Genetic Manipulation of Human Intestinal Enteroids Demonstrates the Necessity of a Functional Fucosyltransferase 2 Gene for Secretor-Dependent Human Norovirus Infection

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ABSTRACT Human noroviruses (HuNoVs) are the leading cause of nonbacterial gastroenteritis worldwide. Histo-blood group antigen (HBGA) expression is an important susceptibility factor for HuNoV infection based on controlled human infection models and epidemiologic studies that show an association of secretor status with infection caused by several genotypes. The fucosyltransferase 2 gene (FUT2) affects HBGA expression in intestinal epithelial cells; secretors express a functional FUT2 enzyme, while nonsecretors lack this enzyme and are highly resistant to infection and gastroenteritis caused by many HuNoV strains. These epidemiologic associations are confirmed by infections in stem cell-derived human intestinal enteroid (HIE) cultures. GII.4 HuNoV does not replicate in HIE cultures derived from nonsecretor individuals, while HIEs from secretors are permissive to infection. However, whether FUT2 expression alone is critical for infection remains unproven, since routinely used secretor-positive transformed cell lines are resistant to HuNoV replication. To evaluate the role of FUT2 in HuNoV replication, we used CRISPR or overexpression to genetically manipulate FUT2 gene function to produce isogenic HIE lines with or without FUT2 expression. We show that FUT2 expression alone affects both HuNoV binding to the HIE cell surface and susceptibility to HuNoV infection. These findings indicate that initial binding to a molecule(s) glycosylated by FUT2 is critical for HuNoV infection and that the HuNoV receptor is present in nonsecretor HIEs. In addition to HuNoV studies, these isogenic HIE lines will be useful tools to study other enteric microbes where infection and/or disease outcome is associated with secretor status.

IMPORTANCE Several studies have demonstrated that secretor status is associated with susceptibility to human norovirus (HuNoV) infection; however, previous reports found that FUT2 expression is not sufficient to allow infection with HuNoV in a variety of continuous laboratory cell lines. Which cellular factor(s) regulates susceptibility to HuNoV infection remains unknown. We used genetic manipulation of HIE cultures to show that secretor status determined by FUT2 gene expression is necessary and sufficient to support HuNoV replication based on analyses of isogenic lines that lack or express FUT2. Fucosylation of HBGAs is critical for initial binding and for modification of another putative receptor(s) in HIEs needed for virus uptake or uncoating and necessary for successful infection by GI.1 and several GII HuNoV strains.

KEYWORDS fucosyltransferase 2, glycobiology, histo-blood group antigens, isogenic enteroids, isogenic organoids, noroviruses, secretor status
Human noroviruses (HuNoVs) are a leading cause of nonbacterial gastroenteritis worldwide. From the first recognition in 1968 that a virus caused an outbreak of HuNoV gastroenteritis in an elementary school in Norwalk, Ohio (1), until 2016, there was no in vitro culture system of HuNoV in intestinal epithelial cells. A novel HuNoV culture system using human intestinal enteroids (HIEs) generated from stem cells isolated from human small intestinal crypts is now available and is being used worldwide to study virus replication, inactivation, and neutralizing antibodies (2–9).

Secretor status is highly associated with infection and disease caused by many HuNoV strains based upon findings from human experimental infection studies and evaluation of acute gastroenteritis outbreaks (10–14). Fucosyltransferase 2 (FUT2) is an enzyme expressed in human epithelial cells that catalyzes α1,2-fucosylation of the terminal galactose, preferentially on glycan type 1 chain precursors. Glycosyltransferases coded for by FUT2 along with FUT3 and ABO genes determine the histo-blood group antigens (HBGAs) found on the epithelial cell surface (15, 16). Persons who lack functional FUT2 alleles do not express ABH HBGAs on epithelial cells, are designated nonsecretors, and are highly resistant to gastroenteritis caused by some HuNoV strains such as GII.4 viruses. Similarly, HIEs derived from nonsecretor individuals are not susceptible to GII.4 HuNoV infection (2). HIEs and gastrointestinal epithelial cells of secretor-positive individuals also bind norovirus virus-like particles (VLPs) (17, 21). However, while HIEs from secretors are permissive to HuNoV replication, FUT2 expression in conventional cancer-derived cell lines is not sufficient to make cells susceptible to HuNoV replication, suggesting that HBGAs function primarily as an initial attachment factor (18, 19). These data also lead to questions on whether there are additional genetically determined differences in HuNoV susceptibility in addition to secretor status. In this study, we evaluated the direct association between FUT2 function and HuNoV infectivity using HIEs with or without FUT2 expression in the same genetic background.

RESULTS

Generation and characterization of isogenic HIE lines. We previously showed that GII.4 viruses can replicate in HIEs derived from secretor-positive individuals and a GII.3 strain can replicate in both secretor-positive and some secretor-negative lines, recapitulating observations seen in epidemiologic studies (2). We determined the FUT2 (Se) and FUT3 (Le) genotypes of jejunal HIE lines and selected two lines (J2 and J4); both lines express FUT3, eliminating Lewis status as a variable in these studies (Table 1). The J2 cell line is heterozygous wild type (Se/Se428) ([J2Fut2+/−]); because the Se gene is autosomal dominant, the cells are secretor positive. The J4 line has a homozygous se428/se428 ([J4Fut2−/−]) recessive mutation and is secretor negative (20). No other mutations associated with loss of or decreased FUT2 enzymatic function were observed in the J2 or J4 line. To better understand the importance of FUT2 in HuNoV infection, we generated isogenic knockout (KO) and knock-in (KI) HIE lines. A lentivirus-delivered CRISPR/Cas9 construct with a guide RNA targeting the Fut2 coding region was used to knock out Fut2 from J2 ([J2Fut2−/−]). Single cell selection under puromycin pressure was used to isolate a clonal population, and deletions of the gene in both alleles were confirmed by sequencing. To generate the KI Fut2 line, we transduced a functional Fut2 coding sequence driven by a cytomegalovirus (CMV) promoter in a lentivirus into J4 ([J4Fut2−/−/FUT2]) cells.

First, we confirmed the phenotype of the Fut2 KO and KI lines by evaluating HBGA expression using an enzyme-linked immunosorbent assay (ELISA) (Fig. 1). J2Fut2+/− cells expressed the secretor-positive glycans, Leb and B, as expected for this secretor-positive, Lewis-positive O8 HIE line. KO of Fut2 altered the J2 phenotype such that the Leb and B glycans were no longer present, and only Lea was detected. J4Fut2+/− cells expressed Lea but not Leb or other secretor glycans, confirming the secretor-negative genotype of this line. KI of Fut2 in J4 cells led to Leb instead of Lea expression.

We next used immunofluorescence microscopy with fluorescently labeled Ulex europaeus agglutinin-1 (UEA-1 lectin) to detect polarized cell surface expression of
Both J2Fut2+/− and J4Fut2−/−/FUT2 cells had apical staining of UEA-1 lectin. This apical staining was lost in J2Fut2−/− and J4Fut2−/− cells (Fig. 2A). These findings indicate that, as expected, terminal fucosylation of the glycan precursor in HIEs depends on the expression of the FUT2 gene. In the secretor-negative lines, unexpected internal cellular UEA-1 staining was present that was not observed in the secretor-positive lines. To confirm that the internal staining was due to specific UEA-1 recognition of α1,2-fucose and not off-target binding due to loss of the specific ligand, we preincubated the UEA-1 lectin with L-fucose prior to staining. When UEA-1 was preincubated with 10 mM L-fucose, staining was not detected in the secretor-negative lines and was significantly reduced in the secretor-positive lines (Fig. 2B). After preincubation with 100 mM L-fucose, UEA-1 staining was not detected in either the secretor-negative or secretor-positive lines (Fig. 2C). Together, these data indicate that the apical and the internal staining observed with UEA-1 is due to specific interaction with α1,2-fucose.

Table 1 summarizes the genotypic and phenotypic findings for the parental, KO and KI cell lines.

**Effect of FUT2 expression on replication of HuNoV strains in isogenic HIE cell lines.** To assess whether these isogenic cell lines support HuNoV replication, we evaluated virus binding and subsequent replication of GII.4, GII.3, GII.17, and GI.1 HuNoV strains that we previously demonstrated replicate in HIEs (2). Since all of these viruses but GII.4 require bile for replication, all infections were performed in the presence of glycochenodeoxycholic acid (GCDCA), a bile acid that supports replication (21).

First, we assessed the effect of KO of FUT2 from J2 cells with the four HuNoV strains (Fig. 3). After 1 or 2 h postinfection (hpi) and washing off the inoculum, GII.4, GII.3, and

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**TABLE 1 Genotyping and phenotyping of HIE lines used in this study**

| HIE line | HIE line modificationa | Genotyping resultsb | Phenotyping results |
|----------|-----------------------|--------------------|--------------------|
| J2Fut2+/− | Not modified          | Se, se^a28         | Le, Le             | Positive, B, Le^b |
| J2Fut2−/− | CRISPR-Cas9 deletion | se^e6, se^Δ        | Le, Le             | Negative, Le^a   |
| J4Fut2−/− | Not modified          | se^e28, se^e28     | Le, le^202,314, Le | Negative, Le^a   |
| J4Fut2−/−/FUT2 | LV Td CMV expression | se^e28, se^e28, Se^CMV | Le, le^202,314, OO | Positive, Le^b   |

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aLV Td, lentivirus transduced.
bSe, secretor; Le, Lewis; Δ, deletion. Se^CMV indicates that the Se gene was expressed under a CMV promoter. Specific mutations in secretor and Lewis genes are indicated by the superscripts.

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**FIG 1** HBGA expression phenotype by ELISA. Five-day differentiated HIEs were evaluated for HBGA expression with primary antibodies against either the Le^a, Le^b, A, or B epitope and HRP-conjugated secondary antibodies. Mean absorbance values from two ELISA replicates are plotted. Error bars denote standard deviations (SD). The threshold of detection is indicated by a dashed line at absorbance of 0.1.
GI.1 binding was significantly reduced in the J2Fut2−/−/H11002 cells compared to the parental J2Fut2+/+/H11001 cells. GII.17 binding was slightly, but not significantly, reduced in several experiments. Viral replication was completely abrogated at 24 hpi for GII.4, GII.17, and GI.1 in the J2Fut2−/−/H11002HIEs. GII.3 was able to replicate in the J2Fut2−/−/H11002 line, consistent with our previous findings that GII.3 is able to replicate in some secretor-negative lines (2). These findings support the requirement for functional FUT2 and secretor-positive HBGA as an initial binding factor for several strains of HuNoV that is critical for replication.

Next, we assayed the J4Fut2−/−/FUT2 line to determine whether a genetically resistant HIE line can become susceptible with the expression of FUT2. We observed increased HuNoV binding at 1 or 2 hpi in the J4Fut2−/−/FUT2 line compared with the parental nonsecretor J4Fut2−/− line for all four HuNoV strains (Fig. 4); however, for GII.4, the increase was not significant. J4Fut2−/−/FUT2 was permissive to infection by all four viruses, based upon increases in viral genome equivalents (GEs) at 24 hpi. Interestingly, although GE levels at 1 to 2 hpi were similar, we observed greater increases in GEs for GII.17 and GI.1 HuNoVs in J4Fut2−/−/FUT2 cells than in J2Fut2−/− cells at 24 hpi. This suggests that additional factors that vary between individual HIE lines, aside from secretor status, may influence the replication of these two strains. Taken together, these results show that functional FUT2 is sufficient and critical for replication of multiple HuNoV strains and that GI.3 is capable of infection in some secretor-negative HIE lines.

DISCUSSION

Previous studies demonstrated that the HuNoV genome is capable of productive infection in transformed cell lines (e.g., Huh-7) following transfection. Overexpression of the FUT2 gene increased virus binding to the cells but did not make them susceptible...
These data indicated the presence of a block between virus binding and entry. The development of the HIE HuNoV infection model identified that enterocytes in these cultures are infected with HuNoVs and allowed the performance of studies to evaluate factors important for virus infection. The current study demonstrates that FUT2 alone is necessary and sufficient for the infection and replication of secretor-dependent GI.1, GII.4, and GII.17 HuNoVs in HIE cells.

Virus attachment and entry into the infected cell are complex processes, and the exact role of fucosylated molecules in HuNoV entry remains to be determined. It is likely that initial binding of secretor-dependent HuNoVs to HIEs is regulated by fucosylation, with the attachment factors being fucosylated. However, it is unclear whether the fucosylated molecules serve only as the initial attachment factor that then facilitates interaction with a virus-specific receptor or whether the fucosylated molecules function as a virus receptor themselves; examples of both mechanisms have been described previously with nonfucosylated glycans. For example, reovirus binding to its proteinaceous JAM-A receptor is enhanced by initial binding to sialic acid residues, while human coronaviruses OC43 and HKU1 bind to 9-O-acetylated sialic acid receptors, and influenza viruses bind to terminal sialic acid receptors. If fucosylated molecules serve only as attachment factors, then the virus receptor(s) for secretor-dependent HuNoVs is present in both secretor-positive and secretor-negative HIEs and is not expressed in other nonsusceptible, secretor-positive cancer-derived or trans-FIG 3 Knocking out FUT2 prevents infection of GI.4, GI.17, and GI.1 HuNoV strains in J2 HIEs. HIE monolayers were inoculated with GI.4, GI.3, GI.17, or GI.1 HuNoV stool filtrate in 500 μM GCDCA-containing Intesticult medium for 1 h (GI.4, GI.3, and GI.17) or 2 h (GI.1) at 37°C. After two washes with CMGF(-) medium, the cells were cultured in the presence of GCDCA for 24 h at 37°C. Total well RNA was extracted, and genome equivalents (GEs) were determined by RT-qPCR. Each data bar represents the mean for three wells of inoculated HIE monolayers. Error bars denote SD. Each experiment was performed two or more times, with three technical replicates in each experiment. Data from a representative experiment are shown in this figure. Numbers above the bars indicate log10 fold change comparing GEs at 24 h postinfection (hpi) to 1 or 2 hpi. Significance was determined by two-way ANOVA with post hoc analysis using Tukey’s test (*, P < 0.05; n.s., not significant).
formed cell lines. On the other hand, if the fucosylated molecules function as the virus receptor, it is unclear why secretor-positive cultured transformed cells derived from cancer patients are not susceptible to infection. It is possible that transformed lines do not properly express the needed receptors or glycans or other signaling pathways essential for infection. Transformation may abrogate proper glycosylation in non-susceptible cell lines compared to nontransformed HIE lines (26).

We observed a striking phenotypic difference in FUT2-expressing HIE cells compared to cells not expressing FUT2. UEA-1 detection of α1,2-fucose was observed exclusively at the apical surface in FUT2-positive cells. Unexpectedly, in the secretor-negative lines, there was internal cellular staining by UEA-1. The staining was due to UEA-1-specific interactions with α1,2-fucose, indicating that these glycan structures are present but unable to transit to the surfaces of the secretor-negative HIE cultures. There is evidence that FUT2 expression leads to surface expression of α1,2-fucosylated molecules. This was shown in specific-pathogen-free mice, where the lumen of the small intestine is mostly lacking in surface fucosylation, and intraperitoneal injection of lipopolysaccharide (LPS) stimulates expression of α1,2-fucosylated molecules at the surface in a FUT2-dependent manner (27, 28). There may be additional fucosyltransferase enzymes present in our HIEs capable of adding fucose to glycoproteins or glycolipids in the absence of FUT2 activity. However, these alternatively fucosylated molecules are unable to transit to the cell surface. Fucosyltransferase 1 (FUT1), expressed in erythroid cells and some other tissues, is also capable of adding α1,2-fucose, although preferentially on glycan chains other than those found in intestinal epithelia. In the parental J2 enteroid line used in this study, FUT1 transcripts are expressed by transcriptome sequencing (RNA-seq), and the fragment per kilobase per million (FPKM) value is
Table 2: Primers used in this study

| Gene       | Primer | Template strand | Sequence (5’→3’)                          | Reference          |
|------------|--------|-----------------|------------------------------------------|--------------------|
| FUT2       | FUT2-280F | +               | AGCCCTCAACATCAAGGCGACTGGGA               | Saxena et al. (35) |
|            | FUT2-564R | –               | AACCAGTCAGGACGCTTGTA                     | Saxena et al. (35) |
|            | FUT2-97F  | +               | ATGGGCCCACCTTCTATCC                     | This study         |
|            | FUT2-1095R | –              | TTATGCTTGTAGAAGGGGAG                 | Ito et al. (36)    |
|            | BO-119F   | +               | GGCATGGAAGAGATCAAG                      | This study         |
|            | BO-120R   | –               | TCTTCAGGTCTGAGTTC                       | This study         |
|            | FUT3      | FUT3-505F       | +                                       | This study         |
|            | FUT3-888R | –               | TCAGCGCCTCTGAGTACG                      | This study         |
|            | FUT3-840F | +               | CAACTGGAAGCCAGACTCA                     | This study         |
|            | FUT3-1485R | –            | CAGCAAGCTCCCTCGAGGG                     | This study         |
| ABO exon 6 | ABO-4522F | +               | CAGAGCTGAGTGGGATTC                    | This study         |
|            | ABO-6R    | –               | CTGGTGGAGATGCGAAGTTG                    | Muro et al. (37)   |
| ABO exon 7 | ABO-6037F | +               | TCCTCAGAGGGCTTTATCA                     | This study         |
|            | ABO-6378R | –               | AGCACTTGGTGGGTT                        | This study         |

~60-fold lower than that of FUT2. In bovine coronary venular endothelial cells, a lotus lectin (LTL) that also recognizes α1,2-fucosylated molecules detects cytoplasmic tubule structures, and this staining is lost after depletion of both FUT1 and FUT2 (29). Further studies showed that LTL coostained with a Golgi marker in newly derived primary human fibroblasts from oral mucosa, but this Golgi colocalization was lost and LTL was instead detected in tubule structures as the fibroblasts were passaged. In our system, the presence of some FUT1 expression may allow for α1,2-fucose-containing structures intracellularly. Future studies will determine the identity and subcellular location of the fucosylated molecules present in the FUT2-negative HIE lines and whether FUT1 is required for their presence.

HIEs provide an excellent tool for future studies on intestinal enzymes involved in glycosylation and how glycosylation alters glycoprotein localization. An association of enteric commensals and pathogens with host secretor status has led to increased recognition of secretor glycans being susceptibility factors important in infection and disease outcomes (30, 31). The exact role played by the glycans in these infections is not fully understood. Our isogenic, physiologically active HIE lines should be helpful to determine whether fucosylation plays a role in microbe binding, entry, or postentry processes that may affect the epithelial responses to infection, and it will be interesting to understand different outcomes of infection with the different microbes.

Materials and Methods

Plasmid constructs. The CDNA of FUT2, obtained from J2 human intestinal enteroid (HIE) total RNA, was amplified with specific gene primers (Table 2) and cloned into the lentiviral expression vector pLVSin-ires-puromycin (32) using In-Fusion cloning kit (TaKaRa-Clontech) according to the manufacturer’s instructions. A Cas9-expressing lentiviral vector (lentiCas9-BLAST; plasmid no. 52962) and single guide RNA (sgRNA) expression vector (lentiGuide-puro; plasmid no. 52963) were purchased from Addgene. The sgRNA sequence targeting the human FUT2 gene (5’-CCACCCAGCTCAGGGGATG-3’) was cloned into the lentiguide-puro vector following the manufacturer’s protocol.

Lentivirus packaging and production. A third-generation lentivirus carrying the FUT2 gene (Lv-FUT2) was produced by cotransfecting HEK293T cells with a combination of a lentivirus plasmid (pLVsin-FUT2-ires-puro, lentiCas9-BLAST, or lentiGuide puro) and three packaging plasmids (pMDLg/prRE [plasmid no. 12251; Addgene], envelope plasmid pMD2.G [plasmid no. 12259; Addgene], and pRSV-Rev [plasmid no. 12253; Addgene]) at a ratio of 3:5:2:1:1, respectively, using polyethylenimine HCl Max molecular weight (MW) 40,000 (Polysciences) (32). The culture supernatant was harvested 60 to 72 h posttransfection, passed through a 0.45-μm filter, concentrated by using Lentix-concentrator (TaKaRa-Clontech) according to the manufacturer’s protocol, and suspended in CMGF(+) medium for transduction.

Lentiviral transduction of HIEs. A cell suspension was prepared from three-dimensional (3D) undifferentiated jejunal HIEs cultivated as previously described (2, 33). After trypsinization and pelleting of the cells at 300 × g, the resulting cell pellet was suspended at a concentration of 3 × 10^5 cells per ml of concentrated lentivirus supplemented with 10 μM Rho-associated protein kinase (ROCK) inhibitor Y-27632 (catalog no. Y0503; Sigma) and 8 μg/ml Polybrene (catalog no. TR-1003-G; EMD Millipore). The mixture was plated in one well of a 48-well plate. The plate was then centrifuged for 1 h at 300 × g at room temperature (RT). After spinoculation, the lentivirus solution was removed, the cells were washed once with CMGF(−) medium, centrifuged again, embedded in 30-μl Matrigel plug, and incubated at 37°C.
and 5% CO₂ in the presence of CMGF(+) medium with ROCK inhibitor for recovery. Five days posttransduction, the cells were treated with puromycin (2 µg/ml) or Blasticidin S (5 µg/ml) until mock-treated cells were completely dead. Single cells were isolated by serial dilution in 96-well plates for sgRNA-transduced HIEs, and deletion of the gene was confirmed by sequencing of genomic DNA from each single cell clone using primers BO-119 and BO-120 (Table 2) that amplified the portion of the FUT2 gene targeted by the sgRNA.

HIE HBGA phenotyping. Differentiated HIE cells were suspended in phosphate-buffered saline (PBS) and boiled for 5 min. After incubation of the boiled supernatants on vinyl, flat-bottomed, 96-well plates (ThermoFisher Scientific) for 4 h at RT and blocking with 10% Carnation nonfat dry milk overnight at 4°C, anti-Le⁺ Gamma-clone (Immucor), anti-Le⁻ (BG-6; Biolegend), anti-A type Gamma-clone (Immucor), or anti-B type Gamma-clone (Immucor) was used as the primary antibody, and horseseradish peroxidase (HRP)-conjugated goat anti-mouse IgG (Sigma) was used as the secondary antibody. 3,3',5,5′-Tetramethylbenzidine (TMB) (KPL) was used as a HRP substrate, and the reaction was stopped with 1 M phosphoric acid (Fisher Scientific) after a 10-min incubation at RT. The absorbance was determined at 450 nm with a Spectramax 190 plate reader (Molecular Devices). Pooled saliva samples from persons who collectively express each histo-blood group antigen (HBGA) type was evaluated using a positive control (34). Saliva from an individual negative for the HBGAs was used as a negative control.

HIE HBGA genotyping. DNA was extracted from HIEs and primers (Table 2) targeting exons 6 and 7 of the ABO gene, the FUT2 (secretor) gene, and the FUT3 (Lewis) gene were used to generate amplicons that were purified using the GeneJET PCR purification kit (ThermoFisher Scientific) and sequenced (GeneWiz). Chromatograms were examined to identify single nucleotide polymorphisms associated with loss of function (for FUT2, A385T, G428A, C571T, and C628T; for FUT3, T59G, T202C, C314T, G484A, G508A, G667A, G808A, and T1067A or A, B, or O genotype (for exon 6, nucleotides 261 and 297; for exon 7, nucleotides 657, 703, 771, 796, 803, 829, and 930).

Immunofluorescence and quantitation. HIE monolayers were grown in glass bottom plates (catalog no. 655892; Greiner Bio-One), differentiated for 5 days, and fixed with 4% paraformaldehyde (PFA) for 20 min at RT. HIE monolayers were permeabilized and blocked with 5% bovine serum albumin (BSA) in 0.1% Triton X-100 in PBS for 30 min at RT. All the subsequent steps were performed in PBS plus 0.1% Triton X-100. HBGAs and cell boundaries were detected after overnight incubation at 4°C with rhodamine-labeled Ulex europeus agglutinin-1 (UEA-1) (1:1,000) (catalog no. RL-1062; Vector Laboratories) and Alexa Fluor 647 phalloidin (ThermoFisher Scientific), respectively. For the fucose inhibition assay, the UEA-I was incubated with different concentrations of α-fucose at RT for 1 h prior to staining. Nuclei were stained with 4,6-diamidino-2-phenylindole (DAPI) (300 nM) for 5 min at RT. Orthogonal 5-µm-thick sections of the sample were captured using a Zeiss LSM 510 confocal microscope. For quantifying fluorescence intensity, five fields per well were analyzed. The fluorescence threshold in these images was set in Image J. Mean fluorescence data from 15 identical regions of interest (ROIs) per field were collected. Comparisons between treatment groups were made using a Student’s t test. P values of <0.05 were considered statistically significant.

In vitro HuNoV infection. Jejunal HIE monolayers in 96-well plates were differentiated for 5 days in commercial Intesticult (INT) human organoid growth medium (Stem Cell Technologies) and inoculated with the indicated positive HuNoV stool filtrates at 37°C for 1 h (GI/Hu/US/2012/GI.4 Sydney [P31]/TCH12-580, 9 × 10⁶ genome equivalents [GEs]/well; GI/Hu/US/2004/GI.3 [P21]/TCH04-577, 4.3 × 10⁶ GEs/well; GI/Hu/US/2014/GI.17 [P38]/TCH14-385, 1.8 × 10⁶ GEs/well) or for 2 h (GI/Hu/US/2006/GI.1 Norwalk [P11]/BCM/723-02, 6.9 × 10⁵ GEs/well) as described previously (2). Each infection was performed in triplicate wells for each time point, and conditions were tested in at least two independent experiments. Inocula were removed, and monolayers were washed twice with CMGF(−) medium to remove unbound virus. Differentiation INT medium (100 µl supplemented with 500 µM glycochenodeoxycholic acid [GCDCA] (Sigma)) was then added to each well, and the cultures were incubated at 37°C for the indicated time points. RNA was extracted from each well using the KingFisher Flex Purification system and MagMAX-96 viral RNA isolation kit. RNA extracted at 1 to 2 hpi was used as a baseline to determine the amount of input virus that remained associated with cells after the infected cultures were washed to remove unbound virus. Replication of virus was determined by HuNoV RNA levels, which were quantified using a standard curve based on a recombinant HuNoV RNA transcript, and replication of virus was determined by assessing changes in virus GE levels at 24 hpi from baseline.

Statistics. All statistical analyses were performed on GraphPad Prism (GraphPad Software, La Jolla, California USA). Samples with RNA levels below the limit of detection of the reverse transcription-quantitative PCR (RT-qPCR) assay were assigned a value that is one-half the limit of detection of the assay. Comparisons between infection time point groups or infected cell lines were made using two-way analysis of variance (ANOVA) with Tukey’s test for post hoc analyses. P values of <0.05 were considered statistically significant.

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