Intestinal development and homeostasis require activation and apoptosis of diet-reactive T cells

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The impact of food antigens on intestinal homeostasis and immune function is poorly understood. Here, we explored the impact of dietary antigens on the phenotype and fate of intestinal T cells. Physiological uptake of dietary proteins generated a highly activated CD44+Helios+CD4+ T cell population predominantly in Peyer patches. These cells are distinct from regulatory T cells and develop independently of the microbiota. Alimentation with a protein-free, elemental diet led to an atrophic small intestine with low numbers of activated T cells, including Tfh cells and decreased amounts of intestinal IgA and IL-10. Food-activated CD44+Helios+CD4+ T cells in the Peyer patches are controlled by the immune checkpoint molecule PD-1. Blocking the PD-1 pathway rescued these T cells from apoptosis and triggered proinflammatory cytokine production, which in IL-10-deficient mice was associated with intestinal inflammation. In support of these findings, our study of patients with Crohn’s disease revealed significantly reduced frequencies of apoptotic CD4+ CD44+ T cells in Peyer patches as compared with healthy controls. These results suggest that apoptosis of diet-activated T cells is a hallmark of the healthy intestine.

Introduction

Immunological tolerance to environmental factors such as intestinal bacteria and food antigens is an active, multilayered process, and its breakdown leads to hyperactivation of mucosal T and B cells and subsequent development of intestinal pathologies (1–3). Successful dietary intervention therapies in patients suffering from celiac disease, food allergies, and inflammatory bowel disease (IBD) demonstrate that diet contains immunologically relevant antigens (4–6). Yet, our understanding of diet-specific immune reactivity is largely derived from T cell receptor (TCR) transgenic mice and model antigens where tolerance mechanisms comprising clonal anergy, deletion, and induction of regulatory T cells (Tregs) have been shown (7–9). These data reveal that the contribution of clonal anergy/deletion versus active suppression by Tregs depends on type, dose, and frequency of applied antigens. Even though the vast majority of peripheral Tregs (pTregs) in the lamina propria (LP) of the small intestine is induced by normal food, oral tolerance is still intact if functional Treg levels are strongly reduced (10, 11).

Although the LP is not considered a typical site of immune induction, Peyer patches (PPs) are specialized lymphoid follicles known for effective stimulation of B and T cells by luminal antigens. Surprisingly, little attention has been paid to their role in tolerance against dietary antigens. This might be due to controversial findings showing, on the one hand, that tolerance occurs in PP-deficient mice and, on the other hand, that PPs are required for oral tolerance (12–15).

Here, we investigated the occurrence of diet-activated T cells in various tissues and characterized their phenotype and fate. We observed that recognition of dietary antigens is essential for the cellular and functional maturation of the small intestine, including generation of Tfh cells and IgA production. In the healthy murine and human gut, food-activated CD4+ T cells in PPs exhibited a proapoptotic phenotype characterized by the transcription factor Helios and the programmed cell death protein 1 (PD-1). In contrast, low Helios expression and reduced apoptosis were observed in PP CD4+ T cells of patients with IBD, suggesting that activation and subsequent death of food-reactive T cells is a hallmark of intestinal homeostasis.

Results

PPs harbor a population of activated Helios+Foxp3+ CD4+ T lymphocytes. First, we addressed the question of whether normal dietary antigens leave a signature of immune activation in nontransgenic animals. To differentiate between stimulation by microbial and
T cells for their regulatory function (18, 19). Although the usage of TCR Vβ chains was broad and similar to Tregs, these cells lacked expression of CD25, CTLA-4, and IL-10, all typically expressed by Tregs (Figure 1, F and G).

Moreover, Helios+Foxp3–CD4+ PP T cells were not able to suppress proliferation of effector T cells (Figure 1H), did not secrete IL-4, IFN-γ, or IL-17 after polyclonal stimulation, and were distinct from a subgroup of latency-associated peptide–expressing (LAP-expressing) Tregs, shown to produce IL-10 and TGF-β (Supplemental Figure 3) (20).

Helios+Foxp3–CD4+ T cells in PPs are activated by dietary antigens. As food is the major source of antigens in GF mice, and thus might explain the high frequency of Helios+Foxp3–CD4+ T cells, we assumed that this population should be affected if animals were kept on an antigen-free, elemental diet (ED). Low proliferation of PP CD4+ T cells in ED mice resulted in a vastly decreased frequency and number of Helios+Foxp3–CD4+ T cells as compared with animals kept on conventional diet (ConvD). Much less pronounced, but still affected by ED, was the number of Tregs in PPs despite unaltered frequencies (Figure 2, A–D).

With exception of the sucking period, the occurrence of Helios’ Foxp3 CD4+ T cells in ConvD and their decline in ED mice was age independent (Supplemental Figure 4, A–C).
To study kinetics of food antigen–dependent expansion of Foxp3 Helios+ and Foxp3-CD4+ T cells, ConvD was replaced by ED for 4 weeks and then switched to ConvD again. A gradual decline in the frequencies of Helios+Foxp3 CD4+ T cells in PPs was observed with the duration of antigen-free alimentation. These levels returned to normal after ED was replaced by ConvD. In contrast, frequencies of Foxp3+ Tregs were not altered by a change of diet (Figure 2D). Flow cytometric analysis of PP CD4+ T cells indicated that the frequency of activated CD4+ Helios+Foxp3- T cells was affected by ED (Figure 2, E and F). The effect of dietary antigens on total PP CD4+ T cells was also seen in GF animals. GF mice fed ED showed further declined frequencies and total numbers of Helios+Foxp3-CD4+ T cells in the PPs, as compared with GF mice kept on ConvD. Of note, despite the unaltered frequency of the Foxp3+ population, the low intestinal cellularity in GF mice kept on ConvD. Of note, despite the unaltered frequency of activated CD4+Helios+Foxp3- T cells in the GC and diminished frequencies of IgA+ B lymphocytes. This explains the low amounts of secretory IgA in the small intestines of ED mice (Figure 3, F and G, Supplemental Figure 8). Importantly, normal thymic T cell development, together with unaltered T and B cell numbers in the spleen, demonstrated that food antigen deficiency does not disturb systemic lymphocyte development (Supplemental Figure 9) but specifically affects expansion and survival of lymphocytes within the small intestine.

**Helios induction by sustained TCR-restricted antigen stimulation.** To investigate whether dietary antigens are recognized in a TCR-specific manner, we analyzed RAG-sufficient OT-II mice in which the vast majority of T cells expresses the OVA-specific Vα2+ TCR and a minor population of T cells, including diet-reactive T cells, expresses an endogenous Vα2+, polyclonal TCR that preferentially accumulated in PPs (Figure 4A). In the absence of OVA, most Vα2+ T cells were Helios-, whereas more than 10% of the Vα2+ CD4+ T cells revealed a signature of food antigen activation in PPs but not in spleen or mLNs (Figure 4B). Correspondingly, OVA-supplemented drinking water triggered the expansion of Helios- Foxp3 Vα2+ T cells in PPs (Figure 4C), but also in mLNs and spleen (data not shown). OVA-stimulated Vα2+ PP T cells differentiated into Helios+Foxp3 or Helios-Foxp3 phenotypes, demonstrating that the majority of Vα2+ T cells expressed Helios independently of Foxp3 and only a small fraction differentiated into Helios+Foxp3 Tregs (Figure 4D). In vitro studies further confirmed that Foxp3-independent induction of Helios expression requires recognition of cognate antigen (Supplemental Figure 10).
These results show that induction of the Helios<sup>+</sup>Foxp3<sup>−</sup> phenotype requires TCR-mediated recognition of dietary antigens, with Va2<sup>+</sup> T cells being specifically activated by OVA but not by other dietary antigens, whereas Va2<sup>−</sup> T cells react in the opposite way.

**Dietary antigen impact on the transcriptional profile of PP CD4<sup>+</sup> T cells.** Since stimulation with food antigens is crucial for the maturation of the intestinal immune system, we investigated the mRNA profile of PP CD4<sup>+</sup> T cells in response to dietary antigens and their activation state. We wondered whether continuous exposure to dietary antigens leads to a phenotype of exhaustion similar to that described for chronic viral infections. In contrast to CD4<sup>+</sup> T cells from chronic viral infection, the transcriptional signature of ConvD-activated CD4<sup>+</sup> T cells was slightly skewed toward exhaustion. CD4<sup>+</sup> T cells from ConvD mice expressed increased mRNA for Lag3, Icos, Pdcd1, and Tigit as compared with ED mice (Figure 5A).

In accordance with our phenotypic analyses, the expression pattern of ConvD animals was markedly biased toward a Tfh cell signature, including transcripts for IL-21, Cxcr5, Bcl6, Pdcd1, and Ascl2. While Tfh cell differentiation may be controlled by a Bcl6- or Ascl2 gene-centered network, the latter network seems to apply to Tfh cell polarization by dietary antigens, as illustrated by positive gene association upon ConvD treatment (Figure 5B).
Finally, we compared the transcriptional profile of activated CD44hiCD62L+Foxp3+ versus naive CD44loCD62L-Foxp3- CD4+ T cells in the PPs of ConvD mice. Gene enrichment analysis revealed a marked upregulation of genes that are associated with apoptosis and oxidative phosphorylation in activated CD4+ T cells as compared with their naive controls (Figure 5, C and D). In summary, these gene signatures show that not only the microbiota but also food antigens induce their naive controls (Figure 5, C and D). In addition, OX40 and Bcl-2, both essential for long-term survival of CD4+ T cells, were almost absent in food-reactive T cells but were strongly expressed in Tregs of PPs (Figure 6, C and D) (21). As lack of survival molecules is common in cells prone to apoptosis, we performed annexin V/PI staining for T and B cells of SPF and GF mice. Strong staining of annexin V was exclusively seen within CD4+ T but not B cells in PPs of SPF and GF mice. To further examine the fate of food antigen–reactive Helios+Foxp3+ CD4+ T cells, we first analyzed the expression of survival molecules required for cell survival. While the majority of CD4+ T cells in spleen and mLNs expressed the IL-7Rα (CD127), a substantial population of PP T cells, including most food protein–reactive T cells, downregulated this survival receptor (Figure 6, A and B). In addition, OX40 and Bcl-2, both essential for long-term survival of CD4+ T cells, were almost absent in food-reactive T cells but were strongly expressed in Tregs of PPs (Figure 6, C and D) (21). As lack of survival molecules is common in cells prone to apoptosis, we performed annexin V/PI staining for T and B cells of SPF and GF mice. Strong staining of annexin V was exclusively seen within CD4+ T but not B cells in PPs of SPF and GF mice (Figure 6E and Supplemental Figure 11, A and B). The apoptotic signature was mainly confined to food-activated CD44hiHelios+Foxp3+ CD4+ T cells (Figure 6, F and G) and was significantly decreased in animals fed ED (Figure 6H).

To define the pathway of apoptosis, sorted CD44hiCD4+ and CD44loCD4+ T cells from PPs of ConvD animals were analyzed for cleavage of poly (ADP-ribose) polymerase (PARP) and caspase activation. Strong cleavage of PARP together with activation of caspase-3 and -8 but not caspase-9 were seen in CD44hi but not in CD44lo T cells (Figure 6I), suggesting that programmed cell death is the fate of diet-activated T cells in PPs of the healthy intestine.

Ingestion of apoptotic cells has been shown to induce IL-10 and to inhibit proinflammatory cytokine production, thus preventing chronic inflammation or autoimmunity (22, 23). Thus, we tested whether apoptotic cells are linked with the induction of IL-10 in PPs. In contrast to spleen and mLNs, substantial amounts of IL-10 were produced by cultured cells of PPs. Importantly, IL-10 secretion by PP-derived cells was dependent on dietary antigens, as only marginal amounts of IL-10 were detectable in ED animals (Supplemental Figure 12, A and B). Further, cocultures of macrophages with lymphocytes from various organs revealed that uptake of T lymphocytes from PPs but not spleen or mLNs triggered macrophage-derived IL-10 synthesis. Secretion of IL-10 correlated with the amount of dead cells, as staurosporine pretreatment further enhanced this effect (Supplemental Figure 12C).

Apoptosis of Helios+Foxp3+ CD4+ T cells is controlled by PD-1. PD-1 is known to be an important regulator of T cell activation and apoptosis (24). We thus tested the expression of PD-1 on both Helios+ and Helios- T cell populations in the PPs. Flow cytometric analysis revealed selective expression of PD-1 in Helios+ but not Helios- PP T cells (Figure 7, A and B). Further, PD-1 ligands PDL-1 and PDL-2 were found to be predominantly expressed on CD11b+ and CD11c+ APCs of the PPs (Figure 7C).

Next, we determined whether PD-1 signaling controls the response of PP CD4+ T cells. To this end, we either injected PD-1-blocking antibodies or isotype controls into normal C57BL/6 mice i.p. or added anti-PD-1 antibodies directly into PP cell cultures. Polyclonal stimulation of PP cell suspensions revealed that anti-PD-1 treatment but not isotype control rescues PP CD4+ T cells from apoptosis to become IFN-γ-secreting effector cells (Figure 7, D–F).

In inflammatory bowel disease, such as Crohn’s disease (CD), a variety of models have been used, including the IL-10–KO mouse which develops inflammatory reactions closest to CD (25). We wondered whether modulation of the PD-1 pathway leads to disease.
Figure 5. Transcriptional profiling of food antigen–dependent CD4+ T cells in PPs. (A) CD4+ T cells from PPs were sorted and analyzed by RNA sequencing. The gene expression signature of T cell exhaustion was compared between ConvD and ED mice. Each dot represents mean transcripts per million (TPM) ± SD for 6 mice per group. (B) Key gene TPM associated with Tfh function were transformed to Z scores and compared between ConvD and ED mice (6 mice per group). ASCL2 and Bcl6 target genes were investigated by Gene Set Enrichment Analysis (GSEA). (C and D) Genes identified by RNA sequencing of CD44+CD62L–Foxp3–CD4+ and CD44–CD62L+Foxp3–CD4+ T cells from PPs of ConvD mice were investigated by GSEA for enrichment of the oxidative phosphorylation pathway and apoptosis as defined in the KEGG database. NES, normalized enrichment score; FDR, false discovery rate.
progression in IL-10–deficient mice. In contrast to WT controls, IL-10–KO mice treated with anti–PD-1 mAb developed severe colitis, accompanied by enhanced cellularity of T cells and increased numbers of IFN-γ+CD4+ T cells in the PPs (Supplemental Figure 13, A–D).

Although these data suggest that PD-1 immune checkpoint blockade in the absence of IL-10 favors colitis development, the precise contribution of PP Helios+CD4+ T cells remains unclear.

Reduced apoptosis of PP T cells in patients with IBD. Finally, we investigated the apoptotic behavior of human PP CD4+ T cells in healthy controls and patients with CD. Biopsies from PPs of the terminal ileum were taken with written consent from persons who had undergone routine intestinal health surveillance or patients with clinically diagnosed CD. The latter group was selected for discrete inflammation of the colon and/or ileum, whereas none of the controls showed signs of inflammation (Supplemental Table 1). To assure that T cells were derived from PPs, lymphocytes were stained and gated for expression of the germinal center marker GL7 prior to further analysis. While approximately 40% of the CD4+ GL7+ PP T cells from healthy controls were annexin V−, patients with CD revealed significantly lower frequencies of apoptotic PP T cells (approximately 22%) (Figure 8, A and B). Increased survival of PP T cells in CD was reflected by enhanced expression of Bcl-2 in viable cells (Figure 8C). Consistent with the finding that Helios marks CD4+ T cells undergoing programmed cell death with subsequent downregulation of Bcl-2 (26, 27), the expression of Helios in Foxp3−CD4+ T cells from PPs of patients with CD was significantly reduced as compared with healthy controls (Figure 8D). These data suggest that high apoptosis of PP CD4+ T cells is a homeostatic mechanism of the healthy gut.

Discussion

Although the value of nutritional treatment in patients with IBD was first reported over 30 years ago, a poor understanding of immunological mechanisms hinders both acceptance and use of medical diets (28, 29). From an immunological point of view, diet contains components of potential antigenicity that the immune system should not actively challenge. Accordingly, several studies have demonstrated that exposure to dietary antigens results in the generation of Tregs. However, the phenotype and fate of food-reactive T cells in the inductive tissue sites have received little attention.
Antigen sampling of the luminal content of the gut is a prerequisite for monitoring both harmless and harmful antigens, and for responding appropriately. In the small intestine, efficient encounter between APCs and antigen-specific T cells takes place in specialized secondary lymphoid organs, that is, mLN s and PPs. Particularly, M cells and the follicle-associated epithelium (FAE) of PPs have been shown to efficiently take up and transport antigens from the gut lumen to immune-inductive sites of PPs (30, 31).

Our search for diet-activated T cells in gut-associated and systemic lymphoid tissues revealed an unusually high frequency of activated CD44¹Helios¹Foxp3–CD4⁺ T cells exclusively in PPs of SPF and GF mice, indicating that activation was mediated by dietary rather than microbial antigens. This finding is in accordance with a previous observation that antigen-experienced T cells naturally accumulate in PPs but not mLN s with age, suggesting that these cells received antigenic stimulation directly in PPs and not in other sites of the intestine. Only a small proportion of these PP-resident CD62L⁺CD4⁺ cells were follicular helper T cells (Tfh) or Tregs (32). In contrast, Tregs became a prominent population in the siLP after weaning of GF animals to ConvD, and alimentation with ED prevented the development of Tregs in the siLP (10).

Preferential expansion of pTregs in the siLP and Helios¹Foxp3–CD4⁺ T cells in PPs seems to reflect the distinct morphology of both tissue sites. In PPs, the majority of DCs express CCR6 and recruit antigen from the subepithelial dome (SED) region before clustering with antigen-specific CD4⁺ T cells in the PPs but not in LP, supporting their selective role in intestinal-inductive sites (33). In contrast, the majority of LP DCs express CXCR1, are CD103⁺, and capture antigen from the intestinal lumen via transepithelial dendrites (34). In the presence of TGF-β and retinoic acid, LP-derived CD103⁺CD11b⁺ DCs have been shown to efficiently induce Foxp3⁺ Tregs (35, 36). Surprisingly, their role in mediating oral tolerance still remains enigmatic, as oral tolerance was intact in CD11b– DC-deficient animals with strongly reduced pTregs (11).

We here describe a population of ConvD-activated CD44⁺Helios⁺Foxp3–CD4⁺ T cells in PPs that is distinct from pTregs, although they express Helios, a marker recently used for differentiation between natural and inducible Foxp3⁺ Tregs (37). Since transient expression of Foxp3 has been described (38), there is a possibility that food-reactive CD44⁺Helios⁺Foxp3–CD4⁺ PP T cells may have originated from an “exFoxp3” population. Such reprogramming is highly unlikely as the frequency of Foxp3⁺ Tregs in PPs was not affected by the type of diet, whereas the Helios⁺Foxp3– population expanded or shrank in the presence or absence of dietary antigens, respectively.

The mechanism of Foxp3-independent Helios expression and its function in activated PP CD4⁺ T cells remains unresolved. Autonomous expression of Helios has been described during differentiation and activation of CD4⁺ T cells, for Tfh cells and autoreactive T cells undergoing apoptosis (27, 39). It is notable that oral antigen uptake preferentially induces IL-27 and IL-10 in PP DCs, both cytokines exhibiting antiinflammatory properties (40, 41). Correspondingly, we observed that lack of dietary protein vastly decreased the number of Helios⁺CD4⁺ T cells and the amount of IL-10 in PPs. Although we were unable to directly track food-reactive T cells in normal mice, analysis of OT-II animals revealed TCR-specific recognition of dietary antigens. Oral uptake of OVA induced the Helios⁺Foxp3⁻ phenotype within the TCR-transgenic Vα2⁺ population, whereas normal chow induced Helios in the endogenous Vα2 CD4⁺ T cell population within PPs.
An important consequence of food antigen recognition by the intestinal immune system is the cellular and functional maturation of the small intestine. In the absence of food antigens, a low amount of IgA together with reduced numbers of B and T cells, including Th cells, assigns a new and essential aspect to dietary proteins aside from nutrition; that is, their antigenicity is required for normal development and function of the small intestine.

Because the accumulation of diet-specific T cells in PPs might cause severe immunopathology if their activity and lifespan remain uncontrolled, we investigated the fate of these cells. In accordance with the downregulation of survival molecules, an exceptionally high level of apoptotic cells was found within the PPs. This increase was confined to the activated CD44+Helios+Foxp3−CD4+ T cell population and was decreased in ED animals, suggesting that caspase-3–mediated apoptosis of diet-activated T cells is a default population and was decreased in ED animals, suggesting that caspase-3–mediated apoptosis of diet-activated T cells is a default mechanism, which warrants intestinal homeostasis.

Helios+Foxp3−CD4+ PP T cells expressed high levels of PD-1, a molecule known to fine tune the fate and the function of T cells (42). We observed that anti-PD-1 treatment of PP T cells triggered IFN-γ production and simultaneously reduced the frequencies of apoptotic CD4+ T cells, indicating that PD-1 blockade rescues dysfunctional T cells from cell death. Interestingly, in the absence of IL-10, blocking of PD-1 favored development of colitis in these mice. Numerous reports show that colitis development remains a complication after treatment of cancer patients with an anti-PD-1 checkpoint inhibitor (43–45).

In line with our murine data, we found that PP CD4+ T cells from healthy donors but not patients with CD revealed a substantially increased frequency of apoptotic T cells with high expression of Helios and low expression of Bcl2. Although we have not yet identified the precise mechanisms for enhanced survival of PP CD4+ T cells in patients with IBD, disturbed apoptosis might explain food antigen–specific immune responses in patients with CD, which were ameliorated by restricting dietary antigens (3). In this context, clinical observations have suggested lymphoid follicles as the initial site of inflammation in patients with CD, and that peak frequency of CD correlates with peak number of PPs at an age between 15 and 25 years (46–48).

Here, we present an approach that combines murine and human studies for characterization of intestinal immune responses against dietary antigens. Our data suggest that apoptosis of PP lymphocytes is a hallmark of the healthy intestine, which eliminates potentially harmful food-reactive T cells and triggers the production of local IL-10. Reduced apoptosis of PP T cells in patients with CD might explain the beneficial effects of ED by depriving these T cells of their antigenic stimulus. These questions underline the importance of further investigating the molecular and cellular mechanisms of increased survival of PP T cells in patients with CD.

**Methods**

*Animals.* C57BL/6 mice were purchased from Charles River Laboratories and kept under specific pathogen–free (SPF) conditions. DEREG mice were provided by Tim Sparwasser (Institute of Infection Immunology, Hannover, Germany). IL10−/− mice, OT-II, and GF C57BL/6 mice were bred at the animal facilities at the University of Marburg. GF mice generated via rederivation through cesarean section were a gift of Petra Kirsch (University of Ulm). GF animals were kept in plastic isolators (Metall and Plastik) with autoclaved food, bedding, and water. Sterility of animals was checked biweekly by culturing feces in thioglycollate broth medium under aerobic and anaerobic conditions for at least 10 days. All handling procedures for GF mice were conducted in a laminar flow hood under sterile conditions.

*Antibiotic sterilization of mice.* To deplete the commensal gut microbiota, 6- to 8-week-old ConvD mice were transferred to sterile cages and orally treated with antibiotic cocktail (1 g/l ampicillin, 0.5 g/l vancomycin, 1 g/l metronidazole, 0.25 g/l imipenem, and 0.2 g/l ciprofloxacin) for 8 weeks according to a standard protocol (49). Bottles with antibiotic-containing drinking water were replaced once per week. The intestinal microbial status was tested weekly by culturing fecal samples under aerobic and anaerobic conditions in thioglycollate broth.

*Animal nutrition.* C57BL/6 mice (SPF and GF) at 6–20 weeks of age were used. The ConvD group were kept on a conventional chow (LASQCDietRod16, LASvendi) and food antigen–free, ED mice were kept on an amino acid–containing diet for up to 5 generations. Pellets of ED diet (ssniff, S7242-E014/-E714) contained all essential vitamins, minerals, trace elements, fat, dextrin, sucrose, and free amino acids.
equimolar to the protein content of normal rodent chow. In summary, the ED pellets contained 20% corn starch, 44% sucrose, 7% cellulose, 17.5% amino acid mixture, 1% vitamin premix w/o choline, 5% mineral trace element premix, 0.2% choline CI, 0.2% sodium bicarbonate, 0.02% butylated hydroxytoluene, 5% corn oil.

Oral administration of OVA. The drinking water was supplemented with 1 mg/ml OVA (Sigma-Aldrich) for low-dose OVA application. OVA-supplemented water was given to OT-II animals for 5 or 7 days prior to analysis.

Isolation of lamina propria mononuclear cells. Single-cell suspensions were performed from spleen, mLNs, and PPs by mechanical disruption and passage through filter (Milteny Biotech). Lamina propria mononuclear cells (LPMCs) of the colon and small intestine were isolated after enzymatic digestion of intestinal tissue pieces with collagenase D and collagenase VIII (MilliporeSigma, 0.4 mg/ml each). Cells were purified by Percoll gradient (40%/70%). LPMCs were collected from the interphase, washed, and resuspended in complete medium.

Small intestinal IgA. The complete small intestine was flushed with 5 ml PBS/2% milk powder, and the intestinal liquid was collected, homogenized, and centrifuged (850g, 30 minutes). Supernatants were used for IgA ELISA. In brief, flat-bottom 96-well plates (NUNC) were coated with rabbit-anti-mouse IgA (Rockland) overnight. After blocking with PBS/10% FCS, 50 μl intestinal liquid was incubated for 3 hours at 4°C, washed twice, and IgA was detected by adding the secondary rabbit-anti-mouse IgA HRP and TMB (R&D Systems). The enzymatic reaction was stopped by the addition of sulphuric acid and absorption was measured with an ELISA Reader (FLUostar Omega, BMG Labtech).

IL-10 ELISA. Single-cell suspensions from PPs, mLNs, and spleen of Conv and ED mice were cultured at a density of 2 × 10⁶ cells/200 μl RPMI medium in round-bottom 96-well tissue-culture plates for 24 hours. The next day, cell-culture supernatants were collected and IL-10 secretion was measured using a murine IL-10 ELISA kit (BD Biosciences) according to the manufacturer’s instructions.

Antibodies, reagents, and staining methods. The following antibodies were obtained from eBioscience unless otherwise stated: anti-CD4 (RM4-5, BioLegend), anti-CD25 (PC61.5), anti-Vβ1, anti-CD279 (anti–PD-1, clone J43) for in vivo screening panel (BD Biosciences), anti-Bcl2 (10C4), anti–PD-L1 (MIH7, BioLegend), and anti–CXCR5 (2G8, BD Biosciences), TCR Vβ (B20.1), anti-B220 (RA3-6B2, BD Biosciences), anti-CD69 (H1.2F3), anti–CTLA-4 (UC10-4F10-11, BD Biosciences), anti–OX40 (OX-86), anti–CXC5R (2G8, BD Biosciences), TCR Vβ screening panel (BD Biosciences), anti–Bcl2 (10C4), anti–PD-L1 (MIH7, BioLegend), and anti–PD-L2 (catalog 122). Anti-CD279 (anti–PD-1, clone J43) for in vivo application was purchased from Hölzelagnostika. Detection of apoptotic cells was performed with annexin-V–FLUOS staining kit (Roche Applied Science). For intracellular cytokine staining, PP cells were stimulated for 4 hours with PMA (50 ng/ml) and ionomycin (750 ng/ml in the presence of Brefeldin A (5 μg/ml, MilliporeSigma). Subsequently, cells were stained with anti-CD4, fixed, and permeabilized in 0.3% saponine buffer before staining for cytokines. Intracellular staining was performed for anti–IL-10 (JESS-16E3). Foxp3- and Helios-expressing cells were routinely detected by Foxp3 staining kit (eBioscience), anti–Foxp3 (FJK-16s), and anti–Helios (22F6). Cells were analyzed by flow cytometry using the FlowJo software (Tree Star).

RNA-Seq and bioinformatical analysis. RNA was purified from sorted PP CD4+ T cells of ConvD and ED mice with the RNeasy Plus Micro Kit according to the manufacturer’s protocol (Qiagen). RNA was quantified with a Qubit 2.0 fluorometer (Invitrogen) and the quality was assessed on a Bioanalyzer 2100 using an RNA 6000 Pico chip (both from Agilent). Samples with an RNA integrity number (RIN) of greater than 8 were used for library preparation. Barcoded mRNA-Seq cDNA libraries were prepared from 10 ng total RNA using NENxt Poly(A) mRNA Magnetic Isolation Module and NENxt Ultra II RNA Library Prep Kit for Illumina (New England Biolabs) according to the manual. Quantity was assessed using Invitrogen’s Qubit HS assay kit and library size was determined using Agilent’s 2100 Bioanalyzer HS DNA assay. Barcoded RNA-Seq libraries were onboard clustered using HiSeq Rapid SR Cluster Kit v2 using 8 μm, and 59 bps were sequenced on the Illumina HiSeq2500 using HiSeq Rapid SBS Kit v2 (59 cycle). The raw output data of the HiSeq were preprocessed according to the Illumina standard protocol. After initial quality assessment with the FastQC software (available online at http://www.bioinformatics. babraham.ac.uk/projects/fastqc/), raw output fastq sample files were trimmed and mapped with the Qiagen CLC Workbench v.10.0.1, using the murine genome version GRCh38. Total read counts were further processed in R, using the DeSeq2 package to compute differential gene expression and adjusted P values. Normalized read counts were ranked and investigated for pathway-specific enrichment by GSEA (available online at http://software.broadinstitute.org/gsea/index.jsp) and were used for graphic representation of differential expression.

The RNA-Seq microarray data were deposited in the NCBI’s Gene Expression Omnibus database (GEO GSE124286) and are available at https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE124286.

Anti–PD-1 treatment of mice and cell cultures. WT and IL10–deficient mice (C57BL/6 background) were administered 100 μg anti–PD-1 Ab (clone J43, Hölzelagnostika) i.p. on days 0, 4, 8, and 12. As isotype control, animals were injected with 100 μg polyclonal Armenian hamster IgG (Hölzelagnostika). On day 15, mice were analyzed for signs of inflammation and intestinal CD4+ T cells were collected and analyzed for apoptosis and cytokine expression by flow cytometry.

In vitro studies, 3 μg/ml anti–PD-1 mAb together with 2.5 μg/ml anti–CD3/CD28 were added to single-cell suspensions from PPs. After 3 days of culture, cells were stimulated for 4 hours with PMA/ ionomycin in the presence of Brefeldin A and analyzed by FACS.

Human Peyer patch T cells. Cells from PP biopsies were isolated by mechanical dissociation through 100-μm and 40-μm cell strainers. The strainers were washed with RPMI 1640 plus 1% Pen/Strep plus 10% FCS (Life Technologies, Gibco) and cells were centrifuged at 400g for 10 minutes at 4°C and resuspended in PBS (Life Technologies, Gibco) for flow cytometric analyses. For surface staining, fluorochrome-conjugated antibodies CD4 (RPA-T4) and GL7 were added and incubated for 15 minutes at 4°C. For the annexin V staining, cells were resuspended in binding buffer and stained with annexin V and 7-AAD (Annexin V Apoptosis Detection Kit, eBioscience). For intracellular staining of cytokines and transcription factors (Bcl2 [100/ D5+124], Foxp3 [PCH101], and Helios [22F6]), cells were fixed, permeabilized, and stained using the Foxp3 staining kit (eBioscience). Flow cytometry analysis was performed on a FACS Canto II (BD Bioscience). Data were analyzed with FlowJo analysis software.

Treg suppression assay. Suppressive properties of Tregs and CD4+Helios Foxp3 cells from PPs were compared. CD4+ GFP responder T cells from spleen and enriched Helios-expressing CD4+ GFP CD44+ CD62L+ T cells were sorted from PPs of DEREG mice. As positive control, iTregs were generated from sorted naive CD4+ T cells
of DEREG mice. In brief, cells were activated by plate-bound anti-CD3 (5 μg/ml; 145-2CD11) and soluble anti-CD28 (1 μg/ml; 37.51) in the presence of IL-2 (50 U/ml; Novartis), TGF-β1 (2 ng/ml, R&D Systems), anti-IL-4 (10% culture supernatant of clone 1B11), and anti-IFN-γ (5 μg/ml, XM1G-2) for 3 days. Responder T cells were labeled with CFSE and cocultured for 3 days with Tregs or Helios-expressing T cells from PPs in the presence of 1 μg/ml soluble anti-CD3 (clone 145-2CD11) and irradiated T cell-depleted splenocytes. Proliferative responses of responder cells were analyzed by FACs.

BrdU incorporation assay. To label proliferating cells in vivo, 1 mg BrdU (Sigma-Aldrich) in 200 μl PB was injected i.p. and animals were subsequently supplied with BrdU (0.8 mg/ml) in drinking water for 5 days. Cell suspensions were stained with anti-CD-4 and anti-BrdU (eBioscience). The percentage of BrdU+ T cells was assessed by FACs.

Western blot analysis. CD44+ and CD44- T lymphocytes were sorted from PPs of C57BL/6 mice and lysed directly in 4× Laemmli sample buffer (with 5% β-mercaptoethanol). Proteins were separated by SDS-PAGE and transferred to nitrocellulose membrane. The following antibodies were used for immunoblot analysis: anti-GAPDH as a loading control, mouse anti-PARP, rabbit anti– caspase-3, mouse anti-caspase-8, and rabbit anti–caspase-9 (Cell Signaling Technology).

Immunofluorescence analysis. Tissue samples were snap frozen in methylbutane cooled with liquid nitrogen. Cryosections of about 5 μm were made with a Frigocut 2800E (Leica), air-dried, and incubated for 20 minutes in ice-cold acetone and stored at -20°C or washed 3 times with PBST for immunostaining. Following incubation in antibody diluent (Dako), samples were incubated with primary antibodies for 2 hours at room temperature followed by washing 3 times in PBST. The following primary antibodies were used: anti-mouse CD4–Alexa Fluor 646 (BioLegend) and anti-mouse GL7-FITC (BD Biosciences). Confocal images of cryosections were acquired by fluorescence microscopy on a Leica TCS SP2 microscope (Leica Microsystems).

Statistics. Statistical analysis was performed using Prism software (GraphPad Software). Results are generally expressed as the mean ± SEM. Data were analyzed using the Student’s t test or Mann-Whitney U test as indicated. Values with P less than or equal to 0.05 were considered statistically significant.

Study approval. All patients gave written consent after approval by the Charité ethics committee (EA4/097/14). All animal experiments were conducted in accordance with German animal protection laws and were approved by the RP-Giessen (MR 20/6 1/2018 and EX7-2015).

Author contributions
US and AV directed and designed the experiments and analyzed the data. PWB, OP, ML, and BS gave scientific advice and critically read the manuscript. AV, SH, ML, HR, RR, and FF performed the animal experiments and in vitro studies. RJ and AP performed the histological analysis. HM, WB, and MK performed and evaluated RNA-Seq experiments. KR performed the molecular and biochemical studies. YRS, RG, and BS conducted the human studies. US and AV wrote the manuscript.

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