Regulatory T Cells Restrict Permeability to Bacterial Antigen Translocation and Preserve Short-Chain Fatty Acids in Experimental Cirrhosis

Oriol Juanola,1,2 Paula Piñero,1 Isabel Gómez-Hurtado,1,3 Esther Caparrós,2 Rocío García-Villalba,4 Alicia Marín,4 Pedro Zapater,1,3,5 Fabián Tarín,1 José M. González-Navajas,1,3 Francisco A. Tomás-Barberán,4 and Rubén Francés1,2,3

Intestinal permeability to translocation of bacterial products is increased in cirrhosis. Regulatory T cells (Tregs) remain central to the interplay between the host and microbial milieu. We propose that Tregs are involved in promoting gut barrier integrity and a balanced interaction with gut microbiota–derived short-chain fatty acids (SCFAs). Carbon tetrachloride cirrhosis was induced in wild-type and recombination activating gene 1 (Rag1)<sup>−/−</sup> mice. Naive T cells and Treg cells were transferred into <span class="caps">Rag1</span><sup>−/−</sup> mice. Intestinal permeability was assessed <em>in vivo</em> after lipopolysaccharide (LPS) oral administration, and bacterial DNA presence was evaluated in mesenteric lymph nodes. Transcript and protein levels of tight-junction (TJ) proteins were measured in colonic tissue. Intestinal T helper profile in response to <em>Escherichia coli</em> (<em>E. coli</em>) was determined by flow cytometry. SCFAs were measured by gas chromatography–mass spectrometry in colonic content before and after <em>E. coli</em> challenge. <span class="caps">Rag1</span><sup>−/−</sup> mice showed significantly increased permeability to LPS and bacterial DNA translocation rate compared with control mice. Naive T and Treg cotransfer significantly reduced gut permeability to bacterial antigen translocation and restored TJ protein expression in <span class="caps">Rag1</span><sup>−/−</sup> mice. Naive T and Treg replenishment in <span class="caps">Rag1</span><sup>−/−</sup> mice restrained proinflammatory differentiation of intestinal lymphocytes in response to <em>E. coli</em>. The main SCFA concentration resulted in significant reduction in <span class="caps">Rag1</span><sup>−/−</sup> mice after <em>E. coli</em> administration but remained unaltered after naive T and Tregs cotransfer. The reduced expression of SCFA receptors induced by <em>E. coli</em> was reestablished following naive T and Treg reconstitution in <span class="caps">Rag1</span><sup>−/−</sup> mice. Conclusion: The restriction of gut permeability, local inflammatory differentiation, and loss of bacteria-derived SCFAs foster the value of Tregs in preventing bacterial translocation in cirrhosis. (Hepatology Communications 2018;2:1610-1623).

Patients with cirrhosis are frequently threatened with bacterial infections that aggravate their prognosis.<sup>(1,2)</sup> The increased exposure to bacterial antigens is due to a pathological translocation from the gut.<sup>(3,4)</sup> Although living microorganisms may translocate and be responsible for severe

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**Abbreviations:** BT, bacterial translocation; CCL<sub>19</sub>, carbon tetrachloride; DCs, dendritic cells; <em>E. coli</em>, <em>Escherichia</em> coli; FFAR, free fatty acid receptor; FITC, fluorescein isothiocyanate; HBSS, Hank’s balanced salt solution; IFN, interferon; IHC, immunohistochemical; IL, interleukin; LPS, lipopolysaccharide; MLN, mesenteric lymph node; mRNA, messenger RNA; PBS, phosphate-buffered saline; PCR, polymerase chain reaction; <span class="caps">Rag1</span>, recombination activating gene 1; SCFAs, short-chain fatty acids; TJ, tight-junction; Tregs, regulatory T cells; WB, western blot; WT, wild type; ZO-1, zonula occludens.

Received July 5, 2018; accepted October 3, 2018.

Additional Supporting Information may be found at onlineLibrary.wiley.com/doi/10.1002/hep4.1268/supplinfo.

Supported by Instituto de Salud Carlos III, Madrid, Spain (PI16/0967); Consellería Educación, Generalitat Valenciana, Valencia, Spain (PROMETEO/2016/001); and FEDER funds (EU).

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DOI 10.1002/hep4.1268

Potential conflict of interest: Nothing to report.
infections such as spontaneous bacterial peritonitis,\(^5\) bacterial products have also been demonstrated to cross the gut barrier and be associated with poor disease outcomes.\(^6,7\)

The immunological response to increased bacterial antigen translocation rates in cirrhosis and other settings involves a decompensation toward a proinflammatory profile in which dendritic cells (DCs), monocytes, and neutrophils are recruited to the injured area.\(^8,9\) As a consequence of this polarization, the tolerogenic response that balances out the inflammation is likely compromised. In fact, the immune homeostasis is lost during progression of cirrhosis. The so-called cirrhosis-associated immune dysfunction helps induce intestinal inflammation, suggesting a plausible role for regulatory immune cells and anti-inflammatory cytokines in this context.\(^10\) Intestinal inflammation leads to increased intestinal permeability, which has been demonstrated in cirrhosis,\(^11-13\) and is key in the translocation of small bacterial antigens that, in turn, are able to induce and/or perpetuate a sustained inflammatory environment.\(^14,15\) In this regard, tight-junction (TJ) proteins, a family of more than 50 members, play an important role in preventing any paracellular breach\(^16\) by which these bacterial antigens may enter the sterile intraepithelial area.

It is well established that normal bacterial colonization is needed for the development of the intestinal barrier.\(^17,18\) Short-chain fatty acids (SCFAs) are produced by the bacterial fermentation of undigested dietary carbohydrates in the intestine and act as the main energy source of colonocytes. SCFA products of commensal microorganisms are important functional players in the maintenance of intestinal barrier integrity. In particular, butyrate is a considerable TJ regulator due to its ability to increase expression of zonula occludens (ZO-1) and claudin-1 as well as influence occluding redistribution \textit{in vitro}.\(^19\) These metabolic intermediaries also work as communicators between the intestinal microbiota and the immune system by promoting extrathymic differentiation of regulatory T cells (Tregs).\(^20\) SCFA receptors, G protein–coupled receptor (GPR)43/free fatty acid receptor 2 (FFAR2) and GPR42/FFAR3, are expressed on the innate immune cells and participate in regulation of the inflammatory response.\(^21,22\)

Treg cells are an essential immune population that interferes with effector T lymphocytes and DCs and provides an immunosuppressive counterpart.\(^23-26\) We have described in the past that the cellular proinflammatory response in patients with cirrhosis receiving norfloxacin was modulated by an interleukin (IL)-10-mediated mechanism.\(^27\) Trying to understand the role of IL-10 in the cirrhotic context, we also reported that IL-10 was required in experimental cirrhotic models to restore normal gut barrier permeability and to lower luminal free endotoxin absorption.\(^28\) Among adaptive IL-10–producing cells, Treg cells are especially relevant because of their homeostatic role in the host–microbe interaction. We have identified increased rates of Treg cells in patients on selective intestinal decontamination, their correlation with serum norfloxacin concentrations, and their role as relevant IL-10–secreting cells in these patients,\(^29\) supporting their implication in norfloxacin-derived down-regulation of serum proinflammatory levels in cirrhosis observed in the past.\(^14\) In addition, this cell type is implicated in gut homeostatic functions, as their role in modulating SCFAs concentration in the gut microbiota has been reported.\(^30\)

**ARTICLE INFORMATION:**

From the 1Instituto ISABIAL-FISABIO, Hospital General Universitario de Alicante, Alicante, Spain; 2Dpto. Medicina Clínica, Universidad Miguel Hernández, San Juan, Spain; 3CIBERehd, Instituto de Salud Carlos III, Madrid, Spain; 4CEBAS-CSIC, Campus de Espinardo, 30100, Murcia, Spain; 5Dpto. Farmacología, Pediatría y Química Orgánica, Universidad Miguel Hernández, San Juan, Spain.

**ADDRESS CORRESPONDENCE AND REPRINT REQUESTS TO:**

Rubén Francés, Ph.D.  
Instituto ISABIAL-FISABIO  
Hospital General Universitario de Alicante  
Avda. Pintor Baeza 12  
03010 Alicante, Spain  
E-mail: frances_rub@gva.es  
Tel.: +34965913928  
Fax: +34965913922
In the present study, our specific aim is to evaluate whether the Tregs effect on reduced bacterial antigen translocation might be associated with an improvement in the gut barrier integrity. The purpose of the study, therefore, was to evaluate the contribution of Tregs on gut barrier integrity in the context of experimental cirrhosis. We propose that Treg cells help maintain a reduced gut permeability by promoting the barrier integrity and favoring a balanced interaction with gut microbiota anti-inflammatory products such as SCFAs in response to episodes of bacterial antigen translocation.

We performed adoptive transfer experiments in a recombination activating gene 1 (Rag1\(^{-/-}\))-deficient mice model of carbon tetrachloride (CCl\(_4\))-induced cirrhosis to evaluate gut barrier integrity markers, gut permeability, bacterial translocation (BT) to mesenteric lymph nodes (MLNs), and the immunological status of intestinal lymphocytes as a key cell population in maintaining gut homeostasis.

**Animals and Methods**

**MICE**

Male C57Bl/6J wild-type (WT) and Rag1\(^{-/-}\) mice on C57Bl/6J background (Jackson Laboratories, Bar Harbor, MN) were included in 16-week cirrhosis-induced protocols. Immunodeficient Rag1\(^{-/-}\) mice have a defective VDJ (variable, diversity, and joining) recombination, so the immune adaptive response is altered by the lack of production of mature T cells and B cells. All of the animals were subjected to a week of quarantine before the cirrhosis protocol. We selected Rag1\(^{-/-}\) mice, which are a T cell–free mouse model, to induce liver toxicity with a classical CCl\(_4\) protocol. The cirrhotic Rag1\(^{-/-}\) mice let us evaluate the activation of sorted naive T cells and Tregs from WT animals. Mice were fed a standard rodent chow and kept at a constant room temperature of 21°C in a 12:12 light/dark cycle. The cirrhosis protocol began with the treatment of animals with 0.25 mmol/L phenobarbital in tap water that was maintained along the study protocol. After 4 weeks, the animals received 2 weekly weight-controlled doses of CCl\(_4\) by oral gavage for 12 weeks. The first CCl\(_4\) dose was 100 μL/kg (2 μL per mouse) in mineral oil, and subsequent doses were adjusted based on changes in weight 48 hours after the previous dose, up to 100 μL per mouse. A group of naive WT mice not subjected to the CCl\(_4\) protocol was used as a negative control for evaluating fibrosis.

Sample size was calculated according to preliminary TJ protein gene expression levels obtained by our group and published in abstract format (Juanola et al. *Hepatology* 2016;64:83A). Assuming an α error of 0.017, a β error of 0.2, an equal distribution of subjects between groups (0.5 each), and an observed mean difference (E) of colonic occludin gene expression between groups of 1 with a SD of 0.3, 4 animals per group per protocol were required. Typically, mortality rates in our work are about 20%. We increased the number of Rag1\(^{-/-}\) mice compared with control mice to prevent possible cell-transfer failures or an increased mortality rate. We included 40 to 45 mice in each protocol (30-35 Rag1\(^{-/-}\) and 10 WT) to make sure we were able to work with at least 5 to 6 mice per group.

Rag1\(^{-/-}\) cirrhotic mice were subjected to adoptive transfer experiments 48 hours before laparotomies, as described.\(^{(29)}\) A group of 24 male C57Bl/6 WT mice not subjected to cirrhosis induction protocol were used for isolating spleen-derived sorted naive T (CD4\(^+\)CD25\(^-\)CD45RB\(^{high}\)) cells and Treg (CD4\(^+\)CD25\(^+\)CD45RBlow) cells for these experiments. Postsort purity was typically more than 98%. Rag1\(^{-/-}\) cirrhotic animals were classified into three groups according to transferred cells: (1) mice not receiving sorted cells; (2) mice transferred with naive T cells (2 × 10\(^5\) cells per mouse); and (3) mice receiving naive T and Treg cells (1 × 10\(^5\) each per mouse). Twenty-four hours prior to laparotomies, a subgroup of cirrhotic Rag1\(^{-/-}\) animals in each condition received oral *Escherichia coli* (*E. coli*), as described subsequently.

Laparotomies were performed under anesthesia with isoflurane. Whole blood was obtained from the cava vein for gut permeability experiments before liver perfusion. Livers were then perfused in situ with 6 mL of Hank’s balanced salt solution (HBSS) without Ca\(^{2+}\) and Mg\(^{2+}\) (Life Technologies Corp., Grand Island, NY) at 37°C at a rate of 1.5 mL per minute. Detectable MLNs were removed, and the liver, spleen, complete small intestine, colon, and content from cecum were collected.

Animals received care according to the criteria outlined in the Guide for the Care and Use of Laboratory Animals. The study was approved by the Animal Research Committee of Universidad Miguel Hernández (Alicante, Spain).
**Fluorescein Isothiocyanate–LPS Intestinal Permeability Assay**

A subgroup of animals in each condition was intragastrically administered 100 µg of fluorescein isothiocyanate (FITC) conjugated LPS from *E. coli* 0111:B4 or dextran (Sigma-Aldrich, St. Louis, MO). Gut permeability was evaluated 2 hours after FITC-LPS or FITC-dextran administration as the percentage of recovery measured by fluorometry at 488 nm in 100 µL of serum samples collected from the cava vein. Dilutions of FITC-LPS and FITC-dextran in phosphate-buffered saline (PBS) were used for standard curves.

**Bacterial DNA Translocation Assessment in MLNs**

MLNs were harvested with sterile instruments in sterile PBS at laparotomies. All tissues were disrupted by using the Tissue Lyser LT (QIAgen, Hilden, Germany), and DNA was immediately isolated using the QIAamp DNA Mini Kit. Bacterial DNA was detected by performing a broad-range polymerase chain reaction (PCR) and partial sequencing analysis of the 16S ribosomal RNA gene, as described.

**Gene Expression Analysis**

Total cellular RNA was isolated using the RNeasy Mini Kit (QIAgen). The qScript One-Step SYBR Green quantitative real-time PCR Kit (Quanta BioScience, Gaithersburg, MD) was used to perform the gene expression of claudin-1, claudin-2, ZO-1, occludin, collagen type I alpha-1 chain (*Col1a1*), tumor growth factor beta (*TGF-β*), matrix metalloproteinase 2 (*MMP2*), tissue inhibitor of metalloproteinase 1 (*TIMP1*), interleukin 2 (*IL2*), and interleukin 10 (*IL10*) in an IQ5 RT-PCR (Bio-Rad, Hercules, CA). *β2-microglobulin* was used as a housekeeping gene in all gene expression analyses. Primer pairs used in the study can be followed in Supporting Table S1.

**Immunohistochemistry Analysis**

Immunohistochemical (IHC) assays were carried out in 4-µm sections of paraffin-embedded colon tissue, and processed following standard procedures. The slides were incubated with primary antibodies claudin-1, claudin-2, occludin, and ZO-1 (Abcam PLC, Cambridge, United Kingdom). As secondary antibodies, we incubated the sections with the correspondent α-goat or α-rabbit biotinylated antibodies (Palex Medical SA, Sant Cugat del Vallés, Spain). Slides were incubated with avidin-biotin complex (Vector Laboratories Inc., Burlingame, CA) and revealed with peroxidase substrate 3,3′-diaminobenzidine (Vector Laboratories Inc.). Nuclei were stained by incubating the sections in Harris hematoxylin (Leica Biosystems Richmond Inc., Richmond, IL). As a negative control, staining was carried out in the absence of a primary antibody. Images were obtained in a camera-assisted optic Leica DM2000 LED microscope (Leica Biosystems, Richmond Inc.). All represented panels are shown at original magnification ×20. A semiquantitative analysis of protein expression was performed using the ImageJ software (https://rsbweb.nih.gov).

**Western Blot Analysis**

Colon homogenates were lysed with radioimmunoprecipitation assay buffer and protein concentration determined by Bradford protein assay (EMD Millipore Corp., Billerica, MA). Twenty micrograms of protein extracts were resolved under reducing conditions on 6% to 15% sodium dodecyl sulfate–polyacrylamide gels and transferred to Immobilon-P membranes (EMD Millipore Corp.). Primary antibodies used are β-actin (Sigma-Aldrich), ZO-1 (Thermo Fisher Scientific, Waltham, MA), and claudin-1, claudin-2, and occludin (Abcam PLC). Finally, membranes were incubated with the appropriate horseradish peroxidase (HRP)–conjugated secondary antibody (Cell Signaling Technology, Leiden, Netherlands). Immobilon Western Chemilum HRP Substrate (EMD Millipore Corp.) was used to detect the activity of the membrane-attached peroxidase, and images were obtained in ChemiDOC XRS+ operated by Image Lab software (Bio-Rad). Protein bands were quantified by densitometry using Scion Image software (Scion Corp., Frederick, MD). Band densities were expressed relative to total β-Actin protein.

**E. coli Culture Conditions and Administration to Mice**

*E. coli* (CECT45) was grown in 9 mL of thio-glycolate broth with resazurin (Biomerieux, Marcy
l’Etoile, France) and incubated overnight at 37°C. Serial dilutions of incubated bacteria were performed before plating them in McConkey agar (Biomérieux) overnight at 37°C. Colony-forming units (CFUs) were then counted, and cells were resuspended in PBS. Each animal was administered $1 \times 10^6$ CFUs orally.

**INTESTINAL LYMPHOCYTE ISOLATION AND $T_h$ DIFFERENTIATION**

The whole small intestine from E. coli–challenged cirrhotic Rag1$^{-/-}$ mice was obtained. Peyer’s patches, fatty tissue, and mesentery were removed from all intestines. The gut was cleaned using cold PBS without Ca$^{2+}$ and Mg$^{2+}$ (Euroclone, Milano, Italy), opened longitudinally, and cut into 1-cm pieces. Intraepithelial lymphocytes (IELs) were obtained by incubating the small intestine pieces twice with HBSS Ca$^{2+}$ and Mg$^{2+}$ free (Life Technologies Corp.) supplemented with 5 mM ethylene diamine tetraacetic acid (Bio-Rad), 1 mM dithiothreitol (Sigma-Aldrich), and 1% penicillin-streptomycin (Life Technologies Corp.) in Incubating Orbital Shake (VWR, Llinars del Vallès, Spain) for 30 minutes at 37°C. Thereafter, the tissue pieces were washed with PBS and then incubated three times in HBSS with 0.5 mg/mL collagenase D (Roche Diagnostics GmbH, Mannheim, Germany), 3 mg/mL dispase II (Sigma-Aldrich), 1 mg/mL DNase I (Roche Diagnostics GmbH), and 1% penicillin-streptomycin with orbital agitation for 30 minutes at 37°C to collect the lamina propria lymphocyte (LPL) cells. All supernatant fractions were filtered with 70-µm nylon cell strainers (Corning Incorporated Life Sciences, Oneonta, NY), washed with PBS 3% fetal bovine serum (FBS) (Life Technologies Corp.), and centrifuged to harvest cell suspensions. IELs and LPLs were pooled and constituted the intestinal lymphocyte content. Live lymphocytes were obtained by preparing a 40%/80% Percoll (Sigma-Aldrich) gradient and then cultured in advanced Roswell Park Memorial Institute (RPMI)-1640 (Life Technologies Corp.) supplemented with 10% FBS, 1% L-glutamine (Life Technologies), and 1% penicillin-streptomycin. Cells were treated and marked according to manufacturer’s instructions in the Mouse $T_h$1/$T_h$2/$T_h$17 Phenotyping Kit (BD Biosciences, San Diego, CA). To induce a complete T-cell activation, E. coli was administered according to the conditions described here. $T_h$ differentiation was evaluated as intracellular production of interferon (IFN)-γ, IL-4, and IL-17. Data acquisition and analysis of marked intestinal lymphocytes were performed using a FACSCanto II flow cytometer operated by FACSDiva software (BD Biosciences).

**GAS CHROMATOGRAPHY–MASS SPECTROMETRY ANALYSIS OF SCFAS IN INTESTINAL CONTENT**

Intestinal content samples with PBS were homogenized, and 300 µL were mixed with 300 µL of 0.5% phosphoric acid. Intestinal content suspensions were sonicated in an ultrasonic bath for 5 minutes and centrifuged for 10 minutes at 17,949g. The aqueous supernatant was extracted with the same volume of methyl tert-buthyl ether for 5 minutes and centrifuged in the same conditions. The upper organic phase was transferred into a tube and 4-methyl valeric acid added as internal standard at a final concentration of 500 µM. Gas chromatography–mass spectrometry (GC-MS) system and chromatographic and mass parameter conditions were followed, as reported.(31)

**STATISTICAL ANALYSIS**

Continuous variables are reported as mean ± SD and categorical variables as frequency or percentages. Quantitative data were analyzed using the Mann-Whitney U test for simple comparisons with the post hoc Bonferroni correction for multiple comparisons. Differences in qualitative variables were analyzed using the $\chi^2$ test. All reported $P$ values are 2-sided, and $P$ values less than 0.05 indicate significance. All calculations were performed using SPSS Statistics 19 (IBM, Chicago, IL).

**Results**

**CIRRHOSIS INDUCTION PROTOCOLS AND MICE GROUPS**

Five independent protocols of CCl$_4$-induced cirrhosis were run to complete the different experiments performed. Each protocol included a group of WT mice and three groups of Rag1$^{-/-}$ mice (nontransferred
**TABLE 1. PROTOCOLS AND NUMBER OF MICE IN DIFFERENT EXPERIMENTAL PROCEDURES DISTRIBUTED BY STUDY GROUPS**

| Protocol | Experiments                                                                 | Number of CCl₄ mice/group | Total number of CCl₄ mice/protocol |
|----------|------------------------------------------------------------------------------|---------------------------|-----------------------------------|
| 1        | FITC-LPS permeability assays                                                 | WT  6 | Rag₁⁻/⁻  8 | Rag₁⁻/⁻ + naive T  8 | Rag₁⁻/⁻ + naive T + Treg  8 | 30 |
| 2        | FITC-dextran permeability assays                                             | WT  6 | Rag₁⁻/⁻  8 | Rag₁⁻/⁻ + naive T  8 | Rag₁⁻/⁻ + naive T + Treg  8 | 30 |
| 3        | TJ proteins mRNA and WB in colonic tissue                                    | WT  6 | Rag₁⁻/⁻  8 | Rag₁⁻/⁻ + naive T  8 | Rag₁⁻/⁻ + naive T + Treg  8 | 30 |
| 4        | FFARs mRNA in colonic tissue (except in C57Bl/6)                               | WT  6 | Rag₁⁻/⁻  8 | Rag₁⁻/⁻ + naive T  8 | Rag₁⁻/⁻ + naive T + Treg  8 | 30 |
| 5        | Th differentiation in colonic ILS in response to E. coli                     | WT  8 | Rag₁⁻/⁻  8 | Rag₁⁻/⁻ + naive T  8 | Rag₁⁻/⁻ + naive T + Treg  8 | 24 |

**GUT PERMEABILITY TO LPS IS REDUCED IN CIRRHTIC RAG₁⁻/⁻ MICE COTRANSFERRED WITH TREG CELLS**

We aimed to study in vivo gut permeability to orally administered FITC-LPS. The results of this functional test are described in Fig. 1A. Rag₁⁻/⁻ mice showed a significantly increased permeability to FITC-LPS compared with WT mice. The recovery rate of LPS...
in $\text{Rag1}^{-/-}$ mice was not reduced after injection with spleen-derived naive T cells. On the contrary, only the coinjection of naive T and Treg cells significantly reduced gut permeability to FITC-LPS. In agreement with this, the translocation of bacterial DNA to MLNs of $\text{Rag1}^{-/-}$ mice cotransferred with naive T and Treg cells was significantly down-regulated to levels present in WT cirrhotic mice (Fig. 1B).

**TREG CELLS ARE ASSOCIATED WITH A RESTORATION OF TJ PROTEIN EXPRESSION IN CIRRHOTIC $\text{Rag1}^{-/-}$ MICE**

The observed reduction in gut permeability led us to study the status of TJ proteins in cirrhotic mice. The expression of TJ proteins ZO-1, occludin, claudin 1, and claudin 2 was measured by gene expression, IHC, and western blot (WB) in all study groups (Fig. 2). Messenger RNA (mRNA) levels of all studied genes were significantly decreased in cirrhotic $\text{Rag1}^{-/-}$ mice either nontransferred or transferred with naive T cells compared with cirrhotic WT mice. The cotransfer with naive T and Treg cells restored transcript expression of all genes to levels shown in cirrhotic WT mice (Fig. 2A).

Cirrhotic $\text{Rag1}^{-/-}$ mice without any transferred cells showed a significant reduction in the protein expression of the different TJ proteins compared with the cirrhotic WT, as shown by IHC and WB analyses (Fig. 2B,C, respectively). None of the TJ protein levels significantly increased when these mice were transfused with naive T cells. However, the cotransfer with naive T and Treg cells restored all TJ protein expression to the levels shown in WT. Visually observed differences between groups in TJ protein expression were shown to be statistically significant after a semiquantitative analysis of IHC and WB images.

**TRANSFERRED TREG CELLS MODULATE IL PROINFLAMMATORY T$_h$1 AND T$_h$17 RESPONSES TO ORAL E. COLI ADMINISTRATION IN CIRRHOTIC $\text{Rag1}^{-/-}$ MICE**

Because Treg cells are functionally required to maintain gut barrier integrity and reduced permeability in CCl$_4$-cirrhotic mice, we were interested in evaluating whether transferred Treg cells in cirrhotic $\text{Rag1}^{-/-}$ mice restrained the induced proinflammatory T$_h$1 differentiation as a possible explanation for the function of Treg cells in gut barrier improvement. Figure 3A shows the flow cytometry gating strategy for determining T$_h$1 subpopulations in ILs of $\text{Rag1}^{-/-}$ mice in response to oral E. coli administration. $\text{Rag1}^{-/-}$ mice showed a low T$_h$1 differentiation from residual intestinal CD4$^+$ cells, as expected (Fig. 3B). The injection of T naive cells induced exacerbated proinflammatory T$_h$1 and T$_h$17 responses that were significantly down-regulated by the cotransfer with Treg cells. The T$_h$2 profile was not altered in any experimental condition.

Because Treg cells exert their action through IL-10 production, we set out to determine transcript expression levels of IL-10 in intestinal lymphocytes. IL-10 levels in cirrhotic $\text{Rag1}^{-/-}$ mice cotransferred with naive T and Treg cells were significantly increased in response to E. coli compared with the rest of the groups, providing a counterbalance for the observed T$_h$1 proinflammatory polarization (Fig. 3C).

**TREG CELLS CONTRIBUTE TO SUSTAIN SCFA LEVELS IN THE INTESTINAL CONTENT OF CIRRHOTIC $\text{Rag1}^{-/-}$ MICE AFTER EXPOSURE TO E. COLI**

Microbiota products such as SCFAs also play a role in maintaining gut homeostasis. We evaluated whether Treg cells are also relevant in SCFA modification in response to E. coli, as a model of induced BT, in the intestinal content of cirrhotic $\text{Rag1}^{-/-}$ mice. The concentration of all measured SCFAs resulted in a significant reduction after E. coli administration, but for acetic acid in nontransferred $\text{Rag1}^{-/-}$ mice. The concentration of all SCFAs remained unaltered in the intestinal content of cirrhotic $\text{Rag1}^{-/-}$ mice cotransferred with naive T and Treg cells except for valeric and isovaleric acids (Fig. 4A). The gene expression levels of receptors FFAR2 and FFAR3 for these SCFAs were measured in colonic tissue homogenates (Fig. 4B). Both receptors were significantly reduced in response to E. coli in cirrhotic $\text{Rag1}^{-/-}$ mice either nontransferred or transferred with naive T cells. The cotransfer with naive T and Treg cells was able to reestablish FFAR2 and FFAR3 levels shown in cirrhotic $\text{Rag1}^{-/-}$ mice not challenged with E. coli.
Discussion

In the present study, we show that Treg cells help maintain reduced gut permeability to bacterial antigen translocation in experimental cirrhosis by improving the barrier integrity. Additionally, in response to an induced bacterial challenge, Treg cells restrict an exacerbated proinflammatory Th commitment and preserve the levels of the most representative anti-inflammatory SCFAs and their receptors. These results give rise to the possibility of designing new interventions aimed at promoting Treg cell differentiation to prevent increased permeability and bacterial antigen translocation episodes in advanced cirrhosis.

BT is a common accepted mechanism for the development of significant bacteria-related complications in cirrhosis, and its prevention must be considered as a permanent goal during disease progression. In the past, we have reported that Treg cells are relevant to the inverse correlation observed between bacterial antigen translocation and norfloxacin through the anti-inflammatory cytokine IL-10 in CCl₄-induced cirrhosis. In the present study, we show in vivo that gut permeability to bacterial antigens is sensitive to the Treg cell population in cirrhotic mice, as orally administered LPS recovered in blood is significantly increased in cirrhotic mice lacking adaptive immunity, and it is down-regulated when these mice are cotransferred with spleen-derived naive T and Treg cells from WT donors (Fig. 1). As a result, these mice also show significantly decreased rates of bacterial DNA translocation in MLNs compared with nontransferred and naive T transferred Rag1⁻/⁻ mice.

It is important to outline that, although T naive transfer into Rag1⁻/⁻ mice is frequently used as an experimental model of spontaneous colitis, which normally takes several weeks to develop, our interest in this model was its intrinsic lack of adaptive immunity. We used it to study the short-term effect (48 hours before laparotomies) of T-cell replenishment in a liver damage–induced protocol. Under these conditions, Rag1⁻/⁻ mice do not develop colitis.

We have proposed that inflammation precedes BT, despite the fact that they feedback each other, in cirrhosis. In fact, systemic inflammation has been described in the absence of BT in experimental cirrhosis, and the reduction of inflammatory environment and improvement of gut barrier integrity markers by means of the Farnesoid X receptor agonist obeticholic acid has been associated with a significant reduction of BT in cirrhotic rats. Our results, in line with this evidence, would point to a progressive depletion of the intestinal Treg population that controls local proinflammatory differentiation as a possible mechanism to explain increased BT observed in advanced cirrhosis.

We provide solid data on the Treg cell involvement in the expression of TJ proteins in colon samples of cirrhotic mice (Fig. 2). The important function of these proteins in preventing paracellular movement across the barrier suggests an effect on expression and distribution of these proteins as a relevant mechanism in the association between Treg cells and a reduced permeability to small bacterial products. Although the transcellular passage of bacterial antigens cannot be discarded, the widening of intercellular spaces has been described among the structural changes of the intestinal mucosa, reviewed by Wiest et al. A decreased expression of ZO-1 and occludin in colon samples of bile duct–ligated mice has been described, and reduced expression of TJ proteins has also been reported in duodenal biopsies from patients with cirrhosis. Considering that the improved expression of different TJ proteins has been reported in experimental cirrhosis with the use of probiotic strains, some of which are known to stimulate Treg cell expression in intestinal mucosa upon interaction with DCs, it is plausible that the progressive, cirrhosis-related inflammatory environment in which T₇₇ and T₈ CD4⁺ T cells are differentiated is accompanied by the
reduction of the Treg population and the decreased expression of TJ proteins. This likely favors increased permeability and BT.

Functionally, the inflammatory control established by Treg cells in response to an oral bacterial challenge is clearly shown in cirrhotic Rag1°/° mice (Fig. 3).
lack of Treg cells is associated with an increased percentage of intracellular IFN-γ and IL-17 expressing cells. Inversely, intestinal lymphocyte–derived IL-10 mRNA expression is significantly increased in Treg-cotransferred Rag1−/− mice. T-cell activation requires the expression of DCs’ costimulatory molecules.\textsuperscript{(42)}
IL-10-increased levels may compromise DCs’ costimulation, switching their phenotype and decreasing effector T-cell activation. In fact, CD80 and CD86 have shown to be down-regulated in DCs from norfloxacin-treated CCl₄ mice only after restoration of the Treg cell population.(29) The evidence that IL-10-deficient cirrhotic mice show increased permeability to bacterial products such as LPS or DNA(28) also supports the implication of Treg cells as relevant IL-10-producing cells(43) in providing the balanced tissue environment to prevent BT episodes in cirrhosis. Whether Tregs can modulate other immune cell populations remains to be addressed in this context and may represent a limitation in this study.

Treg cells stand central to the interplay between the host and microbial milieu. (44) Cirrhosis progression is associated with intestinal bacterial overgrowth, and the gut microbiota dysbiosis toward an increment of Proteobacteria and the reduction of SCFA-producing species. (32,34) Accordingly, the concentration of propionic, butyric, and isobutyric acids in cirrhotic colonic tissue. The improvement of the intercellular sealing likely reduces paracellular translocation of antigenic bacterial products. Treg cell participation in controlling local inflammatory differentiation and bacterial-derived metabolic products such as SCFAs, which help provide homeostatic differentiation, fosters the value of this population in restricting BT episodes in cirrhosis. More studies are needed to establish whether restoring the number of Tregs may have any advantages such as fewer BT episodes or less portal hypertension and liver failure in cirrhosis.

Authors’ contributions: O.J., P.P., I.G.H., E.C.: experimental work, data acquisition, and manuscript writing; R.G.V., A.M., F.T.: short-chain fatty acids measurements; P.Z., J.M.G.N.: statistical analysis and interpretation of results; F.T.: flow cytometry analysis; and R.F.: study concept and design, manuscript writing.

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Supporting Information

Additional Supporting Information may be found at onlinelibrary.wiley.com/doi/10.1002/hep4.1268/suppinfo.