Goblet cells deliver luminal antigen to CD103+ dendritic cells in the small intestine

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The intestinal immune system is exposed to a mixture of foreign antigens from diet, commensal flora and potential pathogens. Understanding how pathogen-specific immunity is elicited while avoiding inappropriate responses to the background of innocuous antigens is essential for understanding and treating intestinal infections and inflammatory diseases. The ingestion of protein antigen can induce oral tolerance, which is mediated in part by a subset of intestinal dendritic cells (DCs) that promote the development of regulatory T cells2–9, and CD103+ dendritic cells (DCs) (with features of macrophages), which promote tumour necrosis factor-α (TNF-α) production, colitis, and the development of Th17 T cells5–7,10. However, the mechanisms by which different intestinal LP-DC subsets capture luminal antigens in vivo remains largely unexplored. Using a minimally disruptive in vivo imaging approach we show that in the steady state, small intestine goblet cells (GCs) function as passages delivering low molecular weight soluble antigens from the intestinal lumen to underlying CD103+ LP-DCs. The preferential delivery of antigens to DCs with tolerogenic properties implies a key role for this GC function in intestinal immune homeostasis.

We examined the in vivo antigen acquisition behaviour of intestinal LP-DCs in fluorescent DC-reporter mice using two-photon microscopy (Supplementary Fig. 1a). The intestine is imaged within the peritoneal cavity and images can be acquired from either the intact intestinal serosa or from the luminal surface through a small longitudinal incision in the intestine (Fig. 1a, b and Supplementary Movie 1, upper panels). The preparation is sufficiently stable to permit three-dimensional imaging of DC behaviour deep within intestinal tissues (Fig. 1c and Supplementary Movie 1, lower left panel) and preserves blood flow and epithelial barrier integrity for more than 4 h of continuous imaging.

We assessed antigen distribution by two-photon microscopy following the intraluminal injection of 10 kDa rhodamine dextran as a model antigen. Dextran coated the surface of the epithelium and filled the space between villi and crypts (Fig. 1a, b). In addition, we observed cylindrical dextran columns approximately 5 μm in diameter and about 20 μm long projecting through the villus epithelium and into the LP, when imaging from either serosal or luminal orientations (Fig. 1a–c, Supplementary Fig. 1b and Supplementary Movie 1, upper panels and lower right panel). Transepithelial dextran columns were common throughout the small intestine from duodenum to ileum, but did not cause a general disruption of the epithelial barrier as shown by the exclusion of dextran from the LP (Fig. 1a–c). We did not detect transepithelial dextran columns in the stomach, caecum, or colon, with the exception of the epithelium overlying the caecal patches (Supplementary Fig. 1c–e and Supplementary Movie 2). Confocal microscopy revealed that dextran columns were a subset of epithelial cells containing intracellular dextran, which had a continuous border of e-cadherin on their basolateral surface and were often in contact with yellow fluorescent protein (YFP1) cells in the LP of CD11c-YFP reporter mice20 (Fig. 1d, e and Supplementary Movie 4, right panel).

Periodic acid–Schiff staining of mucin in sections of the small intestine (Fig. 2a) produced a goblet cell (GC) staining pattern similar in frequency, distribution and dimensions to the dextran columns identified by two-photon microscopy (Fig. 2b, c). Furthermore, in contrast to the acellular and impermeable discontinuities seen in the small intestine epithelium11, dextran columns were associated with a nucleus (Fig. 2d, Supplementary Fig. 2a and Supplementary Movie 3). To determine if the dextran-filled cells were in fact GCs, sections of intestine from mice given lysine-fixable dextran were stained with antibodies to mucin 2 (MUC2) and cytokeratin 18, which are both highly expressed by GCs12. Dextran columns showed near perfect colocalization with MUC2+ and cytokeratin 18+ epithelial cells displaying GC morphology (Fig. 2e, f). Therefore, we term this phenomenon ‘goblet-cell-associated antigen passages’ (GAPs). To address the possibility that GAPs are apoptotic GCs, we co-stained for various markers of apoptosis, including cleaved cytokeratin 18, cleaved caspase 3 and TdT-mediated dUTP nick end labelling (TUNEL; Supplementary Fig. 3a–i). In all cases, we found no association between apoptotic GCs and GAPs. Moreover, GAPs are distinct from villous M cells, because they did not colocalize with the M-cell marker glycoprotein 2 (GP2) (Fig. 2g)13. The frequency and distribution of GAPs assessed by two-photon microscopy was similar in all strains of specific-pathogen-free (SPF) mice examined (Supplementary Fig. 2b–d), with a non-significant trend toward more GAPs detected in the terminal ileum (Supplementary Fig. 2h). GAPs were also evident in human jejunum resection specimens (Fig. 2h, i), suggesting that GAPs are a general phenomenon of the healthy small intestine. We examined the frequency of GAPs in C3H/HeJ/Bir IL-10−/− mice14, which develop spontaneous intestinal inflammation with GC loss, and in germ-free mice that lack normal gut flora. The number of GAPs and GCs correlated strongly; GAPs and GCs were significantly more numerous in germ-free mice (Supplementary Fig. 2e, g) and significantly fewer in IL-10−/− mice (Supplementary Fig. 2f, g).

Previous studies have shown that LP-DCs can extend transepithelial dendrites (TEDs) between intestinal epithelial cells to sample luminal contents and microbiota15–18. However, these studies used ex vivo and exteriored tissue preparations that involved removing the luminal contents and mucous before imaging. Using our in vivo imaging preparation and confocal microscopy, we found that although LP-DCs probed the epithelium actively with their dendrites (Fig. 1c and Supplementary Movie 1, lower left panel, and Supplementary Movie 4), they did not extend TEDs into the intestinal lumen to capture fluorescent antigen in healthy mice (based on over 50 independent intravital imaging experiments examining all regions of the small intestine from the tip of the
of luminal dextran or beads on the basis of fluorescence colocalization (Supplementary Fig. 4a and Supplementary Movie 5, upper left panel).

Next, we examined whether paracellular leak would serve as a major source of luminal antigen for small intestine LP-DCs. The intraluminal injection of 5 mg of 10 kDa dextran produced a faint ‘feather’-like staining pattern between villous epithelial cells, consistent with paracellular leak (Supplementary Fig. 4b and Supplementary Movie 5, upper right panel). Time-lapse two-photon imaging showed that dextran collected at the base of the epithelium, but was flushed out of the villi efficiently during contraction and did not remain colocalized with LP-DCs even in areas of extensive paracellular leak (Supplementary Fig. 4b and Supplementary Movie 4, left panel, and Supplementary Movie 5, upper right panel). In addition, we did not detect paracellular leak around GAPs in fixed sections by confocal microscopy, despite the more permissive tight junctions of GCs. However, we cannot exclude the possibility that LP-DCs capture low levels of antigen via paracellular leak, because this process might be below the level of detection of our imaging approach.

In contrast to TEDs and paracellular leak, two-photon time-lapse imaging provided direct evidence that GAPs are a source of luminal antigen for LP-DCs. In addition to dextran (Fig. 3a), GAPs were capable of transporting protein antigens (Fig. 3b). Although most GAPs remained visible for the duration of our imaging experiments, they were a dynamic phenomenon (Supplementary Movie 5, lower panels). Moreover, the manner in which LP-DCs interacted with GAPs varied. In some cases, DCs made stable contacts and slowly collected antigen over several minutes (Fig. 3a and Supplementary Movie 5, lower left panel) whereas in others, DCs actively probed GAPs and captured clumps of antigen (Fig. 3b and Supplementary Movie 5, lower right panel). We assessed the molecular weight exclusion limit of GAPs and found that beads ranging from 0.02 to 1.0 in size did not enter GAPs (Supplementary Fig. 4c). In contrast, GAPs filled rapidly with 10 kDa dextran (Supplementary Fig. 4d) and dextran colocalized with CD11c-YFP LP-DCs 2 h after intraluminal injection (Fig. 3a and Supplementary Movie 5, lower left panel). GAPs also filled with larger dextrans (Supplementary Fig. 4d); however, capture by LP-DCs was markedly reduced with 70 kDa dextran (Fig. 3d) and undetectable with 2,000 kDa dextran during our 4-h imaging window.

The small intestine LP contains two prominent DC populations: CX3CR1+CD103+ DCs with tolerogenic potential and CX3CR1+CD103+ DCs that have features of macrophages and have been implicated in intestinal inflammation. We visualized LP-DC subsets in vivo using dual-reporter mice created by crossing CD11c-YFP mice with CX3CR1-GFP knock-in mice (ref. 28) in which CD103-CD103+LP-DCs can be distinguished by the presence or absence of CX3CR1-GFP expression, respectively (Fig. 3c and Supplementary Fig. 5). We frequently observed CD11c-YFP+CX3CR1-GFP+LP-DCs sampling GAPs by two-photon microscopy, but this behaviour was rare in CD11c-YFP+CX3CR1-GFP+LP-DCs (Fig. 3c and Supplementary Fig. 6c); out of 50 total LP-DCs containing dextran, 49 were CD11c-YFP+CX3CR1-GFP+ (Fig. 3d). Furthermore, when we directly observed antigen transfer from GAPs to DCs (20 out of 500 GAPs examined), GAPs delivered antigen exclusively to CD11c-YFP LP-DCs, often stained positive for cytokeratin 18. In contrast to TEDs and paracellular leak, two-photon time-lapse imaging confirmed that the cells interacting with GCs were CD103+CD11c+LP-DCs and not B220+ plasmacytoid DCs (pDCs) (Supplementary Fig. 6a). Moreover, flow cytometry showed that luminal antigen was captured preferentially by CD103+DCs at a proportion of roughly 10:1 over CD103+DCs (Fig. 3e) and rarely colocalized with pDCs (Supplementary Fig. 6b), consistent with the sampling bias observed by two-photon imaging. Control experiments demonstrated that antigen uptake during cell isolation was negligible (Supplementary Fig. 7), thus the flow cytometry results measure the in vivo antigen acquisition capacity of different LP-DC subsets. Interestingly, CD103+LP-DCs, but not CD103+LP-DCs, often stained positive for cytokertin 18 (Fig. 3f and g), which is highly expressed by villous GCs (Fig. 3h).
Because cytokeratin 18 expression is undetectable in LP-DCs by quantitative real-time PCR (Fig. 3h), this suggests that CD103+ LP-DCs interact selectively with GAPs and can capture GC-derived proteins.

In two-photon imaging experiments, LP-DCs captured fluorescent luminal ovalbumin (Ova) readily from GAPs (Fig. 4a and Supplementary Movie 6), similarly to fluorescent dextran and BSA. To determine if LP-DCs could process and present luminal antigen, we administered Ova in vivo, then sorted LP-DCs and co-cultured them with OTI T cells\textsuperscript{11}. T-cell proliferation was assessed on day 3 by both CFDA (dye) dilution (Fig. 4b, c) and by counting T cells after culture (Fig. 4d). Total LP-DC populations (CD45\textsuperscript{+}, CD11c\textsuperscript{+}, MHCII\textsuperscript{+}) were capable of inducing modest OTI T-cell proliferation that was significantly greater than controls (Fig. 4b, d). CD103\textsuperscript{+} LP-DCs stimulated significant OTI T-cell proliferation, whereas CD103\textsuperscript{−} LP-DCs from the same mice did not (Fig. 4d). The failure of CD103\textsuperscript{−} LP-DCs to stimulate T cells was not due to a lack of intrinsic antigen presentation capacity, because they were capable of inducing comparable levels of OTI T-cell proliferation to CD103\textsuperscript{+} LP-DCs (81.4% to 88.2%, respectively) when Ova was added to the cell culture (Fig. 4c). These findings suggest that CD103\textsuperscript{+} LP-DCs have cross-presentation capacity, similar to their mesenteric lymph node counterparts\textsuperscript{11}.

In some two-photon time-lapse recordings we observed GAPs forming or disappearing (Supplementary Movies 3 and 5, lower panels), suggesting that GAP formation could be related to changes in GC function or secretion. To test whether GC secretion was associated with GAP

**Figure 2** | GCs are associated with the transepithelial passage of luminal material. a, b, Periodic acid–Schiff (PAS) staining of GCs (a) and dextran columns (b) visualized by two-photon microscopy. c, PAS and dextran stained structures have similar morphology and dimensions. d, Dextran columns were often associated with a nucleus (white arrow). e, f, Dextran columns (Dex, red) colocalized with GC markers cytokeratin 18 (Cyt18, white) (e) and MUC2 (f). g, Cytokeratin 18-positive cells (white arrows) did not colocalize with the M-cell marker GP2 (yellow arrows). h, i, Dextran columns were present in healthy human small intestine (h) and stained positive for cytokeratin 18 (i). Scale bars, 30 μm (a and b, h), 20 μm (d, e, g and i), and 10 μm (f). Error bars, s.d.

**Figure 3** | GAPs deliver soluble antigen to CD103\textsuperscript{+} LP-DCs in the steady state. a, b, Time-lapse two-photon imaging of model antigens (red) dextran (a) and bovine serum albumin (BSA) (b) delivered by GAPs to LP-DCs (green, CD11c-YFP). Antigen from GAPs colocalized with LP-DCs (yellow arrows) over time. c, Two-photon imaging of CD11c-YFP CX3CR1-GFP mice. GAPs (red) delivered antigen preferentially to CD103\textsuperscript{+} LP-DCs (CD11c-YFP\textsuperscript{+} CX3CR1\textsuperscript{−} GFP\textsuperscript{−} ; green) over CD103\textsuperscript{−} LP-DCs (CD11c-YFP\textsuperscript{+} CX3CR1\textsuperscript{+} GFP\textsuperscript{+} ; cyan) LP-DCs. d, Enumeration of GAPs and GAPs delivering antigen to LP-DC subtypes in CD11c-YFP CX3CR1-GFP mice in two-photon recordings. e, Flow cytometry of LP cells showed that CD103\textsuperscript{−} DCs captured more luminal dextran than CD103\textsuperscript{+} DCs. FITC, fluorescein isothiocyanate. SSC, side scatter. f, Cytospins on sorted LP-DCs stained with DAPI (blue) and the GC marker cytokeratin 18 (red). g, Significantly more CD103\textsuperscript{+} LP-DCs than CD103\textsuperscript{−} LP-DCs stained cytokeratin 18-positive per high-powered field (P = <0.001). h, Neither DC population expressed detectable cytokeratin 18 mRNA. Scale bar, 15 μm (a and b), 50 μm (c), 25 μm (f). Data in g taken from nine or more high-powered fields (representing more than 150 cells) Data in h performed in triplicate and is representative of two experiments. Error bars, s.e.m. nd, not detected.
formation, we administered the cholinergic agonist carbamylcholine (CCh) to stimulate GC secretion and imaged the small intestine by two-photon microscopy. After CCh administration, the frequency of T cells after co-culture with LP-DCs (Fig. 4f) increased compared to LP-DCs from SPF housed mice. CD103− LP-DCs presented luminal antigen significantly better than CD103+ LP-DCs from germ-free mice. Two-photon imaging of intestines in M1KO mice. M1KO mice lacked antigen delivery in germ-free mice, indicating that GAPs are a source of luminal antigen for CD103+ LP-DCs. Two-photon imaging revealed a concomitant loss of GAPs, with the epithelium forming a tight barrier to luminal dextran (Fig. 4j and Supplementary Fig. 9d). Moreover, LP-DC populations isolated from M1KO mice failed to stimulate OTI T cells above background levels (Fig. 4k and Supplementary Fig. 9e). The lack of antigen presentation by CD103+ LP-DCs from M1KO mice was not due to an intrinsic defect, because these LP-DCs were capable of inducing robust T-cell proliferation when exogenous Ova was added to the T-cell cultures (Supplementary Fig. 9f). Taken together, these findings indicate that GAPs are a major mechanism for delivering luminal antigens to LP-DCs in the steady state.

Understanding how the balance between tolerance and immunity is achieved at the intestinal mucosa is crucial for oral vaccine development and the treatment of chronic intestinal inflammatory diseases. This study identifies GAPs as a mechanism by which CD103+ LP-DCs can acquire innocuous antigens from the intestinal lumen in the steady state. How preferential antigen delivery is achieved is the focus of ongoing studies. Biased delivery could be a natural consequence of CD103+ LP-DC sampling the epithelium more actively or perhaps being recruited selectively by chemotactants released by or near GCs. GC deficiency or dysfunction in mice and humans has been linked to the development of intestinal inflammation23–27. Although this association has been attributed to the loss of mucins and other biologically active GC products, our findings suggest that GCs could play a key role in promoting intestinal immune homeostasis by delivering luminal antigen to tolerogenic LP-DCs.

**METHODS SUMMARY**

**Mice and human specimens.** All mice were of a C57BL/6 background unless otherwise specified. Healthy human jejunal sections were obtained from bariatric surgeries, placed in PBS containing dextran for 1 h before imaging. Procedures and protocols were carried out in accordance with the institutional review board at Washington University School of Medicine.
Two-photon microscopy. The intravital imaging preparation used in this study is similar to previously described methods with the following differences: imaging is performed with the tissue within the peritoneal cavity, faecal material is not scraped from the mucosal surface and in some experiments atropine (1 mg kg\(^{-1}\)) was injected subcutaneously to dampen peristaltic movement of the small intestine. At this dose atropine did not affect the formation of TEDs or the detection of GAPs. Model fluorescent antigens, dextran (2–5 mg), Ova (2 mg), BSA (2 mg) and FluoSpheres (1 ml undiluted) (all from Invitrogen) were injected into the intestinal lumen approximately 2h before imaging. Human resection specimens were incubated in 10 µg ml\(^{-1}\) of dextran at room temperature for 1h before imaging.

Full Methods and any associated references are available in the online version of the paper at www.nature.com/nature.
METHODS

Mice and human specimens. C57BL/6 mice, Rag-/- mice, Balb/c mice, IL-10/-/- knockout mice on the C3H/HeJ/Bir,129 background25, OTI T cell receptor transgenic mice2, CX3CR1-GFP knock-in mice28, Math1fl/fl mice22 and Villin Cre transgenic mice27 were purchased from The Jackson Laboratory. CD11c-YFP transgenic mice28 were a gift from M. Nussenzweig and LysM-GFP mice31 were a gift from K. Ley. Gnotobiotic mice were obtained from the Washington University Digestive Disease Research Core Center murine models core. Animals, other than gnotobiotic mice, were housed in a specific-pathogen-free facility and fed routine chow diet. Animals were 8 to 16 weeks of age at the time of analysis. Intestines from mice receiving 230 rads of gamma irradiation and killed 6 h later served as positive controls for apoptosis markers. All mice were on a C57BL/6 background unless otherwise specified. Healthy human jejunal sections were obtained from bariatric surgeries, placed in PBS containing dextran for 1 h before imaging. Procedures and protocols were carried out in accordance with the institutional review board at Washington University School of Medicine.

Intravital two-photon microscopy. The intravital imaging preparation used in this study is similar to previously described methods10,15 with the following differences: the intestine is not exteriorized during imaging, faecal material is not scraped from the mucosal surface and in some experiments atropine (1 mg kg-1) was injected subcutaneously to dampen peristaltic movement of the small intestine. Mice were anaesthetized with isoflurane and a small vertical incision was made in the abdominal wall to expose the peritoneal cavity and contents. The intestine is secured to the bottom of a glass coverslip on the upper chamber plate using a thin ring of Vetbond tissue adhesive (3M). Because the coverslip sits directly over the incision in the mouse abdomen the tissue remains in the peritoneal cavity for imaging. No additional manipulations were performed to image from the serosal surface. The imaging chamber was maintained at 37 °C using a dual-channel heating system (Warner Instruments). To image from the luminal surface, a small longitudinal incision was made in the intestine taking care to avoid large blood vessels. Model fluorescent antigens, dextran (2–5 mg), Ova (2 mg), BSA (2 mg) and FluoroSpheres (1 ml undiluted) (all from Invitrogen) were injected into the intestinal lumen ~2 h before imaging. Human resection specimens were incubated in 10 µg ml-1 of dextran at room temperature for 1 h before imaging. Time-lapse imaging was performed with a custom-built two-photon microscope running ImageWarp acquisition software (A&B Software). For time-lapse imaging, we averaged 15 video-rate frames (0.5 s per slice) during the acquisition. Each plane represents an image of 220 × 240 µm in the x and y dimensions, 21 to 31 sequential z-steps (2.5 µm each) were acquired to form a z-stack. In our experiments epithelial integrity was assessed by dextran and DAPI staining, which in healthy tissue, demarcates the luminal surface of the epithelium (aside from GAPs) and shows an ordered arrangement of DAPI-stained nuclei, respectively. In some experiments mice were given 108 Salmonella typhimurium strain 3716 orally 24–72 h before imaging. Flow cytometry, immunohistochemistry and confocal microscopy. Flow cytometry and the staining of intestine sections were performed as previously described4,28. Reagents used for flow cytometry include anti-CD11c, anti-CD45, anti-CD4, anti-MHCII, anti-CD3, anti-CD8a, anti-PDCA-1 (all from eBioscience), 7-aminoactinomycin D (7-AAD), anti-CD103 and anti-B20 (both from BD Biosciences). Data was acquired with a FACScan cytometer (BD Biosciences) retrofitted with additional lasers. Data acquisition was performed using CellQuest (BD Biosciences) and Rainbow (Cytek) or FlowJo software (Tree Star). Data analysis was performed on a Macintosh computer running FlowJo software.

To immobilize the lysine-fixable fluorescent dextran, intestinal sections were treated with 2% paraformaldehyde immediately after two-photon imaging. Reagents used for immunohistochemistry include 4’,6-diamidino-2-phenylindole (DAPI; Sigma-Aldrich), anti-CD11c, anti-CD103, anti-e cadherin (all from BD Biosciences), anti-cytokeratin 18 and anti-MUC2 (both from abcam), anti-GFP (MBL International), anti-cleaved cytokeratin 18 (Enzo Life Sciences), and anti-cleaved caspase 3 (Cell Signaling Technology). The terminal deoxynucleotidyl transferase-mediated dUTP nick end labelling (TUNEL) assay (Trevigen) was performed as per the manufacturers recommendations. Confocal microscopy was performed using a Zeiss LSM510 Meta laser scanning confocal microscope (Carl Zeiss) equipped with a ×63, 1.4 numerical aperture Zeiss Plan Apochromat oil objective. Images were obtained using the Zeiss LSM510 software.

T-cell proliferation assays. Mice were anaesthetized and the small intestine injected intraluminally with 2 mg of Ova (Sigma-Aldrich) dissolved in phosphate buffered saline (PBS), or PBS alone (controls) as in the two-photon imaging experiments. In some experiments mice received 3 µg of carbamylcholine (Sigma-Aldrich) subcutaneously 20 min after the administration of luminal Ova. Two hours after the administration of Ova, cell populations were isolated from the intestinal lamina propria as previously described22 and sorted by flow cytometry into total LP-DC populations (7-AAD-, CD45+, MHCI+, CD11c+), CD103+ DC populations (7-AAD-, CD45+, MHCI+, CD11c+ CD103+), or CD103− DC populations (7-AAD-, CD45+, MHCI+, CD11c+ CD103−). Sorted DC populations were cultured with sorted CFDA (Invitrogen)-labelled OTI splenic T cells at a ratio of 1:10 DC to T cells. As a positive control, 20 µg of Ova was added to cultures of DC populations isolated from mice receiving luminal PBS, unless otherwise stated. After 3 days cultures were evaluated for CFDA dilution and the number of T cells by flow cytometry and cell counting.

Real-time PCR. RNA isolation, cDNA synthesis, standard curve construction and real time PCR were performed as previously described22. The following primers were used: 18S, forward, 5’-CGGCTACCATCCTAAGGAA-3’ and reverse, 5’-GCTGGAATTACCGCGGCT-3’; cytokeratin 18, forward, 5’-CAGCCACGGCTCTATGACAGG-3’ and reverse, 5’-CTTTCTCGGTCTGATTCCAC-3’. Small intestine epithelial cell populations were stained with cytokeratin 18 and AlexaFluor 488 (Invitrogen)-labelled lectin from Ulex europaeus (UEA-I; Sigma-Aldrich). Goblet cell populations were isolated by flow cytometric sorting as UEA-1+ cytokeratin 18+ cells.

Statistical analysis. Data analysis using Student’s t test or a one-way ANOVA with a Tukey’s post test were performed using GraphPad Prism (GraphPad Software).