan School of Public Health) for assistance with flow cytometry and helpful discussions. This work was supported by the Defense Advanced Research Projects Agency (Cooperative Agreement W911NF-12-2-0036), Food and Drug Administration (Grant HHSF223201310079C), and the Wyss Institute for Biologically Inspired Engineering at Harvard University.