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Although defects in intestinal barrier function are a key pathogenic factor in patients with inflammatory bowel diseases (IBDs), the molecular pathways driving disease-specific alterations of intestinal epithelial cells (IECs) are largely unknown. Here, we addressed this issue by characterizing the transcriptome of IECs from IBD patients using a genome-wide approach. We observed disease-specific alterations in IECs with markedly impaired Rho-A signaling in active IBD patients. Localization of epithelial Rho-A was shifted to the cytosol in IBDs, and inflammation was associated with suppressed Rho-A activation due to reduced expression of the Rho-A prenylation enzyme geranylgeranyltransferase-I (GGTase-I). Functionally, we found that mice with conditional loss of Rhoa or the gene encoding GGTase-I, Pggt1b, in IECs exhibit spontaneous chronic intestinal inflammation with accumulation of granulocytes and CD4+ T cells. This phenotype was associated with cytoskeleton rearrangement and aberrant cell shedding, ultimately leading to loss of epithelial integrity and subsequent inflammation. These findings uncover deficient prenylation of Rho-A as a key player in the pathogenesis of IBDs. As therapeutic triggering of Rho-A signaling suppressed intestinal inflammation in mice with GGTase-I-deficient IECs, our findings suggest new avenues for treatment of epithelial injury and mucosal inflammation in IBD patients.

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Although defects in intestinal barrier function are a key pathogenic factor in patients with inflammatory bowel diseases (IBDs), the molecular pathways driving disease-specific alterations of intestinal epithelial cells (IECs) are largely unknown. Here, we addressed this issue by characterizing the transcriptome of IECs from IBD patients using a genome-wide approach. We observed disease-specific alterations in IECs with markedly impaired Rho-A signaling in active IBD patients. Localization of epithelial Rho-A was shifted to the cytosol in IBDs, and inflammation was associated with suppressed Rho-A activation due to reduced expression of the Rho-A prenylation enzyme geranylgeranyltransferase-I (GGTase-I). Functionally, we found that mice with conditional loss of Rhoa or the gene encoding GGTase-I, Pagg1b, in IECs exhibit spontaneous chronic intestinal inflammation with accumulation of granulocytes and CD4+ T cells. This phenotype was associated with cytoskeleton rearrangement and aberrant cell shedding, ultimately leading to loss of epithelial integrity and subsequent inflammation. These findings uncover deficient prenylation of Rho-A as a key player in the pathogenesis of IBDs. As therapeutic triggering of Rho-A signaling suppressed intestinal inflammation in mice with GGTase-I–deficient IECs, our findings suggest new avenues for treatment of epithelial injury and mucosal inflammation in IBD patients.

Introduction

Forming a mechanical boundary against pathogens, antigens, and toxins, the monolayer of intestinal epithelial cells (IECs) represents the human body’s largest contact area with the environment (1). From their early development in the crypt bottom until shedding of aged cells at the villus tip, IECs follow a continuous turnover process (2–4). Under physiological conditions, a complex interaction between cytoskeleton rearrangement and tight junction proteins guarantees that the cell shedding itself does not mean a disturbance of epithelial integrity (5). However, alterations of epithelial integrity may lead to development of gut inflammatory disorders (1, 6), such as inflammatory bowel diseases (IBDs) (7). IBDs are associated with marked alterations of IECs, leading to increased tight junction permeability, altered cytoskeletal rearrangement, and induction of epithelial cell death with a subsequent loss of barrier function (1, 6). In vivo studies in IBD patients demonstrated that loss of IEC integrity even preceded the clinical relapse of disease (8), suggesting that alteration of barrier function is crucial in the pathogenesis of IBDs. However, the exact IEC intrinsic mechanisms involved in the initiation of barrier dysfunction remain enigmatic.

On a subcellular level, Rho GTPases represent signaling molecules centrally involved in arrangement of cytoskeletal proteins and epithelial cell dynamics (9–13). Since their activation preferentially takes place when GTPases are associated with cellular membranes (14, 15), function of Rho proteins crucially depends on their intracellular localization (16). Recruitment of small GTPases is governed by prenylation, a posttranslational process consisting of the attachment of hydrophobic isoprenoids to the C-terminal CAAX motif of the protein (17). In the intestinal epithelium, it has been shown that apoptotic as well as physiological cell extrusion requires Rho-mediated signaling pathways (18, 19). Moreover, Rho-A/ROCK/Myosin II interaction is known to be important for morphogenesis of the primitive gut tube in Xenopus (20). However, little is known about regulation of Rho-A signaling in the intestinal epithelium and its impact on gut homeostasis in vivo.

Using gene expression profiling of IECs from inflamed versus uninfamed tissue, we identified altered Rho-A signaling in IBD patients. Although Rho-A was expressed at normal levels, cytosolic accumulation of Rho-A in IECs suggested impaired Rho-A activation via prenylation upon inflammation. Consistently, genetic deletion of Rhoa or Pagg1b (encoding for the prenylation-catalyzing enzyme geranylgeranyltransferase-I [GGTase-I] driving Rho-A activation) in murine IECs impacted on cytoskeleton and cell shedding, resulting in epithelial injury and spontaneous gut inflammation. Our data demonstrate that injury in GGTase-I–
information of included patients is summarized in Supplemental Table 1 (supplemental material available online with this article; doi:10.1172/JCI80997DS1). Comparative genomic expression analysis of IECs from inflamed versus uninflamed gut areas of CD patients showed that samples within each group clustered, and we were able to identify 129 genes differing significantly (fold-change ≥ 1.5; \( P \leq 0.05 \)) between both groups (Figure 1A). Interestingly, gene ontology analyses indicated inflammation-associated down-regulation of pathways involved in epithelial cell dynamics and IEC extrusion, such as Rho-A signaling, epithelial adherens junctions, and regulation of actin-based motility by Rho (Figure 1B). Notably, pathways involving Rho GTPases other than Rho-A, such as Cdc42 signaling, showed no significant changes (Supplemental Figure 1A).

Although our data implicated a relation between impaired Rho-A function in IECs and inflammation in human IBDs, neither protein nor mRNA expression levels of Rho-A were found to be reduced in IECs from inflamed gut areas of CD or ulcerative deficient epithelium was driven by Rho-A dysfunction and that this phenotype could be successfully reversed by Rho activation. Together, our study highlights for the first time to our knowledge the relevance of functional Rho-A signaling and its regulation via prenylation for maintenance of epithelial integrity. Thus, prenylated Rho-A in epithelium emerges as a therapeutic target in IBDs.

### Results

**A crucial role of epithelial Rho-A in intestinal inflammation.** Intestinal erosions and ulcers caused by epithelial injury represent clinical hallmarks of IBDs, but the underlying pathogenesis of epithelial disruption in the context of gut inflammation is not fully understood. To identify alterations in signaling pathways in intestinal epithelium upon inflammation, we used a heuristic approach and performed gene expression arrays in purified IECs from Crohn’s disease patients (CD patients) (array data accessible through NCBI’s Gene Expression Omnibus [GEO GSE72780]). Clinical results suggested that gene expression analysis of IECs from inflamed versus uninflamed gut areas of CD patients showed samples within each group clustered, and we were able to identify 129 genes differing significantly (fold-change ≥ 1.5; \( P \leq 0.05 \)) between both groups (Figure 1A). Interestingly, gene ontology analyses indicated inflammation-associated down-regulation of pathways involved in epithelial cell dynamics and IEC extrusion, such as Rho-A signaling, epithelial adherens junctions, and regulation of actin-based motility by Rho (Figure 1B). Notably, pathways involving Rho GTPases other than Rho-A, such as Cdc42 signaling, showed no significant changes (Supplemental Figure 1A). Although our data implicated a relation between impaired Rho-A function in IECs and inflammation in human IBDs, neither protein nor mRNA expression levels of Rho-A were found to be reduced in IECs from inflamed gut areas of CD or ulcerative
ular, the small intestine of Rho-AΔIEC mice could be characterized by crypt abnormalities, villous blunting, reduced number of goblet cells (Figure 3, C and D, and Supplemental Figure 2A), marked infiltration of CD4+ T cells and neutrophils (Figure 3E and Supplemental Figure 2B), and significantly elevated levels of Tnfa compared with Rhoa fl/fl littermates. Colon tissue turned out to be less affected, and the increased numbers of CD4+ T cells and increased mucosal TNF-α levels in the colon of Rho-AΔIEC mice did not reach statistical significance as compared with WT animals (Supplemental Figure 2, C and D). Interestingly, the observed intestinal phenotype of Rho-AΔIEC mice was not derived from a developmental problem during embryogenesis, since neonates from Rho-AΔIEC mice appeared to be healthy, showed no reduced body weight, and did not differ from Rhoafl/fl littermates with regard to intestinal tissue structure (Supplemental Figure 2, E–G). Our data therefore identified epithelial Rho-A as a key regulator in the maintenance of intestinal homeostasis and epithelial cell architecture. The altered epithelial structure in Rho-AΔIEC mice was associated with a significantly increased intestinal permeability (Figure 3G), suggesting that epithelial Rho-A deficiency leads to impairment of intestinal barrier function.

colitis patients (UC patients) compared with uninflamed tissue (Figure 2A and Supplemental Table 2). However, while — in IECs of control patients — Rho-A was detected almost exclusively at the plasma membrane, IECs in inflamed areas of gut were characterized by a prominent cytosolic accumulation of Rho-A (Figure 2B, Supplemental Figure 1, B and C, and Supplemental Table 3), suggesting a predominance of inactive Rho-A in IECs of IBD patients. Similarly, cytosolic Rho-A enrichment upon inflammation could also be confirmed in IECs from dextran sulfate sodium–exposed (DSS-exposed) mice (Figure 2C and Supplemental Figure 1D), suggesting a link between dysfunction of epithelial Rho-A and gut inflammation. To further address the functional consequences of impaired epithelial Rho-A, we generated mice with an IEC-specific deletion of Rhoa (Rho-AΔIEC mice) by crossbreeding mice carrying the LoxP-flanked Rhoa gene (21) with mice expressing Cre-recombinase under the control of the epithelium-specific villin promoter (22). Lack of Rho-A in IECs resulted in decreased body weight (Figure 3A) and spontaneous mucosal inflammation in the small intestine, as indicated by high resolution colonoscopy (Figure 3B). Evaluation of inflammatory tissue alteration by histologic scoring confirmed small intestinal pathology in Rho-AΔIEC mice. In particular, the small intestine of Rho-AΔIEC mice could be characterized by crypt abnormalities, villous blunting, reduced number of goblet cells (Figure 3, C and D, and Supplemental Figure 2A), marked infiltration of CD4+ T cells and neutrophils (Figure 3E and Supplemental Figure 2B), and significantly elevated levels of Tnfa (Figure 3F) compared with Rhoafl/fl littermates. Colon tissue turned out to be less affected, and the increased numbers of CD4+ T cells and increased mucosal TNF-α levels in the colon of Rho-AΔIEC mice did not reach statistical significance as compared with WT animals (Supplemental Figure 2, C and D). Interestingly, the observed intestinal phenotype of Rho-AΔIEC mice was not derived from a developmental problem during embryogenesis, since neonates from Rho-AΔIEC mice appeared to be healthy, showed no reduced body weight, and did not differ from Rhoafl/fl littermates with regard to intestinal tissue structure (Supplemental Figure 2, E–G). Our data therefore identified epithelial Rho-A as a key regulator in the maintenance of intestinal homeostasis and epithelial cell architecture. The altered epithelial structure in Rho-AΔIEC mice was associated with a significantly increased intestinal permeability (Figure 3G), suggesting that epithelial Rho-A deficiency leads to impairment of intestinal barrier function.

Figure 2. Rho-A profile in human and murine colitis. (A) Rho-A protein expression in IECs isolated from human gut; blots are representative of 2 experiments (top); and Rhoa mRNA expression (bottom) in IECs isolated from human gut. Mean values ± SEM are shown (controls, n = 4; IBD uninflamed, n = 9; CD, n = 5; UC, n = 4). No statistical significance. 1-way ANOVA with LSD multiple comparisons test. (B) Representative pictures and quantification from Rho-A immunostaining (red) in human gut samples. Sections were counterstained with EpCAM (green) and Hoechst (blue). Arrows indicate cytosolic accumulation of Rho-A. Bars show percentage of cells with cytosolic Rho-A (2 villi or crypts/sample; 30 IEC/sample). Mean values ± SEM (controls, n = 5; IBD uninflamed, n = 7; CD, n = 3; UC, n = 3). ††P ≤ 0.001 vs. control; *P ≤ 0.05 vs. IBD-uninflamed; 1-way ANOVA with LSD multiple comparisons test. Original magnification, ×63, zoom ×4. (C) Western blot analysis of Rho-A in cytosolic and total proteins from colonic IECs from DSS-exposed and unchallenged mice. Blots are representative of 2 experiments (n = 4/group). Con, control; un, uninflamed; infl, inflamed.
Deficient prenylation in the inflamed intestinal epithelium of CD and UC patients. Since impaired Rho-A signaling in IBDs could not be explained by decreased Rho-A expression (Figure 2A), we postulated an impaired activation of epithelial Rho-A under inflammatory conditions via prenylation, a posttranslational modification necessary for membrane affinity of this GTPase (23). Notably, expression analysis of the Rho-A–associated prenylation-catalyzing enzyme GGTase-I indicated significantly diminished protein and mRNA levels in IECs of CD and UC patients during mucosal inflammation (Figure 4, A and B, Supplemental Figure 3A, and Supplemental Tables 3 and 4). The association between intestinal inflammation and diminished GGTase-I expression could be further confirmed in the experimental model of DSS-induced colitis in mice. In this model, significantly decreased GGTase-I expression went along with accumulation of nonprenylated (Np) proteins in IECs upon inflammation, indicated by an increased detection of Np small GTPase Rap1A (Np-Rap1A) (Figure 4, C and D, and Supplemental Figure 3, B and C). Thus, deficient prenylation in IECs upon inflammation might be responsible for altered subcellular localization and signaling of Rho-A in IBDs.

To gain further insights into the signaling mechanisms controlling Ppget1b expression in gut epithelium, epithelial organoids were cultured in the presence of different proinflammatory cytokines. Use of gut organoid technique allowed us to perform our analyses in the absence of extraepithelial factors (24). Interestingly, exposure to the IBD hallmark cytokine IL-6 (25) resulted in a signifi-
differenced in Rhoa expression could not be observed when we analyzed total gut tissue (Supplemental Figure 4B), our data furthermore suggest that regulation of intestinal Rhoa expression specifically occurs within the epithelial compartment.

Deficient prenylation within IECs results in alteration of cell shedding and loss of gut homeostasis. To further characterize the impact of prenylation on Rho-A signaling and intestinal inflammation in vivo, we created a mouse line with inducible conditional Ppargt1b deficiency in IECs. Breeding of Ppargt1bfl/fl mice (26) with villin-Cre-ERT2 mice (22) (tamoxifen induces ERT2-dependent Cre-recombinase expression) led to the generation of Ppargt1bΔIEC mice, which developed normally and showed no signs of intestinal pathology before tamoxifen treatment (data not shown). Tamoxifen-induced specific deletion of Ppargt1b within IECs caused significantly decreased Ppargt1b expression in epithelial organoids (Figure 5A), suggesting that pathologically increased IL-6 signaling in the inflamed gut of IBD patients contributes to downregulation of Ppargt1b expression and subsequent impairment of Rho-A activity in IECs. By comparing IECs in colon and small intestine, we observed contrary expression patterns for Il6 and Ppargt1b with high Il6 (Supplemental Figure 4A) and relatively low Ppargt1b expression (Figure 5B) in the small intestine. Our data therefore implicate that regional as well as inflammation-induced alteration of prenylation levels in IECs might be augmented by IL-6. Besides relatively low levels of Ppargt1b expression, IECs in small intestine could also be characterized by a decreased expression of Rhoa mRNA compared with IECs in colon (Figure 5B), making the small intestine potentially more susceptible to Rhoa and Ppargt1b inhibition. As regional differences in Rhoa expression could not be observed when we analyzed total gut tissue (Supplemental Figure 4B), our data furthermore suggest that regulation of intestinal Rhoa expression specifically occurs within the epithelial compartment.

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**Figure 4. GGTase-Iβ profile in human and murine colitis.** (A) GGTase-Iβ protein expression in IECs isolated from human gut; blots are representative of 2 experiments. The same samples are shown in Figure 2A (top); Ppargt1b mRNA expression (bottom) in IECs isolated from human gut. (control, n = 3; IBD uninfamed, n = 7; CD, n = 3; UC, n = 4). *P ≤ 0.05 vs. control; **P ≤ 0.05 vs. IBD uninfamed. (B) Representative pictures and quantification of GGTase-Iβ immunostaining (red) in human gut samples. (control, n = 6; IBD uninfamed, n = 7; CD, n = 7; UC, n = 5). Bars show percentage of cells expressing GGTase-Iβ (2 villi or crypts/sample; 60 IECs/sample). **P ≤ 0.01 vs. control; *P ≤ 0.05 vs. IBD uninfamed. Original magnification, ×63. (C) GGTase-Iβ expression in IECs from DSS-exposed mice: mRNA expression, measured by qPCR (n = 8/group). †††P ≤ 0.0001 vs. control (top); representative blot out of 3 experiments (bottom). (D) GGTase-Iβ immunostaining (red) in colon from DSS-treated mice. Mean intensity quantification (n = 8/group). Original magnification, ×63. *P ≤ 0.05 vs. control. Immunostainings were counterstained with EpCAM (green) and Hoechst (blue) in B and D. Mean values ± SEM are shown in A–D. One-way ANOVA with LSD multiple comparisons test were used in A and B. Independent samples t test was used in C and D. Con, control; un, uninflamed; infl, inflamed.
a decrease of epithelial \( \text{Pggt1b} \) mRNA (Supplemental Figure 5, A and B) and GGTase-I protein expression and, subsequently, an enrichment of Np proteins (Figure 6, A and B, and Supplemental Figure 5C). Such deletion resulted in body weight loss (Figure 7A), while \( \text{Pggt1b}^{\text{fl/fl}} \) control mice remained unaffected. Macroscopic, endoscopic, and histological analyses identified chronic intestinal inflammation and injury in \( \text{Pggt-I}^{\beta \Delta \text{IEC}} \) mice (Figure 7, B–E). The colon showed signs of moderate inflammation that did not reach statistical significance, whereas significant marked destruction of the villus-crypt architecture and loss of epithelial integrity could be observed in the small intestine of \( \text{Pggt-I}^{\beta \Delta \text{IEC}} \) mice as compared with floxed control mice (Figure 7E). Immunofluorescence and biochemistry analyses of small intestine tissue demonstrated that inflammatory injury of small intestine in \( \text{Pggt-I}^{\beta \Delta \text{IEC}} \) mice went along with a massive infiltration of CD4+ T cells, macrophages, and neutrophils (Figure 7F) and significantly increased levels of proinflammatory cytokines, such as IL-1\( \beta \), IL-6, and TNF-\( \alpha \) (Figure 7G and Supplemental Figure 5D). Colon tissue of \( \text{Pggt-I}^{\beta \Delta \text{IEC}} \) mice showed only moderate alterations with regard to inflammatory cell infiltration and cytokine production (Supplemental Figure 5, E an F). To investigate whether deficient prenylation in IECs might compromise the gut barrier, we next subjected mice to FITC-dextran oral gavage. We noted a significantly higher increase of FITC serum levels in \( \text{Pggt-I}^{\beta \Delta \text{IEC}} \) mice as compared with control mice, suggesting a breakdown of intestinal barrier function (Figure 7H). Time course studies furthermore indicated that development of epithelial injury and impairment of gut barrier after \( \text{Pggt1b} \) deletion even preceded induction of proinflammatory cytokine production (Supplemental Figure 6). The loss of intestinal barrier function therefore appeared as a primary defect in \( \text{Pggt-I}^{\beta \Delta \text{IEC}} \) mice.

To investigate whether prenylation might be an essential process for epithelial cell survival, we next studied epithelial development in organoids (24) from \( \text{Pggt-I}^{\beta \Delta \text{IEC}} \) mice. \( \text{Pggt1b} \) deletion impaired the maintenance of complex epithelial organization and resulted in destruction and death of ex vivo cultured and tamoxifen-exposed organoids from \( \text{Pggt-I}^{\beta \Delta \text{IEC}} \) mice (Figure 8 and Supplemental Figure 7), suggesting that GGTase-I activity acts as an epithelial-intrinsic factor controlling cell survival. These data clearly demonstrated that gut epithelial organization and integrity directly depend on prenylation within IECs.

Dysregulation of apoptosis and necroptosis in IECs has been previously described to cause intestinal inflammation via loss of barrier function (4). To investigate whether excessive epithelial cell death might be a key driving factor of the severe phenotype occurring in \( \text{Pggt-I}^{\beta \Delta \text{IEC}} \) mice, we determined whether apoptosis or necroptosis might play an important role. However, neither the additional IEC-specific deletion of \( \text{Casp8} \), nor that of the necrop-

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**Figure 5. Regulation of GGTase-I\( \beta \) and Rho-A expression in intestinal epithelium.** (A) \( \text{Pggt1b} \) and \( \text{Rhoa} \) mRNA expression (measured by qPCR) in cytokine-treated intestinal organoids from unchallenged WT mice (20 ng/ml of the cytokine, 8 hours stimulation) (\( n = 3 \)/group). \( ^{\dagger} P \leq 0.05 \) vs. control; independent samples \( t \) test. (B) \( \text{Pggt1b} \) and \( \text{Rhoa} \) mRNA expression in IECs isolated from different gut segments from unchallenged WT mice (\( n = 4 \)/group). \( ^{\dagger} P \leq 0.05 \); 1-way ANOVA with LSD multiple comparisons test. Bars show mean ± SEM in A and B.

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**Figure 6. Validation of tamoxifen-induced \( \text{Pggt1b} \) deletion in IECs (\( \text{Pggt-I}^{\beta \text{IEC}} \) mice).** \( \text{Pggt-I}^{\beta \text{IEC}} \) mice were treated for 3 consecutive days with tamoxifen by i.p. injection. Day 0 was defined as the day of the first tamoxifen injection. (A) Western blot of GGTase-I\( \beta \) and Np-Rap1A in isolated IECs; blots are representative of 3 independent experiments. (B) Representative pictures of GGTase-I\( \beta \) (red) and Np-Rap1A (red) immunostainings in duodenum (\( n = 4 \)/group). Sections were counterstained with EpCAM (green) and Hoechst (blue). Original magnification, ×63.
tosis inducer Rip3k, rescued the lethal consequence of Pggt-IβTiΔIEC deletion (Supplemental Figure 8, A–D). Additionally, proliferation of IECs along the crypt axis was not reduced in the absence of epithelial GGTase-I (Supplemental Figure 8E), excluding the possibility that cell proliferation arrest is a key underlying mechanism for tissue injury in Pggt-IβTiΔIEC mice.

Our finding of severe barrier dysfunction in the absence of alterations in key homeostatic mechanisms finally led us to study epithelial cell shedding, a process crucial for the regulation of epithelial turnover in the steady state. Notably, using in vivo confocal microscopy of the gut (5), we observed marked alterations of the monolayer due to the occurrence of epithelial gaps in Pggt-IβTiΔIEC mice but not control mice (Figure 9A). Under physiological conditions, such gaps are formed by the shedding of epithelial cells and sealed by neighboring cells (3, 5, 27) to avoid breakdown of barrier function. To determine whether barrier function and cell shedding were altered in Pggt-IβTiΔIEC mice, as recently observed in IBD patients (8, 28, 29), we performed in vivo rhodamine leakage assays. These studies revealed an increased passage of topically applied dextran from the lumen to the lamina propria in Pggt-IβTiΔIEC mice.
Electron microscopic analysis of GGTase-I–deficient IECs revealed disruption of the epithelial architecture and altered cytoskeleton rearrangement (Figure 10A). Consistently, localization of actin fibers and myosin IIA within IECs was no longer restricted to the apical cell membrane in the absence of GGTase-I, but it spread along lateral membranes and also in the cytosol (Figure 10B and Supplemental Figure 10A). Cytoskeleton rearrangement was analyzed by kinome array of GGTase-I–deficient IECs.

as compared with control mice (Figure 9B and Suplemental Figure 9A). Villus tips in Pggt-IβΔIEC mice showed marked accumulation of leakage points and appearance of dextran-permeable IECs (Figure 9C), suggesting altered epithelial cell integrity. Furthermore, cell shedding was increased in Pggt-IβΔIEC mice (Supplemental Figure 9B), consistent with the barrier loss and suggesting shedding alterations as key processes mediating the phenotype of Pggt-IβΔIEC mice.

Figure 8. Development and survival of GGTase-Iβ–deficient organoids. Small intestinal crypts isolated from control and Pggt-IββΔIEC mice were treated with tamoxifen in vitro. (A) Representative microscopic pictures out of 3 independent experiments. Original magnification, ×10. (B and C) Cell death staining measured by propidium iodide incorporation (red) as described in Methods. Nuclei were counterstained with Hoechst (blue). Representative pictures; original magnification, ×20 (n = 17) (B). Quantification of normalized mean fluorescence intensity (red/blue). Bars show mean values ± SEM (n = 17) (C). †P ≤ 0.05 vs. control; independent sample t test.

Figure 9. Epithelial turnover and integrity within GGTase-Iβ–deficient epithelium. (A) Representative in vivo images of small intestinal villi stained with acriflavine (green) (3 independent experiments). Arrows indicate epithelial gaps. Original magnification, ×40. (B) Live imaging of cell shedding in the small intestine from control and Pggt-IββΔIEC mice; acriflavine (green), and luminal rhodamine-dextran (red). Representative pictures out of 2 experiments. Original magnification, ×20. (C) Representative pictures and quantification of leakage entry points (red arrow) and permeable IECs (white arrow). Original magnification, ×40. Mean ± SEM; n = 8/group. †P ≤ 0.05 and †††P ≤ 0.0001 vs. control mice; independent samples t test.
Phosphorylation of 19 out of 64 cytoskeleton-related proteins was modified, and many of these proteins are known to be involved in Rho-A regulation (Supplemental Figure 10B). Since cytoskeleton rearrangement and tight junction redistribution represent one of the early events occurring in cell shedding (27), we hypothesized that cell shedding was arrested in Pggt-βTmice. Hence, we quantified the number of IECs showing redistribution of actin fibers (funnel-like structures, arrested events) versus cells that have been already shed into the lumen (nucleus does not belong to the monolayer, completed events) (Supplemental Figure 10C). Indeed, the accumulation of arrested (early) versus completed (late) shedding events in GGTase-β-deficient small intestine (Figure 10C) indicated an impaired completion of cell extrusion. Accumulated half-shed cells might subsequently impair the normal cytoskeletal mechanisms to seal the resulting gap.

Mucosal inflammation in the absence of GGTase-I is driven by blockade of Rho-A signaling. In order to further analyze the interplay between deficient prenylation, impaired Rho-A signaling, and epithelial injury, we next performed a more profound characterization of molecular alterations in GGTase-β-deficient IECs. Quantitative proteomic and gene expression assays (array data accessible through NCBI’s GEO [GSE72781]) revealed that the steady-state levels of 322 proteins and the expression of 725 genes were regulated in IECs of Pggt-βTmice as compared with control mice (fold-change ≥ 2; P ≤ 0.05) (Figure 11, A and B). Gene ontology analysis from GGTase-I–deficient IECs implicated an association between GGTase-I prenylation and cell viability/cell death induction, cytoskeleton rearrangement, and cell morphology (Figure 11, C and D, and Supplemental Table 5), consistent with our mechanistic observations and previous findings on potential Rho targets (30). Interestingly, when we compared the gene expression profile in IECs from IBD patients and GGTase-I–deficient IECs, Rho-A signaling could be identified as one of the only 2 pathways significantly regulated in both conditions (Figure 11E). Beside Rho-A signaling, only the aryl hydrocarbon receptor (AHR) pathway (31) was regulated in both settings, but it seemed to be of minor functional relevance, since the expression of AHR repressor (AHRR) (32) turned out to be unchanged in IECs from Pggt-βTmice (Supplemental Figure 11). Furthermore, Rho-A- and GGTase-I-deficient IECs resembled each other in the accumulation of arrested cell shedding events (Figure 10, B and C, and Figure 12, A and B) and in their inability to form regular organoid structures (Figure 12C). Together, these data strongly suggested that impairment of Rho-A signaling represents the pivotal consequence of deficient GGTase-I activity in IECs, relevant for epithelial integrity and IBD pathogenesis.

Figure 10. Cytoskeleton rearrangement and cell shedding within GGTase-β-deficient epithelium. (A) Representative electron microscopic pictures from duodenum of Pggt-βTmice and control mice. Arrows indicate invaginations (red), apical network fibers (blue), and filamentous ultrastructure (yellow) (2 independent experiments). Original magnification, ×5,000; insets ×20,000. (B) Representative pictures from phallolidin staining of F-actin fibers (green) in colon and duodenum from Pggt-βTmice. Nuclei were counterstained with Hoechst. Arrows indicate redistribution of actin fibers. Original magnification, ×63. (C) Quantification of arrested vs. completed shedding events, expressed as percentage of total shedding events. Mean values ± SEM; n = 11 (colon); n = 7 (duodenum).
Furthermore pointed to a potential therapeutic applicability of Rho activators for treatment of epithelial-derived gut pathology.

Discussion

Epithelial injury represents an important but not fully understood aspect of IBD pathogenesis (33). While recent studies have shown that epithelial injury in IBDs can be initiated by mechanisms regulating cell death in IECs, such as apoptosis and necroptosis (34), we here identified a new mechanism of epithelial damage in IBDs induced by impaired Rho-A signaling with subsequent cytoskeletal alterations of IECs and altered cell shedding. Such abnormal cell shedding may cause intestinal erosions and ulcer formation in IBDs and favors impaired barrier function that may clinically precede the development of acute flares in IBD patients (8, 27–29).

In line with this suggestion, GGTase-Iβ-deficient IECs showed a substantial decrease of membrane-bound and an accumulation of cytosolic Rho-A (Figure 13, A and B, and Supplemental Figure 12A), while other small Rho GTPases did not show this profile (Supplemental Figure 12B). Reduced membrane-localization of Rho-A in IECs of Pggt-IβTiΔIEC mice was paralleled by inefficient GTPase activation, resulting in reduced levels of active GTP-bound Rho-A (Figure 13C), and decreased phosphorylation of the downstream Rho-A target protein myosin light-chain 2 (MLC2) (Figure 13D). Thus, we confirmed that Rho-A activation and signaling is markedly impaired in IECs in the absence of GGTase-I. On a functional level, we wondered whether activation of Rho-A signaling would rescue the intestinal pathology in Pggt-IβTiΔIEC mice and, thus, treated these mice with Rho activators. Analysis of Pggt-IβTiΔIEC mice exposed to the specific Rho activator CN03 indicated that successful triggering of Rho-A activity in IECs in vivo (Supplemental Figure 12C) resulted in a significant amelioration of mucosal damage in endscopy (Figure 14, A and B), a clear tendency toward decreased histological damage score ($P = 0.054$) (Figure 14, C and D), and significantly diminished expression of $Ilb$ and $Iil6$ in the duodenum (Figure 14E) as compared with untreated mice. Rho-A blockade via impaired prenylation thus emerged as a key driver of mucosal damage in Pggt-IβTiΔIEC mice. The observed in vivo effect of CN03 furthermore pointed to a potential therapeutic applicability of Rho activators for treatment of epithelial-derived gut pathology.

In the inflamed gut, epithelial-extrinsic mediators, such as proinflammatory cytokines derived from activated lamina propria immune cells (TNF-α, IL-6, and IL-13) (35–37), are potent inducers of IEC apoptosis and pathological cell shedding with subsequent cytoskeletal alterations of IECs and altered cell shedding. Such abnormal cell shedding may cause intestinal erosions and ulcer formation in IBDs and favors impaired barrier function that may clinically precede the development of acute flares in IBD patients (8, 27–29).

In the inflamed gut, epithelial-extrinsic mediators, such as proinflammatory cytokines derived from activated lamina propria immune cells (TNF-α, IL-6, and IL-13) (35–37), are potent inducers of IEC apoptosis and pathological cell shedding with subsequent impairment of epithelial barrier function (35, 37–40). However, an increased occurrence of abnormal intestinal permeability in healthy relatives from IBD patients (41), and the finding that loss of barrier function in IBDs often preceded the clinical relapse of disease (8, 28), strongly argue against the assumption that IBD-associated epithelial barrier defects rep-
noted in the large intestine. This could be due to the compensatory effects of other small GTPases in the large intestine. The identification of these GTPases will require further studies in the future.

Our findings indicate that epithelial integrity, intestinal architecture, and homeostasis critically depend on sufficient prenylation of Rho proteins within IECs. IEC-specific deletion of the enzyme GGTase-I resulted in intestinal disease characterized by accumulation of inactive cytosolic Rho-A, epithelial injury, and mucosal inflammation. Likewise, abrogation of Rho-A in IECs led to marked epithelial alterations and intestinal inflammation. Our findings suggested that impairment of GGTase-I–mediated prenylation in IECs causes accumulation of Np cytosolic Rho-A molecules that are functionally inactive. This subsequently results in impaired regulation of cytoskeleton rearrangement (42) with instability of cell shape and alteration of cell shedding. Accumulation of arrested cell shedding events within the epithelial monolayer finally resulted in loss of barrier function and pathologically increased intestinal permeability. Altered intestinal permeability may subsequently cause chronic intestinal inflammation. Indeed, time-course studies in inducible GGTase-I–deficient mice revealed that altered barrier function precedes mucosal inflammation, suggesting a primary barrier defect in such animals. The potential relevance of these findings for humans was underlined by the observation that significant downregulation of GGTase-I levels and cytosolic Rho-A accumulation was seen in the gut epithelium of IBD patients with active disease. Thus, in addition to the role of Rho-A signaling in cytokine TNF-α. The phenotype of these mice was particularly prominent in the small intestine, while little inflammation was present a pure secondary phenomenon of chronic colitis. Instead, epithelial-intrinsic alterations may play a primary role in IBD pathogenesis, and their modulation might open a field for potential therapeutic or preventive interventions. In this context, the here described requirement for sufficient Rho-A prenylation in gut epithelium represents an interesting new aspect, which significantly improves our understanding of intracellular regulation in IECs. In particular, the observation that Pggt1b deletion in ex vivo organoids impaired the maintenance of complex epithelial organization, even in the absence of any immune cell–derived factor, strongly supported the independency of Rho-A signaling cascades from epithelial-extrinsic mediators. On the other hand, the observed regulatory impact of IL-6 on GGTase-1β expression clearly demonstrated the high complexity of the interplay between epithelial and epithelial-extrinsic mediators in pathogenesis of IBDs. Our observation that IECs in small intestine of unchallenged WT mice showed as significantly decreased expression of Rhoa mRNA compared with IECs in colon might implicate augmented susceptibility of the small intestine to Rho-A and GGTase-1β inhibition. It will therefore be important to investigate in future studies whether levels of epithelial Rho-A expression or activation along the gastrointestinal tract might indeed impact on regional differences in disease manifestations in IBD patients.

The functional relevance of Rho-A and GGTase-I–mediated Rho-A activation for functional integrity of the intestine was highlighted by studies in mice with conditional gene targeting in which deletion of Rhoa or Pggt1b in IECs caused spontaneous chronic intestinal inflammation with accumulation of granulocytes and CD4+ T cells and augmented production of the proinflammatory cytokine TNF-α. The phenotype of these mice was particularly prominent in the small intestine, while little inflammation was...
tion of the Rho-associated kinase (ROCK), as well as deletion of the Rho-A target MLC kinase (MLCK), has been shown to result in incomplete cell extrusion (27). Microscopically, ROCK and MLCK inhibition resulted in a predominant appearance of funnel-like structures (arrested shedding events) built up by reorganized actin filaments and tight junction proteins along lateral membranes of shedding cells. However, the final consequence of the accumulation of arrested shedding events has not been considered previously (27). Such accumulation of epithelial funnel-like structures could also be observed in the intestine of Pggt-IβΔIEC mice, indicating that redistribution of tight junction proteins takes place, but complete extrusion and resolution of shedding does not occur. Altered cell shedding in the absence of appropriate Rho-A function finally resulted in epithelial instability and loss of barrier function, leading to chronic intestinal inflammation. Hence, our observations confirmed that blockade of cell shedding due to cytoskeleton alterations, mediated by Rho-A among others, is an essential mechanism for disturbances of epithelial integrity.

The newly gained insights into Rho-A–dependent IEC intrinsic mechanisms involved in the initiation of intestinal barrier dysfunction open new avenues for potential interventional or translational approaches. Therapeutically triggered induction of Rho-A signaling via specific Rho activators or induction of Rho downstream signaling molecules such as ROCK might therefore represent a novel strategy for maintenance of intestinal integrity and prevention or healing of mucosal damage in IBDs and other inflammatory disorders of the intestine.

**Methods**

**Animal models.** Mice carrying LoxP-flanked Pggt1b (Pggt1bΔ/Δ) (26), caspase 8 (Casp8Δ/Δ) (4), Rhoa alleles (RhoaΔ/Δ) (21), as well as Rip3k KO mice (46), were described previously. Pggt1bΔ/Δ mice were provided by Martin O. Bergö (Sahlgrenska Cancer Center, Institute of Medicine, University of Gothenburg, Gothenburg, Sweden). RhoaΔ/Δ mice were provided by Cord Brakebusch (Biotech Research and Innovation Centre, University of Copenhagen, Copenhagen, Denmark). To generate specific deletion of Pggt1b or Rhoa genes in IECs, Pggt1bΔ/Δ or RhoaΔ/Δ mice were crossbred with villin-Cre or villin-Cre-ERT2 mice (22). Pggt1bΔ/Δ mice were crossbred with Casp8Δ/Δ or Rip3kΔ/Δ in order to impair apoptosis or necroptosis, respectively. Ethanolic tamoxifen (Sigma-Aldrich) solution was emulsified in sunflower oil and administered for 3 consecutive days by i.p. or intrarectal routes (1 mg/mouse/day). Littermates carrying loxP-flanked target genes but not the Cre-recombinase were used as control. Genotyping was performed by PCR in ear or tail genomic DNA. Animal studies were conducted in a sex- and age-matched manner using littermates for each experiment. Both male and female animals were used at the age of 6–10 weeks. All mice were kept under specific pathogen–free conditions. Except for DSS-induced colitis, animals from different experimental groups were cohoused in every experiment. Mice were routinely screened for pathogens according to Federation for Laboratory Animal Science Associations (FELASA) guidelines.

DSS-induced colitis. Colitis was induced in C57BL/6J mice by daily supplementation of drinking water with 2% w/v dextran sulfate sodium (MP Biomedicals) for 7 days (47, 48); DSS was withdrawn on day 7, when animals received normal drinking water. Animals were sacrificed 10 days after starting of the DSS treatment.

Mouse endoscopy. Gut status of mice was monitored by high-resolution mouse video endoscopy as previously described (49). Mice were anesthetized with 2%–2.5% isoflurane in oxygen during endoscopy. Any colitis was scored basing on the following 5 parameters: thickening of the colon wall, changes in the normal vascular pattern, presence of fibrin, mucosal granularity, and stool consistency. Endoscopic scoring of each parameter was performed (0 to 3), leading to a cumulative score between 0 (no signs of inflammation) and 15 (very severe inflammation) (50).
oral gavage (4,000 g/mol average molecular weight of FITC-Dextran). Serum was collected 4 hours after gavage, and FITC concentration was determined spectrophotometrically using FITC-Dextran as standard.

In vivo cell shedding. In short (5, 27), mice were anesthetized by ketamin/xylazin i.p. injection, and the intestine was exteriorized and opened. The mucosa was topically stained with acriflavine (Sigma-Aldrich) and rhodamine-dextran (10,000 g/mol) as luminal dye (Invitrogen). Surgical preparation was pinned up on a coverslide and mounted in a chamber for perfusion of saline solution. Time-sequential Z-stacks images of the villus tip were taken with a confocal microscope (Leica Microsystems).

Quantification of epithelial cell shedding. Epithelial cell shedding was quantified by several techniques. On one side, the cell shedding rate was quantified by the counting of cell shedding events detected by

Figure 14. Treatment of Pggt-IpβTiΔIEC mice with the Rho activator CN03. Three independent experiments. (A) Representative colonic endoscopy pictures. (B) Endoscopy score. (C) Histologic analysis: representative pictures from H&E staining. Original magnification, ×20. (D) Damage score quantification. (E) Iltb and Il6 expression in duodenum measured by qPCR. Mean values ± SEM from 3 independent experiments; (control, n = 4; Pggt-IpβTiΔIEC, n = 6; CN03, n = 6 per group) in B, D, and E. †P ≤ 0.05 vs. control mice; *P ≤ 0.05 vs. Pggt-IpβTiΔIEC mice; 1-way ANOVA with LSD multiple comparisons test in B, D, and E.
in vivo cell imaging. The number of cell shedding events was normalized by the length of the basal membrane on the individual pictures and the time of image acquisition. A total number of 10 animals were analyzed. On the other hand, we have also counted the number of leakage and “permeable cells” (our terminology for cells with dextran in the cytoplasm) in 2 single in vivo images from each specimen (n = 8 animals). Finally, a similar method was performed in order to quantify the number of arrested versus completed cell shedding events. However, in this case, we took advantage of actin staining with phalloidin, which allowed us to detect the cytoskeleton protein redistribution occurring in early stages of the cell shedding process. We thus quantified the overall cell shedding rate, the distribution of different stages of cell shedding, and the appearance of dextran-permeable cells.

IEC isolation. Murine gut tissues and human surgical specimens were freshly used for IEC isolation. Therefore, tissue was incubated in a solution containing DTT and EDTA (34). Purification of IECs was performed by Percoll gradient.

Crypt isolation and organoid culture. Small intestine was aseptically isolated, washed, and cut. Intestinal fragments were washed, incubated in chelation buffer (containing EDTA), and vigorously resuspended in order to isolate intestinal crypts. Crypts were embedded in Matrigel (BD Biosciences) on ice and seeded in surface cell culture plates. After polymerization of Matrigel, basal culture medium was added to the culture (24). The entire medium was changed every 2 days, and organoids were passaged weekly. Organoids were treated in vitro with 0.5 µg/ml tamoxifen; the caspase inhibitor Z-VAD (10 µg/ml) (Bachem) or Necrostatin-1 (30 µM) (Sigma-Aldrich), and cytokines (20 ng/ml).

Quantification of cell viability and death in crypt organoid cultures. We used a cell viability imaging kit (Roche Diagnostics) in order to quantify the death/destruction of organoids after in vitro treatment of organoids with tamoxifen and subsequent deletion of Pgt1b gene. This Kit includes 3 different dyes in order to identify nuclei (blue dye), viable cells (green dye), and dead cells (red dye). Identification of dead cells is based on permeability of those cells due to destruction of the cell membrane. Detection of cell viability is based on metabolic activity, which allows the cleavage of the substrate, resulting in green fluorescence. Nuclear staining (blue) enabled normalization of results based on the overall number of detected cells. Quantification of the mean intensity of dead cell signal (red signal) and nuclear signal (blue signal) was performed using ImageJ software.

Histology, IHC, and electron microscopy. Formalin-fixed paraffin-embedded samples were used for H&E staining and histopathological analysis. The score of the histological damage in colon, ileum, and duodenum took into account the 2 parameters epithelial erosion and distorted tissue architecture. Each parameter was scored (none, 0; mild, 1; pronounced, 2; severe, 3), and both results were summed afterward (maximum score of 6). Scoring was performed in a blinded fashion. H&E was performed in cryosections, using TSA Cy3 (PerkinElmer). Images were acquired by fluorescence microscopy (Olympus). IHC was performed using the online tool Database for Annotation, Visualization, and Integrated Discovery (DAVID) and the platform Ingenuity.

Gene expression assay (human; array data accessible through NCBI’s GEO [GSE72780]). More than 30 samples were collected from the gut of patients suffering from CD or control patients. IECs were isolated from patient specimen as described. RNA was isolated using the RNeasy Lipid Tissue Mini kit according to manufacturer’s instructions (QIAGEN). Nine samples were chosen for GeneChip analysis according to the following inclusion criteria: histopathological confirmation of inflammation status, acceptable RNA quality as confirmed by a bioanalyzer, and comparable expression of the epithelial marker villin.

GeneChip analysis. Genome-wide expression analysis was performed in the Genomics core facility of the Institute of Human Genetics of the University Erlangen-Nuremberg using the Affymetrix GeneChip Human Genome U133 Pls 2.0 Arrays. For multiple gene expression analysis, we used a cell viability imaging kit (Roche Diagnostics) in order to quantify the death/destruction of organoids after in vitro treatment of organoids with tamoxifen and subsequent deletion of Pgt1b gene. This Kit includes 3 different dyes in order to identify nuclei (blue dye), viable cells (green dye), and dead cells (red dye). Identification of dead cells is based on permeability of those cells due to destruction of the cell membrane. Detection of cell viability is based on metabolic activity, which allows the cleavage of the substrate, resulting in green fluorescence. Nuclear staining (blue) enabled normalization of results based on the overall number of detected cells. Quantification of the mean intensity of dead cell signal (red signal) and nuclear signal (blue signal) was performed using ImageJ software.

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Gene expression assay (mouse; array data accessible through NCBI’s GEO [GSE72781]). Gene-chip experiments were performed by the Erlangen University core facility (n = 3 per group). RNA quality was first assessed by Eukaryotic Total RNA Nano (Agilent Technologies), and gene expression was measured using the Affymetrix mouse 430 2.0 chip. For multiple gene array testing, including differential expression analysis, the software package FlexArray (Michal Blazejczyk, Mathieu Miron, and Robert Nadon, 2007, Genome Quebec, Montreal, Canada; http://www.qginnovationcenter.com/services/bioinformatics/index.aspx?l=e) was used. GO-based analyses were performed using the online tool Database for Annotation, Visualization, and Integrated Discovery (DAVID) and the platform Ingenuity.

Gene expression assay (human; array data accessible through NCBI’s GEO [GSE72780]). More than 30 samples were collected from the gut of patients suffering from CD or control patients. IECs were isolated from patient specimen as described. RNA was isolated using the RNeasy Lipid Tissue Mini kit according to manufacturer’s instructions (QIAGEN). Nine samples were chosen for GeneChip analysis according to the following inclusion criteria: histopathological confirmation of inflammation status, acceptable RNA quality as confirmed by a bioanalyzer, and comparable expression of the epithelial marker villin.

GeneChip analysis. Genome-wide expression analysis was performed in the Genomics core facility of the Institute of Human Genetics of the University Erlangen-Nuremberg using the Affymetrix GeneChip Human Genome U133 Pls 2.0 Arrays. For multiple gene
array evaluations, including advanced significance analysis, clustering, and gene ontology-based gene class testings, ANOVA of the software package Partek Genomics Suite were used. In brief, GC-RMA algorithm was applied to CEL-files of all microarrays. Next, log scale transformation, grouping, and baseline definition were performed. For advanced significance analysis, the following parameters were applied: unpaired t test, \( P < 0.05 \), fold-change > 1.5, K-Means Clustering. Redundancies in gene list (e.g., cluster probe test lists) were eliminated, and gene ontology enrichment analysis was performed, allowing the detection of statistically overrepresented protein functions (GO terms) in a subset of genes. More than 54,000 expression tags were analyzed, from which 129 significantly (\( P < 0.05 \)) differed (fold-change > 1.5) between IECs from inflamed and uninflamed gut areas of CD patients.

**Immunoblotting.** Protein extracts were obtained by suspension and incubation in mammalian protein extraction reagent (Thermo Scientific) supplemented with protease and phosphatase inhibitor tablets (Roche Diagnostics). For membrane/cytosol fraction separation, lysis buffer (RIPA) with and without detergents and high-speed centrifugation steps were used. Protein extract was cleared by centrifugation, protein concentration was measured by Bradford assays, and denaturation was performed by boiling in LDS sample buffer (Thermo Scientific). Afterward, proteins were separated by SDS-PAGE gels, blotted onto nitrocellulose or polyvinylidene fluoride (PVDF) membranes, and blocked with 5% nonfat milk. Membranes were incubated overnight at 4°C with the following primary antibodies: GGTase-I, 1:2,500 (catalog WH000529M2, Sigma-Aldrich); Np-Rap1A, 1:1,000 (catalog sc-1482, Santa Cruz Biotechnology Inc.); Rho-A, 1:500 (catalog sc-179, Santa Cruz Biotechnology Inc.); phospho-MLC2, 1:1,000 (catalog 3671, Cell Signaling Technology); Cdc42, 1:1,000 (catalog 2466, Cell Signaling Technology); Rac-1, 1:1,000 (catalog sc-95, Santa Cruz Biotechnology Inc.); Rho-B, 1:1,000 (catalog 2098, Cell Signaling Technology); Rho-C, 1:1,000 (catalog 3430, Cell Signaling Technology); and AHRR, 1:1,000 (catalog sc-138745, Santa Cruz Biotechnology Inc.). Convenient HRP-linked secondary antibodies were used (incubation for 90 minutes at RT), and chemiluminescence was detected by using ECL Western blotting substrate (Thermo Scientific). Protein expression was compared with the level of actin, 1:2,000 (catalog 4967, Cell Signaling Technology). Inclusion of a positive control sample (protein lysate from testis) allowed reliable identification of the 38–43 kDa protein GGTase-I. **Proteomic assay.** Of the total protein from purified IECs, 20 µg were digested using a modified filter aided sample preparation (FASP) method (52). Three biological replicates were included, except where indicated. Reversed-phase nano-UPLC separation of tryptic peptides and mass spectrometric analysis using an ion-mobility enhanced data-independent analysis workflow were performed as previously described (52). All samples were analyzed in 3 technical replicates. **Cytoskeleton protein activation array.** Activation of 64 cytoskeleton-related proteins was assessed by measuring their phosphorylation status using Cytoskeleton II Phospho Antibody Array (catalog PCPI41, Cytoskeleton Inc.), following manufacturer’s instructions.

**Rho-A activation assay.** Rho-A activation status was measured by using Rho-A G-LISA Activation Assay (catalog BK214, Cytoskeleton Inc.), following manufacturer’s instructions.

**Rho activator treatment.** Mice were randomly distributed into 2 groups (Pggt-Iβflx/lacZ ± CN03) and Pggt-Iβ littermates were used as controls. Mice were treated with the Rho Activator II (catalog CN03, Cytoskeleton Inc.) before onset of intestinal disease (oral, 0.3 µg/mouse; intrarectal, 0.5 µg/mouse). Tamoxifen was administered intrarectally. Animals were sacrificed 10 days after starting tamoxifen induction.

**Statistics.** Significance analysis of normally distributed data was performed using 2-tailed Student’s t test. Error bars represent ± SEM. For multiple comparisons, data were analyzed by 1-way ANOVA and LSD post-hoc test (IBM SPSS Statistic 21). \( P \) values of less than 0.05 and 0.001 were considered significant and highly significant.

**Study approval.** Animal protocols were approved by the Institutional Animal Care and Use Committee of the University of Erlangen. Animal studies were performed in accordance with German law and with approval from Regierung Mittelfranken (Ansbach, Germany). Collection of human samples was approved by Ethics Committee of the Department of Medicine of the University of Erlangen-Nuremberg (Erlangen, Germany) and Humanitas Research Hospital. Each patient gave written informed consent.

**Author contributions**

RLP, C Becker, IA, and MFN designed the research. RLP and VT performed the majority of the experiments. CG performed gene expression analysis in human IECs. ST performed quantitative proteomic analysis. KA performed electron microscopy. UB, RA, GF, SV, and SD collected human material for the study. AJMW was involved in performance of live animal imaging. SW was involved in discussion and design of experiments. C Brakebusch and MB provided us with genetically modified mouse strains. RLP, IA, MFN, AJMW, and C Becker analyzed the data and wrote the paper. ABE performed Gene-chip experiments. All the authors discussed the results and commented on the manuscript.

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