Intelectins are ancient carbohydrate binding proteins, spanning chordate evolution and implicated in multiple human diseases. Previous GWAS have linked SNPs in ITLN1 (also known as omentin) with susceptibility to Crohn’s disease (CD); however, analysis of possible functional significance of SNPs at this locus is lacking. Using the Ensembl database, pairwise linkage disequilibrium (LD) analyses indicated that several disease-associated SNPs at the ITLN1 locus, including SNPs in CD244 and Ly9, were in LD. The alleles comprising the risk haplotype are the major alleles in European (67%), but minor alleles in African superpopulations. Neither ITLN1 mRNA nor protein abundance in intestinal tissue, which we confirm as goblet-cell derived, was altered in the CD samples overall nor when samples were analyzed according to genotype. Moreover, the missense variant V109D does not influence ITLN1 glycan binding to the glycan β-D-galactofuranose or protein–protein oligomerization. Taken together, our data are an important step in defining the role(s) of the CD-risk haplotype by determining that risk is unlikely to be due to changes in ITLN1 carbohydrate recognition, protein oligomerization, or expression levels in intestinal mucosa. Our findings suggest that the relationship between the genomic data and disease arises from changes in CD244 or Ly9 biology, differences in ITLN1 expression in other tissues, or an alteration in ITLN1 interaction with other proteins.
“omentin” is sometimes used.

Orthologs of human ITLN1 are induced during intestinal parasitism and related Th2-type immune responses, including asthma.

Human ITLN1 was found to recognize β-D-galactofuranose, a galactose isomer incorporated into the glycans of microorganisms, including protozoa, fungi, and bacteria, but not mammalian cells. Within β-D-galactofuranose, ITLN1 binds to the exocytic vical 1,2-diol, a chemical moiety present on other microbial carbohydrate-containing structures, such as Kdo2 (Di[3-deoxy-D-manno-octulosonic acid]) of lipopolysaccharides. This carbohydrate specificity suggests microbial pattern recognition binding for ITLN1, in its otherwise elusive role in innate immunity.

Genome-wide association studies (GWAS) have identified several single nucleotide polymorphisms (SNPs) present at the ITLN1 locus as risk alleles for Crohn’s disease, a main subtype of IBD, which is characterized by dysbiosis, epithelial barrier dysfunction, and immune function perturbations; processes consistent with an innate immune dysfunction.

Herein, we further characterize the ITLN1 locus in the context of IBD-associated SNPs, examine the intestinal expression and glycan binding of ITLN1 associated with risk and non-risk alleles, and localize ITLN1 protein expression to goblet cells of the human small and large intestine.

Results

The intelectin-1 locus and Crohn’s disease. To gain insight into the potential influence of IBD-associated allelic variants on expression and protein function, we genotyped surgical specimens from a previously reported cohort of individuals requiring surgical resection for Crohn’s disease or ulcerative colitis (n = 134). Accordingly, we found by direct sequence analysis of PCR products that the GWAS-identified SNP, rs2274910 (T/C), located in intron 3 of ITLN1, was associated with rs2274907 (A/T), which encodes a missense variant in exon 4, where valine is substituted with aspartic acid at amino acid position 109 (V109D).

Utilizing the 1000 Genomes Project Phase 3 of the Ensembl database we found rs2274910 and rs2274907 are in linkage disequilibrium (LD) across human populations, consistent with findings of association in our U.S. cohort (Supplemental Table 1). We sequenced the entire coding region of ITLN1 cDNA in all specimens of this cohort, exploring the potential for additional exon variants in IBD subpopulations. We found rs2274908 (C/T: H89H/Q) exclusively as a synonymous (silent) variant associated with rs2274907. No additional missense variants were identified.

To display allelic variants associated with disease, we constructed a locus map of ITLN1 (Fig. 1A) annotated with published SNPs as outlined by the National Human Genome Research Institute—European Bioinformatics Institute (NHGRI-EBI) Catalog and NCBI-LitVar database, including variants within ITLN1 and its flanking regions intergenic to CD244 (3’) and ITLN2 (5’). ITLN1 and CD244 (rs1333062 and rs4656940, respectively) have been annotated together as risk loci for IBD, due in part to their close proximity (~14,000 nt). The SNP rs133062 is located in the 3′-flanking region of ITLN1, 46 nt intergenic to exon 8, where rs4656940 is located within intron 1 of CD244. The odds ratio (OR) for Crohn’s disease of rs4656940 is approximately equivalent (1.15; 95% confidence interval [CI] = 1.09–1.21) to that of rs2274910 (1.14; 95% CI = NR) of ITLN1.

Several annotated SNPs in CD244 (rs4656942 and rs11265498) and in the adjacent gene Lyn (rs540254 and rs560681), but not any in neighboring SLAMF7, are in LD with GWAS-identified rs2274910 and rs4656940 (Supplemental Table 2). In addition, rs4656958 located in the 5′ flanking region of ITLN1 (2004 nt 5′ to exon 1) has also been identified by GWAS as a risk allele for IBD (OR = 1.10; 95% CI = 1.03–1.19). Thus, GWAS-identified IBD-associated SNPs are located within ITLN1 and its flanking regions, and these SNPs are in LD with SNPs in CD244 and Lyn, which are genes located 3′ to ITLN1. Data in the GTEx browser revealed that quantitative trait loci (QTL) for expression (eQTL) in blood for ITLN1 and CD244 are both affected by several disease risk SNPs, including rs1333062, rs2274907 and rs2277539 (Supplemental Table 3). In addition, QTL for splicing (sQTL) of USF1 were similarly affected by these SNPs, as was eQTL for Lyn by rs465940. No significant eQTLs or sQTLs were identified in small intestine or colon tissue. Although other genes located at 1q23.3 may be responsible for dysbiosis, epithelial barrier dysfunction, and immune function perturbations; processes consistent with an innate immune dysfunction.

Further review of ITLN1 rs2274910 (C/T) revealed a distinction in the frequency of the GWAS-identified risk allele (C) between African (28%) and non-African (67%) superpopulations (SP) across the context of different SP groups. The study populations for most IBD-focused GWAS are comprised primarily of non-African SP, including investigations identifying ITLN1 as a risk locus. Using the Ensembl database, we performed pairwise LD analyses of disease-associated ITLN1 variants (Fig. 1A, Table 1) in African and European SP using rs2274907 (A/T: V109D) as a focus point simply due to it being a missense variant. In addition to being in complete LD (D' = 1.0, r2 = 1.0) with rs2274910, rs2274907 (V109D) was found also to be in LD with GWAS-identified alleles across the ITLN1 locus, where risk alleles are the major alleles in European SP and minor alleles in African SP (Table 1, Fig. 1B).

The allele genotype frequencies in European SP: VV (7.6%), VD (49.3%), and DD (43.1%) contrasts sharply with that in the African SP VV (45.7%), VD (43.1%), and DD (11.2%) (Supplemental Fig. 1). In our small cohort of surgical specimens, allele frequencies were not significantly different for those diagnosed as either Crohn’s disease or ulcerative colitis as compared to respective controls, but there was a suggestion of skewing in specimens diagnosed as ileal compared to colonic Crohn’s disease, worthy of follow-up analysis in a larger cohort (Supplemental Table 1). Our findings clarify that rs2274910 (T, D109), which is in complete LD with GWAS-identified rs2274910 risk allele (C), is the major allele in non-African SP; in Crohn’s disease multiple GWAS report a further skewing of this high frequency of the risk allele observed in the studied populations.

Intestinal intelectin-1 expression, glycan-binding, and oligomerization is unaltered by missense variant rs2274907 (A/T, V109D).

Di Narzo, et al. examined blood and intestinal eQTLs from a...
Table 1. European and African superpopulation haplotype analysis at the ITLN1 locus. LD, linkage disequilibrium; AF, allele frequency; Abs, absent. D’ and r² values represent co-inheritance range (i.e., lowest and highest values) for each sub-population within African and European superpopulations.

| Location: Chromosome 1q23.3; GRCh38:CM000663.2 (Forward strand) | European LD: rs2274907 (D/V) | African LD: rs2274907 (D/V) |
|----|----|----|
| Location: Chromosome 1q23.3; GRCh38:CM000663.2 (Forward strand) | D', r² | D', r² |
| Location: Chromosome 1q23.3; GRCh38:CM000663.2 (Forward strand) | AF | AF |

Figure 1. ITLN1 polymorphisms. (a) The physical arrangement of ITLN1 relative to centromere/telomere orientation and flanking genes on chromosome 1q23.3 is diagramed, along with single nucleotide polymorphisms associated with human disease, including IBD: rs465694023, rs1126550158, rs1205871759, rs1157877060, rs465695361, rs133306224, rs227490730, rs227490830, rs22749101, rs229755925, and rs465695825–27. Red text highlights GWAS-identified SNPs. (b) 1000 Genomes Project Phase 3 super population allele frequency (percentages) of rs2274907 (A/T, V109D): AFR, African; AMR, American, EAS, East Asian; EUR, European; SAS, South Asian. (c) 3D structure of human trimeric ITLN1 bound to β-D-galactofuranose (black). The distance between V109 residue (red) and coordinating calcium (green) are indicated on each monomer (PDB: 4WMY). (d) Amino acid sequence adjacent to coding variant rs2274907 (V109D) in orthologs of human ITLN1 (“—” identical residue present ITLN1).
therapy-resistant Crohn's disease cohort. Their analysis indicated eQTLs for ITLN1 in intestine (but not blood) are linked to GWAS-identified risk alleles rs4656958 and rs2724910, and eQTLs for CD244 in blood (but not intestine) are linked to the risk alleles rs4656958, rs4656940 and rs2724910. In both cases, a statistically significant decrease in expression is associated with the GWAS-identified SNPs. To better clarify expression patterns of ITLN1 within intestinal tissue, we generated an ITLN1-specific quantitative real-time PCR assay to enumerate the absolute mRNA transcript levels. ITLN1 mRNA was abundant in each anatomic segment across both small and large intestine (Supplemental Fig. 2a). Similarly, high levels of ITLN1 gene expression were observed in surgical specimens of both ileum and colon (Fig. 2a). Crohn's disease diagnosis did not alter transcript levels in either small or large intestine (Fig. 2a). Of note, ITLN1 expression in the colon was elevated (p < 0.05) in specimens from individuals diagnosed with ulcerative colitis (Fig. 2a). To determine whether ITLN1 expression is modified by risk-associated haplotype, we stratified the data by genotype. In ileum, colon, and combined sample sets, the risk associated haplotype (queried by rs2274907 genotype) did not modify ITLN1 expression (Fig. 2b and Supplemental Fig. 2b). Together, these findings suggest that the mucosal transcript level of ITLN1 is unaltered in Crohn's disease. Moreover, the risk-associated haplotype does not modify mucosal ITLN1 expression (Table 1).

The distinguishing chemical properties of valine and aspartic acid provided impetus to further explore the potential of missense variant V109D to modify ITLN1 oligomeric structure or function. Using PyMOL to visualize the published crystal structure of human ITLN1 (PDB: 4WMY), residue 109 was found to be located at the protein surface; however, it is not located at the sites annotated for glycan binding and/or calcium coordination (Fig. 1c). Bio-layer interferometry analysis found that recombinant V109-ITLN1 reversibly bound to immobilized β-D-galactofuranose with kinetics and affinity that mirror recombinant D109-ITLN1 (117 nM versus 107 nM, respectively) (Fig. 1c). Without finding discernable influence on glycan binding, we next tested if the V109D change altered ITLN1 oligomerization. For other secreted lectins (e.g., ficolins and mannose-binding lectin), oligomerization is important for avidity in binding cell-surface glycans as well as subsequent effector functions. Previous studies report that human ITLN1 forms stable homotrimers, consisting of ~35 kDa monomers, connected through intermolecular disulfides (C31 and C48). Analysis of human ileal lysates confirmed that ITLN1 forms a stable homotrimer at approximately 110 kDa; however, a consistent ITLN1 band near the ~220 kDa marker was also identified using a gradient SDS-PAGE (4–20%). This suggests that human ITLN1 forms higher molecular weight oligomers (likely dimers of homotrimers), which are stable under denaturing (but not reducing) conditions (Fig. 2d). These findings agree with those reported for an amphibian ITLN1 ortholog, where dimers of homotrimers were shown to mediate bacterial agglutination. We also performed immunoblots of ileal specimens isolated from V109 and D109 homozygote individuals to determine the potential influence on oligomer formation. Neither VV or DD genotype affected the formation of homotrimers or hexamers (Fig. 2d). These data suggest that it is unlikely that the role of V109D variation in Crohn's disease pathogenesis results from alterations in ITLN1 glycan binding or oligomerization.

Alternatively, we reasoned that D109 might influence protein stability, and that such differences would be observed as changes in protein abundance. In human ileal samples, we observed that under reducing conditions ITLN1 was present primarily as a monomer at ~40 kDa, with the exception of a minor fraction migrating at ~70 kDa (Fig. 2d). This minor band may represent a covalent non-reducible dimer but further study is needed to fully characterize this entity. Human ITLN1 possesses a single glycosylation site at N163, which slightly increases the molecular weight from that predicted (33 kDa) by the deduced amino acid sequence. Ileal samples from individuals diagnosed with Crohn's disease demonstrated equivalent relative proportions of protein oligomers and no additional degradation products compared to control specimens (Fig. 2d). Thus, ITLN1 was found to be an abundant protein in the small intestine, where expression was comparable in specimens from control and Crohn's disease biopsies, consistent with mRNA expression data (Fig. 2a, d).

To further investigate the possible significance of V109D variation in Crohn's disease, we performed a sequence analysis (NCBI BLAST) of the neighboring amino acid region across various ITLN1-like orthologs. In all Eutherian mammals queried (n = 63), aspartic acid (D) was reported as the amino acid at the corresponding position (Fig. 1d). Even amongst primates, valine at position 109 is unique to Homo sapiens. Nevertheless, other chordates were found to encode hydrophobic amino acids like valine (e.g., leucine) at this position, including ancient ascidians (Fig. 1d). While aspartic acid marks the risk-allele associated with Crohn's disease, the V109D variation is not a "mutation", despite its location in a conserved domain of the protein and differences in chemical properties of these residues. Furthermore, V109, or the genetic variants in LD with this allele, could be considered "protective" in the context of Crohn's disease.

Human intelectin-1 is a goblet cell product. In mice, where mammalian intelectins were first discovered, Int1 is expressed in Paneth cells. In contrast, a recent single-cell RNA sequence analysis investigation identified ITLN1 mRNA in goblet cells of the ileum and colon. We sought to localize ITLN1 protein in these tissues to confirm and extend this cell-type expression pattern in humans. Both ITLN1 and ITLN2 are expressed in the human small intestine and cannot be distinguished using most commercially available antibodies; therefore, we generated paralog-specific antibodies to clarify the cellular origin of ITLN1. Colon biopsies of non-IBD subjects, preserved in fixatives to retain mucus (either Carnoys fixative or Histochrome), were incubated with either antisera targeting a domain highly conserved in intelectins (ITLNCONS) or targeting the N-terminal specific domain (ITLN1NTERM). Both ITLNCONS and ITLN1NTERM immunoreactivity were localized to colonic goblet cells (Fig. 2e, Supplemental Fig. 2c). To confirm goblet cell localization, samples were stained with a mucin-2 (MUC2) antibody generated against a peptide sequence located within the C-terminal domain (adjacent to the autocatalytic sequence) of the mature, glycosylated MUC2 protein (MUC2C). This antibody targets goblet cell granules and secreted protein. While MUC2C stained goblet cells most strongly in their secretory.
Figure 2. ITLN1 mRNA and protein expression in the small intestine and colon. (a) ITLN1 mRNA expression in surgical specimens of ileum and colon according to diagnosis. The mRNA copy number per 10 ng total RNA was determined with quantitative real-time RT-PCR using external cDNA standards (see methods). (b) ITLN1 mRNA expression by V109D genotype (excluding ulcerative colitis samples). Box plots present the median and first and third quartiles, whiskers present the range. Statistical analysis of ITLN1 mRNA expression was performed using either a Kruskal–Wallis test with Dunn's multiple comparisons or two-tailed Mann–Whitney U test; *p < 0.05. (c) Representative real-time bio-layer interferometry responses of recombinant ITLN1 V109 and D109 binding to immobilized β-D-galactofuranose (upper). Responses at 490 s were plotted against ITLN1 concentration and fit to a single site binding equation to determine Kd (lower). Data is representative of three separate experiments. (d) SDS-PAGE (4–20% gradient) immunoblots of human ileum by V109D genotype (left, n = 2 per genotype) and disease state (right). (e) Immunohistochemistry of human colon using the following antibodies: ITLN conserved (ITLNCONS), ITLN1-specific (ITLN1NTERM), MUC2 mature (MUC2C3), deglycosylated MUC2 (MUC2APO, PH1900). Tissue counterstained by DAPI (blue). (f) Immunofluorescence of human colon using ITLNCONS-FLOR (anti-chicken; Alexa Flour 647) and MUC2C3 (anti-rabbit; Alexa Flour 488). (g) Immunohistochemistry of human ileum; ITLN2-specific (ITLN2NTERM) antibody. Red box indicates Paneth cells identified with the ITLNCONS antibody. CTRL = control (non-Crohn's disease and non-ulcerative colitis), CD = Crohn's disease, UC = ulcerative colitis. Images in (e–g) are representative of multiple (> 6 fields) from 9 specimens. Light microscopy scale bars: 20x (200 µm), 40x (100 µm), and 100x (50 µm); fluorescence microscopy scale bars: 20x (100 µm) and 63x (50 µm).
granule, the cellular staining pattern of ITLN1 was different. Specifically, punctate staining of ITLN1 adjacent to the granule and relatively weak staining within the granule was a feature of all ITLN1-positive samples tested (n = 6 colon, n = 3 ileum). To further characterize the staining pattern of ITLN1 within goblet cells, we used another MUC2 antibody that targets the tandem repeat region of the apo-MUC2 protein (PH1900, MUC2APO). This antibody identifies non-O-glycosylated MUC2, which resides within the endoplasmic reticulum region of goblet cells. MUC2APO staining of the cytoplasmic region in goblet cells mimicked the staining pattern for ITLN1TERM (Fig. 2e, Supp. Fig. 2C). At higher magnification, ITLN1TERM staining was identified in both cytoplasmic and granule compartments, and readily apparent in both longitudinally and transversely oriented tissue sections (Fig. 2e). To better localize ITLN1 within goblet cells, we generated a chicken anti-ITLN antibody (conserved, ITLNCONS-FLOR) for dual labeling with the rabbit-derived MUC2C3 reagent in fluorescence microscopy. As observed by conventional microscopy, ITLN1 and MUC2 largely colocalized within colonic goblet cells, where ITLN1 was also found occasionally in the cytosolic compartments adjacent to nuclear (DAPI) staining (Fig. 2f). Immunohistochemistry of human ileum using the ITLN1TERM antibody also identified goblet cells, where the aforementioned cytosolic and granular staining pattern was also evident (Fig. 2g). It is worth noting that as a feature distinct from findings in mice, human Paneth cells remained unstained with the ITLN1TERM antibody (Supp. Fig. 2D). Although staining of human ileum with the ITLNCONS antibody identified Paneth cells in addition to goblet cells, paralog-specific antibodies determined Paneth cell staining to be human ITLN2 (Fig. 2g, and Nonnecke, et al.: unpublished observations). Together, our results demonstrate that human ITLN1 is expressed in goblet cells, both in the ileum and colon.

Discussion

Various GWAS investigations have identified SNPs at the ITLN1 locus as risk alleles for Crohn's disease. We observed that these SNPs are in LD, forming a risk haplotype that extends into the adjacent CD244 and Ly9 genes. Given the abundant expression of ITLN1 in intestinal tissues and the likely role of this lectin in innate immunity, we focused our investigation on ITLN1 in the intestine. The first GWAS-identified SNP at this locus (rs2274910) is located in intron 3 of ITLN1. We determined that this variant was associated with an adjacent SNP in exon 4 (rs2274907), which encodes a valine to aspartic acid amino change at residue 109. Although valine is the most common allele in African SP, aspartic acid is both the most common allele and the disease-associated allele in European SP—the primary population studied in IBD GWAS. Our results clarify that V109D is in LD with other GWAS-identified Crohn's disease-associated risk alleles of rs2274910 (T/C), and other alleles forming the risk haplotype are common, where T and C alleles of rs2274910 are present at 43% and 57%, respectively across Phase 3 Individuals. Similarly, the Crohn's disease risk allele of autophagy-related 16-like 1 (ATG16L1) rs2241880 (A/G) is common (60:40%) and associated with a modest OR (1.45; CI = 1.27—1.64). Further investigation is required to determine the mechanism by which genetic variation at this locus affects risk for Crohn's disease. In this study, we tested relevant possibilities, and our data support that it is unlikely to be due to ITLN1 expression in intestinal goblet cells. It is possible that expression of ITLN1 in another tissue could underlie disease risk. Moreover, expression of other genes at this locus, notably either CD244 or Ly9, which include SNPs in LD with the risk haplotype might explain disease risk. Further investigation is required to determine the mechanism by which genetic variation at this locus affects risk for Crohn's disease. In this study, we tested relevant possibilities, and our data support that it is unlikely to be due to ITLN1 expression in intestinal goblet cells. It is possible that expression of ITLN1 in another tissue could underlie disease risk. Moreover, expression of other genes at this locus, notably either CD244 or Ly9, which include SNPs in LD with the risk haplotype might explain disease risk. Further investigation is required to determine the mechanism by which genetic variation at this locus affects risk for Crohn's disease. In this study, we tested relevant possibilities, and our data support that it is unlikely to be due to ITLN1 expression in intestinal goblet cells. It is possible that expression of ITLN1 in another tissue could underlie disease risk. Moreover, expression of other genes at this locus, notably either CD244 or Ly9, which include SNPs in LD with the risk haplotype might explain disease risk.

Methods

Human specimens and ethics statement. All methods were carried out in accordance with relevant guidelines and regulations. Surgical specimens were obtained at The Cleveland Clinic Foundation (Cleveland, OH) with informed consent from all patients, and deidentified under protocols approved by the Institutional Review Board at The Cleveland Clinic Foundation as previously described. These specimens were collected from subjects undergoing surgery for Crohn's disease (CD), ulcerative colitis (UC), or non-CD/UC-related (control, CTRL) surgical indications (i.e., colon cancer, bowel obstruction, and familial adenomatous polyposis). Race was recorded for these specimens based on medical record reporting. Diagnoses were based on standard criteria using clinical, radiological, endoscopic and histopathological findings. Exclusion criteria included the diagnoses of backwash ileitis, indeterminate colitis, concurrent CMV or C. difficile infection. Samples were immediately snap frozen in liquid nitrogen (for RNA, DNA, and protein analysis) or placed into tissue fixative for histology. The mucosa, utilized for mRNA and protein expression, was dissected away from the intestinal/colonic serosa and associated adipose tissue. The overview of regional mRNA expression (Supp. Fig. 1) used commercially-obtained RNAs of human tissue (Biochain, Newark, CA).

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Databases. Single nucleotide polymorphisms (SNPs) referenced to human ITLN1 were identified using the NHGRI-EBI Catalog of human genome-wide association studies (https://www.ebi.ac.uk/gwas/home) and NCBI-LitVar (https://www.ncbi.nlm.nih.gov/CPBresearch/Lu/Demo/LitVar/#!!query = ) databases3,24. Further investigation of identified SNPs, including descriptions of putative function and population frequency, was procured from Ensembl and the 1000 Genomes Project (Phase 3)31,32. Allele (i.e., nucleotide) associations and linkage disequilibrium were studied using the NCBI's dbSNP (LDpair) database (https://ldlink.nci.nih.gov/?tab=help+Ldpair)35. QTL analysis was queried at The Genotype-Type Expression (GTEx) Portal (https://www.gtexportal.org/home/)36. Missense variant rs2274907 (V109D) was applied to the referenced crystal structure of human ITLN1 (PDB ID: 4WMQ) using PyMOL software (The PyMOL Molecular Graphics System, Version 1.8.2.1 Schrödinger, LLC., https://pymol.org/2/) to orientate location.

Genotyping. Genomic DNA was isolated and purified using the QIAamp DNA minikit (Qiagen, Germantown, MD). Exon variants in each of our IBD cohort were explored by Sanger sequencing of cDNA corresponding to the entire coding region of ITLN1 amplified using specific primers (2 s: 5'-GGAGCGTTTGGAGAAAGCTGCA-3' and 1147a: 5'-ATTAAACATTCTAGCTACTGGTG-3'). The sequenced PCR products were aligned to the NCBI-ITLN1 reference (NM_017625.3), identifying rs2274907 (A/T: V109D) and rs2274908 (C/T: H89H/Q) in some specimens. Association of single nucleotide polymorphisms rs2274910 (T/C) and rs2274908 (C/T) was identified by PCR amplification from genomic DNA across adjacent regions in multiple samples, including both homozygous, as well as heterozygous (T/C and A/T) genotypes using specific oligonucleotide primers (1062 s: 5'-ACAAGGAGCCCTTTCGGCCATG-3'; 1431a: 5'-AGTTGGGACCAACCTTGTAGTC-3'). The PCR reactions were initiated by denaturation of the DNA template at 95 °C for 10 min followed by 45 cycles consisting of 95 °C for 1 s, a -1 °C per two-cycle 'touchdown' annealing temperature for 5 s and extension at 72 °C for 10 s. Following the cycle runs, samples were denatured to establish the melting temperature(s) of the PCR product. The sample melt temperatures were compared to that of the internal standard to confirm template specificity. Absolute quantification of target gene within each sample was obtained by inclusion of an external cloned cDNA standard of known concentration, where crossing point (i.e., cycle number) is matched to a standard curve. Data was plotted in Prism version 9.0.0 (GraphPad).

RNA isolation. The general procedures for RNA isolation and synthesis of cDNA were previously described by our group31,32. For the current study, RNA was freshly isolated for all specimens from frozen tissue25. Briefly, frozen tissue was dispersed using Brinkmann Polytron homogenizer in guanidinium thiocyanate (GTC) buffer (4–5 M GTC, 0.5% sodium-N-lauryl sarcosine, 10 mM EDTA, 20 mM Tris–HCl, 0.1 M 2-mercaptoethanol). Homogenates were layered above 5.7 M CsCl/EDTA in diethyl pyrocarbonate (DEPC) H2O in polyallomer tubes (11 × 34 mm; Beckman, Indianapolis, IN). Using a TLS-55 swinging bucket rotor (Beckman, Indianapolis, IN) samples were centrifuged for 3 h at 55,000 rpm in a TL-100 ultracentrifuge (Beckman). Following centrifugation, excess GTC and CsCl were removed by aspiration and the RNA pellet was solubilized with RNA sample buffer (0.5% N-lauroylsarcosine, 5 mM EDTA, 5 mM tris, and 5% 2-mercaptoethanol). The RNA buffer solution was transferred to tubes containing an equal volume of 50:50 phenol–chloroform (Calbiochem/EMD Millipore, San Diego, CA), vortexed, and centrifuged for 3 min at 13,000 rpm. The resulting supernatant was washed with 100% chloroform (Sigma-Aldrich, St. Louis, MO), centrifuged, and added to tubes containing 100% ETOH and 0.3 M sodium acetate. Following overnight incubation at -80 °C, RNA was precipitated by centrifugation for 20 min at 10,000 rpm and washed once with 80% ETOH. The resulting RNA pellet was re-suspended in DEPC H2O, and concentration was determined by ultraviolet absorbance spectroscopy (260 nm)31,32.

cDNA synthesis. RNA (2–5 μg) was precipitated under centrifugation, washed once with 80% ETOH and re-suspended in DEPC H2O. cDNA synthesis was performed according to the procedures outlined by the SuperScript III First-Strand cDNA Synthesis System (Invitrogen, Thermo Fischer Scientific, Waltham, MA). The cDNA product was purified using a column QIAquick PCR Purification Kit (Qiagen, Germantown, MD).

Quantitative real time PCR (qRT-PCR). Primers specific for human ITLN1 (211 s: 5'-CTTCGTCTCTCATTCTGCCCCC-3'; 297a: 5'-CTGGTAGATAAACACCTTCTC-3') and ACTB (128 s: 5'-TGATGTTGCGGTA TGGGTCA-3'; 220a: 5'-GGTGCTCGATGGGTACCTTCAG-3') were designed using MacVector version 16 Software. Real-time PCR was performed on cDNA template corresponding to 10 ng of RNA using a Roche Diagnostics Lightcycler 2.0 (Roche, Indianapolis, IN) as previously described25,75. Briefly, each reaction included 4 mM MgCl2, 0.5 μM forward and reverse primers, and 1× LightCycler FastStart SYBR Green I mix (Roche, Mannheim, Germany). A control without cDNA template was included in the procedural assay. The PCR conditions were: initial denaturation at 95 °C for 10 min, followed by 45 cycles with each cycle consisting of denaturation, 95 °C for 1 s annealing at 60 °C for 5 s; and extension at 72 °C for 10 s. Following the cycle runs, samples were denatured to establish the melting temperature(s) of the PCR product. The sample melt temperatures were compared to that of the internal standard to confirm template specificity. Absolute quantification of target gene within each sample was obtained by inclusion of an external cloned cDNA standard of known concentration, where crossing point (i.e., cycle number) is matched to a standard curve. Data was plotted in Prism version 9.0.0 (GraphPad).

Rabbit anti-ITLN1 polyclonal antibodies. Analysis of the pre-ITLN1 deduced protein sequence using the SignalP4.0 algorithm indicated signal cleavage would occur between G16 and W1753. The adjacent 19 amino acids of ITLN1 (WSTDEANTYKFEWCTSSS) comprised a completely unique sequence compared to the corresponding region of the human ITLN2 sequence (AAASSEMLESREFETCAFSFS). Adjacent to this discrimi-
nating 19-amino acid sequence (W17 to P35) of ITLN1 is a domain beginning at amino acid S36 that has very high sequence identity to ITLN2 as well as other intelectin orthologs. BLASTp search of the non-redundant protein database with the 19-amino acid W17 to P35 peptide sequence yielded numerous primate ITLN1 orthologs with highly similar sequence (>80%), but no other mammalian proteins. This 19-amino acid peptide was synthetically prepared, purified and characterized to confirm proper mass, as previously described. The peptide was coupled to ovalbumin (OVA, Thermo Fischer Scientific, Waltham, MA) at a ratio of 20 molecules peptide per molecule OVA using two strategies. First, one equivalent volume of 0.2% glutaraldehyde was added to the 20:1 peptide/ovalbumin mixture in PBS; the mixture was then stirred for 1 h at room temperature. 1 M glycine was added to a final concentration of 200 mM and stirred for an additional 1 h. Second, the peptide was reacted with maleimide-activated-ovalbumin (Imject™ Maleimide-Activated Ovalbumin, Thermo Fischer Scientific, Waltham, MA) according to the suppliers’ protocol. Briefly, the peptide was mixed with the activated ovalbumin at a 20:1 molar ratio in PBS and allowed to react at room temperature for 2 h. Each of the two peptide/OVA conjugates were dialyzed against PBS for 24 h at 4 °C, changing dialysis buffer every 6–8 h. The protein concentration was adjusted to 750 μg/ml with PBS and then equal portions of the two peptide/OVA conjugates were mixed to then use as antigen. In a separate set of reactions, the peptide was coupled to keyhole limpet hemocyanin (KLH, Thermo Fischer Scientific, Waltham, MA) at a ratio of 200 molecules peptide per molecule KLH using the identical protocols (glutaraldehyde and maleimide-activated-KLH (Imject™ Maleimide-Activated KLH, Thermo Fischer Scientific, Waltham, MA) and equal portions of the resulting two peptide/KLH conjugates were mixed to use as antigen. Rabbits were immunized by Antibodies Inc. (Davis, CA) with the ITLN1 peptide/OVA antigen on days 1, 14, 21, and then boosted by immunization with ITLN1 peptide/KLH antigen on days 35 and 49 to produce the N-terminal polyclonal antiserum (termed ITLN1NTERM). Preimmunization sera was collected and used as control antisera. All methods for these procedures were in accordance with relevant guidelines and regulations, and approved by the U.S. Department of Health and Human Services Public Health Service Animal Welfare Assurance Committee (Assurance ID D16-00576). Rabbit polyclonal antiserum generated by immunization with human ITLN1-derived synthetic peptides was described and characterized previously (termed ITLNCONS)\textsuperscript{11,15}

### Chicken anti-ITLN1 IgY.

The ITLN1 protein sequence was analyzed using the online algorithm for design of peptide-directed antibodies provided by the NHLBI (https://hpcwebapps.cit.nih.gov/AbDesigner/)\textsuperscript{36}. A 28 aa peptide (CTVGDRWSSQQSKADYPEGDGNWANYN) was identified as an attractive antigen candidate. This peptide spanned the V108D polymorphic site, and was highly similar in sequence to human ITLN2 and C57BL/6 mouse Itln1 (27/28 identical residues to both of these orthologs). This peptide was synthetically prepared, purified and characterized\textsuperscript{34}. In two separate reactions, the peptide was coupled to KLH, using the two chemical coupling strategies (glutaraldehyde and maleimide) and protocols, as described above. Each reaction product was dialyzed against PBS and protein concentration adjusted to 750 μg/ml. Equal portions of the two peptide/KLH conjugates were mixed to use as antigen. Leghorn chickens were given a primary inoculation with the KLH conjugate in complete Freund’s adjuvant, followed by two booster inoculations at 3 and 6 weeks (with incomplete Freund’s adjuvant) by Aves Labs, Inc. (Davis, CA). An enriched IgY fraction from egg yolk of immunized chickens (termed ITLNCONS-FLOR\textsuperscript{37}) was obtained by organic extraction de-lipidation, high-salt precipitation and then extensive dialysis against PBS.

### Bio-layer interferometry.

Dissociation constants for recombinant ITLN1 with biotin-β-D-galactofuranose were determined using a bio-layer interferometry OctetRed instrument (ForteBio, Fremont, CA). Biotin-β-D-galactofuranose was diluted to 5 μM in PBS, pH 7.5, and loaded (120 s) onto Streptavidin biosensors (ForteBio). The sensor was washed in PBS for 60 s, and then a baseline (60 s) established in (20 mM HEPES, pH 7.4, 150 mM sodium chloride, 0.1% Tween-20, 0.1% BSA). The shake rate was set to 1000 rpm and binding assays were performed at room temperature. Data was plotted in Graphpad Prism software (version 9.0.0, Graphpad Software, San Diego, CA). The response at equilibrium (490 s) was plotted against concentration of ITLN1 and curves fit to a one-site total non-linear regression equation to determine equilibrium dissociation constants.

### Immunoblotting.

Iideal non-CD and CD specimens, genotyped for ITLN1 rs2274907 (V109D), were homogenized in assay buffer (150 mM NaCl, 1% Triton X-100, 0.5% sodium deoxycholate, 0.1% SDS, 50 mM Tris, pH 8.0) containing a cocktail (1:100) of protease inhibitors (Protease Inhibitor Cocktail III: 4–(2-aminoethyl)benzenesulfonyl fluoride (AEBSF, 1 mM), aprotinin (600 nM), bestatin (50 µM), E-64 (15 µM), leupeptin (20 µM), pepstatin (10 µM); Calbiochem/EMD Millipore, Burlington, MA). Protein concentration was determined by bicinechonic acid (BCA) assay (Peirce, ThermoFisher Scientific). Following heating at 95 °C for 30 min (with or without 2.5% v/v 2-mercaptoethanol for reducing conditions), protein (25–50 μg/lane) was loaded to SDS–polyacrylamide gels (10% and 4–20% acrylamide) and electrophoresed for 40–50 min at 200 V, prior to wet transfer (Towbin Buffer: 25 mM Tris, 192 mM glycine, 20% v/v methanol, pH 8.3) to nitrocellulose membranes (0.2 μm pore, Bio-Rad Laboratories, Hercules, CA) at 350 mA for 60–80 min. Estimated molecular sizes of ITLN1 monomers and higher order oligomers were deduced using the MagicMark™ XP Western
Protein Standard (ThermoFisher Scientific). Membranes were blocked in PBS-T containing 5% w/v skim milk for 30 min, and incubated with primary antibodies at 4 °C. Following overnight incubation, membranes were washed with PBS-T for 1 h, probed with donkey anti-rabbit IgG horseradish peroxidase (HRP) linked secondary antibody (GE Healthcare Amersham, Pittsburgh, PA) for 3 h at room temperature, rinsed in PBS-T, and visualized using Femto and ECL West chemiluminescent substrates (ThermoFisher Scientific). Chemiluminescent signal was detected using a Biospectrum AC Imaging System (UVP, Upland, CA).

**Immunohistochemistry (IHC).** Surgical tissue of human ileum and colon, fixed in Carnoy’s fixative or HistoChoice™ Tissue Fixative (Sigma Aldrich, St. Louis, MO), were paraffin embedded and sectioned (4–5 μM) and mounted on X-tra™ positive-charged slides (Leica Biosystems, Buffalo Grove, IL). Sections were deparaffinized, rehydrated in gradient ethanol, H2O, and PBS and blocked with 5% donkey serum (in 1× PBS) for 30 min prior to overnight incubation at 4 °C with antibody51. Following incubation, slides were washed, and probed with Vectastain ABC kit goat anti-rabbit/goat secondary antibodies (Vector Laboratories, Burlingame, CA), activated with 3,3-diaminobenzidine peroxidase (Vector Laboratories, Burlingame, CA), and counterstained in 0.4% light green (Sigma Aldrich, St. Louis, MO). 0.2% glacial acetic acid for 5 min prior to dehydration and mounting51. Samples were visualized using an Olympus BX51 microscope (Center Valley, PA) and images were processed using Adobe Creative Suite software (version 25, Adobe Inc., San Jose, CA).

**Fluorescence IHC.** Specimen slides were prepared as before. Following overnight incubation with primary antibodies, samples were incubated with two secondary antibodies (goat anti-chicken IgY Alexa Fluor Plus 647 [Thermo Fischer Scientific, Waltham, MA] and goat anti-rabbit IgG Alexa Fluor 488 [Abcam, Cambridge MA]), rinsed, and then stained with DAPI using the TrueVIEW Autofluorescence Quenching Kit (Vector Laboratories, Burlingame, CA) as outlined by the manufacturer. Images were captured using a Leica SP8 STED 3× microscope (Leica Microsystems Inc., Buffalo Grove, IL). Acquired Z-stacks were further processed to generate figure images using Fiji ImageJ software (version 1.0, https://imagej.net/Fiji)57.

**Statistical analysis.** Statistical analysis was performed using Graphpad Prism software (version 9.0.0, Graphpad Software, San Diego, CA). Specific details are outlined in figure and table legends.

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**Author contributions**

E.B.N., B.L., and C.L.B. designed research; E.B.N., P.A.C., A.E.D., F.A. and W.Y. performed research; C.A.D.L.M., W.Y., W.L., B.S., and M.E.V.J. contributed new reagents/analytic tools; E.B.N., P.A.C., A.E.D., M.A.U., C.A.D.L.M., B.S., M.E.V.J., L.L.K., E.J.H., B.L., and C.L.B. analyzed data; and E.B.N. and C.L.B. wrote the paper. All authors reviewed the manuscript.

**Competing interests**

The authors declare no competing interests.

**Additional information**

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