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An intercrypt subpopulation of goblet cells is essential for colonic mucus barrier function

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Abstract

Introduction—An intricate balance with our intestinal microbes is pivotal to health. A key interface of host-microbial interactions occurs in the mucus covering the intestinal epithelial surface. In the colon, the mucus layer serves as a barrier inhibiting direct epithelial contact with the dense population of microbes. Defects in this system are a hallmark of colitis. The mucus layer is structurally dependent on the polymeric mucin MUC2, and it is synthesized by goblet cells (GCs), specialized secretory cells classically viewed as a homogenous cell type. Studies identifying divergent functional features in GC subpopulations, including differential mucus biosynthesis rate and responses to bacteria, suggests GC populations may in fact be heterogeneous.

Rationale—In the present study, we characterized intestinal GC expression diversity, and defined how a specific GC subtype, localized in the intercrypt surface epithelium, functionally contributes to the formation of the mucus barrier.

Results—Using mCherry-MUC2 transgenic mice to sort and isolate GCs, we generated transcriptomic and proteomic profiles to characterize the GC expression landscape in both the small intestine and colon. Single cell transcriptomic analysis revealed several unique GC clusters in each tissue regions, which segregated into two separate trajectories. One trajectory had enriched expression of known GC-specific genes (e.g., *Clca1*, *Fcgbp*), and was designated as canonical GCs. Conversely, the other trajectory was enriched for expression of genes typically associated with enterocytes (e.g., *Dmbt1*, *Gsdmc4*), which we designated as non-canonical GCs.

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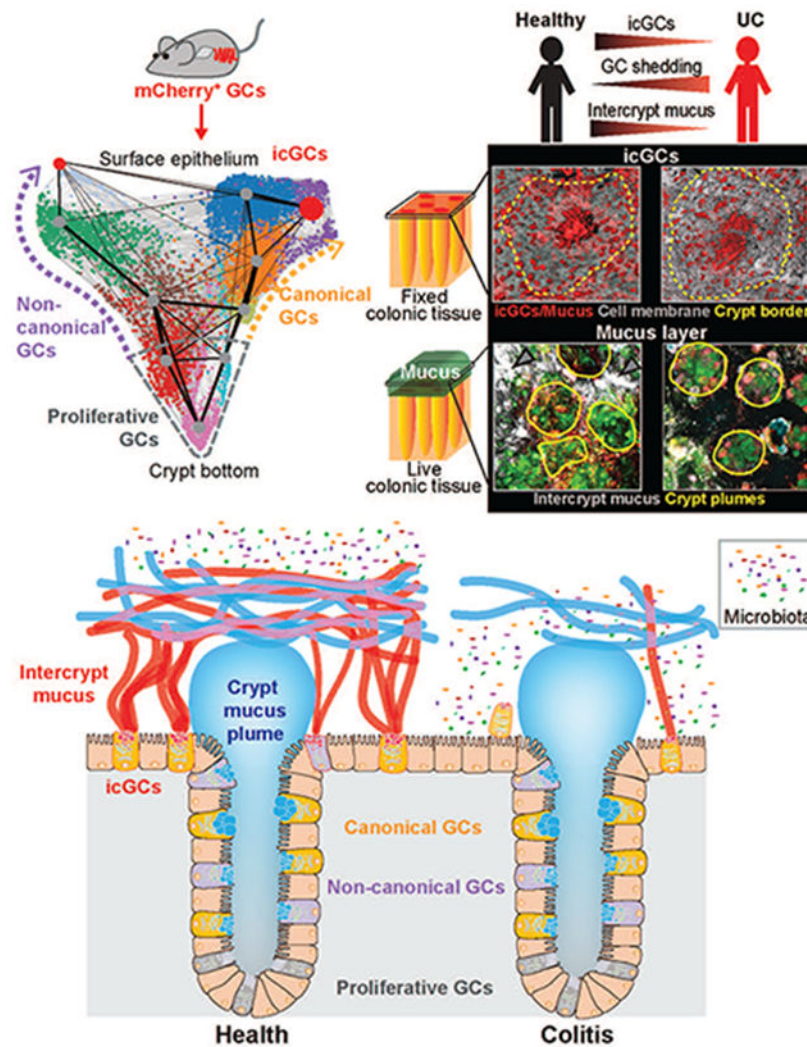
In the colon, the most differentiated GCs are the high mucus turnover cells localized to the surface epithelium between crypts and were thus designated as intercrypt GCs (icGCs). These cells had distinct expression profiles from crypt-resident GCs, thus we investigated their role in forming the mucus barrier. We exploited the lectin binding features of mucus to resolve the three-dimensional organization of mucus in live tissue explants. Results demonstrated that icGCs secreted distinct mucus that filled the spatial regions between mucus plumes secreted from crypt openings. The intercrypt mucus was impenetrable to bacteria-sized beads; however, it was more penetrable to smaller molecules compared to crypt plume mucus. Penetrable surface mucus may be important for absorption of ions, and other compounds, while denser mucus within the crypt compartment contributes to the shielding of the stem cell niche.

Both mucus subtypes appear to be important for the overall protective function of mucus, as a barrier impenetrable to bacteria was formed by the mixed net-like organization of intercrypt and crypt plume mucus. A mouse model with dysfunctional icGCs lacking normal intercrypt mucus exhibited an inadequate mucus barrier and was more susceptible to both chemically-induced and spontaneous (age-dependent) colitis, thus arguing for the indispensable role of icGCs in maintaining a functional mucus barrier. Furthermore, biopsies from patients with ulcerative colitis, also in remission, exhibited increased GC shedding and reduced icGC numbers. These characteristics was associated with structural defects in the mucus barrier, including gaps in the intercrypt mucus that exposed areas of the surface epithelium.

Conclusion—The current study identifies GCs as a heterogeneous population of cells with diverse functional features that indicate a dynamic cellular system. GCs at different locations along the crypt-surface axis contribute to a functional mucus barrier that protects the epithelium from microorganisms. The icGCs possess a unique role in mucus organization, where their malfunction is associated with colitis in both mice and humans.

Diversity of GC expression and function

Intestinal GCs can be divided into several distinct clusters that form canonical and non-canonical trajectories, and gene expression profiles outline their topographical position. The specialized surface GC type, the icGCs, secrete mucus crucial for forming a protective barrier between bacteria and the epithelium. Depletion of icGCs impairs mucus function, a phenomenon observed in ulcerative colitis (UC) patients.



Abstract

The intestinal mucus layer, a vital element of epithelial protection, is produced by goblet cells. Intestinal goblet cells are assumed to be a homogenous cell type. Herein, we delineated their specific gene and protein expression profiles and identified several distinct goblet cell populations forming two differentiation trajectories. One distinct subtype, the intercrypt goblet cells (icGCs), located at the colonic luminal surface, produced mucus with properties that differed from mucus secreted by crypt residing goblet cells. Mice with defective icGCs had increased sensitivity to chemically induced colitis and manifested spontaneous colitis with age. Furthermore, alterations in mucus and reduced number of icGCs were observed in patients with both active and remissive ulcerative colitis, highlighting the importance of icGCs in maintaining functional protection of the epithelium.

Keywords

Single cell RNA sequencing; Spdef; Muc2; mucin; epithelium

Introduction

Mucus secreted from the intestinal epithelium forms a continuous layer throughout the intestinal tract (1, 2). A primary function of mucus is to protect the epithelium against luminal microbes. In the distal colon, this is achieved by an inner mucus layer on the tissue surface, which creates a physical barrier between the bacteria and the epithelium (3-5). The inner mucus layer is modified to form an expanded outer mucus that is more accessible to bacteria (6); mucus along the whole intestine facilitates fecal transport (6). In the small intestine, a single layer of permeable mucus fortified by antimicrobial peptides protects the epithelium (1, 2).

The primary structural component of the mucus, MUC2, is a large multimeric and heavily *O*-glycosylated protein. The exposed terminal glycans form an outer surface on MUC2 that can be recognized by bacteria and lectins (7). Upon secretion, oligomers of MUC2 and other mucus components form a sieve-like structure, allowing passage of small molecules whilst hindering translocation of microorganisms. To date, a comprehensive understanding of mucus organization remains to be elucidated.

The vital importance of a functional mucus barrier is demonstrated by mice lacking *Muc2*, which develop spontaneous colitis (8). Furthermore, several observations have correlated mucus defects with the pathogenesis of ulcerative colitis (UC) and with experimental mouse models of colitis (9-11). Although mucus disruption is observed prior to intestinal inflammation and colitis induced by dextran sodium sulfate (DSS) in mice (12), it remains uncertain whether mucus dysfunction in humans is a cause or consequence of colitis-associated inflammation.

Mucus constituents, including MUC2, are secreted by goblet cells (GCs), and their high expression and complex structure put a high demand on the cellular biosynthesis machinery. GCs express specific proteins involved in these processes as *AGR2* and *ERN2* (13, 14) (all gene and protein names are listed in Table S1). As with all intestinal epithelial cells, GCs are derived from crypt-residing intestinal stem cells. GCs differentiation is promoted by several transcription factors, including *ATOH1* and *SPDEF*, whereas they are negatively regulated by *HES1* (15-18). Global single cell analysis of the intestinal epithelium has often classified GCs as a single population (19-21). However, a subpopulation of GCs only detected in active UC has recently been described in humans (22), suggesting dynamic differentiation of GCs during inflammation.

In the present study, we characterized GC transcript and protein profiles in both small and large intestine. Single cell analysis identified several clusters, forming two distinct differentiation trajectories. GCs in the intercrypt region of the colonic surface epithelium, intercrypt GC (icGCs), possessed a unique transcript profile and produced distinct, more penetrable mucus that filled the spatial regions between mucus plumes secreted from crypt-resident GCs. *Spdef*^{-/-} mice showed secretory defects in the icGC population and altered mucus architecture. These changes contribute to increased susceptibility to DSS-induced colitis, as well as increased incidence of spontaneous colitis. In a majority of UC patients, both in remission (UCr) and with active disease (UCa), a reduced number of icGCs

and an altered mucus organization was observed. Thus, intestinal GCs are remarkably heterogeneous, with colonic icGCs constituting an essential subpopulation for normal formation and function of the mucus barrier.

Goblet cell specific transcript and protein expression

We isolated mCherry-MUC2 positive cells by FACS from the distal colon and the most distal small intestine of transgenic RedMUC2^{98trTg} mice (23) (Fig. S1, A and B). GCs comprised 17% and 5% of viable epithelial cells in the colon and small intestine, respectively (Fig. S1C). Bulk RNA sequencing (RNA-seq) and proteomics by mass spectrometry were performed comparing GCs to the non-GC pool (Table S2 and S3). Principal Component Analysis (PCA) grouped samples based both on cell type and origin (Fig. S1D). Protein and mRNA showed good overlap and Pearson's correlation coefficient comparing mRNA and protein was 0.57 in the colon and 0.63 in the small intestine (Fig. S1, E and F). Comparison of GC and non-GC populations identified approximately 500 proteins and 800 transcripts significantly enriched in GCs (Fig. 1, A and B). Many of the GC enriched genes were shared between the small intestine and colon, but some showed site-specificity (Fig. 1C). Several known mucus components such as Muc2 (endogenous), Clca1, Fcgbp and Zg16 were among the 50 most abundant GC enriched genes and proteins. Gene Ontology (GO) enrichment analysis primarily classified the GC expression profile into secretory related processes but also identified lipid metabolism, immune response, proteolysis, and protein glycosylation (Figs. 1D and S2).

Goblet cells comprise several different clusters

To study the intrinsic variability of GCs under basal conditions, we performed single cell RNA-seq (scRNA-seq) on sorted GCs from RedMUC2^{98trTg} mice. Colonic GCs grouped into eight different clusters (Fig. 2, A and B, and Table S4). Clusters 6 and 4 were enriched for the proliferation marker *Mki67*, whereas clusters 7 and 8 were enriched for *Mxd1*, a gene expressed by fully differentiated cells (24). *Mxd1*⁺ cells were further separated by more specific markers, *Sln4* and *Aqp8*. Furthermore, GCs showed two main gene expression profiles; Clusters 1, 3, 5, and 7 exhibited a higher expression of typical GC-associated genes such as *Clca1* and *Fcgbp* (1, 25), whereas clusters 2 and 8 were enriched in expression of *Gsdmc4*, *Dmbt1* or *Aqp8*; all more abundant in the non-GC population according to bulk RNA-seq (Fig. 2, A and C). PCA analysis showed a higher diversity within the clusters associated with non-GC genes and with proliferation (Fig. 2B). A cluster tree revealed four main groups of GCs: typical, proliferative (*Mki67*⁺), non-typical, and fully differentiated (*Mxd1*⁺) (Fig. 2D).

Small intestinal GCs grouped into nine clusters (Fig. S3, A and B, and Table S4) when Muc2 expressing Paneth cells were removed (26) (Fig. S4). Proliferative genes were abundant in clusters 2 and 9 (*Mki67*⁺). Fully differentiated GCs (*Mxd1*⁺) expressed *Ido1* and were further divided into clusters 5 and 8. Typical GC-associated genes (*Clca1* and *Fcgbp*) were most highly expressed in clusters 1, 3, 5 and 8, while cluster 4 showed a more pronounced non-GC profile (*Sis*, *Dmbt1*, *Gsdmc4*) and was more diverse in the PCA (Fig.

S3, A to D). Expression of typical Paneth cell products such as *Defa24* and *Lyz1* was also noted in several clusters, but highest in cluster 3 and 6 (Fig. S3, A and C).

Maturation trajectories of intestinal goblet cells

To investigate whether GC populations follow distinct trajectories throughout their differentiation and maturation, we performed a pseudotime analysis (Table S5). Colonic GCs revealed two trajectories originating from proliferative cells (*Mki67*⁺) at the lower region of the crypt. Typical GC markers (*Atoh1*, *Fcgbp* and *Cla1*) were enriched in one trajectory designated as canonical GCs. Expression of these genes were found in *Muc2*-expressing cells along the crypt (Fig. 3, A to C). A second trajectory comprised cells with a non-GC profile and was designated non-canonical GCs as markers (*Hes1*, *Gsdmc4* and *Dmbt1*) for this trajectory were more abundant in *Muc2*-negative cells (Fig. 3, A to C). The terminals of the canonical and non-canonical GC trajectories ended in two small populations of *Mxd1*⁺ GCs highly expressing *Sfn4* or *Aqp8* that were localized at the surface epithelium (Fig. 3, A to C). Similar results were obtained analyzing individual datasets, and the same GC expression pattern of non-canonical GC markers, as seen in RedMUC2^{98trTg} mice, was also observed in wild type (WT) mice (Fig. S5, A and B). These trajectories were further supported by expression of different transcription factors. Specifically, *Atoh1* was found in most GCs, especially in the canonical GCs, whereas *Hes1* was expressed in some GCs along the crypt following the non-canonical trajectory (Fig. 3, B and C).

We next compared functions of genes associated with canonical and non-canonical GC trajectories in the colon (Table S6). Gene ontology (GO) enrichment analysis of differentiated colonic GCs (*Mki67*⁺) revealed that processes related to secretion, glycosylation, and endoplasmic reticulum stress were enriched in the canonical GCs, whereas non-canonical GCs demonstrated enrichment of genes associated with lipid and amino acid metabolism, detoxification, and intestinal absorption (Fig. 3D). Expression levels of *Ii18* and *Nlrp6*, genes associated with sentinel GCs (senGCs) (23), were higher in the non-canonical population. *Wfdc2* and *Areg*, genes whose expression is altered with colitis (22, 27), were also enriched in the non-canonical GCs (Fig. 3E).

Both pooled and individual datasets from the distal small intestine reveal a trajectory starting from the proliferative pool of GCs that leads to a branching structure (Figs. S6, A and B, and S5C). Unlike colon, this trajectory was dominated by canonical GCs with a later bifurcation made up of non-canonical GCs. The most mature canonical GCs present at the tip of the villi, were marked by the expression of *Ido1*. *Defa24*⁺ cells were spread along the canonical trajectory and identified in several GCs of the crypt, whereas *Lyz1* was limited to fewer GCs (Fig. S6, B to D)

Intercrypt goblet cells

The two colonic clusters of fully differentiated GCs (cluster 7 and 8, *Mxd1*⁺) expressed many genes in common, but differed as cluster 8 expressed several genes shared with non-GC epithelial cells (Fig. 2, A and B). Markers for these GCs were only detected at the surface epithelium in the intercrypt area (Fig. 3C). Functional profiling of these cells

compared to non-proliferative crypt GCs revealed increased expression of genes involved in cellular response to stress and bacteria, likely related to their luminal facing location (Fig. 4A and B, S7A, and Table S6). Additionally, processes such as programmed cell death and negative regulation of differentiation were upregulated as a consequence of their late stage of differentiation. They were also enriched in protein transport-associated genes (Fig. 4, A and C). Several genes involved in protein folding and quality control (*Agr2*, *Pdia4-6*) were expressed at lower levels in these cells compared to crypt GCs (Fig. 4D).

The possibility to use lectin staining to differentiate GCs at different locations in the epithelium was determined. The wheat germ agglutinin (WGA) lectin (N-acetylglucosamine/sialic acid-binding) detected crypt-residing GCs in fixed tissue, whereas the *Ulex europaeus* agglutinin I (UEA1) (fucose-binding) readily detected GCs of the upper crypt and surface epithelium (Fig. 4E). Lectin specificity is dependent on the combined effect of the glycosyltransferases expressed and glycan presentation, differential lectin staining could thus not be fully explained by glycosyltransferase expression (Fig. S7B). UEA1⁺/WGA⁻ GCs were restricted to the intercrypt surface epithelium. Accordingly, these cells were designated as intercrypt GCs (icGCs). In addition to specific lectin reactivity, icGCs were also strongly stained by antiserum to the non-O-glycosylated Muc2 precursor found in the endoplasmic reticulum (Fig. 4E). However, they were barely detectable using AB/PAS or immunostaining of mature Muc2, which instead primarily labelled the theca in crypt-residing GCs (Fig. 4F). This difference was likely due to the rapid secretion and reduced storage of mature Muc2 in icGCs compared to crypt-residing GCs(28).

The icGCs, as defined by their staining, shape and location include all intercrypt surface GCs, but appear to be primarily comprised of the most differentiated canonical GCs (cluster 7, high *Slfn4* expression) in line with their high expression of mucus associated genes, such as *Muc2*, *C1cal* and *Fcgbp* (Fig. 4G). They show a functional profile with response to stress, cell differentiation, apoptosis and protein transport (Fig. S7C). For example, expression of the constitutive secretory regulator *Stxbp1* was high, while the regulated secretory equivalent *Stxbp2* was predominantly expressed in crypt GCs. Higher expressions of the endoplasmic reticulum stress related gene *Ern2* combined with altered expression of proteins involved in folding indicated functional alterations in protein biosynthesis (Fig. 4D), and a subset of icGCs also expressed the immunomodulatory gene *Ido1* (Fig. S7, D and E). Although icGCs mainly consist of canonical GCs, the contribution of non-canonical surface GCs (cluster 8) cannot be excluded. The transcription factor *Spdef*, thought to drive terminal GC differentiation, was expressed by most GCs, but was found to be more abundant in the canonical lineage clusters, including icGCs (Figs. 4H, and S7F).

Unlike in the colon, expression of the icGC marker *Slfn4* was not restricted to GCs in the distal small intestine, suggesting that cells identical to icGCs are absent in the distal small intestine. Nevertheless, the cluster of *Mxd1*⁺/*Ido1*⁺ GCs present at the villi could potentially represent a similar subset; however, additional research is required for confirmation (Fig. S8).

Colonic mucus organization and properties

Using fluorescent-conjugated UEA1 and WGA, either alone or in combination, to stain secreted mucus in flushed colonic tissue *ex vivo*, revealed two distinctive staining patterns: WGA stained plume-like structures positioned on top of crypt openings (crypt plumes), whereas UEA1 primarily stained mucus material filling the space between plumes (intercrypt mucus) (Figs 5, A and B, and S9, A to C, and Movie S1). Intensity histograms and low co-localization coefficients in tissue proximate mucus confirmed that UEA1 and WGA primarily stained different material (Figs. 5C, and S9, D and E). Isosurface mapping demonstrated that UEA1-positive intercrypt mucus emerged from icGCs, whereas WGA-positive material emerged from the crypt (Fig. 5D). The UEA1- and WGA-stained fractions intermixed on top of the plumes and merged to form a dense net-like arrangement at the mucus surface (Fig. 5B), as demonstrated by an increased co-localization coefficient in the mix zone and at the surface (Fig. S9E). Mucus not subjected to flushing had a staining pattern mirroring that observed in flushed mucus; however, the mix zone appeared thicker, and the plumes smaller (Fig. S9F). Formations resembling the crypt-covering mucus plumes could also be observed by staining of fixed colonic tissue sections (Fig. S9G). However, the mucus plumes were also intensely stained by UEA1, indicating different lectin binding depending on method. Consequently, the observed mucus organization observed *ex vivo* is also likely present *in vivo*.

Bacteria-sized 1 μm beads settled on the mucus surface (Fig. S10A). On the contrary, 0.2 μm beads penetrated the mucus surface and intercrypt mucus and reached the epithelium. However, crypt plume mucus excluded even the 0.2 μm beads (Fig. 5E, and S10B). To rule out that intrinsic properties of the beads affected their distribution, FITC-conjugated dextran with a molecular mass of 2,000 kDa (FD2000) was used. Similar to the 0.2 μm beads FD2000 penetrated down to the epithelium in the intercrypt regions but did not fully penetrate the crypt plumes (Fig. S10C). These findings suggest that the crypt plumes have more stringent barrier properties compared to intercrypt mucus.

Inducing ion and liquid flow by forskolin treatment, or reducing it using *Cftr*-deficient mice did not dramatically alter the overall mucus structure, although the volume of the crypt plumes appeared to increase with forskolin and decrease with impaired *Cftr* activity (Fig. S10, D and E). The ability of the crypt plumes to exclude 0.2 μm beads remained with deficient *Cftr* function. Thus, liquid secretion does not seem to determine overall mucus structure.

Altered icGCs in *Spdef*^{-/-} mice

The transcription factor *Spdef* is thought to be involved in the terminal differentiation of GCs (16, 17), and as it is expressed in the most differentiated GCs, the icGCs, we investigated GC effects in *Spdef* deficient mice. Colonic GCs from 6 week-old *Spdef*^{-/-} (crossed with RedMUC2^{98trTg}) mice showed altered expression of several GC-cluster marker genes, primarily in the canonical lineage, but also in the proliferative cells, compared to WT GCs (Fig. 6A). Furthermore, several of the genes either 2-fold up- or down-regulated in *Spdef*-deficient compared to WT GCs were more abundant in the icGCs, but

genes encoding the main mucus components (5) were unaltered (Fig. 6B, S11A, and Table S7). Many icGC genes were markedly reduced in *Spdef*^{-/-} mice, whereas *Ido1*, expressed by few icGCs, initially showed a slight increase in expression and later expanded to more cells (Figs. 6 A and B, and S11, B and C). Thus, the absence of *Spdef* induce several alteration but appeared to have a pronounced effect on icGCs.

Lectin-based investigation of icGCs in colonic whole-mounted tissue from 7 and 13 week-old *Spdef*^{-/-} mice showed normal icGC and crypt numbers, but revealed age-dependent icGC morphological alterations (Figs. 6, C and D, and S12, A to C). At 7 weeks of age *Spdef*^{-/-} icGCs appeared largely unaffected, whereas in 13 week-old *Spdef*^{-/-} mice the apical membrane staining of icGCs was more intense, and they contained less UEA1 stained material. Transmission electron microscopy (TEM) further revealed gross morphological changes in icGCs, which had increased number of microvilli at the apical membrane, enlarged ER, and reduced theca size, in line with previous observations (17), whereas other GCs appeared normal (Figs. 6E, and S12D). This can explain the difference in staining patterns observed (Fig 4F). The alterations of the apical membrane in icGCs of *Spdef*^{-/-} mice were even more evident by scanning electron microscopy (SEM), resulting in quantitatively lower number (Fig. 6F and S12E).

To investigate defects in mucus secretion and turnover, *in vivo* labeling of *O*-glycans with N-azidoacetylgalactosamine (GalNAz) was performed (28). Mucus turnover in icGCs and crypt GCs of *Spdef*^{-/-} mice was similar to WT mice at 13 weeks of age; however, *Spdef*^{-/-} icGCs, but not crypt GCs, contained less mature mucus compared to their WT counterparts (Figs. 6G, and S12 F and G). The combined results reveal dramatic effects on icGCs in *Spdef*^{-/-} mice.

Altered mucus phenotype and susceptibility to colitis in *Spdef*^{-/-} mice

We next investigated the contribution of icGCs to mucus layer organization and function. At 13 weeks of age, *Spdef*^{-/-} mice largely lacked distinct intercrypt mucus. UEA1 instead primarily stained the WGA-positive crypt plumes (Fig. 7, A and B), which was further confirmed with both isosurface overlay and intensity histogram analysis (Fig. 7, C and D, and S13A). Closer examination of the *Spdef*^{-/-} mucus surface mesh indicated that it mainly originated from the crypt plumes and appeared less dense than in WT explants (Fig. 7B and 5B). Using immunostaining for the Muc2 non-*O*-glycosylated precursor as an icGC marker, UEA1-stained material could be detected in icGCs in 13 week-old *Spdef*^{-/-} mice, indicating that lectin reactivity of mucus produced by *Spdef*^{-/-} icGCs was retained but decreased (Fig. S13B).

Mucus penetrability by 1 μm beads was unaltered in 7 week-old *Spdef*^{-/-} mice compared to WT but the mucus was found to be thinner and more penetrable in 13 week old animals (Fig. 7E and F, and S13C). Nevertheless, crypt-openings remained protected by the plumes (Fig. S13D). A deficient mucus barrier in 13 week-old *Spdef*^{-/-} animals was confirmed by fluorescent *in situ* hybridization (FISH). Gut-residing bacteria were separated from the epithelium by a continuous mucus layer in WT and 7 week old *Spdef*^{-/-} mice, whereas defects in the mucus layer was observed in the older *Spdef*^{-/-} mice (Fig. 7G).

Spdef^{-/-} and WT mice at 7 weeks of age responded similarly to DSS-induced colitis, whereas colitis was more severe in the 13 week old *Spdef*^{-/-} mice (Fig. 7H, and S13E). Notably, *Spdef*^{-/-} mice required euthanasia at 6 days of DSS treatment according to ethical guidelines (Fig. S13F). Additionally, at 26 weeks of age almost all *Spdef*^{-/-} mice exhibited signs of spontaneous colitis by histopathological examination of colonic tissue sections (Fig. 7, I and J, and fig. S13G). FISH revealed bacteria in proximity to the epithelial surface and at the bottom of the crypts in *Spdef*^{-/-} mice, whereas bacteria were clearly separated from the tissue in WT colon (Fig. 7K). These results indicated that there is an age-dependent loss of colonic mucus barrier integrity, in *Spdef*^{-/-} mice where icGC abnormalities are likely a contributing factor

Fewer icGCs and mucus alterations in ulcerative colitis patients

In human sigmoid colon tissue sections, icGCs were strongly stained by the *Lycopersicon esculentum* lectin (LEL) (N-acetylglucosamine-binding), but stained only weakly or negative for *Lotus tetragonolobus* lectin (LTL) (fucose-binding). Conversely, crypt-residing GCs were intensely stained by LTL and largely negative or weakly stained for LEL. The WGA lectin stained GCs throughout the crypt, with increasing intensity towards the icGCs. Thus, human icGCs could be broadly distinguished as WGA⁺/LEL⁺/LTL⁻ GCs (Fig. 8A). Staining of the non-*O*-glycosylated MUC2 precursor was as prominent in human colonic icGCs as in the mouse (Fig. 8B).

Staining for T-antigen (Galβ1-3GalNAc) by Jacalin, which labels all GCs, was used for GC detection in whole-mounted sigmoid colon biopsies from healthy controls and UC patients (Fig. 8C, and Table S8). Individual GCs were mapped and assigned as crypt-opening GCs or icGCs depending on their location (Fig. S14A). A tendency towards fewer observable GCs per crypt was noted in both UC_r and UC_a patients compared to controls, where the reduction of icGCs was further pronounced (Fig. 8D, and S14B). This difference was not due to smaller crypt surface area (Fig. S14C). The frequency of icGCs at different relative positions along the crypt center-to-border axis was used to identify crypt-opening and intercrypt zones, based on the presence of icGCs in each area (Fig. S14D). The number of GCs at intercrypt zone was reduced in UC_r and UC_a patients whereas the number of GCs in the crypt-opening zone was unaltered (Fig. 8E). The difference in number of GC was especially pronounced at the outer border of the crypt (Fig. 8E and S14E). These results indicate that icGCs were lost in the intercrypt zone in UC_a and UC_r patients.

GC shedding was quantified as mucus-containing cells above the epithelial surface and was scored from 0 to 3 (Figs. 8, F and G, S14F, and Table S9). Both UC_r and UC_a samples showed more GC shedding. There was a negative correlation between the number of icGCs and the GC shedding score, indicating that the reduced number of icGCs was a consequence of increased cell shedding (Fig. S14K).

The *ex vivo* mucus organization of healthy human colonic biopsies was similar to that observed in mouse, where crypt covering plumes primarily stained with LTL, and intercrypt mucus stained with both LEL and WGA (Fig. 8H, and Movie S2). The bead penetration

assay showed similar results with 1 μm beads settling on the mucus surface, while 0.2 μm beads penetrated the intercrypt mucus (Fig. 8I).

Analysis of biopsies from UC patients revealed a loss of intercrypt mucus in a majority of the patients (Fig. 8J, and Movie S3). Some patients with absent intercrypt mucus maintained a fairly intact mucus surface, whereas others had a very weak mucus surface staining. Overall both UCr and UCa specimens had more discernable mucus defects than controls, based on scores grading the presence/absence of intercrypt mucus and the density of surface mucus from 0 (normal) to 5 (all mucus lost) (Figs. 8, J and K, S14G, and Table S10). Jacalin did not stain the mucus gaps in UC, but gave more pronounced mucus plume staining than in control patients (Fig. S14H). Furthermore, no increase in Jacalin or T-antigen binding Peanut agglutinin (PNA) staining was observed in icGCs in colonic sections from UC (Fig. S14I).

Plotting the mucus defect score to the number of icGCs showed separate clustering of control and UC patients (Fig. S14J), with a trend towards a positive correlation between the mucus defect score and GC shedding score (Fig. S14K). No correlations could be found between available clinical information (number of active episodes during the last two years, time to last active episode, medication), mucus and icGC scores (data not shown).

Our findings revealed that many UC patients with active disease and in remission have reduced numbers of icGCs, increased GC shedding, and altered mucus organization.

Discussion

The protective intestinal mucus is produced by GCs, and their mRNA and protein expression profiles were now determined. Overall, GCs have cellular pathways that focus on protein production, folding, glycosylation, vesicle transport, and secretion, highlighting mucus production as their main task. Yet, in contrast to the current understanding, single cell analysis of GCs identified distinctive canonical and non-canonical differentiation trajectories, comprising several GC clusters in both small and large intestine. While both trajectories were clearly identifiable in the colon, the non-canonical trajectory was less prominent in the distal small intestine. The non-canonical GCs trajectory is characterized by transcripts in common with enterocytes. Enterocyte markers can also be observed in the GC clusters of other single cell datasets of all intestinal epithelial cells when carefully examined (19, 21, 22, 29). The expression of genes involved in the activation of senGCs, as well as genes dysregulated during colitis are found in non-canonical GCs suggesting that these cells could be crucial for maintaining intestinal homeostasis (22, 27, 30). The GC diversity identified indicate an unpredicted degree of plasticity within the GC population.

In the small intestine, epithelial-derived microbial defense mechanisms are largely attributed to the Paneth cells; however we observed GCs with a defense profile, expressing antimicrobial genes indicating that GCs are a more intrinsic part of the innate immune system than previously considered. The co-secretion of antimicrobials and mucus may represent a strategy to increase the concentration of defense proteins close to the epithelium.

The icGCs located at the colonic surface epithelium are the most differentiated GCs. They possess a distinct morphology, a specific gene expression profile, are detected by specific lectin combinations and non-*O*-glycosylated Muc2 antiserum, and rapidly produce and secrete intercrypt mucus. Based on this we propose that icGCs are a novel, functionally distinct GC subtype, in a similar fashion to the previously described senGCs (23). The high expression of several mucus genes as well as protein transport-associated genes in icGCs is in line with the rapid biosynthesis and secretion of mucus previously observed in these cells (28). Additionally, *Stxbp1*, enriched in icGCs, has been associated with baseline secretion in lung GCs, whereas *Stxbp2*, which is more prominent in crypt GCs, is associated with regulated secretion (31). This suggests that secretion by icGCs is continuous at baseline, while secretion from crypt GCs is more responsive to stimulation (32). Furthermore, enhanced mucus production increases the cellular demand of protein folding. However, several chaperones, including the mucin biosynthesis specific *Agr2* (13), were expressed at lower levels in icGCs, which might indicate that the continuous secretion from icGCs is different from "plume secretion" from crypt GCs. The increased expression of the GC specific gene *Em2*, regulating *Muc2* mRNA levels, also indicate different control in these cells (14).

Ex vivo visualization of mucus on live biopsies provided new insight into mucus organization, revealing different properties of the mucus emerging from icGCs and crypt GCs. The stratified mucus layer seen in immunostainings of fixed colonic tissue likely represents the net-like structures seen at the mucus surface *ex vivo*. The mucus plumes and intercrypt mucus likely correspond to the most recently secreted material, originally described as more MUC2-intense and often associated with the tissue surface in sections (2, 5). The contrasting appearance of mucus by the two methods might arise from different conditions in fixed and live tissue, especially shrinkage upon fixation of the highly hydrated mucus and static pressure exerted by the luminal content.

The enhanced barrier function of the mucus plumes covering the crypt openings provides an extra barrier against microbial intruders, protecting the stem cell niche at the bottom of the crypt. More penetrable intercrypt mucus secreted from icGCs at the epithelial surface interconnects the crypt plumes and while it provides a barrier to microbes, it may enable liquid, ion and nutrient uptake, consistent with the absorptive functions of the surface epithelium. These findings may also explain why motile bacteria can penetrate the mucus at these specific locations (33). Although yet to be resolved, the delineating features of intercrypt mucus and crypt plumes may be a result of cell-specific forms of MUC2, or co-secretion of other factors known to modulate mucus properties, such as proteases, cross-linkers, ions, or fluid volumes (34-37).

The effect of *Spdef* deficiency was prominent in icGCs, based on altered gene expression, morphology and reduced mucus production and secretion of the surface epithelium. These mice spontaneously developed colitis, but the intercrypt mucus defects were evident prior to the onset of inflammation, supporting a fundamental role of intercrypt mucus. Nevertheless, contributing effects of *Spdef* ablation on other GCs or cells causing additional functional alterations cannot be ruled out.

Previous studies have shown that patients with UC have mucus barrier defects; however, such impairments are primarily described in the context of active disease (9, 30). Herein, we have shown that a majority of UC patients, not only those with active disease but also those in remission, have alterations in their mucus organization. The explants from UC patients exhibited enhanced GC shedding on the surface epithelium and reduced icGC numbers, resulting in structural defects in the mucus barrier such as gaps in the intercrypt mucus and exposed areas of surface epithelium. Taken together, alterations of icGCs and intercrypt mucus may represent an early and sustained indicator of disease that could contribute to disease initiation.

The results from mouse and human distal colon suggest comparable mucus organization at the epithelial surface between species. The intercrypt mucus contributes to the mucus barrier organization and is important for limiting bacterial penetration. The common denominator identified in inflammatory initiation is the reduced number or function of icGCs resulting in local mucus barrier defects. The mechanisms likely differ between mice and humans and is more diverse among patients. The experimental limitations to specifically target icGCs and the few patients possible to include preclude detailed understanding, but pave the way for further studies and potential treatment strategies.

In summary, in-depth mRNA/protein expression profiling of intestinal GCs unveils striking heterogeneity in this cell population, which points to a markedly nuanced orchestration of the protective mucus system in the gut. Pseudotime analysis not only identified the anticipated canonical trajectory, but also revealed an additional non-canonical path consistent with a complex differentiation network. Plasticity has been observed in several studies of the intestinal stem cell-proliferative cell niche (38), and in post-mitotic cells (39). Our results support dynamic shaping of the GC population that may reflect both intraepithelial communication and influence from environmental cues. We further identified icGCs as a new functionally distinct GC subtype localized at the apical surface of the colonic epithelium. The icGCs secrete mucus that differs from mucus produced by crypt-residing GCs, and both are required to provide a protective mucus barrier. Deficient intercrypt mucus secretion or reduced icGC numbers leads to disruption of the mucus layer with an impaired organization at the mucus surface, correlating with increased colitis susceptibility. In UC patients, reduced icGC numbers and deficient mucus is an early marker as observed not only in active disease patients but also during remission.

Material and methods

A full description of the methods can be found in the supplementary materials

Animals

All animal experimental procedures were in compliance with Swedish animal welfare legislation and approved by the Swedish Laboratory Animal Ethical Committee in Gothenburg, Sweden (number: 280-12, 73-15, 74-15, 2285-19 and 2292-19). C57BL/6, RedMUC2^{98trTg} mice (23, 40), *Spdef*^{-/-} (17), crossed to the RedMUC2^{98trTg} strain, and *Cftr*^{tm1Unc} mice (41) were bred in house in SPF conditions and all experiments used age-

matched mice of both genders. DSS-colitis was induced by administration of 3% w/v DSS and disease activity index was scored daily (12, 42).

Human samples

Sigmoid colon biopsies were obtained from patients referred for colonoscopy at Sahlgrenska University Hospital (Gothenburg, Sweden) (Table S8) in compliance with the local human research ethical committee (136-12) and the Declaration of Helsinki as described previously (43). Patients with normal intestine, evaluated macroscopically, were included as controls. Patients with full Mayo score 0 were considered as remission, and 1 as active UC.

Intestinal epithelial cell isolation and sorting

Intestinal epithelial cells were isolated from freshly collected distal colon and distal small intestine from RedMUC2^{98trTg} mice and *Spdef*^{-/-} RedMUC2^{98trTg}. GC and non-GC were sorted with FACS Jazz (Becton Dickinson) according to the presence or absence of mCherry signal with dead cells and doublets removed. The purity of sorted cells was > 96%.

RNA extraction and sequencing

Sorted cells were pelleted and total RNA extracted using RNeasy kit and QIAshredder columns (Qiagen). The cDNA libraries were prepared using the TruSeq Stranded Total RNA Sample Preparation kit with Ribo Zero Gold (Rev. E; Illumina) and sequenced via paired-end with the NextSeq500 platform (Illumina). The reads were mapped against the mouse reference genome mm10 from Ensembl using STAR (44). The number of mapped reads was calculated using HTseq (45) and different expression and statistical analysis with DESeq2 in R (46).

Protein extraction

Sorted cells were used and digested with with LysC and trypsin. Samples were analyzed with an EASY-nLC 1000 system connected to a Q-Exactive mass-spectrometer (Thermo Fisher). MS raw files were processed with MaxQuant software (47) by searching against mouse UniProt database combined with an in-house database containing all human and mouse mucin sequences, including mCherry MUC2 (<http://www.medkem.gu.se/mucinbiology/databases/>) using Label-Free Quantification (LFQ) method. Statistics were analyzed with Welch's two-sample t-test, and the Benjamini-Hochberg test for adjusted p-values.

Data analysis for RNA-sequencing and mass spectrometry.

Volcano plots were created using EnhancedVolcano (48). The top 50 genes and proteins were clustered using the heatmap.2 function from gplots. Venn diagrams were made with VennDiagram. All these packages were run in R. Fold change ≥ 2 and adjusted p-value ≤ 0.05 (proteins) or ≤ 0.01 (RNA) were used to select enriched proteins and genes. Gene Ontology enrichment analysis was performed using the DAVID tool (49, 50).

Single cell preparation, sequencing and analysis

GCs from distal colon and distal small intestine were sorted and droplet-based scRNA-seq was performed using the chromium single cell 3' v2 platform (10x Genomics). Library sequencing was performed using the NextSeq500 platform. The data processing was performed using Cell Ranger software (10x Genomics) and Seurat (51, 52) in R. Immune, enteroendocrine, Paneth cells, and cells with low expression of *Muc2* (<3) were removed. We identified 15392 genes across 6123 cells for the colon and 14794 genes across 3552 cells for the small intestine. Statistical analysis was performed using Wilcoxon Rank Sum test and adjusted p-values Bonferroni correction.

Pseudotime analysis was performed using Scanpy (53) in Python resulting in 14447 genes across 3402 cells for the small intestine and 14892 genes across 5936 cells for the colon. GCs were clustered using the Lovain algorithm. Statistically analysis used Wilcoxon Rank Sum test and the adjusted p-values Benjamini-Hochberg test.

Immunostaining, *in situ* hybridization and colonic whole-mounted tissue

Paraffin-embedded methanol-Carnoy-fixed sections were dewaxed, rehydrated and antigen-retrieved for immunostaining with different combinations of anti-Ido1 (Cell Signaling), anti-Epcam (Abcam), chicken monoclonal anti-Defa24, raised against a well characterized peptide(54), anti-Lysozyme (Genway) or anti-apoMUC2 (detecting the non-O-glycosylated mucin) antibodies (PH497 and PH1900 for mouse and human respectively (55, 56). Bacterial FISH was performed as previously described (57) with counterstaining using fluorescently conjugated lectins. For RNA-*in situ* hybridization we used the RNA scope technique (Advanced Cell Diagnostics) according to manufacturer's instruction on formalin-fixed tissue sections. GCs in formalin-fixed whole-mounted tissue from mouse or human biopsies were stained by lectins or an anti-human apoMUC2 and counterstained with CellMask Deep Red plasma membrane stain (Thermo Fisher Scientific). In all cases, DNA was counterstained by Hoechst-34580 (Thermo Fisher Scientific). Images were recorded with an LSM700 Axio Examiner Z.1 confocal microscope (Zeiss) and further processed using the Imaris software (Bitplane) or ImageJ v1.52k (58). Scoring criteria for GC shedding are given in table S9.

GalNAz labeling of mice and turnover of mucus

WT and *Spdef*^{-/-} mice at 13 weeks of age were labeled with GalNAz by intraperitoneal injection with subsequent detection using by Alkyne-Tetramethylrhodamine (TAMRA) (both from Thermo Fisher Scientific) on fixed colonic tissue sections as previously described (28).

Ex vivo mucus analysis

Procedure, materials and equipment for *ex vivo* examination of colonic mucus were employed as previously described (43) with visualization of mucus by fluorescently conjugated lectins and investigation of mucus barrier properties using fluorescent beads or FITC-conjugated dextran (2000 kDa). Quantification of bead penetrability was performed as previously described (59, 60). Z-stacks were acquired using LSM700 with a Plan-Apochromat x 20/1.0 DIC water objective (Zeiss) and the ZEN 2010 software. Image

analysis was performed using Imaris. Mucus defect scoring was done in a blinded fashion according to table S10.

Electron microscopy

Colonic tissue was excised, flushed and fixed in modified Karnovsky's fixative, cut and further processed for either TEM or SEM using a Zeiss Leo 912AB Omega electron microscope or a Zeiss Gemini DSM 982 microscope respectively. GCs were manually counted in a blinded fashion for quantification.

Statistical analysis

Statistical analysis and graphical illustrations were performed using GraphPad PRISM 7.03 or 8.3.1 (GraphPad Software). Data are presented as mean \pm standard deviation (SD) or median with range, as indicated. Statistical tests were applied as indicated. For all statistical analyses: * $p < 0.05$, ** $p < 0.01$, *** $p < 0.0001$, **** $p < 0.00001$, ns = non-significant.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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Data availability

MS proteomics data have been deposited to the ProteomeXchange Consortium (<http://proteomecentral.proteomexchange.org>) via the PRIDE (61) partner repository with the dataset identifier PXD011527. All raw data from the bulk RNA-seq and scRNA-seq are deposited to the GEO database (GSE144436).

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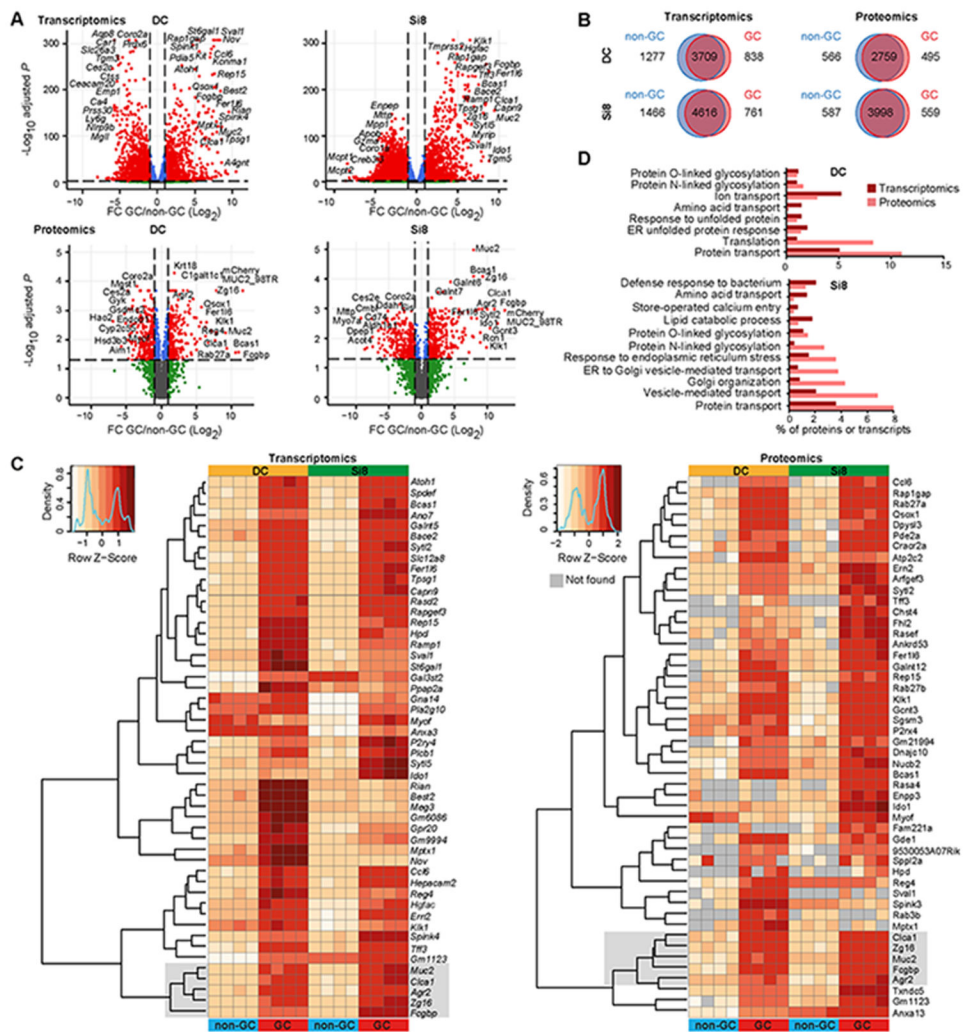


Fig. 1. Multi-omics analysis of purified GCs from the distal colon (DC) and the distal small intestine (Si8).
(A) Volcano plots of transcriptome and proteome in GC and non-GC populations. **(B)** Venn diagram indicating the number of genes and proteins associated with each population using fold change (FC) values ≥ 2 and adjusted-p value ≤ 0.05 (protein) or ≤ 0.01 (mRNA). **(C)** Heat maps of the 50 genes and proteins most differentially expressed in the GCs compared to non-GCs. **(D)** Biological processes upregulated in GCs according to Gene Ontology (GO) enrichment analysis. Four replicates (pool of two mice per replicate).

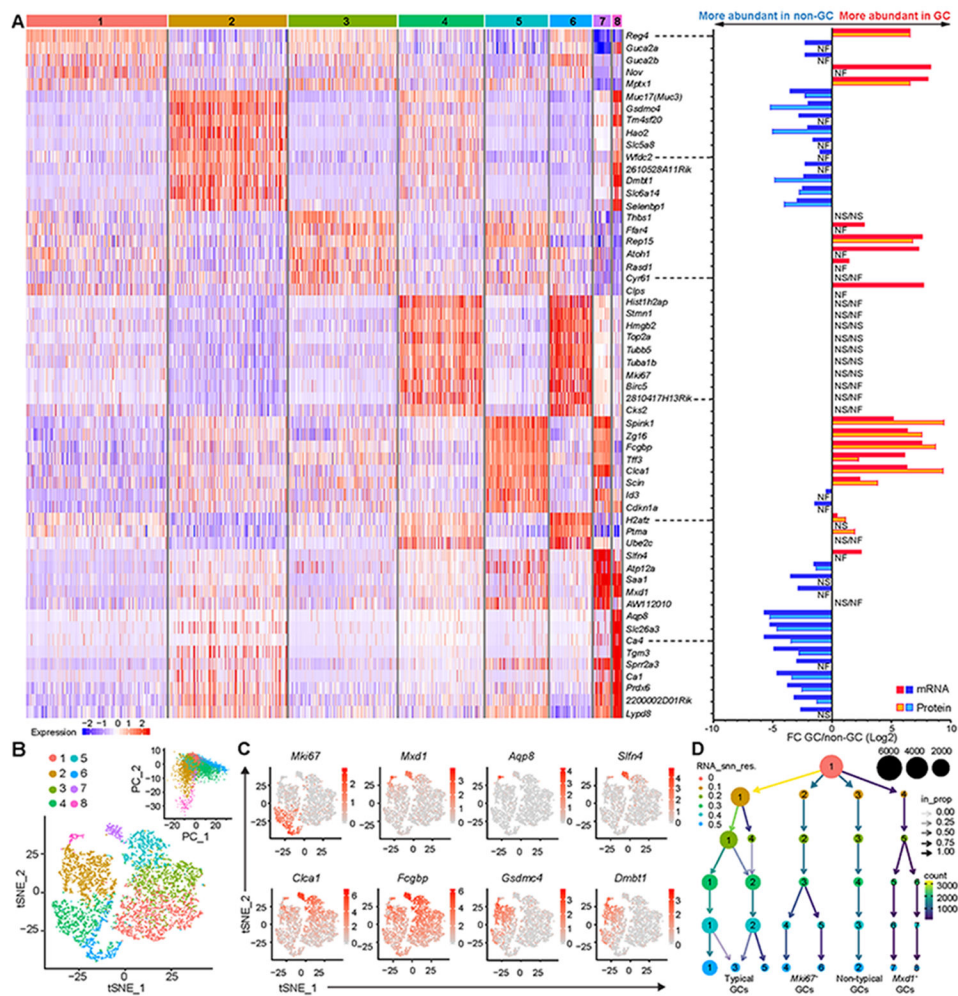


Fig. 2. Different clusters identified among the *Muc2*⁺ cells in the distal colon by single-cell analysis. (A) Left panel: Heat map of up to 10 enriched genes per cluster; Right panel: mRNA and protein FC of GC to non-GC genes displayed in the heat map. (B) tSNE plot of colonic GCs grouped in eight clusters and principal component analysis (PCA) of cluster diversity. (C) tSNE plots highlighting expression and distribution of selected genes. (D) Cluster tree showing the four main subgroups of GCs during the clustering process. Two replicates (two mice per replicate) were combined for the analysis. FC: Fold change. NS: Not significant; NF: Not found.

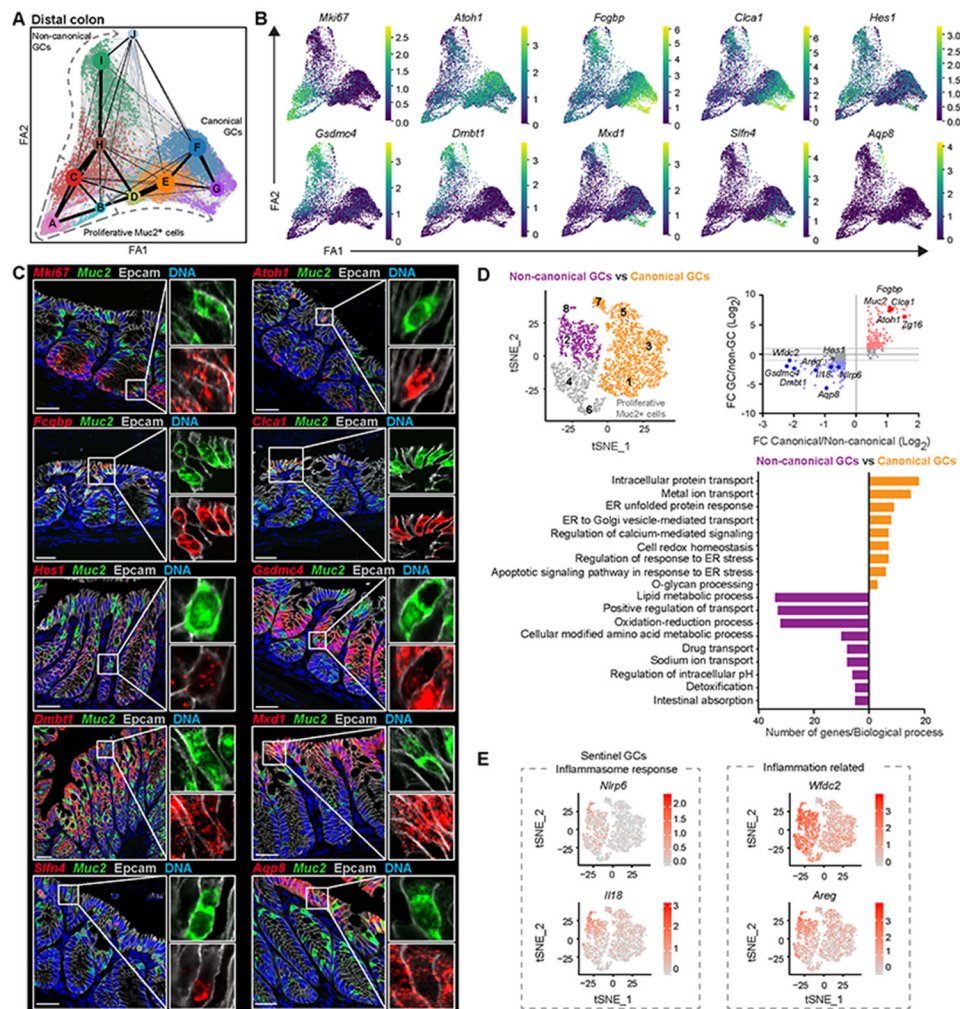


Fig. 3. Pseudotime analysis unveils two trajectories of GCs in the distal colon.

(A) Trajectory and Partition-based graph abstraction of colonic GCs. Two replicates (2 mice per replicate) were combined. (B) Expression of selected genes along the described trajectories. (C) mRNA-*in situ* hybridization of genes enriched in different clusters counterstained with *Muc2*, Epcam for cell borders and Hoechst for DNA. Scale bars = 25 μ m. (D) tSNE plot, scatter plot of the bulk RNA-seq data of genes significantly enriched in the canonical and non-canonical GC populations, and biological processes upregulated in canonical GCs compared to non-canonical GCs according to GO enrichment analysis. (E) tSNE plots highlighting expression of sentinel GC signaling genes and genes dysregulated during inflammation.

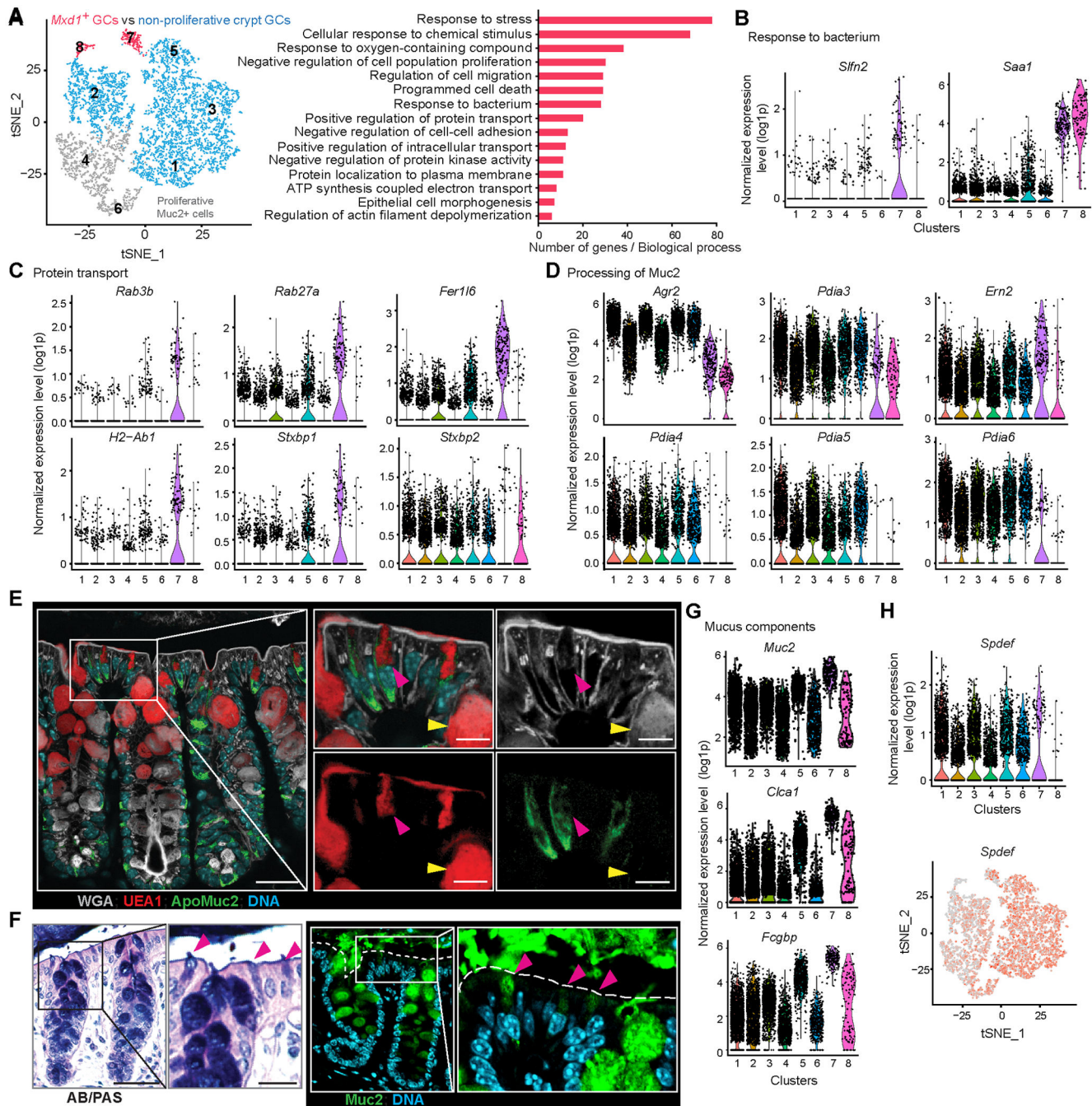


Fig. 4. Gene expression in icGCs.

(A) Biological processes upregulated in the surface located GCs (*Mxd1*⁺) compared to crypt-residing GCs according to GO enrichment analysis. (B, C, D) Violin plots of gene expression related to response to bacteria (B) protein transport (C) and processing of Muc2 (D) in the different clusters. (E) Staining with WGA, UEA1, the non-*O*-glycosylated Muc2 precursor and Hoechst for DNA in a colonic cross section. Arrowheads indicate crypt-opening GC (yellow) and an icGC (pink). Scale bars = 20 μ m, and 10 μ m in magnification; n = 24. (F) Mouse colonic section stained with AB-PAS or mature Muc2 with icGCs indicated

(pink arrowheads). Epithelial surface (dashed line), scale bars = 50 and 20 μm (left and right panel in AB-PAS). **(G)** Violin plots of mucus gene expression in the different clusters. **(H)** Violin and tSNE plots showing expression of *Spdef* in GCs.

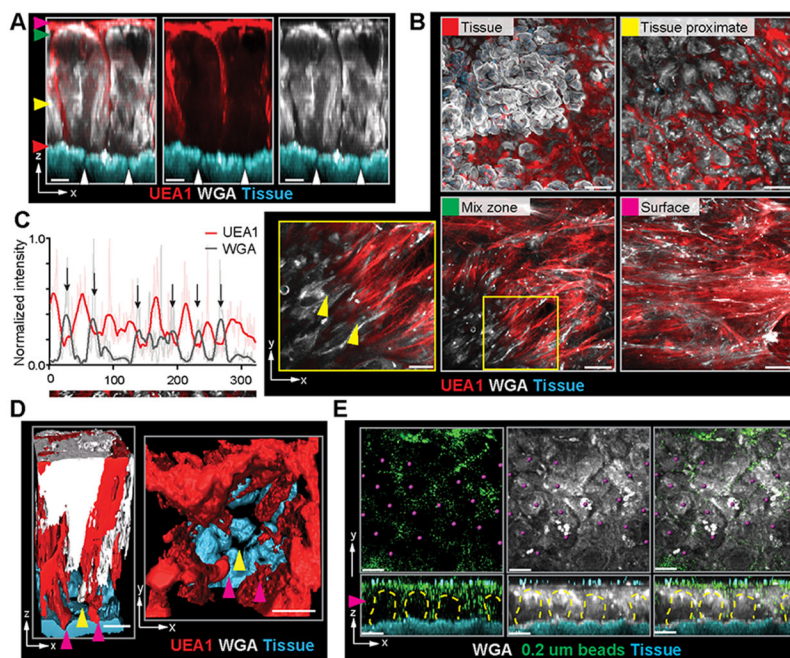


Fig. 5. Colonic mucus organization and penetrability.

(A) UEA1 and WGA-stained flushed mouse colonic mucus in x/z -projection with marked (white arrowheads) crypt-openings. Color code arrowheads match images in B. (B) x/y projections at different z -stacks levels as indicated in A. Magnification of the mixed zone mucus (yellow arrowheads mark WGA and UEA1 fusion points); $n = 30$. (C) Intensity profile of UEA1 and WGA along a straight line in tissue proximate mucus (Fig. S9). Normalized raw data in pale color, crypt-openings (arrows); $n = 17$. (D) Isosurface overlay of UEA1 and WGA-stained mucus secreted from icGCs (pink arrowheads) or a crypt-opening (yellow arrowhead). (E) x/y and x/z projections of $0.2 \mu\text{m}$ fluorescent beads in WGA-stained colonic mucus. Areas with low bead concentration (dashed line), underlying crypt-openings (purple spheres), z -position for x/y projections (arrowhead); $n = 6$. Scale bars: $50 \mu\text{m}$ (B, E), $25 \mu\text{m}$ (A, B magnification) and $20 \mu\text{m}$ (D).

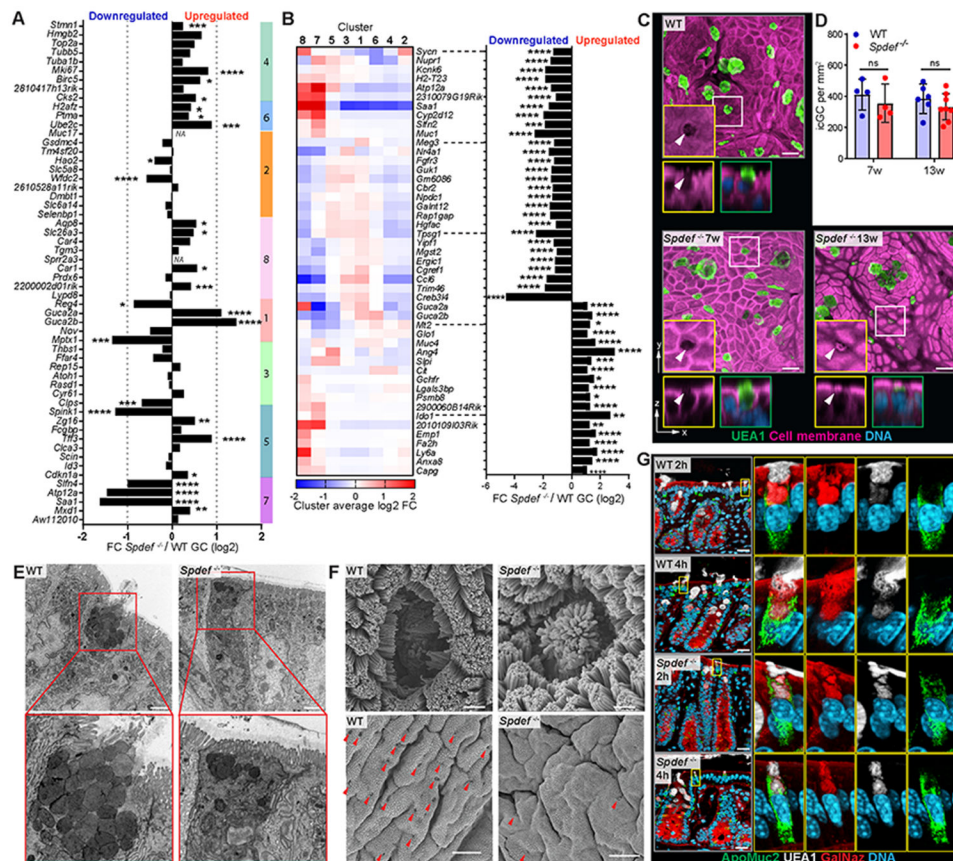


Fig. 6. icGC alterations in *Spdef*^{-/-} colon.

(A) Fold change (FC) expression of single GC cluster (color bars) marker genes in *Spdef*^{-/-} versus WT with adjusted p-values. (B) FC expression among the previously defined single cell clusters (left) of the most up- and down-regulated genes in *Spdef*^{-/-} compared to WT GCs obtained by bulk RNA seq with adjusted p-values (right). (C) UEA1, cell membrane and DNA staining of whole-mounted colonic tissue from 7 and 13 week-old WT and *Spdef*^{-/-} mice. Magnified areas (white boxes), apical surface of icGCs (arrowheads), scale bars = 20 μ m. (D) Quantification of the number of icGCs per mm² in whole-mounted colons from WT and *Spdef*^{-/-} mice. Mean \pm SD, analyzed by 2-way ANOVA with Sidak's multiple comparison test; n = 4 - 8. (E) TEM micrographs of icGCs in WT and *Spdef*^{-/-} colon with magnifications. Scale bars = 2 μ m; n = 4. (F) SEM scans of the icGCs and the epithelial surface in WT and *Spdef*^{-/-} distal colon. Identified GC openings (red arrowheads), scale bars = 1 μ m (upper) 20 μ m (lower), n = 4. (G) *In vivo* labeling for 2 and 4 h by GalNAz detected by Alkyne-TAMRA on colonic sections of 13-week old WT and *Spdef*^{-/-} mice. Scale bar = 20 μ m; n = 4.

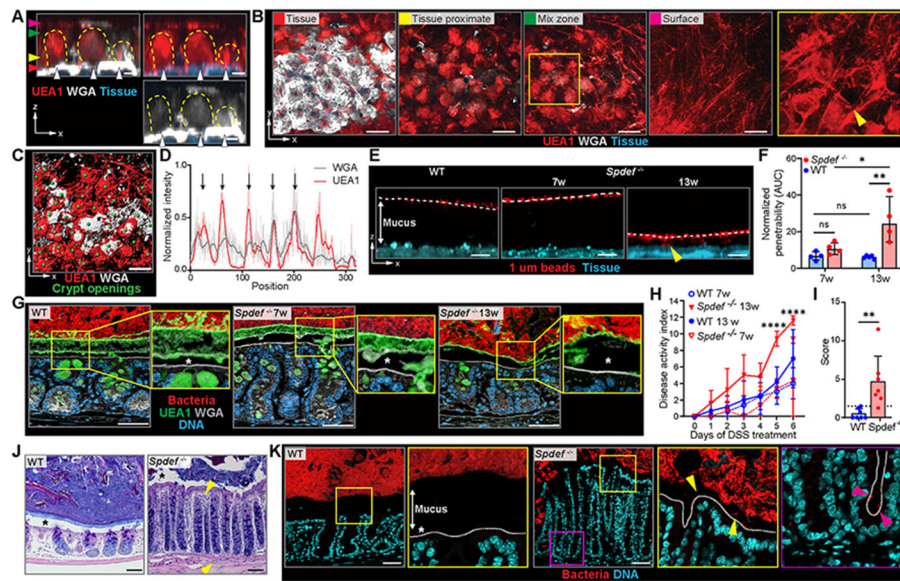


Fig. 7. Age-dependent loss of intercrypt mucus and colitis in *Spdef*^{-/-} mice.

(A) UEA1- and WGA-stained *Spdef*^{-/-} colonic mucus in x/z-projection. Crypt-openings (white arrowheads), crypt plumes (dashed line). Color code arrowheads match images in B. (B) x/y projections at different z-stack levels as indicated in A. Magnification of mucus from the crypt plumes expanding to form the mucus surface mesh (arrowhead); n = 12. (C) Isosurface overlay of UEA1 and WGA. Underlying crypt-openings marked by green spheres. (D) Intensity profile of UEA1 and WGA in tissue proximate mucus (Fig. S12). Normalized raw data (pale color), crypt openings (arrows); n = 3. (E) 1 µm beads on WT and *Spdef*^{-/-} colonic mucus. Mucus surfaces (dashed lines), beads at tissue (arrowhead); n = 4 - 5. (F) Normalized bead penetrability; n = 4 - 5. (G) Bacteria FISH, UEA1 and WGA in sections from WT and *Spdef*^{-/-} colon; n = 7. (H) Colitis score of *Spdef*^{-/-} and WT mice treated with 3% DSS; n = 3 - 13 (I) Histological colitis score in 26 week-old WT and *Spdef*^{-/-} mice. Normal score range of 6-19 week-old WT (n = 14, data not shown) was set as the colitis cut-off (dashed line). (J) AB-PAS stained colonic sections from 26 week-old WT and *Spdef*^{-/-} mice. Infiltrating inflammatory cells (arrowheads); n = 7. (K) Bacteria FISH in sections from 26 week-old WT and *Spdef*^{-/-} animals. Epithelial surface (dashed line), bacteria at the epithelium (arrowheads); n = 4. Scale bars = 50 µm. * in images indicate shrinkage artifacts due to fixation. Graphs shows mean ± SD analyzed by 2-way ANOVA with Tukey's multiple comparison test (F and H) or unpaired Student's t-test (I).

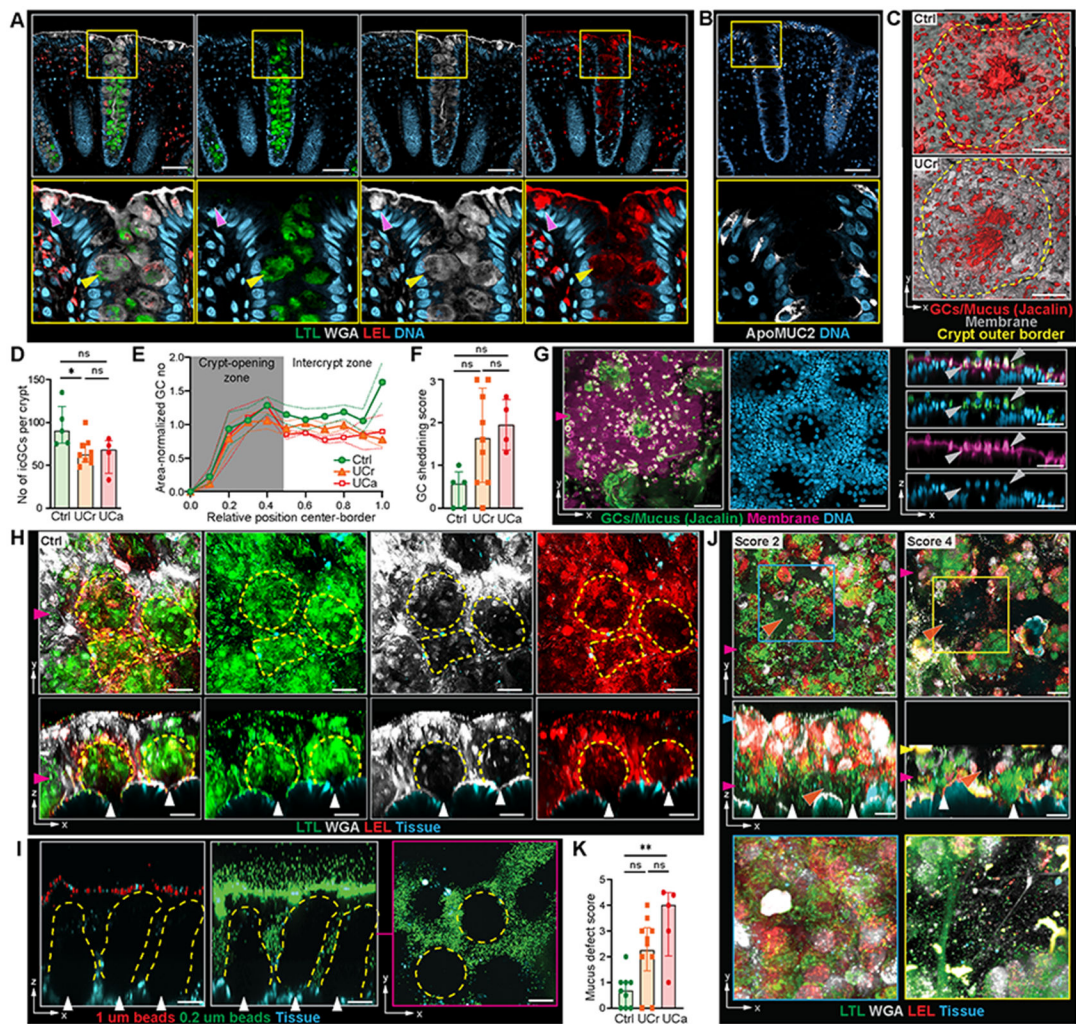


Fig. 8. Mucus alterations and loss of icGCs in UC patients.

(A, B) Lectin (A) and non-*O*-glycosylated MUC2 immunostaining (B) in a human sigmoid colon cross section. Arrowheads: crypt-opening GC (yellow) and icGC (pink); $n = 12$. (C) Whole-mounted colonic tissue from control (Ctrl) and UC patients. The outer border of the crypt (dashed yellow line). (D, E, F) Number of icGCs per crypt (D), number of GCs (normalized to the relative area for each segment) at different relative positions along the crypt center-border axis (E), and GC shedding score (F) in Ctrl, UCr and UCa colonic biopsies; $n = 4 - 9$. (G) GC shedding (gray arrowheads) in a fixed whole-mounted UCr biopsy. Position for x/z projection (pink arrowhead). (H) Biopsies from human sigmoid colon stained *ex vivo* with LTL, WGA and LEL. Crypt plumes (dashed lines), level of corresponding projection (arrowhead), $n = 9$. (I) Mucus penetrability of 1.0 and 0.2 μm fluorescent beads. Mucus without 0.2 μm beads (dashed line), crypt-openings (arrowheads); $n = 3$. (J) Mucus defect scores images in biopsies from UC patients. Mucus gaps (orange arrowhead), crypt openings (white arrowheads), and z-positions for x/y projections (color coded arrowheads). (K) Mucus defect score in Ctrl, UCr and UCa patients. Scale bars =

50 μm . Bars indicate median \pm interquartile range. Statistical analysis was performed by Kruskal-Wallis analysis with Dunn's multiple comparison test.

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