Characterisation of proguanylin expressing cells in the intestine – evidence for constitutive luminal secretion

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Guanylin, a peptide implicated in regulation of intestinal fluid secretion, is expressed in the mucosa, but the exact cellular origin remains controversial. In a new transgenic mouse model fluorescent reporter protein expression driven by the proguanylin promoter was observed throughout the small intestine and colon in goblet and Paneth(-like) cells and, except in duodenum, in mature enterocytes. In Ussing chamber experiments employing both human and mouse intestinal tissue, proguanylin was released predominantly in the luminal direction. Measurements of proguanylin expression and secretion in cell lines and organoids indicated that secretion is largely constitutive and requires ER to Golgi transport but was not acutely regulated by salt or other stimuli. Using a newly-developed proguanylin assay, we found plasma levels to be raised in humans after total gastrectomy or intestinal transplantation, but largely unresponsive to nutrient ingestion. By LC-MS/MS we identified processed forms in tissue and luminal extracts, but in plasma we only detected full-length proguanylin. Our transgenic approach provides information about the cellular origins of proguanylin, complementing previous immunohistochemical and in-situ hybridisation results. The identification of processed forms of proguanylin in the intestinal lumen but not in plasma supports the notion that the primary site of action is the gut itself.

Guanylin is a small peptide produced in the intestinal mucosa from the \textit{Guca2a} gene which has established roles in intestinal fluid homeostasis and maintenance of gut physiology. Together with the related peptide uroguanylin (encoded by the \textit{Guca2b} gene), and heat-stable enterotoxin STA, guanylin activates the Guanylate Cyclase C receptor (GC-C), which is encoded by the \textit{Gucy2c}-gene and expressed along the gastrointestinal (GI) tract. Unusually for a gut peptide/receptor combination, however, the peptide appears to access its receptor from the luminal direction\textsuperscript{2–4}. GC-C activation catalyses the generation of cGMP, stimulating transepithelial chloride flux, inhibition of sodium proton exchangers\textsuperscript{5}, and increase in water movement into the intestinal lumen\textsuperscript{6}. Overstimulation or gain of function mutations in GC-C lead to altered GI transit diarrhoea, and loss of function mutations have been associated with meconium ileus in infants\textsuperscript{7–9}.

The reported active 15 amino acid bioactive peptide guanylin derives from the C-terminus of a longer pre-proguanylin peptide, which comprises a signal sequence (residues 1–21) and proguanylin (residues 22–116 in the mouse sequence, 22–115 in the human). A number of reports have suggested that proguanylin is the predominant secreted form of the peptide, but the enzymatic pathway underlying proguanylin processing to generate...
bioactive guanylin remains unclear. Previous studies aimed to address the potential cellular origin of guanylin. Early reports suggested, based on the hormonal function proposed for guanylin peptides, that enteroendocrine cells were the likely source of these peptides, including enterochromaffin cells and somatostatin secreting D-cells. However, other cell types were also reported as sources of guanylin, including goblet cells, Paneth cells, colonic Paneth-like cells, tuft cells and enterocytes, but with some discrepancies between techniques and studies. Stimuli underlying proguanylin secretion remain poorly characterised, with the strongest body of evidence supporting a link to salt consumption, corresponding with an observed increase in Guca2a expression in the intestines of rats fed a high salt diet.

The GC-C signalling axis has been reported to play a role in crypt-villus epithelial proliferation and to act as a tumour suppressor gene. Reduced GC-C receptor signalling was linked to hyperplasia of crypts and villi along the gastrointestinal tract and was associated with increased susceptibility to tumorigenesis. In a cohort of patients with stage I-III colorectal cancer, guanylin mRNA and peptides were lost and/or significantly lower in cancerous tissues compared to healthy adjacent tissues in >85% of cases, and targeting the GC-C pathway at the early stages of colorectal cancer has been proposed as a candidate therapeutic strategy.

Circulating proguanylin levels were lower in people with obesity and raised following Roux-en-Y gastric bypass surgery, suggesting potential links to metabolism or food intake. Corresponding with these findings, mice fed a high fat diet had lower expression and peptide levels of guanylin and forced re-expression of guanylin reduced the rate of obesity associated colorectal cancer. Gucy2c-deficient mice are hyperphagic and heavier compared with wild-type mice. In wild-type mice, food intake was reduced following intravenous administration of proguanylin, but not guanylin, and increased following treatment with a proguanylin antisense and it was speculated local processing in the hypothalamus releases active uroguanylin, suggesting a central role for GC-C signalling. However, another group found that systemic administration of proguanylin-derived peptides modulated food ingestion or glucose homeostasis in mice, despite targeting the same GC-C receptor. GC-C activation has, however, been shown to stimulate secretion of the anorectic peptide glucagon-like peptide-1 (GLP-1) from enteroendocrine cells in the GI tract.

Despite the multiple proposed physiological roles of guanylin peptides and increased interest in their use for treating irritable bowel syndrome or colorectal cancer, the cellular origins of guanylin and the mechanisms underlying its secretion are poorly understood. To address these questions, we generated a transgenic mouse model in which the Guca2a promoter drives the expression of the yellow fluorescent protein Venus. This model was utilised alongside the use of mass spectrometry and a newly-established monoclonal antibody-based immunoassay to measure proguanylin and proguanylin-derived peptides in human plasma, tissues and cell supernatants isolated from preclinical experimental systems.

Results

Proguanylin levels in human plasma. By LC-MS/MS, we detected high levels of proguanylin (22–115) in human plasma, but were not able to detect shorter proguanylin-derived peptides containing the C-terminal active sequence (Fig. 1a). Therefore, this suggests that proguanylin is mostly circulating as the intact proform which we could detect using a newly developed proguanylin immunoassay. Fasting plasma proguanylin levels in healthy humans were 15.2 ± 2.7 ng/mL (mean ± sd), and followed a normal distribution (Fig. 1b). Plasma levels fell slightly after a 75 g or 50 g oral glucose tolerance test but there was no significant effect of a mixed liquid meal (Fig. 1c,d). Lean patients who had undergone total gastrectomy with Roux-en-Y reconstruction for gastric cancer had elevated fasting proguanylin concentrations (26.5 ± 6.9 ng/mL), but no significant change after oral glucose ingestion (Fig. 1d). Although a number of the gastrectomy patients carried risk alleles in CDH1, neither pre- nor post-operative proguanylin levels appeared influenced by the CDH1 genotype. Fasting proguanylin concentrations were elevated in a proportion of subjects who had undergone small bowel transplantation (Fig. 1e). One transplant subject had blood samples collected after removal of the small and large intestine and again after bowel transplantation, revealing very low proguanylin after intestinal removal (~0.2 ng/mL, close to the detection limit of the assay) but a normal level (19.4 ng/mL) after transplantation. Collectively, these results provide strong support for the idea that the gut is the predominant source of proguanylin in human blood, which circulates at relatively high levels compared to other intestinally derived hormones, such as GLP-1.

Identification of proguanylin-derived peptides by LC-MS/MS. Analysis of tissue homogenates from sequential sections along the mouse small intestine, colon and rectum by LC-MS/MS did not detect proguanylin itself, likely because other peptides eluted with similar retention times, but identified several proguanylin-derived peptides (Fig. 2a). Whilst these were barely detectable in homogenates from the proximal duodenum, levels increased along the GI tract before falling again in the rectum. Interestingly, the canonical sequence for the reported active 15 amino acid peptide (position m-102–116) was present at extremely low levels in tissue homogenates and detectable only in jejunum and colon, whereas a form containing one additional amino acid at the N-terminus was readily detected in all samples. Similar analysis of the apical and basolateral solutions from murine jejunum and colon mounted in Ussing chambers (after 120 min each from one colon and one jejunum) identified full length proguanylin (22–116) in both solutions, although at much higher concentrations in the apical compartment (Fig. 2b–d). In the apical compartments of both tissues we also identified several proguanylin derived peptides such as 60–88, 101–116, 103–116 and 104–116 (Fig. 2b,e). The same peptides were detected in the basolateral compartments, but at much lower concentrations, mirroring the lower levels of proguanylin. Immunoassay measurements of proguanylin from human colonic tissues mounted in Ussing chambers also revealed much higher levels released into apical than basolateral compartments (Fig. 2f). Forskolin + IBMX had only a minor effect on proguanylin secretion, although this was a strong stimulus of GLP-1 release into the basolateral compartment, measured in parallel.

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Mapping guanylin-expressing cells along the murine gut axis. We were unable to detect specific immunohistochemical staining with the monoclonal antibodies isolated during the development of the new immunoassay in human intestinal tissue slices or primary mixed epithelial cultures (data not shown). Instead,

Figure 1. Proguanylin detection in plasma. (a) LC-MS/MS evidence of full length proguanylin in human plasma. Right panel shows the total ion chromatogram for an m/z range corresponding to the proguanylin in a charge state of 8 (±0.01 around each isotopic variants), asterisk marks the peak corresponding to proguanylin, left panel is the m/z distribution at retention time corresponding to the peak marked by an asterisk. (b) Data distribution and normality for proguanylin concentration in plasma of fasted healthy volunteers (n = 51). Skewness = 0.37; kurtosis = -0.08; P = 0.69 when evaluated for Shapiro-Wilk test. (c) Effect of liquid meal (n = 28) or 75 g OGTT (n = 12) on plasma proguanylin levels in healthy volunteers after 60 minutes. (d) Effect of 50 g glucose ingestion on plasma proguanylin levels of healthy controls and participants with total gastrectomy after 60 minutes (n = 17, controls: n = 11). Patients with a mutation in the CDH1 are indicated in orange. (e) Serum levels of proguanylin in fasting healthy volunteers and patients following intestinal transplantation. *p < 0.05 and ***p < 0.001 using a paired t-test when comparing intra individual values across two time points or unpaired t-tests when comparing two groups of individuals.
Figure 2. Proguanylin production and secretion in the gut. (a) Measurement of GUCA2A peptides along the mouse intestine from duodenum to rectum (1 to 7 are small intestine regions every 5 cm from duodenum to ileum, and 8 to 10 are proximal colon, distal colon and rectum respectively) analysed by LC-MS/MS. Data are mean ± SEM, n = 4. (b) Apical (dark colours) and basolateral (light colours) secretion of guanylin peptides from mouse colonic (red) and jejunum (blue) tissues. (c,d) Evidence of full length proguanylin (marked by asterisk) in mouse jejunum apical compartment (c) and absence in the basolateral compartment (d) showing chromatogram for the m/z range corresponding to the [M + 8 H]8+ charge state of proguanylin (left panel) and the m/z distribution at the expected retention time. (e) PEAKS peptidomics search results showing peptides (of less than 65 amino acids) matching the proguanylin sequence in the apical side of the Ussing chamber experiment using murine colonic tissue. (f) Apical (blue) and basolateral (green) secretion of proguanylin and basolateral secretion of GLP1 from human colonic tissues using Ussing chambers from 6 tissue preparations from 4 different donors. Data are represented as mean ± SD * and **p < 0.05 using a paired t-test adjusting for multiple testing, comparing apical vs basolateral secretion and unstimulated vs F/I stimulated.
a transgenic mouse model was designed to identify which cell types produce proguanylin, by expression of the fluorescent protein Venus under the control of the guanylin promoter. The line was derived from one founder in which a BAC construct containing the Venus sequence in place of the coding sequence of Guca2a was incorporated into the mouse genome (Fig. 3a). Offspring showed fluorescence in the small and large intestine, enabling the use of fluorescence-assisted cell sorting (FACS) to separate fluorescent (positive) from non-fluorescent (negative) cells separately from the duodenum, jejunum, ileum and colon. RT-qPCR analysis of sorted positive and negative cell populations showed enrichment of Guca2a and Venus expression in the fluorescent populations, thus validating the model (Fig. 3b,c). Interestingly, the fluorescent cell populations were also enriched for the expression of uroguanylin, Guca2b (Fig. 3d).

The proportion of Venus fluorescent cells was highly variable between different intestinal tissues. Using flow cytometry to count the number of cells exhibiting fluorescence above a fixed threshold, we estimated that the

Figure 3. Generation and characterisation of the guanylin-Venus mouse. (a) Schematic depicting the insert of BAC RP23-104G3 and replacement of the Guca2a coding region from exons 1–3 with the yellow fluorescent protein Venus. Lengths (in kb) of the murine genomic sequence 3′ and 5′ of the Guca2a coding region in the BAC are indicated. (b–d) RT-qPCR analysis of FACS sorted positive (green) and negative (blue) populations for each tissue for Guca2a (b), Venus (c) and Guca2b (d) n = 3 each; negative and positive populations were compared using paired t-test *P < 0.05 **P < 0.01. (e–h) duodenum (e), jejunum (f), ileum (g) and colon (h) of Guca2a-Venus mice immunostained for GFP (green) and DNA (Hoechst; purple).
Venus-positive cell frequency was ~3% of the cellular epithelial cell population in the duodenum, ~50% in the jejunum, 30% in the ileum and 20% in the colon. Immunostaining of tissue sections confirmed this disparity in guanylin-expressing cell distribution along the intestinal axis (Fig. 3e–h). Duodenum contained only scarce Venus-positive cells that were scattered along the crypt-villus axis (Fig. 3e). In jejunum and ileum, Venus-positive cells appeared to separate into two distinct populations, one located at the tops of the villi and another in the crypts. In colon, the fluorescent marker was largely restricted to the surface epithelium at the top of the crypts (Fig. 2h). These staining patterns suggest that Guca2a is expressed in mature enterocytes in the jejunum, ileum and colon but not duodenum, as well as in some other epithelial cell types.

Identification of guanylin-expressing cells by transcriptomic profiling. To determine the identity of the fluorescent cells in the small and large intestine of Guanylin-Venus mice, we performed RNA sequencing analysis of FACS-sorted Venus-positive and -negative cell populations from different regions of the GI tract from three mice each. Confirming the RT-qPCR data, both Guca2a and Guac2b were enriched in fluorescent cell populations from all four intestinal regions, and surprisingly, we also detected higher expression of the receptor GC-C (Gucy2c) in fluorescent cells from the jejunum, ileum and colon (Fig. 4d), although not duodenum. By principal component analysis (PCA) of the RNAseq results, the samples were found to cluster both by the tissue of origin and by whether they were fluorescent or non-fluorescent, indicating that Guca2a expressing cells are transcriptomically different from negative cells, and that they vary along the GI tract (Fig. 4a). Analysis of the top 500 most differentially expressed genes across all groups revealed clear separations between Guca2a positive and negative cells, and between Guca2a-expressing cells in the duodenum compared with their counterparts in the jejunum and ileum (Fig. 4b). Analysis of the numbers of Guca2a-cell enriched genes in different intestinal regions also revealed strong similarity between Guca2a-positive cells from the duodenum and jejunum, compared with those from the colon (Fig. 4c). 515 genes were enriched in Guca2a-positive cells compared to negative cells from the same region, in all four tissues, suggesting at least a degree of overlap between the Guca2a cell populations from different regions of the GI tract.

To identify the cell types expressing Guca2a, we examined the expression of known marker genes for different epithelial cell populations: goblet cells, enterocytes, Paneth cells, tuft cells, enteroendocrine cells and proliferative cells (Fig. 4d and Supplementary Fig. 1). Goblet cell markers (Muc2, Tff3, Spik4) were enriched in positive cells across all four tissues. Enterocyte cell markers (Adh6a, Khk, Alpi, Slc16a3, Batf2) were enriched in guanylin-expressing cells, particularly in the jejunum and ileum, and colonic marker cells (Car2, Slc2a1) were also enriched in guanylin-expressing cells. Paneth cell markers (Lyz1 and Defa22) were enriched in positive cells from the small intestine but not colon, consistent with the fact that Paneth cells are not present in the colon, but Paneth-like cells markers (Kit, Egf) were enriched in positive cells of the colon.35 Markers of Tuft cells (Dclk1, Lrmp, Trpm5), enteroendocrine cells (including Gcg, Gip, Ppy, Cck, Chga, Tph1) and proliferative cells (Lgr5, Mki67, Sox4, Stmn1, Foxm1) were relatively depleted in Guca2a-positive compared with negative cells. Overall, this transcriptomic analysis suggests that Guca2a is expressed by several different cell populations in the intestinal epithelium: goblet and Paneth/Paneth-like cells in all 4 intestinal tissues, and enterocytes/colonocytes in tissues distal to the duodenum. Notably, Guca2a labelling cells did not exhibit expression patterns typical of enterocytes in the duodenum, or of enteroendocrine cells, tuft cells or proliferative cells.

Identification of guanylin-expressing cells by immunostaining. We further examined the cellular identity of Guca2a expressing cells by double immunostaining of tissue slices from Guca2a-Venus mice and by RT-qPCR of FACS-purified fluorescent cells. To address the overlap with Paneth cells, we examined expression of Lyz1, and immunostaining for lysozyme (Fig. 5). Confirming the RNAseq data, Lyz1 was enriched in Guca2a-fluorescent cells from different regions of the small intestine (Fig. 5a). Dual staining for lysozyme and Venus (using an anti-GFP antibody) showed double labelled cells at the crypt bases in the duodenum, jejunum and ileum (Fig. 5d–f), with the majority of lysozyme-positive cells being GFP positive: 67% in the duodenum, 91% in the jejunum and 86% in the ileum (Fig. 5b). Paneth cells represented ~50% of all Venus positive cells in the duodenum, but less than 15% in the jejunum and ileum, reflecting the higher overall frequencies of Guca2a-positive cells in the jejunum and ileum (Fig. 5c).

To validate and quantify the localisation of Guca2a in goblet cells, we measured expression of the markers Muc2 by qPCR, and performed double staining for Muc2 and GFP in Guca2a-Venus tissues. Expression of Muc2 was enriched in Venus-positive compared with negative cells (Fig. 6a), with the most marked differential expression evident in the duodenum. Many cells exhibited double-positive staining for Muc2 and GFP in all four tissues (Fig. 6c–g), with ~50% of all goblet cells staining positive for GFP in the duodenum, jejunum and ileum compared with ~20% in the colon (Fig. 6b). In the duodenum, 53% of all GFP positive cells expressed co-staining for Muc2, whereas Muc2-positive goblet cells represented only 9%, 16% and 19% of the total GFP-positive cell populations in jejunum, ileum and colon, respectively, again reflecting the higher overall frequencies of fluorescent cells in these tissues (Fig. 6c).

Altogether, these results show that the Guca2a positive cell population in the duodenum is largely made up of Paneth and goblet cells, each representing about half of all Guca2a cells in this region. In the jejunum and ileum, Paneth and goblet cells together only explain a third of all Guca2a positive population, the remainder being mature enterocytes, as further supported by the expression of the different intestinal cell-type markers by qPCR (Fig. 7). In the colon, only ~20% of Guca2a positive cells stained positive for Muc2, and the remainder appeared to be dominated by mature colonocytes.

Regulation of guanylin secretion. We next investigated whether proguanylin release could be modulated by a panel of candidate stimuli known to stimulate different epithelial cell types. In particular, we tested agents that could potentially target a variety of cell types, including PMA to activate PKC and forskolin/IBMX to
Figure 4. Transcriptome analysis of Guca2a-expressing cells. (a) Principal-Component analysis of the top 500 most variable genes between positive and negative cells from the four regions of the GI tract. Non-coding and mitochondrial genes were excluded. In purple are the Venus-positive cell subpopulations and in grey are the negative population. Tissue of origin is represented by shape (circle: duodenum; triangle: jejunum; square: ileum; cross: colon). (b) Heatmap showing the top 500 most differentially expressed genes between positive and negative cells across all 4 regions. Values are log10 normalised read counts. Genes and samples are grouped via hierarchical clustering based on Euclidean distance and complete linkage. Colon = red; duodenum = yellow; jejunum = light blue; ileum = dark blue. Purple = positive cells; grey = negative cells (c) Venn diagram summarising the overlap of positively-enriched genes in the guanylin-positive cell subpopulation for each region. Positively enriched genes were selected with p-adjusted < 0.05. (d) Heatmap of selected cell markers (rows) between positive and negative cells (columns sorted by cell type and tissue of origin). Values are log fold change compared to mean expression across all samples. Colour coding as for (b).
increase cAMP concentrations; stimuli known to enhance secretion from enteroendocrine cells via cell surface receptors, including bile acids, peptones and bombesin; and stimuli previously reported to modulate guanylin release, including elevated salt concentrations and carbachol. Secretion was first measured from primary human colonic crypt cultures containing a variety of cell types, using LC-MS/MS to provide parallel quantification of a readily detectable proguanylin fragment 22–49 and GLP-1. After 1 h of incubation, guanylin secretion was slightly enhanced by the PKC activator PMA, but was not altered by bombesin, forskolin/IBMX or an agonist of the Gs-protein coupled bile acid receptor GPBAR1, which robustly elevated GLP-1 (Fig. 8a,b).

**Figure 5.** Guanylin expression in Paneth cells. (a) Venus-positive (green) and negative (blue) subpopulations of isolated intestinal cells from the Guca2a-Venus mice were analysed by RT-qPCR for Lyz1 expression. n = 2–3 each; positive and negative populations were compared using a paired t-test *P < 0.05. (b) Proportion of Lyz1 only positive cells (blue) and Venus and Lyz1 double positive (green) compared to total number of Lyz1 positive cells. (c) Proportion of Venus only positive cells (yellow) and double positive to Venus and Lyz1 (green) compared to total number of Venus cells. (d–f) Guca2a-Venus mice sections immunostained for Venus (GFP, green), lysozyme (red) and DNA (Hoechst; blue) from duodenum (d), jejunum (e) and ileum (f).
We next used differentiated human duodenal and colonic organoids as a model of mixed epithelial cell types, to measure proguanylin secretion by immunoassay over 24 h in response to a panel of stimuli including sodium chloride, nutrients, bile acids and neurotransmitters. As expected, undifferentiated organoids had very low levels of GUCA2A expression and did not secrete detectable concentrations of proguanylin, whereas GUCA2A expression and proguanylin secretion increased following organoid differentiation (Fig. 8e,f and Supplementary Fig. 2). Differentiated organoids secreted proguanylin into the medium, and in the duodenum (but not colon) this was marginally stimulated by 50 mM NaCl but not 100 mM mannitol (Fig. 8e,f). No other condition modulated proguanylin secretion in either duodenal and colonic organoids, except for an inhibition seen with PMA, contrasting with the slight increase observed in primary cultures after more acute (1 h) stimulation.

We then assessed candidate mechanisms regulating proguanylin secretion using a human enterocyte cell line model, TC7, a subclone of the Caco-2 cell line. This more homogenous culture system also allowed us to monitor changes in the expression of proguanylin, either after acute (4 h) or more prolonged (24 h) stimulation. Addition of 10 or 50 mM NaCl, or 100 mM mannitol as an osmolarity control, resulted in a reduction in proguanylin secretion into the media after 4 hours, with the higher NaCl and mannitol treated cells also exhibiting a correspondingly lower expression of GUCA2A relative to the housekeeper gene ACTB (Fig. 8g,h). After 24 h incubation, however, proguanylin levels in salt and mannitol-treated wells were similar to those in control wells, and GUCA2A expression was elevated (Fig. 8i,j). We also measured proguanylin release and biosynthesis in TC7 cells
in response to known enteroendocrine stimuli: bombesin, carbachol, peptones, taurodeoxycholic acid, PMA and forskolin/IBMX. After 4 h incubation, the only stimulus that significantly increased the release of proguanylin was 10μM forskolin/IBMX, corresponding with a parallel increase in GUCA2A expression (Fig. 8g,h). After 24 h,
Figure 8. Proguanylin secretion in vitro. (a,b) Effect of different stimuli on proguanylin(22–49) (a) and GLP-1 (b) secretion from primary human colonic cultures, measured by LC-MS/MS after 1 h incubation in the absence (basal) or presence of bombesin (100 nM), forskolin/IBMX (10 μM each), PMA (1 μM) or GPBAR-A (3 μM). (c,d) Quantities of proguanylin in TC7 cell supernatants (c) and lysates (d) at different time points in standard media (blue) with forskolin/IBMX (10 μM each, red) or brefeldin A (5 μg/mL, green). Data are mean ± SD, n = 8. (e,f) Effects of different stimuli on proguanylin secretion, measured by immunoassay, from human duodenal organoids (e) and human colonic organoids (f) Cultures were incubated with 50 mM NaCl, 100 mM mannitol, 100 nM bombesin, 1 mM carbachol, 5 mg/mL peptone, forskolin/IBMX (10 μM each), 1 μM PMA, 100 μM taurodeoxycholic acid (TDCA) or 100 nM aldosterone for 24 h. Data were normalised to total protein content and to the mean basal value. Data are mean ± SD, n = 6–18. (g–j) Secretion and expression of proguanylin in TC7 cells in response to different stimuli. TC7 cultures were incubated with stimuli as in (e). Supernatant proguanylin levels, measured by immunoassay (g,i) as well as transcript levels (h,j) were measured
proguyanlin release and transcript levels were also increased by peptones, and a small reduction in secretion was found in PMA-treated wells (Fig. 8i,j). At this later timepoint with forskolin/IBMX, the GUCA2A transcript levels were not significantly different from controls (Fig. 8).

As we had inconsistent results in these different culture systems, with only forskolin/IBMX apparently increasing proguyanlin release in TC7 cells, we quantified proguyanlin in cell supernatants and cell extracts at different time points over 24 hours. After 4 h, the amount of proguyanlin in the supernatant, measured by immunoassay, was similar to that found in cell lysates, and after longer incubations the supernatant concentrations increased approximately linearly, whereas the lysate content stayed relatively constant (Fig. 8c,d). These results suggest that the cells turn over their entire proguyanlin content in approximately 4 hours. In cells treated with forskolin/IBMX, slightly higher supernatant proguyanlin concentrations were detected after 8 and 24 hours, corresponding with a small increase in the lysate content at 8 h. The slightly higher secretory rate in forskolin/IBMX might therefore reflect a stimulation by cAMP of proguyanlin biosynthesis. When TC7 cells were treated for 4 or 8 hours with brefeldin A, an inhibitor of ER/Golgi transport, proguyanlin secretion was largely abolished, although lysate concentrations remained unchanged (Fig. 8c,d), suggesting that ER/Golgi transport is required for proguyanlin secretion, and that there is very little storage of proguyanlin in post-Golgi compartments.

Discussion

In this study, we applied new tools to investigate the cellular origin and secretion of guanylin peptides, including a new immunoassay for human proguyanlin, LC-MS/MS analysis of proguyanlin-derived peptides, and a transgenic mouse model expressing Venus under the control of the proguyanlin promoter that enabled the identification, isolation and transcriptomic analysis of proguyanlin-expressing cells. We found that proguyanlin is constitutively secreted into the gut lumen, predominantly by mature enterocytes in the distal small and the large intestine, but in the duodenum only by Goblet and Paneth cells.

Guanylin and uroguanylin have been reported to exhibit differential expression along the GI tract, with the mRNA of these two genes mainly expressed in the proximal small intestine and the colon36,38,39. Using the guanylin-Venus transgenic mouse, we observed guanylin-dependent fluorescence in a variety of intestinal cell types and found that the purified fluorescent cell populations were enriched for both Guca2a and Guca2b. The pattern of Venus fluorescence in the duodenum was atypical compared with the rest of the GI tract, as it was largely restricted to Paneth and goblet cells. In the remainder of the gut, these cell types were also fluorescent, but we additionally observed expression in “mature” enterocytes at the small intestinal villus tips and the colonic surface epithelium. These results are in agreement with a recent RNA-seq study16, which found a similar distribution of guanylin and uroguanylin along the murine rostro-caudal and crypt-villus intestinal axes. Unlike this murine RNA-seq analysis, we did not find tuft cell markers in guanylin-Venus cells, but our results are consistent with a recent study16 that failed to detect GUCA2A in human and rat tuft cells97, and another report suggesting that uroguanylin but not guanylin is expressed in this cell type19. In agreement with the RNA-seq studies16,97, we found no evidence for guanylin expression in enteroendocrine cells, contrasting with reports showing expression in human or rat enteroendocrine cells by immunohistochemistry12,13,28. Whilst these previous studies themselves are inconsistent by identifying different enteroendocrine subpopulations (EC-, D- and L-cells) to express guanylin, possibly due to antibody cross-reactivity, we cannot exclude possible species differences.

LC-MS/MS analysis of human plasma readily detected full-length proguyanlin, consistent with our measurement of high circulating proguyanlin concentrations by immunoassay. Shorter proguyanlin fragments were not detectable by LC-MS/MS in human plasma. In media from murine Ussing chamber studies, we detected full-length proguyanlin together with lower levels of a variety of cleaved forms. Our finding that full-length proguyanlin (22–116/115) is the major released form is consistent with previous reports in humans and rats38,40. Whilst we did not detect the originally described canonical sequence for the 15aa long active guanylin (mouse 102–116/human 101–115)41, we regularly and consistently detected peptides that were one amino acid shorter or longer at the N-terminus, as also reported previously for rats and humans38,40 thus suggesting that the originally isolated guanylin 15-mer likely represents an artefact of the acetic extraction method as previously proposed41. In murine tissue extracts, LC-MS/MS confirmed the previously-observed proximal to distal gradient of proguyanlin-fragments, and identified a variety of proguyanlin cleavage products, some of which matched those found to be secreted in the Ussing chamber studies (mouse 101–116 and 103–116). As we were not able to quantify the relative proportion of cleaved to full-length proguyanlin in the tissue extracts, it is not clear how much proguyanlin processing occurs intracellularly and how much depends on enzymatic cleavage in the gut lumen or bloodstream. Although the enzymes responsible for luminal proguyanlin processing remain unknown, we speculate a role for memprins, which are zinc metalloproteinases found highly expressed in the small intestinal epithelial brush border43, with sequence specificity predicted to cleave proguyanlin to release the 16 amino acid long guanylin peptide, but of course future research will be required to investigate if proguyanlin is a substrate for these metalloendo-peptidases.

In agreement with previous studies on rat tissues24,44, we found that proguyanlin is predominantly secreted into the gut lumen rather than at the basolateral side, in sharp contrast to enteroendocrine hormones such as...
GLP-1. This finding was observed in both mouse and human intestine and is consistent with the localisation of guanylin-expression in goblet and Paneth cells, which are specialised to secrete peptides and mucus in the luminal direction. Proguanylin was also, however, detectable in the basolateral compartment of Ussing chamber experiments, albeit at lower levels than in the luminal chamber, consistent with its detection in human plasma. As it seems unlikely that proguanylin arrives in the circulation following its diffusion/leakage out of the gut lumen across the epithelial layer, there should be an additional secretory pathway in the basolateral direction from a cell type that remains unclear. Our finding that proguanylin was undetectable in the plasma of a patient who temporarily lacked an intestine prior to transplantation, argues against the possibility that circulating proguanylin derives from a non-intestinal source. The observation that bowel transplant patients had raised fasting plasma proguanylin levels is interesting and in line with previous reports of elevated proguanylin expression after Roux-en-Y gastric bypass surgery. The variety of operations performed, ranging from isolated small bowel transplants to multivisceral transplants including liver and pancreas, however, prevents currently a more detailed analysis of what might upregulate proguanylin secretion into the plasma after surgery.

We were unable to identify clear stimuli of guanylin secretion in vitro, although these studies were limited by the cellular models available. TC7 cells did not exhibit strong evidence of regulated proguanylin release, but cellular content had a rapid turnover and secretion was blocked by brefeldin A, arguing for a continuous constitutive vesicular secretory pathway. As TC7 cultures would not have contained goblet and Paneth cells, intestinal organoids and primary intestinal cultures were used as an alternative in the hope they might have contained a mixture of cell types, but these also failed to exhibit robust secretion in response to candidate stimuli. PMA inhibited proguanylin secretion from organoid cultures over a 24 hour incubation, whereas it had no effect on secretion from TC7 cells over a similar timeframe and slightly stimulated secretion from primary cultures during a 1 hour incubation. The underlying explanation for the different responses of the two models is unclear but might reflect the different incubation times and the presence of different cell types in organoids and primary cultures, whereas TC7 cells are of an enterocyte lineage. We were also unable to provide convincing evidence in support of previous reports that guanylin levels are linked to increased sodium levels. Our data are best compatible with the idea that proguanylin secretion from enterocytes occurs constitutively, with limited storage of acutely recruitable vesicular pools, as proguanylin accumulated over time in TC7 supernatants, but not in the cells themselves. Previous reports of carbachol-triggered proguanylin release from intact tissue pieces mounted in Ussing chambers and bethanechol-stimulated release from perfused colonic preparations might reflect degranulation of goblet cells, which are known to respond to cholinergic stimulation. Moro et al. also showed that bombesin-stimulated proguanylin release was inhibited by tetrodotoxin, suggesting the involvement of the enteric nervous system (ENS), possibly explaining the lack of a response seen in the here employed culture systems, as these would lack the ENS.

Proguanylin concentrations in human plasma averaged ~15 ng/ml, approximately 1000-fold higher than fasting plasma levels of gut hormones such as GLP-1, and were largely unaffected by food or glucose ingestion. Taken together with the finding that secretion in vitro was largely unresponsive to a range of candidate stimuli, our data do not support the idea that plasma proguanylin acts as a classical hormonal signal related to nutrient intake. Indeed, although elevated proguanylin levels were reported previously in bariatric patients, consistent with our own findings, administration of proguanylin or guanylin in mouse models did not influence food intake or glucose homeostasis. As GC-C is predominantly expressed in the GI tract, it remains uncertain whether circulating proguanylin has any signalling role.

Proguanylin released into the intestinal lumen is cleaved into smaller peptides, and although we did not detect the classical 15 amino acid guanylin peptide, we consistently found slightly longer and shorter peptides that are likely to exhibit activity on the GC-C receptor. It has been hypothesised previously that guanylin secreted by Paneth cells deep inside the crypts might activate CFTR in neighbouring enterocytes thus sustaining luminal fluid flow from the crypt to villus, whilst guanylin secreted by goblet cells could similarly provide water for full mucin hydration. Guanylin released from the villus tips could potentially target nearby or distal enterocytes, providing a paracrine/autocrine feedback mechanism to ensure adequate epithelial hydration. As full length proguanylin has any signalling role.

Future studies will be required to identify specific release pathways involved in secretion from the different cell types producing proguanylin, the mechanisms and enzymes regulating active guanylin generation in the gut lumen, and the physiological roles (if any) of circulating proguanylin.

Materials and Methods

Unless otherwise stated, all chemicals were supplied by Sigma-Aldrich.

Ethical approvals. Ethical approval for collection of human plasma samples was granted by the following National Research Ethics Committees: East of England – Cambridgeshire and Hertfordshire (ref: 13/EE/0195), East of England – Norfolk (ref: 14/EE/1247); and East of England – Cambridge South (ref: 16/EE/0338). Ethical approval for use of human tissue pieces for cell culture, organoid generation and Ussing chamber studies was granted by the East of England – Cambridge Central Research Ethics Committee (ref: 09/H0308/24); surgical specimens were obtained under this ethical approval from Tissue Bank at Addenbrooke’s Hospital (Cambridge, UK). All human studies were conducted in accordance with the ethical standards of the Helsinki Declaration of 1975. All human participants gave written informed consent, either themselves or through their parent and/or legal guardian if they were under the age of 18 years old, and all methods were performed in accordance with the relevant guidelines and regulations.

Animal procedures during the generation of antibodies were approved by the Babraham Institute Animal Welfare and Ethical Review Body. Animal procedures involving the generation and/or use of Guanylin-Venus
mice were approved by the University of Cambridge Animal Welfare and Ethical Review Body. All animal experiments were performed in accordance with the relevant guidelines and regulations under the UK Home Office project licences 70/8084, 70/7824 and PESOF6065 and were carried out in accordance with the Animals (Scientific Procedures) Act 1986 Amendment Regulations (SI 2012/3039).

**Human blood samples.** Healthy volunteers and patients who had previously undergone prophylactic total gastrectomy were recruited for oral glucose tolerance tests (50 or 75g OGTs) or mixed liquid meal (Ensure plus) and blood was collected just before and 60 minutes after ingestion into lithium-heparin coated tubes and centrifuged 10 min at 4°C at 3500 g, plasma was stored at −80°C until analysis⁴⁷. Other blood samples were obtained from patients having received small bowel transplants and healthy volunteers after an overnight fast and serum samples were stored at −80°C until analysis.

**Transgenic mice generation.** BAC constructs containing the sequence of the yellow fluorescent protein Venus driven by the mouse guanylin promoter were made by the RedEt technique⁴⁸. Briefly the sequence in the 3 exons coding for Guca2a (Guanylin) from the start codon to the stop codon in the murine-based BAC RP23-104G3 (Source Bioscience) was initially replaced by a counter-selection cassette RpsL-neo (GeneBridges) and subsequently by the Venus sequence⁴⁹ using Red/ET recombination technology (GeneBridges). The RpsL/Neo (FSD001/FSD002) or Venus (FSD003/FSD004) sequences were amplified by PCR with Phusion-polymerase adding Guca2a-gene specific 3’ and 5’ sequences (Table 1). Positive recombinants were isolated using appropriate antibiotic selection and characterised by PCR. The identity and correct positioning of the introduced Venus sequence was confirmed by direct sequencing using oligonucleotides FSD045-FSD076 (Table 1) and restriction analysis using the Zral enzyme (NEB) for DNA fingerprinting. The BAC was purified using a Maxi-Prep (Qiagen) and diluted to 1 ng/μL into injection buffer (10 mM Tris-HCl (pH 7.5), 0.1 mM EDTA, 100 mM NaCl, 0.03 mM spermine, and 0.07 mM spermidine). The Central Biomedical Services (University of Cambridge) performed the pronuclear injection of ova derived from C57Bl6/J/CBA F1 crosses, which were implanted into pseudopregnant females. Integration of the transgene was screened by PCR on the pups’ ear clips after extracting the DNA by proteinase K digestion using the primers GFPF1 and GFPR1 (Table 1). Experiments were performed on tissue derived from animals backcrossed into C57Bl6J to varying degree (3–8 generations), with no changes having been observed in the expression profile along the intestinal axis during the backcrossing procedure.

**Immunohistochemistry.** Tissues from different regions of mouse gut were fixed in 4% PFA (Alfa Aesar) at room temperature overnight, dehydrated in sucrose (6h 15% followed by 24h 30%) at 4°C. Tissue was embedded in OCT (VWR) and sectioned (7–10 μm) using a cryostat and mounted on Superfrost Plus glass slides (Thermo Fisher Scientific). Tissues sections were blocked in 10% donkey serum and 1% BSA and permeabilized with 0.05% Tween-20. Slides were stained with primary antibodies diluted in blocking solution over night at room temperature (RT) (Table 2). Slides were washed in PBS and 0.05% Tween-20 and then incubated with secondary antibodies conjugated to Alexa-Fluor 488, 555 or 633 (Invitrogen) and Hoechst nuclear stain (1:1300). Sections were mounted in hydromount (National Diagnostics) and images taken using a TCS SP8 confocal microscope using LAS software (Leica).

**Primary murine intestinal single-cell digestion for fluorescence-activated cell sorting.** Four distinct parts of the gastrointestinal tract corresponding to the duodenum (top 3 cm of the small intestine), jejunum (3 cm in the middle small intestine) and ileum (bottom 3 cm of the small intestine) as well as the whole colon and rectum were treated separately and digested into single cells⁵⁰. Each intestine part was opened longitudinally, rinsed in PBS and the outer muscle layer was removed. Each part was incubated in 10 mL solution containing 15 mM EDTA (or 30 mM for colon) and 1 mM DTT in Ca²⁺/Mg²⁺-free PBS for 7 minutes at RT. Tissue pieces were transferred into 15 mL Ca²⁺/Mg²⁺-free PBS with 10 μM ROCK inhibitor y27632 (Tocris) and shaken by hand for 30 seconds to collect epithelial fragments, and tissue transferred back to the EDTA solution. Incubation in EDTA and shaking was repeated 5 times. Epithelial fragments were kept on ice and after final shaking, centrifuged at 500 g at 4°C and media was removed and replaced with 10 mL Trypsin 0.25% EDTA, 100 mg/mL DNAse for 5 minutes at 37°C. 15 mL of Hanks’ buffered salt solution (HBSS) supplemented with 10% PBS and 5 μM y27632 was added and cells triturated before centrifugation at 500 g at 4°C for 5 minutes. Cells were resuspended in 10 mL washing solution (HBBS with 0.1% BSA and 5 μM y27632) and filtered through 100 μm and 50 μm cell filters. Single cells were pelleted again (500 g at 4°C for 5 minutes), and resuspended in 150 μL of the washing solution containing an anti-Cd45-PE antibody (1/500) (Thermo Fisher MA110233) (1/50) on ice. Cells were centrifuged at 500 g at 4°C for 5 minutes, resuspended in 200 μL washing solution containing DAPI and incubated for 5 minutes. Cells were washed twice with 1 mL washing solution and finally resuspended in 500 μL washing solution containing 5 μM DRAQ5 (Fisher Scientific). Single cell suspensions were sorted within 2 h on a FACsJazz (BD) at the Cambridge NIHR BRC Cell Phenotyping Hub. For each tissue, cells with high Venus fluorescence and non-fluorescent cells were sorted directly into lysis buffer (RLT plus, Qiagen) whilst dead cells, nucleus-free remnants and immune cells were excluded on the basis of DAPI, DRAQ5 and CD45 staining, respectively.

**RNA extraction, RNA sequencing and RT-qPCR.** Total RNA from FACS-isolated cells was extracted using RNAeasy Micro Plus kit (Qiagen) according to the manufacturer’s instructions and concentration and quality assessed using a Bioanalyser RNA Pico kit (Agilent). RNA samples with RNA integrity number values between 8.6 and 10 were used. Libraries were prepared and amplified using SMARTer Pico V2 Mammalian prep kit ( Takara) from 10 ng of RNA. Libraries were single-end 50 base pair sequenced at CRUK Cambridge on an Illumina HiSeq. 4000 platform. The average total number of reads in each sample was 32.3 million with an average of 21.4 million mapping uniquely to the mouse genome GRCm38 using STAR⁵¹. Read counts per gene were
Table 1. Table of primers.

| Primer   | Purpose                                      | Sequence (5'-3')          |
|----------|----------------------------------------------|---------------------------|
| FSD001   | Replacement GN by rspl/neoR cassette forward | GCCCCACGTGTATTACCACCAGCAGCTACTGGCCGTGTTCCTCTTG |
| FSD002   | Replacement GN by rspl/neoR cassette reverse | CCTCTAGAGATAGGGGGGCTTCCACATGGGCTGAGAGGAAGG |
| FSD003   | Replacement GN by Venus cassette forward     | GCCCCACGTGTTACACCAGCAGCTACTGGCCGTGTTCCTCTTG |
| FSD004   | Replacement GN by Venus cassette reverse     | CCTCTAGAGATAGGGGGGCTTCCACATGGGCTGAGAGGAAGG |
| FSD045   | To sequence GN-Venus + −5000bp f1          | CTCCAGGAGGAGAAAAGAG       |
| FSD046   | To sequence GN-Venus + −5000bp f2          | CATGAGCTCCAGCTACTGCT      |
| FSD047   | To sequence GN-Venus + −5000bp f3          | CAAACCGTGCCACCTTTATTC     |
| FSD048   | To sequence GN-Venus + −5000bp f4          | CTGGCCTGGCAGAGATGATG      |
| FSD049   | To sequence GN-Venus + −5000bp f5          | CAGCACCTGCCACACTGAG       |
| FSD050   | To sequence GN-Venus + −5000bp f6          | CTTTGACACATGCTCTTCAG      |
| FSD051   | To sequence GN-Venus + −5000bp f7          | CTCTTTCATCGAGGGCTTCC      |
| FSD052   | To sequence GN-Venus + −5000bp f8          | CAGCAGATAGGGCCGAGACTAG    |
| FSD053   | To sequence GN-Venus + −5000bp f9          | CTTCTGGGGTCACTAACTTGC     |
| FSD054   | To sequence GN-Venus + −5000bp f10         | CTTGATCCTCAGCTCTTCCAG     |
| FSD055   | To sequence GN-Venus + −5000bp f11         | CGTAACAGCTCTGCTGCTCAG     |
| FSD056   | To sequence GN-Venus + −5000bp f12         | CTTGAAAGCTGTCCTAGATG      |
| FSD057   | To sequence GN-Venus + −5000bp f13         | GCAACCCAGAAGACATCAAG      |
| FSD058   | To sequence GN-Venus + −5000bp f14         | GAACACTGTCGCTCCAGATG      |
| FSD059   | To sequence GN-Venus + −5000bp f15         | GCAGAGAGATAGGCTGCTG       |
| FSD060   | To sequence GN-Venus + −5000bp f16         | GGAGCTGTCCTAGCTCTGAGT     |
| FSD061   | To sequence GN-Venus + −5000bp f17         | GAGTGTCTCTGTAGTCTG        |
| FSD062   | To sequence GN-Venus + −5000bp f18         | CAATCAGGAGATGACGCTC       |
| FSD063   | To sequence GN-Venus + −5000bp f19         | CTACACCAGCTACTCCTCTCGC    |
| FSD064   | To sequence GN-Venus + −5000bp f20         | CACCTACTGGGACCAAGTGTTC    |
| FSD065   | To sequence GN-Venus + −5000bp f21         | CTTGATGCTTCTCTGCTTGTG     |
| FSD066   | To sequence GN-Venus + −5000bp f22         | CACTCGGAGACTGCTGTACAG     |
| FSD067   | To sequence GN-Venus + −5000bp f23         | CTGGAGGAAAGCTGCTGACC      |
| FSD068   | To sequence GN-Venus + −5000bp f24         | CACAGGATGTTGGAAGAGTAAGG   |
| FSD069   | To sequence GN-Venus + −5000bp f25         | GCAGTGTAGTAGCTCCAGAG      |
| FSD070   | To sequence GN-Venus + −5000bp f26         | CTCTGACAGCTCTACTGCTG      |
| FSD071   | To sequence GN-Venus + −5000bp f27         | GGAAGCCCTCTGATAGAAGAG     |
| FSD072   | To sequence GN-Venus + −5000bp f28         | CTGGAGGAGATGCTGCTGCCAG    |
| FSD073   | To sequence GN-Venus + −5000bp f29         | CTCGAGTGTGGAGGCTGCTG     |
| FSD074   | To sequence GN-Venus + −5000bp f30         | CACATCTCTGCTGGACACGAG     |
| FSD075   | To sequence GN-Venus + −5000bp f31         | GAAATAAGGCTGAGCTGTTG      |
| FSD076   | To sequence GN-Venus + −5000bp f32         | CAGTGAGCTGAGCCTACTTG      |
| GFPF1    | mouse screening                             | GACGTAAAACGCCACAAGT       |
| GFPR1    | mouse screening                             | GGATCTTGAAGGTCCGCTTC     |

determined using STAR and differential gene expression was determined using DESeq. 252. Raw data and generated count table are available with the GEO accession number GSE128117. RNA from homogenised tissues and cell cultures were extracted using TRI Reagent following standard protocols. RNA was treated with DNase I (Ambion), and reverse transcription was performed using the High-Capacity cDNA Reverse Transcription Kit (Applied Biosystems). RNA from sorted cells was reverse-transcribed using SuperScript III (Invitrogen) according to the manufacturer’s protocol. RT-qPCR was performed using the fast Taqman universal mastermix and the Taqman probes listed in Table 3 on a QuantStudio 7 (Applied Biosystems). Gene expression was measured relative to that of β-actin measured in parallel in the same sample using the ΔCt method.

**Mass spectrometry.** Plasma samples were extracted using ACN precipitation further purified using an HLB Prime μElution solid-phase extraction (SPE) plate (Waters, MA, USA) 51, whilst the Ussing chamber samples were extracted using SPE only. Peptides were analysed using a Thermo Fisher Ultimate 3000 LC system coupled to a Q Exactive Plus Orbitrap mass spectrometer (ThermoScientific, San Jose, USA) as previously described. 24. LC-MS/MS data were analysed using Peaks v8.5 to identify peptides sequences on the Mus musculus and Homo sapiens Swissprot databases (downloaded on 26/10/2017) and XCalibur 4.1 (ThermoFisher) to
forms or total protein levels, the cells were lysed in 150 μL of lysis buffer (50 mM Tris- HCl, 150 mM NaCl, 1% IGEPA-L-CA630, 0.5% deoxycholic acid, and one tablet per 50 mL of complete EDTA-free protease inhibitor cocktail (Roche)) on ice for 30 min, centrifuged at 10,000 g for 10 mins at 4 °C and the supernatants analysed.

Table 2. Table of antibodies and antisera used.

| Use                  | Antibody target                               | Company and catalog number | RRID           | Dilution |
|----------------------|-----------------------------------------------|----------------------------|----------------|---------|
| hybridoma            | FITC labelled monoclonal antibody anti-mouse IgG | Jackson ImmunoResearch (115-095-164) | AB_2338600   | 10 μg/mL |
| HTRF assay           | AlexaFluor 647 labelled goat anti mouse Fc IgG | Jackson ImmunoResearch (115-605-164) | AB_2338913   | 7.5 nM  |
| Cell sorting         | anti-CD45-PE Thermo Fisher (MA110233) | AB_11153576          | 1/500         |
| IC                   | Donkey anti-mouse 647 Life technologies (A31571) | AB_162542           | 1/300         |
| IC                   | Donkey anti-goat 488 Life technologies (A11055) | AB_2534102          | 1/300         |
| IC                   | Donkey anti-rabbit 555 Life technologies (A31572) | AB_162543           | 1/300         |
| IC                   | Goat anti-FGF Abcam (Ab5450) | AB_304897            | 1/1000        |
| IC                   | Rabbit anti-Lysozyme Dako (A0099) | AB_2341230          | 1/200         |
| IC                   | Rabbit anti-Muc2 Santa Cruz (sc15334) | AB_2146667          | 1/200         |
| IC                   | Mouse anti-E-cadherin BD (610181) | AB_397580            | 1/200         |

Quantify peptides measuring signal peak areas for specific sets of m/z and retention time corresponding to the identified peptides. Quantification of the guanylin peptides along the mouse intestine was analysed on previously published LC-MS/MS data (ProteomeXchange PXD009788, 10.6019/PXD009788)47.

Antibody generation. The production and purification of soluble proguanylin in E. coli was described by Lauber and collaborators54. This method was used to produce 25 mg of proguanylin for the hybridoma generation, screening and assay development. The purified protein was subjected to endotoxin removal using Polymyxin B resin55. To perform the primary screen of the hybridoma clones and IgG characterisation, 2 mg of proguanylin was enzymatically biotinylated using the BirA enzyme (Avidity LLC #BirA500) according to manufacturer’s instructions. Female CD1 mice aged 6–8 weeks were used for hybridoma generation and injected subcutaneously with 100 μg of recombinant proguanylin followed by 3 boosts of 100 μg each over 21 days. The immunisation, hybridoma generation and cDNA preparation and variable chain sequencing and IgG purification were performed as previously described55. For the clone selection, a Homogeneous Time Resolved Fluorescence biochemical assay was developed to measure the binding of IgGs to the biotinylated targets using Europium cryptate conjugated streptavidin (Cisbio #610SAKLB) as the donor and AlexaFluor 647 conjugated goat anti-mouse Fc IgG (Jackson ImmunoResearch #115-605-164) as the acceptor. Test samples were incubated for 15 h at RT with 2.5 nM biotinylated-Proguanylin premixed with 1 nM Streptavidin-cryptate, and 7.5 nM AlexaFluor 647 conjugated goat anti-mouse Fc IgG. The fluorescence was measured on an EnVision plate reader (Perkin Elmer) using a 320 nm excitation filter and 665 nm and 590 nm emission filters.

Cell culture. TC7 cells (RRID:CVCL_0233, a sub-clone of the caco-2 cell line60) were previously characterized as differentiating epithelial cells61 and were cultured in Dulbecco’s modified Eagle’s medium (DMEM, 4.5 g/L glucose) supplemented with 10% Fetal Bovine Serum (FBS), 100 units/mL penicillin, 100 μg/mL streptomycin and 2 mM L-glutamine at 37 °C, 5% CO2 and passaged every 7 days.

Human organoid lines were generated using a modified protocol60,61. Briefly, fresh surgical specimens of fresh human duodenum and colon were minced, and fragmented with 30 μM EDTA for 3 × 10 mins, with tissue shaken in PBS after each EDTA treatment. The fraction with intestinal crypts was then further digested using TrypLE (Life Technologies) for 5 mins at 37 °C to generate small cell clusters, which were seeded into basement membrane extract (BME, R&D technology), with 20 μL of lysis buffer for 30–60 mins at 37 °C. Organoid medium61 was then overlaid and changed 3 times per week. Organoids were passaged every 10–21 days, with TrypLE digestion for 15 mins at 37 °C followed by mechanical shearing with rigorous pipetting for 30 times to breakup organoids into small clusters which were then seeded as before in BME. To generate differentiated cell types, organoid diameters were washed 3 times in Advanced DMEM/F12 over 1 h. Organoid media lacking mnt3A, SB202190, A83-01 and nicotinamide was then added to the wells and changed every day for up to 10 days.

Secretion assay. Mouse primary cultures were prepared as described62. Briefly, colons from individual mice were harvested and digested with 0.4 mg/mL collagenase XI. Isolated crypts were plated on matrigel overnight and the day after, cells were rinsed with secretion buffer (in mM) 5.6 KCl, 138 NaCl, 4.2 NaHCO3, 1.2 NaH2PO4, 2.6 CaCl2, 1.2 MgCl2, and 10 HEPES, pH 7.4 and 0.001% BSA) 3 times over 30 minutes and incubated in secretion buffer with indicated drugs for 1 h. Supernatant was then collected and centrifuged 5 min at 2,000 g at 4 °C and transferred to a fresh Protein Lobind tube before purification through an SPE plate and analysis by LC-MS/MS as described.

Differentiated TC7 cells (grown at confluency for 9 days) or 8–9 day old differentiated human organoids were used for secretion experiments. The wells were washed 3 times in culture medium. Test agents were dissolved in the same medium and 250 μL applied for TC7 secretions, or 150 μL applied for organoid secretions. TC7 cells or organoids were incubated at 37 °C for 4 h, 8 h, or 24 h. Medium was centrifuged for 10 mins at 350 g at 4 °C, with supernatant used for downstream analysis. For experiments that required cell lysates to measure total protein levels, the cells were lysed in 150 μL of lysis buffer (50 mM Tris-HCl, 150 mM NaCl, 1% IGEPA-L-CA630, 0.5% deoxycholic acid, and one tablet per 50 mL of complete EDTA-free protease inhibitor cocktail (Roche)) on ice for 30 min, centrifuged at 10,000 g for 10 mins at 4 °C and the supernatants analysed.
Ussing chamber secretion experiments were performed with minor modifications as described previously. Human and mouse intestinal tissue was transported to the laboratory in ice-cold L15 medium. Serosa and most of the muscular layers were stripped away with fine forceps and the remaining tissue was mounted in an Ussing chamber (EM-LV5YS-4 system with P2400 chambers and P2405 sliders; Physiologic Instruments). The active epithelial surface area of each segment was 0.4 cm². Both parts of the Ussing chambers were filled with Ringer solution containing 120 mM NaCl, 3 mM KCl, 0.5 mM MgCl₂, 1.25 mM CaCl₂, and 23 mM NaHCO₃, with 10 mmol/L glucose, maintained at 37 °C, and continuously bubbled with 5% CO₂/95% O₂ (vol/vol). The transepithelial potential difference was monitored under open circuit conditions, using a DVC 1000 amplifier (WPI) and recorded through Ag-AgCl electrodes and 150 mM NaCl agarose bridges. After 20–30 minutes, solutions from both sides were exchanged with 1.2 mL of fresh Ringer solution containing 10 mM glucose, 0.1% fatty acid–free BSA, 10 μM amastatin, 500 kIU/mL aprotinin and Infacol (Forest Laboratories, 1:105 dilution, to prevent extensive foam formation). Solutions for LC-MS/MS only contained the additives 10 mM glucose and 0.001% BSA. Samples (120 μL) were taken from both compartments typically at 10, 40 and 85 minutes after solution exchange. IBMX (100 μM) and forskolin (10 μM) were added bilaterally immediately after the second sample was taken at 40 min. Reported secretion was normalized for 1 cm² of the tissue and a 60-minute secretion period. GLP-1 and Proguanylin were measured by MSD immunoassay or LC-MS/MS as indicated.

**MSD assay.** Anti-ProGuanylin antibody AB1000025 was coated using 5 μL of 50 ng/μL on an assay plate (MSD #MA6000) incubated at RT overnight. Plates were blocked using 1% BSA in PBS for 2 h at RT. After washing 3 times with washing buffer (PBS-Tween (0.05%)), 25 μL of samples were added to each well and incubated for 2 h at RT. Following incubation, plates were washed 3 times with washing buffer and 25 μL of a second Sulfo-Tag labelled anti-ProGuanylin antibody AB1000028 prepared at 1 μg/μL in Diluent 41 (MSD) was added per well for 1 h at RT. Once washed with washing buffer, 150 μL of Read buffer 2X was added to each well before reading the plate immediately using a MSD Workbench. A standard curve was constructed by fitting a sigmoidal function to the absorbance obtained from the known proguanylin concentrations and the concentration of experimental samples was assessed by interpolating them to the standard curve using the MSD discovery software. Samples in lysis buffer were diluted 1/20, dilution at which the lysis solution had no effect on measurements. GLP-1 and Proguanylin were measured by MSD immunoassay or LC-MS/MS as indicated.

**Data availability**

The datasets generated during and/or analyzed during the current study, the animal model (Guanylin-Venus) and the immuno-assay developed during this study are not publicly available but are available from the corresponding author on reasonable request. The RNAseq data has been deposited as outlined in material and methods.

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### Table 3. Table of Taqman assays probes.

| Gene                          | Species | TaqMan™ Gene Expression Assay Id |
|-------------------------------|---------|----------------------------------|
| Guanylin (Guca2a)             | mouse   | Mm00433863_m1                   |
| Uroguanylin (Guca2b)          | mouse   | Mm01192051_m1                   |
| Beta act (Actb)               | mouse   | Mm02619580_gl                    |
| Mucin 2 (Muc2)                | mouse   | Mm01276696_m1                   |
| Lysozyme (Lyz1)               | mouse   | Mm00657323_m1                   |
| SAM Pointed Domain Containing ETS Transcription Factor (Spdef gene) | mouse   | Mm00600221_m1                   |
| SRY-Box 9 (Sox9)              | mouse   | Mm00448840_m1                   |
| Villin (Vill)                 | mouse   | Mm00494146_m1                   |
| Glucagon (Gcg)                | mouse   | Mm01269055_m1                   |
| Gastric Inhibitory Polypeptide (Gip) | mouse | Mm00433601_m1                   |
| Peptide YY (Ppy)              | mouse   | Mm00520716_gl                    |
| Cholecystokinin (CcK)         | mouse   | Mm00446170_m1                   |
| Somatomedin (Sot)             | mouse   | Mm00436671_m1                   |
| Doublecortin Like Kinase 1 (Delk1) | mouse | Mm0044950_m1                   |
| Transient Receptor Potential Cation Channel Subfamily M Member 5 (Trpm5) | mouse | Mm01129032_m1                   |
| Chemokine (C-C motif) ligand 9 (Ccl9) | mouse | Mm00441260_m1                   |
| Macrophage Myristoylated Alanine-Rich C Kinase Substrate (Marcksl1) | mouse | Mm00456784_m11                   |
| Leucine Rich Repeat Containing G Protein-Coupled Receptor 5 (Lgr5) | mouse | Mm00438890_m1                   |
| Guanylin (GUCA2A)             | Human   | Hs00157859_m1                   |
| Uroguanylin (GUCA2B)          | Human   | Hs00951189_m1                   |
| Beta act (ACTB)               | Human   | Hs01060665_gl                    |
| Venus (Yfp)                   | Other   | Mr03987581_mr                    |

The datasets generated during and/or analyzed during the current study, the animal model (Guanylin-Venus) and the immuno-assay developed during this study are not publicly available but are available from the corresponding author on reasonable request. The RNAseq data has been deposited as outlined in material and methods.
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Author contributions
F.S.D., P.L., R.K., T.D., J.R., D.A.G., J.L.P.A., T.V., F.F. and B.G.C. performed experiments and analysed the data. C.L.M., S.J.M., R.H.H. and G.P.R designed and ran the clinical study, obtained regulatory approvals, recruited participants, obtained and processed samples from participants, and analysed the clinical data. B.G.C., S.R., M.G., F.M.G. and F.R. supervised the work and guided the project. All authors contributed to writing and/or approved the final manuscript.

Competing interests
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