



# Human Sapovirus Replication in Human Intestinal Enteroids

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**ABSTRACT** Human sapoviruses (HuSaVs), like human noroviruses (HuNoV), belong to the *Caliciviridae* family and cause acute gastroenteritis in humans. Since their discovery in 1976, numerous attempts to grow HuSaVs *in vitro* were unsuccessful until 2020, when these viruses were reported to replicate in a duodenal cancer cell-derived line. Physiological cellular models allowing viral replication are essential to investigate HuSaV biology and replication mechanisms such as genetic susceptibility, restriction factors, and immune responses to infection. In this study, we demonstrate replication of two HuSaV strains in human intestinal enteroids (HIEs) known to support the replication of HuNoV and other human enteric viruses. HuSaVs replicated in differentiated HIEs originating from jejunum, duodenum and ileum, but not from the colon, and bile acids were required. Between 2h and 3 to 6 days postinfection, viral RNA levels increased up from 0.5 to 1.8 log<sub>10</sub>-fold. Importantly, HuSaVs were able to replicate in HIEs independent of their secretor status and histo-blood group antigen expression. The HIE model supports HuSaV replication and allows a better understanding of host-pathogen mechanisms such as cellular tropism and mechanisms of viral replication.

**IMPORTANCE** Human sapoviruses (HuSaVs) are a frequent but overlooked cause of acute gastroenteritis, especially in children. Little is known about this pathogen, whose successful *in vitro* cultivation was reported only recently, in a cancer cell-derived line. Here, we assessed the replication of HuSaV in human intestinal enteroids (HIEs), which are nontransformed cultures originally derived from human intestinal stem cells that can be grown *in vitro* and are known to allow the replication of other enteric viruses. Successful infection of HIEs with two strains belonging to different genotypes of the virus allowed discovery that the tropism of these HuSaVs is restricted to the small intestine, does not occur in the colon, and replication requires bile acid but is independent of the expression of histo-blood group antigens. Thus, HIEs represent a physiologically relevant model to further investigate HuSaV biology and a suitable platform for the future development of vaccines and antivirals.

**KEYWORDS** sapovirus, norovirus, human intestinal enteroids, histo-blood group antigens, enteric virus, enteric viruses

*Sapovirus* (SaV) is a genus of small RNA viruses within the *Caliciviridae* family, which also contains the *Norovirus* (NoV) genus. HuSaV was first discovered in 1976, by electron microscopic observation of fecal samples from patients with gastroenteritis in the United Kingdom (1), but the most studied strain originated in the early 1980s, from samples from children in a Japanese orphanage located in Sapporo (2). HuSaVs contain a single-stranded, positive-sense, poly-A tailed RNA genome of 7.1 kb to 7.7 kb, encoding two open reading frames (ORF) that are packaged in an icosahedral capsid ranging from 30 to 38 nm in diameter (3). ORF1 encodes a polyprotein that is proteolytically cleaved into at least six nonstructural proteins and possibly the major structural protein VP1 (4),

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which self-assembles to form a particle of 180 copies of VP1 (5). ORF2 encodes a minor structural protein, VP2 (6). A subgenomic RNA comprising the coding sequence of VP1 and VP2 has been identified in a porcine strain; thus, VP1 may arise from the cleavage of the polyprotein, or from the translation of this subgenomic RNA (3). *Caliciviridae* share a capsid structure, with VP1 divided in two domains: a S domain, mostly conserved between strains, and a P domain, with a subdomain P2 displaying the most variable region in amino acid sequences, primarily accounting for antigenic and immunogenic diversity (5). VLP production does not require VP2 owing to the self-assembling properties of VP1 dimers, and HuSaV VP2 structure and function are still largely uncharacterized. HuSaVs are classified into four genogroups: GI, GII, GIV and GV (7). Other SaV genogroups have been identified in various animal species such as minks, bats, sea lions, dogs, rodents, or chimpanzees (8).

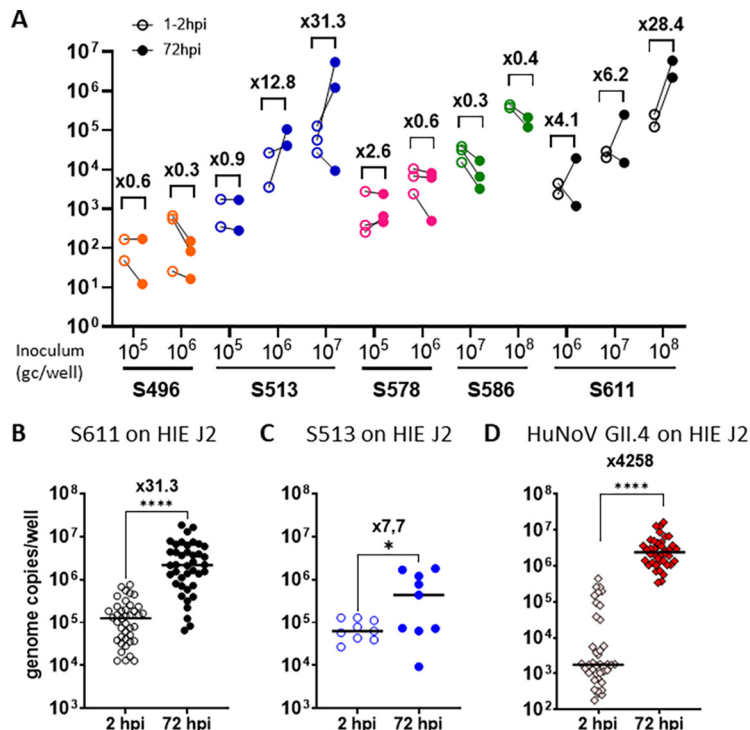
HuSaV infection causes classical acute gastroenteritis (AGE) symptoms that are considered milder or similar to those caused by HuNoV and human rotavirus (RV) (9). Molecular detection is required to properly identify the causative pathogen and is infrequently performed for HuSaVs. Yet, HuSaVs are one of the common causes of viral gastroenteritis globally, causing ~3.4% of cases based on a meta-analysis of 106 studies around the world (10). In recent studies, HuSaVs have been detected more frequently than human RV in countries with childhood vaccination programs targeting RV (11–13). Infection is likely to occur across the human population at a young age, usually before 2 years (14, 15). Viral progeny is excreted in feces, up to  $10^{11}$  copies per grams of stool (16), and HuSaV transmission occurs through the fecal-oral route via contact with HuSaV-positive feces, vomitus, contaminated surfaces, contaminated food and/or drinks. Yet, as opposed to the marked winter seasonality of HuNoV in the Northern Hemisphere, seasonality is less apparent for HuSaV. Studies on different continents show differing seasonal patterns, but correlation with high rainfall, low temperature, and floods is suspected (9). In sewage, HuSaV may be detected year-round (17).

Similar to HuNoV, *in vitro* replication of HuSaV was a challenge for more than 40 years from when the virus was first isolated. A porcine sapovirus (PoSaV), the Cowden strain, can be cultivated in porcine LLC-PK1 cells in a bile-dependent manner (18). More recently, Takagi et al. succeeded in propagating GI.1 and GII.3 HuSaV strains in HuTu80 (duodenum) and NEC8 (ileocecum) cancer cell lines, in the presence of bile acids, further demonstrating the important role of this cofactor in HuSaV infection (19). However, due to the previous lack of a reliable culture system, there is still little information about the HuSaV viral life cycle and tropism.

Studies of tissues obtained from infected piglets suggest that replication of the PoSaV Cowden strain occurs in the small intestine, especially in the duodenum and jejunum (20). PoSaV requires sialic acids linked to O-glycoproteins as receptors, but the HuSaV tropism and receptor(s) remain unknown (21). Regarding HuNoV, epidemiological studies have demonstrated that some HuNoV strains are, at least partially, restricted to people with the secretor phenotype (22), who express histo-blood group antigens (HBGAs) in their intestinal mucosa. Most HuNoV strains bind to HBGAs (23, 24). Conversely, HuSaV has not previously been shown to interact with HBGAs (25).

Human intestinal enteroids (HIEs) have become an essential tool to study HuNoV replication in human cells, overcoming decades of unsuccessful attempts at creating a cellular model allowing culture of these viruses (26). HIEs are derived from LGR5+ stem cells isolated from intestinal crypts, maintained in a 3-dimensional environment, and can be grown to form multicellular structures recapitulating small intestine tissues or as two-dimensional monolayers. Using this physiological model has allowed a better understanding of HuNoV–host interactions and confirmed the dependency of several HuNoV strains for the expression of HBGA to initiate successful infection (27–30). This model also allows replication of enteric viruses, including rotavirus, enterovirus, coronavirus, adenovirus, astrovirus (31).

Here, we undertook the study to evaluate whether HIEs support the replication of HuSaVs, to examine whether replication requires bile acids, and to investigate HuSaV

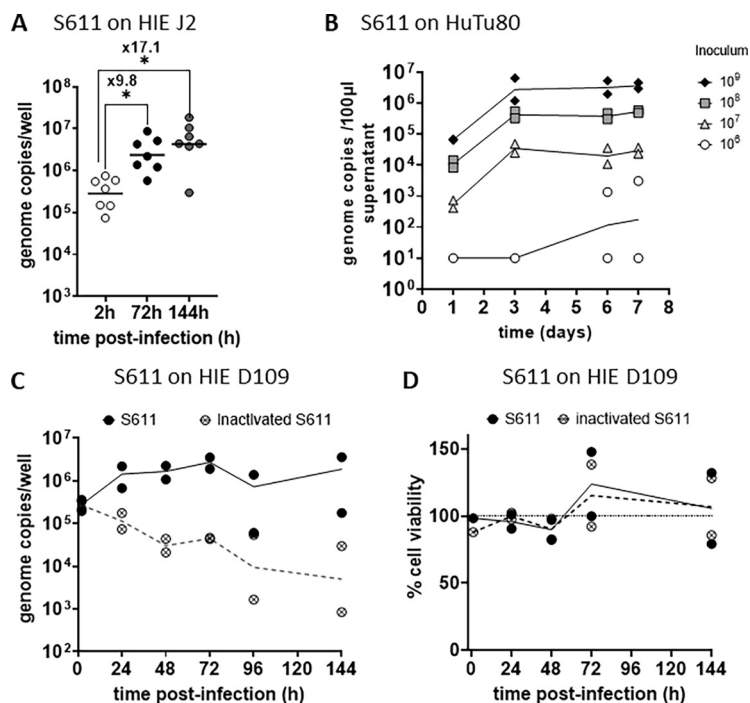


**FIG 1** Two human sapovirus strains replicate in jejunal human intestinal enteroids (HIE). (A) Five HuSaV-positive stools filtrates were used to infect differentiated monolayers of HIE J2 cells in the presence of 500  $\mu$ M bile acid GCDCA, with inocula of  $1.10^5$  to  $1.10^8$  HuSaV genome copies per well (horizontal axis). HuSaV genome quantities in each well (vertical axis) were measured by qRT-PCR at 1 or 2h (empty circles) and 72h (full disks) postinfection (pi). Differences between HuSaV titers at 2 and 72 hpi were not statistically significant,  $n = 2$  to 3 experiments,  $P > 0.05$ , Wilcoxon test. (B) An inoculum of  $1 \times 10^8$  gc/well of HuSaV strain S611 was used in repeated experiments to infect differentiated J2 monolayers with 500  $\mu$ M GCDCA and showed a mean genome fold change of 31.3 ( $n = 40$  experiments,  $P < 0.0001$ , Wilcoxon test). (C) An inoculum of  $1 \times 10^7$  gc/well of HuSaV strain S513 was also used to infect differentiated J2 monolayers with 500  $\mu$ M GCDCA and showed a geometrical mean genome amplification of 7.7 ( $n = 9$  experiments,  $P = 0.012$ , Wilcoxon test). (D) In parallel with HuSaV infections, differentiated J2 monolayers were also infected with  $1.10^5$  genome copies per well of HuNoV GII.4 TCH11-64 as a control with 500  $\mu$ M GCDCA and showed efficient replication with a geometric mean fold increase of 4258 ( $n = 40$  experiments,  $P < 0.0001$ , Wilcoxon test). (A, B, C, D) each dot is the mean of three technical replicates for one independent experiment at 2h (empty circles or pink diamonds) and 72h postinfection (plain disks and red diamonds), horizontal lines are the geometrical mean of virus genome copies per well considering all experiments, and the number above indicates the mean fold change in viral genome between the two time points.

tropism and the requirement of HBGAs for infection using HIEs from different segments of the gut and from different human donors with different HBGA phenotypes. Our results show that the HIE model is fit to investigate HuSaV–host interactions.

## RESULTS

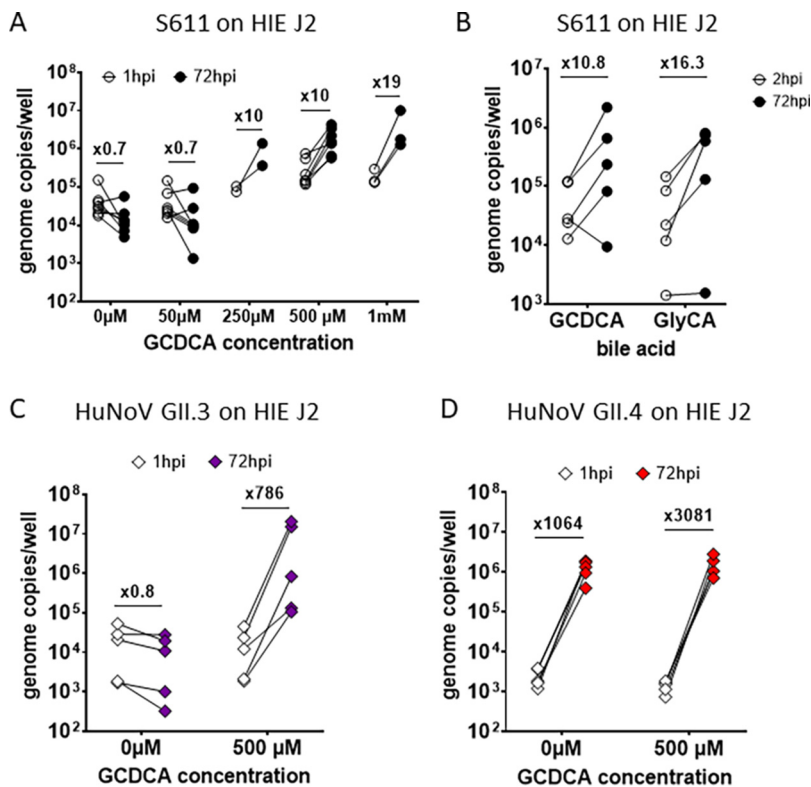
**HIE cultures support HuSaV replication.** To evaluate if HIEs could support HuSaV replication, we inoculated differentiated J2 HIE monolayers with HuSaV-positive stool filtrates at different concentrations in the presence of the bile acid GCDCA, similar to the protocol used previously for HuNoVs (32). Of the five stool filtrates evaluated, two (S611 and S513) showed 0.5 to 1.8  $\log_{10}$ , i.e., 3 to 63-fold, increases in genome copies from 1 or 2 hpi to 72 hpi, respectively, depending on the amount of virus in the inoculum (Fig. 1A). We confirmed that the S611 strain replicated successfully on J2 HIE monolayers more than 40 times over a 2-year period, with a geometric mean fold change in viral genome of 32 between 1 or 2 hpi and 72 hpi ( $P < 0.0001$ , Wilcoxon signed-rank test) (Fig. 1B). Due to a lower titer, S513 was used in 9 experiments on J2 HIE monolayers, and showed replication in 6, with a mean fold change of 7.7 between 1 or 2h and 72 hpi ( $P = 0.01$ , Wilcoxon signed-rank test), lower than S611 in the same



**FIG 2** Kinetics and magnitude of HuSaV S611 replication in HIE. (A) Differentiated J2 monolayers were infected with  $1.10^8$  cg/well of HuSaV S611 with  $500 \mu\text{M}$  GCDCA and the viral genome quantified after 2h, 72h and 144h. Each dots represents the mean of three technical replicates in an experiment ( $n = 7$ ). In comparison with 2 hpi, viral replication was detected at 72 hpi ( $P < 0.05$ , Wilcoxon test) and 144 hpi ( $P < 0.05$ , Wilcoxon test) with geometrical mean fold changes in viral genome of 9.8 and 17.1, respectively. (B) HuTu80 were infected with S611 at four different concentrations from  $1.10^5$  to  $1.10^9$  cg/well with  $500 \mu\text{M}$  GCDCA. Mean viral titers of three technical replicates were measured in the culture supernatants at 1, 3, 6, and 7 days postinfection and are plotted for each experiment ( $n = 2$ ), with lines connecting the geometrical means. (C) Differentiated D109 monolayers were inoculated with  $1.10^8$  cg/well native (plain black circles) or heat-inactivated (crossed circles) S611 with  $500 \mu\text{M}$  GCDCA and the viral genome quantified after 2h, 24h, 48h, 72h, 96h, and 144h. The mean of technical replicates is plotted as a circle for each experiment ( $n = 2$ ) and geometrical means are connected with a plain (native virus) or a dotted (inactivated virus) line. (D) The viability of the differentiated D109 monolayers either infected by native (plain black circles) or heat-inactivated (crossed circles) S611 with  $500 \mu\text{M}$  GCDCA for  $n = 2$  experiments in comparison with an uninfected control (100%, dotted line).

HIE line (Fig. 1C). In most experiments, monolayers were also inoculated with HuNoV GII.4 TCH11-64 strain as a control (Fig. 1D), for which the mean fold change was much higher (4,388 mean fold change;  $P < 0.0001$ , 1h or 2h versus 72 hpi, Wilcoxon signed-rank test) than for the two tested HuSaV strains. Together, these results demonstrate that two strains of HuSaV replicated repeatedly in J2 HIEs, albeit with lower fold changes after 3 days of infection compared to a HuNoV GII.4 strain.

We next investigated the kinetics and magnitude of S611 replication in HIEs. To assess if the low fold changes observed at 72 hpi were due to a more protracted course of infection for HuSaV in HIEs, we measured the fold increase after 6 days (144h) of infection (Fig. 2A). Viral genome concentrations were higher at 72 hpi and 144 hpi than at 2 hpi ( $P < 0.05$  for both, Wilcoxon signed rank test), with fold changes in viral genome of 9.8 and 17.1 at 72h and 144 hpi, respectively, confirming successful viral replication. There was a small but significant difference between the two late time points ( $P$  value = 0.047). We next tested the replication of the S611 strain in the HuTu80 duodenal cell line that was recently shown to support replication of HuSaV GI.1, with various amounts of virus in the inoculum. Like in HIEs, S611 replicated in HuTu80 cultures, with fold changes in viral genome copies of  $\sim 2 \log_{10}$  between 1 day and 3 or 6 days postinfection (Fig. 2B), lower than the fold increase previously reported for another GI.1 strain (19). Of note, we were able once to use the S611 fresh after collection, before any freeze-thaw cycle, on J2 HIEs in a single experiment that showed a high

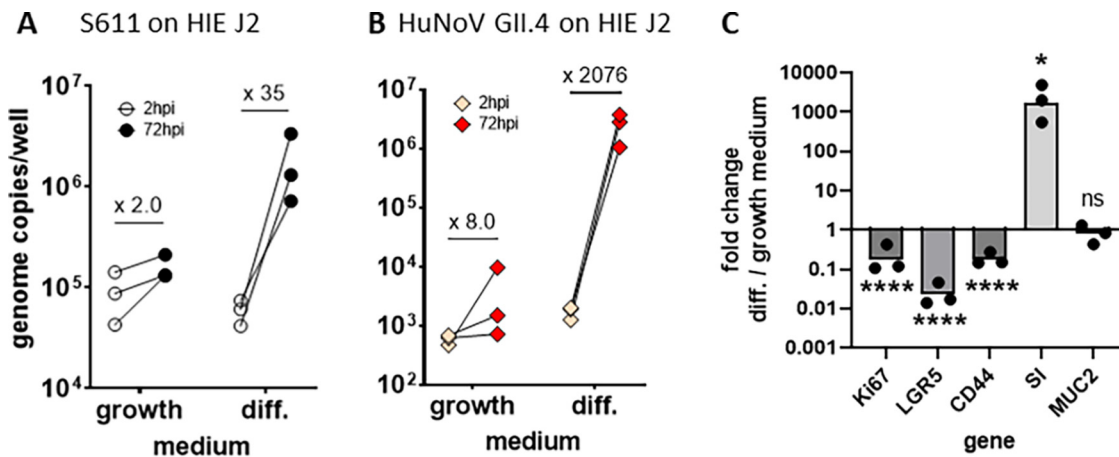


**FIG 3** Bile acids are a necessary cofactor of HuSaV infection in HIE. (A) Differentiated J2 monolayers were infected with  $1.10^8$  cg/well of S611 in the presence of the bile acid GCDCA at concentrations ranging from 50  $\mu$ M to 1 mM, or in the absence of bile acid. The viral titer was measured and is depicted as the mean of three technical replicates for  $n = 2$  to 6 experiments at 1 hpi (white circles) and 72 hpi (black circles) and the geometrical mean fold change between the two points indicated above. (B) The bile acid GlyCA (right side) was also used in comparison to GCDCA (left side) with concentrations of 500  $\mu$ M on differentiated J2 monolayers infected with  $1.10^8$  cg/well of S611,  $n = 5$  experiments. (C) and (D) Experiments in (A) were also conducted with  $1.10^6$  cg/well of HuNoV GII.3 (purple) or  $1.10^5$  cg/well of HuNoV GII.4 (red), with GCDCA concentrations of 500  $\mu$ M or without GCDCA.

fold increase ( $\times 1,115$ ) of the viral genome in 72 hpi (Fig. S1). The kinetics of S611 replication in the D109 HIE line showed an increase in viral genome copies between 1 and 72h postinfection, followed by a plateau until 6 days (144h) postinfection (Fig. 2C). To control for the exposure of HIEs to fecal filtrates, in the absence of viral replication, we inoculated cultures with heat-inactivated virus, where the viral genome copies showed a slow but steady decline as expected (Fig. 2C). In parallel, the viability of the cell culture was monitored using a metabolic assay and was not affected by the HuSaV replication, nor by exposure to the inactivated virus (Fig. 2D). These results show that in our experimental conditions, HuSaV S611 replicates at low levels and does not alter the viability of HIEs.

**Replication of S611 HuSaV strain in jejunal HIE is dependent on bile acid and cell differentiation.** We next assessed whether bile acids were necessary for HuSaV S611 replication in HIEs as previously shown for replication of the PoSaV Cowden strain (33) and in previous work conducted with HuSaV in the HuTu80 human duodenal cell line (19). J2 HIE monolayers were inoculated with HuSaV S611 in the presence of increasing concentrations (from 0  $\mu$ M to 1 mM) of the bile acid GCDCA. We observed that GCDCA was necessary for a productive infection of J2 HIE monolayers by HuSaV S611, with fold changes in viral genome copies  $>3$  appearing when GCDCA concentrations were between 250  $\mu$ M and 1 mM (Fig. 3A). A concentration of 500  $\mu$ M was set for the next experiments.

We also tested the effect of another bile acid, sodium glycocholate (GlyCA), as it was used in the study of HuSaV replication in HuTu80 cells (19). HuSaV S611 replicated similarly with 500  $\mu$ M GCDCA (mean fold change of 10.8 between 2h and 72h pi) or GlyCA (mean fold change of 16.3) (Fig. 3B).



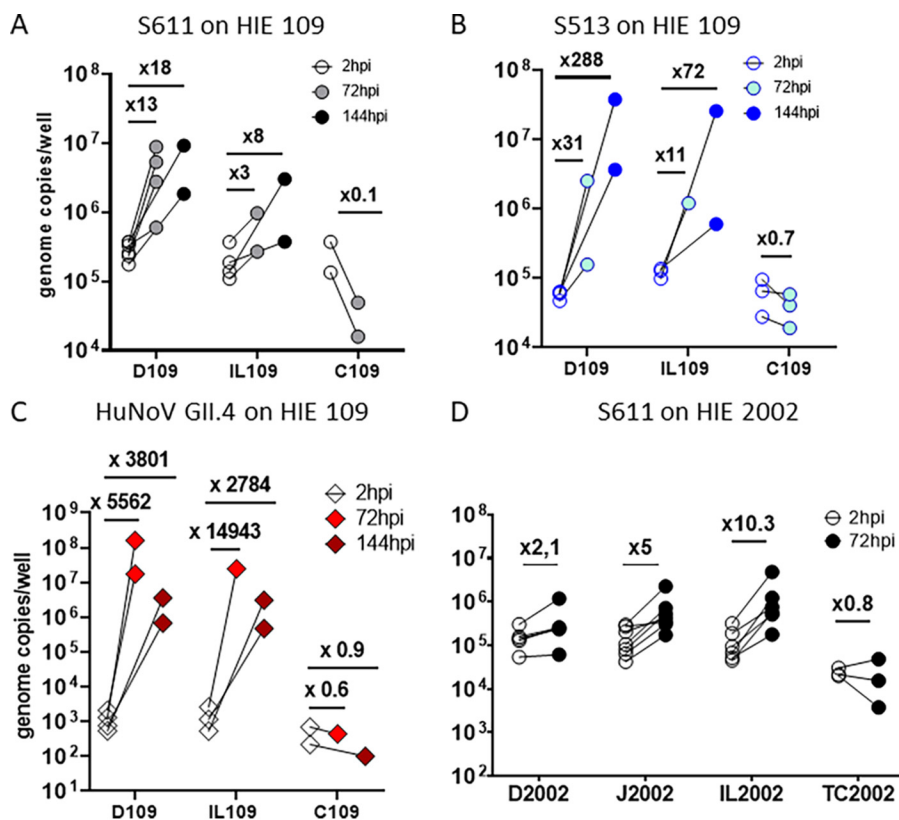
**FIG 4** HuSaV replicate in differentiated jejunal HIE. Comparison of the mean viral titer per well of S611 HuSaV (A) and GII.4 HuNoV (B) at 2 hpi (white bars) and 72 hpi (black bars) in J2 monolayers cultured and infected in growth or differentiation (diff.) medium with  $500 \mu\text{M}$  GCDCA, for  $n = 3$  experiments, 3 technical replicates per experiment. Geometric mean fold changes between the two time points are indicated above for each condition. (C) Fold change in relative expression levels of 5 genes (normalized on GAPDH) between HIE J2 monolayers cultured in differentiation versus growth medium in the experiments presented in (A). \*,  $P < 0.05$ , \*\*\*\*,  $P < 0.0001$ , paired  $t$  test.

HuNoV bile-dependent strain (GII.3) and partially independent strain (GII.4) were also inoculated in parallel as controls and recapitulated results obtained previously (26) (Fig. 3C and D).

We then took advantage of the HIEs to investigate whether HuSaV infects intestinal stem cells and/or differentiated cells, such as enterocytes that are a target of HuNoV. J2 HIEs were plated as monolayers and either infected the following day and kept in stem-cell growth medium (Intesticult OGM) or induced to differentiate using a medium with reduced concentrations of niche growth factors for 4 days before being infected. HuSaV replicated to higher fold changes in differentiated cells compared to in undifferentiated cells (Fig. 4A). Similar results, albeit with higher fold increases, were observed with HuNoV GII.4 (Fig. 4B). The expression of five genes known to be modulated during differentiation was assessed by qRT-PCR. Relative expression of stem cell and proliferation markers (Ki67, LGR5, CD44) was lower under the condition with differentiation medium (mean fold changes of 0.22, 0.036, 0.19, respectively;  $P$  values  $< 0.0001$ , paired  $t$  test). Conversely, Sucrase-isomaltase (SI) was markedly induced (fold change of 2592,  $P$  value  $< 0.05$ , paired  $t$  test), indicating that cells differentiated into enterocytes. The mucin 2 (Muc2) marker of goblet cells was not affected (fold change 0.92,  $P$  value = 0.74, paired  $t$  test). Altogether these results show that HIE differentiation is necessary for a consistent replication of HuSaV.

**HuSaV replicates in cells of the three portions of small intestine from different donors.** To assess the tropism of HuSaV in the human gut, we used HIEs originating from different segments of the intestine of the same donor. In donor 109, we observed the replication of both S611 (Fig. 5A) and S513 (Fig. 5B) in duodenal (D109) and ileal (IL-109) cells, after 72 or 144 hpi, but not in cells from the colon (C109). The HuNoV GII.4 TCH11-64 strain showed the same pattern (Fig. 5C), as previously reported (29). S611 replicated poorly in IL-109, whereas S513 showed higher fold increases in ileal cells. In HIEs from donor 2002 (Fig. 5D), for which cultures from all segments from a single individual were available, S611 replicated in cells from the jejunum (J2002) and the ileum (IL-2002), but not from the duodenum (D2002) or the transverse colon (TC2002). From these results, we conclude that HuSaV can replicate in HIEs from different donors, and in cells from the 3 portions of the small intestine, but that strain-related or donor-related factors may modulate this tropism.

**HuSaV replication in HIE is independent of secretor status.** The tested HuSaV strains replicated in HIE cell monolayers from three different donors: 2, 109 and 2002, all secretor positive (Table 1). We next investigated if secretor phenotype and blood



**FIG 5** HuSaV replicates in human cells from the three segments of the small intestine. Differentiated HIE monolayers from different segments of the small intestine (J for jejunum, D for duodenum, IL for ileum, C for colon and TC for terminal colon) originating from two donors, 109 (A, B, C) and 2002 (D), were infected with  $1.10^8$  cg/well of HuSaV S611 (A, D – white, gray, black circles),  $1.10^7$  cg/well of HuSaV S513 (B – blue circles) or  $1.10^5$  cg/well of HuNoV GII.4 as a control (C – pink and red diamonds) with  $500 \mu\text{M}$  GCDCA. The mean viral titer per well for three technical replicates is depicted as circles for 2h, 72h or 144 hpi, for  $n = 2$  to 6 experiments. Above each condition is displayed the corresponding viral genome fold change between the two time points.

type (HBGA phenotype) could impact HuSaV replication, as it does restrict several HuNoV strains. S611 replicated successfully in 3 of 7 experiments in jejunal HIE J10 (secretor and Lewis negative) and in 6 of 7 experiments in J11 (secretor positive) (Fig. 6A, left panel), whereas HuNoV GII.4 did not replicate in J10 (Fig. 6A, right panel). Interestingly, S611 initial binding and viral genome fold changes in 72 hpi appeared to be lower on the J10 HIE line than in the two secretor positive lines.

Next, we tested the dependency to the secretor phenotype in isogenic cells, using two HIE lines and their genetically modified counterparts: J4, a secretor negative line, with the J4FUT2KI knock-in, and J2, a secretor positive, with the J2FUT2KO knockout. HuSaV S611 was able to replicate in all HIEs independently of the expression of *FUT2* (Fig. 6B and C). As observed previously (26, 34), HuNoV did not show replication in secretor negative J10, J4 and J2FUT2KO HIE lines. These results show that S611 HuSaV strain can replicate in both secretor positive and negative jejunum HIEs from different donors, with 5 secretor positive and 3 secretor negative HIE lines being tested. As expected, GII.4 HuNoV showed no replication in the 3 secretor negative HIE lines (Fig. 6A and B and C, right panels). Of note, most of the tested HIEs are O blood group but S611 also replicated in A and B type group HIEs (D-IL-109 for A group, and J2 for B group, Table 1).

## DISCUSSION

Prior attempts to grow HuSaV, mostly in primary kidney and intestine cells of human or simian origin, have been reported, but many have produced either inconclusive or

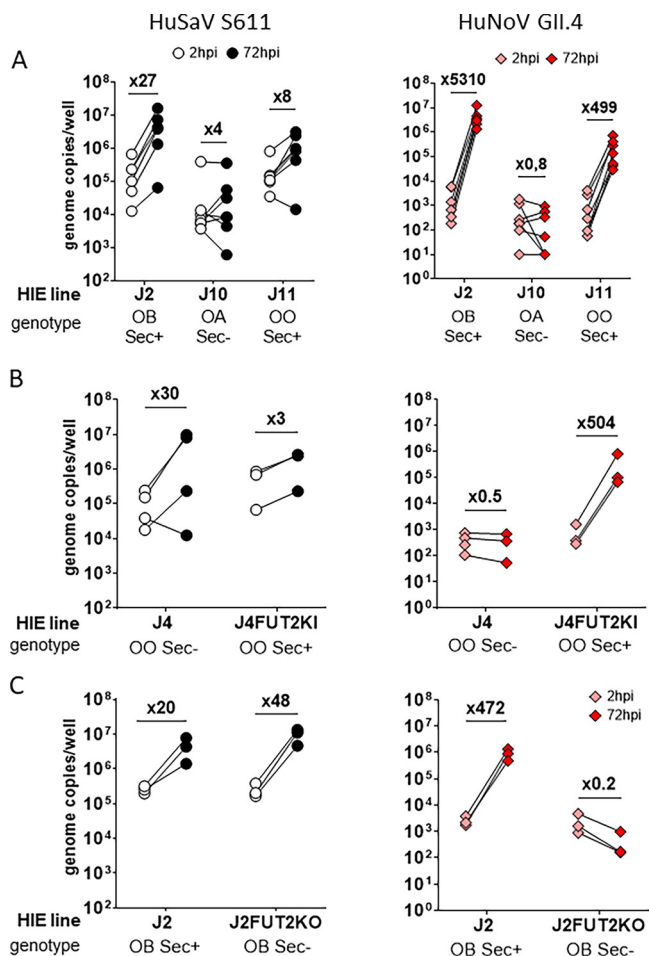
**TABLE 1** List of HIE lines used in this study

HIE line	Tissue of origin	Phenotype		Reference
		Secretor status	ABO	
J2	Jejunum	Sec +	B, Lewis b	Ettayebi et al., 2016 (26)
J11	Jejunum	Sec +	Lewis b	
J2002	Jejunum	Sec +	Lewis b	
D2002	Duodenum			
IL2002	Ileum			Ettayebi et al., unpublished data
TC2002	Transverse colon			
J4FUT2KI	Jejunum	Sec +	Lewis b	Haga et al., 2020 (34)
D109	Duodenum	Sec +	A, Lewis b	Rajan et al., 2018 (35)
IL109	Ileum			
C109	Colon			
J2FUT2KO	Jejunum	Sec -	Lewis a	Haga et al., 2020 (34)
J4	Jejunum	Sec -	Lewis a	Haga et al., 2020 (34)
J10	Jejunum	Sec -	A	Rimkute et al., 2020 (30)

nonreproducible results (35–37). On the other hand, the PoSaV Cowden strain replicates successfully in porcine kidney cells (18, 38), and its replication is dependent on bile acids with sialic acids acting as a binding factor. Recently, Takagi et al. reported efficient replication and passaging of GI.1 and GII.3 HuSaV strains in a HuTu80 adenocarcinoma cell line of duodenal origin in the presence of bile acids, and a reverse genetics system also produced GII.3 virions after transfection of HuTu80 cells (6, 19). Our work focused on HIEs, a physiological model of human intestine in which several enteric viruses can replicate. The genetically diverse range of donors and genetically modified HIE lines also allows investigation of viral tropism and cofactors required for replication.

In this work, we were able to successfully grow two HuSaV strains in HIEs: S611 (GI.1) and S513 (GI.2) (Table 2). These two genotypes cause the majority of cases in Europe (39, 40), and were the most frequently detected in HuSaV-positive fecal samples in a 2021 study in Spain (11). These two strains showed consistent viral replication in the HIE J2 line of jejunal origin, over a time frame of 2 years for S611, characterized by a moderate multiplication. Indeed, the viral genome fold changes within 3 or 6 days postinfection did not reach levels as high as those achieved by GII.4 HuNoV in HIEs (2 to 3  $\log_{10}$ ) or HuSaV in HuTu80 (19) (from 2 to 6  $\log_{10}$ ), which precluded serial propagation experiments in our study. However, in our hands, S611 replication in the HuTu80 cell line was also moderate (1 to 2  $\log_{10}$ ), which suggests that this may be a characteristic of the strains used in this study. Indeed, previous work has shown that some HuNoV strains also display a low or moderate viral replication in HIEs (26, 29, 41). A unique experiment conducted with fresh S611, before any freezing, showed a higher 3  $\log_{10}$ -fold (x1115) increase in the J2 HIE line (Fig. S1), but additional fresh stool samples containing virus were not available to evaluate the repeatability of this observation. Hence, our results showing moderate fold changes in viral genomes for HuSaV S611 and S513 need to be repeated with more strains, and if possible, fresh stool samples, before drawing conclusions on the magnitude of HuSaV replication in HIE in general. In addition, kinetic experiments showing that the viral replication mostly occurs in the first day of infection suggest that HuSaV S611 did not propagate efficiently in HIE cultures, and was limited to one replication cycle, as also observed for several HuNoV strains (42). Finally, although most experiments showed replication of S611 (38 out of 40 experiments) and S513 (6 out of 9), in some cases with similar inoculum and experimental conditions, the viruses did not grow. This has been observed previously for some HuNoV strains with low or moderate amplification (41). Similarly, heterogeneity in viral fold change was also reported for a HuNoV strain across repeated experiments (29). Here, HuNoV GII.4 was included in most experiments to control for the ability of the cell cultures to sustain an infection (see Materials and Methods). Thus, the resulting heterogeneity in HuSaV replication may be linked to a possible heterogeneity of the





**FIG 6** HuSaV replicate in HIE independently of HBGA genotype. Differentiated monolayers of jejunal HIE from different donors were infected with  $1.10^8$  cg/well of HuSaV S611 (white and black) or  $1.10^5$  cg/well of HuNoV GII.4 (pink and red) with  $500 \mu\text{M}$  GCDCA in each case and the mean viral titer per well measured for three technical replicates and  $n = 3$  to  $7$  experiments at 2 hpi (white, pink) and 72 hpi (black, red). (A) Three jejunal HIE cells lines, J2, J10, J11 with contrasted HBGA genotype (OB Sec+, Sec-, and OO Sec+, respectively). (B) J4 cells (OO Sec- genotype) and the corresponding knock-in, J4FUT2KI (OO Sec+). (C) J2 cells (OB Sec+) and the corresponding knockout, J2FUT2KO (OB Sec-).

stool samples where the viral particles may form aggregates or vesicle-cloaked clusters, as reported for HuNoV (43, 44).

Despite these moderate levels of replication, the HIE model was still adequate to investigate HuSaV biology. We first confirmed that bile acids are required for efficient replication of HuSaV S611 in HIEs. Here, GCDCA was mainly used, similarly to the HuNoV replication model (26), but could be replaced by GlyCA (Fig. 3B), as was done on HuTu80 cells (19). Some strains of HuNoV being able to replicate in HIEs independently of bile acid (Fig. 3D) (26), other HuSaV samples should be tested in this model as

**TABLE 2** List of viral strains used to perform infection experiments

Virus	Strain	Titer (gc/mL) <sup>a</sup>	Genotype	Accession number or ref
HuNoV	TCH11-64	$5.8 \times 10^9$	GII.4	(26, 29)
	TCH04-577	$1.07 \times 10^{10}$	GII.3	<a href="#">GU930737</a> (26, 29)
HuSaV	S496	$1.2 \times 10^9$	GIV.1	<a href="#">OP654149</a>
	S513	$5.81 \times 10^8$	GI.2	<a href="#">OP654151</a>
	S578	$2.89 \times 10^8$	GI.2	<a href="#">OP654152</a>
	S586	$7.2 \times 10^{10}$	GII.3	<a href="#">OP654150</a>
	S611	$3.66 \times 10^{10}$	GI.1	<a href="#">OP654153</a>

<sup>a</sup>gc/mL: genome copies/mL of fecal filtrate.

well to generalize this observation. Based on HuNoV work, bile acids are thought to promote viral replication by enhancing endosomal uptake, acidification, and ceramide levels on the apical membrane (32). Work on PoSaV also showed similar effects, as bile acids are required for virions to escape endosomes into the cytoplasm and for the production of ceramide (45).

Likewise, intestinal stem-cell differentiation was necessary to support efficient replication of HuSaV S611 in jejunal HIE (Fig. 4A), as observed previously for HuNoV (Fig. 4B) (29). Moreover, since this differentiation was mostly characterized by a strong induction in SI expression, a marker of enterocytes, rather than Muc2, a marker of goblet cells (Fig. 4C), our results suggest that enterocytes may be the main cell type sustaining HuSaV replication in HIE cultures, although the contribution of other cell types cannot be excluded.

Using the HIE model also allowed us to investigate the potential tropism of HuSaV in the human intestine, with cells originating from jejunum, duodenum, ileum and colon of two different donors. We observed the replication of both HuSaV strains in HIEs derived from the three segments of the small intestine, with variations between the cultures from the two tested donors. D2002 showed only one experiment with >3-fold change in viral genome at 72 hpi, but efficient replication occurred in duodenal cells from the other donor, D109 cells. Conversely, in the colonic HIE (C109 and TC2002), viral titers of HuSaV S611 or S513 decreased over 3 days (Fig. 5A and D), indicating that cells derived from this section of the intestine do not support measurable viral replication. Our data strongly suggest that the tested HuSaV strains replicate in the small intestine but not in the colon. Likewise, PoSaV is believed to mainly infect small intestinal tissues based on histological localization of virus-induced damage in gnotobiotic piglets (46). This is also consistent with the fact that the small intestine is physiologically exposed to bile acids, a necessary cofactor for replication of the PoSaV Cowden strain and the HuSaV strains tested to date.

HBGA polymorphisms in the human population are known to drive sensitivity or resistance to infection with HuNoV and other enteric pathogens (47). Epidemiological data suggest that this is not the case for HuSaV (48, 49), but these studies were conducted on small cohorts (with number of HuSaV infected patients between 33 and 42) and some did not observe the usual secretor-associated trend when looking at HuNoV infections (50). Here, we took advantage of HIEs originating from donors expressing various HBGAs to assess the dependency of HuSaV on secretor or ABO phenotypes. S611 replicated both in secretor-positive (J2, J11, J4FUT2KI) and secretor negative (J10, J4, J2FUT2KO) jejunal HIE cells and in HIE from A, B, and O blood group donors. Some small differences can be noted, such as a reduced viral binding at 2 hpi, less experiments showing viral replication (3 out of 7 experiments) and lower viral replication when infecting the secretor-negative J10 HIEs compared to secretor positive J2. However, this could be due to other unknown differences between J2 and J10, which were derived from different donors and could have different levels of expression of the viral receptor or other cellular cofactors. To exclude these, we then used isogenic cell lines differing only with the expression of FUT2. J4 HIEs, initially secretor negative with homozygous *se428/se428* in the *FUT2* gene, supported S611 replication, with viral binding at 2 hpi similar to that of J2 cells. When knocked-in for *FUT2* (J4FUT2KI), these cultures remained infected by S611. Our results are also in agreement with previous studies made with GI and GV HuSaV VLPs that showed no interactions between VLPs and human HBGAs from saliva or synthetic carbohydrates (25).

Further epidemiological studies conducted on a larger scale could totally eliminate a possible link between secretor status and susceptibility to HuSaV infection, although data presented in this paper and previous studies strongly suggest that HuSaV infection and replication are independent from intestinal HBGAs.

Due to a lower viral concentration and volume of sample, S513 could not be used in all experiments, which would have allowed more comparisons between the two HuSaV GI genotypes. In 4 different cell lines tested with both viruses (J2, D109, IL-109 and

C109), the two strains behaved similarly. Three other tested strains did not replicate on HIE J2 (Fig. 1A), which could be due to their lower titers, especially for S496 and S578. Thus, additional strains will need to be tested to investigate whether other GI genotypes or other HuSaV genogroups also replicate in HIEs.

Initially overshadowed by more pathogenic viral AGE agents, HuSaVs have increasingly been detected in the human population, possibly due to the more widespread use of molecular diagnosis tools and/or to an actual epidemiological change, following RV vaccine implementation. Recent research highlight some unique characteristics of the SaV genus in comparison with the more studied NoV (5, 51). Most of the HuSaV biology remains to be explored, and we show here that HIEs represent a physiologically relevant model for further investigation of HuSaVs replication cycle and virus-host interactions.

## MATERIALS AND METHODS

**Viral strains.** HuSaV-positive or HuNoV-positive 10% fecal suspensions were prepared in PBS, 0.22  $\mu$ m-filtered, aliquoted and stored at  $-80^{\circ}\text{C}$ , as described previously (26), before being used to inoculate cells. Viral titers were determined by quantitative, one-step reverse transcription and PCR (qRT-PCR) on extracted nucleic acids, as described previously for HuNoV (52). Table 2 describes the viral strains used for subsequent experiments. The HuSaV S611 strain was heat-inactivated for 15 min at  $60^{\circ}\text{C}$  before being used to inoculate HIE monolayers in some experiments.

**Viral strain sequencing.** Stool fecal suspensions were incubated with 2000 U of OmniCleave Endonuclease (Epicentre, Madison, US) for 1 h at  $37^{\circ}\text{C}$  to eliminate free nucleic acids, followed by nucleic acid extraction as in (53). A final step of RNA purification and concentration was performed using a Zymo-spin column (RNA Clean & Concentrator, Zymo Research, Irvine, USA). Libraries were prepared by synthesizing cDNA using the Superscript IV kit (Life Technologies) with random hexamers according to the manufacturer's instructions (Life Technologies). cDNA first strand was physically fragmented (Ultrasonicator M220, Covaris) for 2.5 min. and afterwards, the NEBNext Ultra II RNA Library Prep kit for Illumina (E7770L –NEB) was used for Illumina pools. Sequencing was performed on an Illumina MiSeq to generate  $2 \times 250$  bp reads. Viral reads were assembled into full genomes and genotyped using the Genome Detective online tool (<https://www.genomedetective.com/>).

**HIE cells.** HIEs used in this study were obtained from healthy tissues of human donors who gave informed consent, according to a protocol approved by the Baylor College of Medicine Institutional Review Board (BCM IRB). The HIE cultures were maintained as described previously (29) in agreement with the ethical evaluation committee of the French Institute of medical research and Health (CEE). The tissue origin of the stem cells is designated using letters: J for jejunum, D for duodenum, IL for ileum and C for colon, and the number indicates the donor. In addition, a genetically modified J2 HIE line (J2FUT2KO) that is an isogenic J2 line with the *FUT2* gene responsible for the secretor phenotype knocked out and J4 cells genetically modified to express a functional *FUT2* gene (J4FUT2KI), as described previously, were also used for experiments (34).

**HIE culture.** All HIEs were grown in a 30  $\mu$ L dome of Matrigel growth-factor-reduced basement membrane matrix (Corning) surrounded with 500  $\mu$ L of Intesticult Intestinal Organoid Growth Medium (OGM; StemCell) supplemented with antibiotics (0.3% PenStrep) in 24-well plates (Nunclon, ThermoScientific). Cells were incubated at  $37^{\circ}\text{C}$  and 5%  $\text{CO}_2$ . Medium was changed every 2 to 3 days, and cells were passaged every week using trypsin 0.05%- EDTA (Gibco, ThermoFisher) and repeated pipetting to dissociate cells within the enteroids, before being centrifuged, resuspended in Matrigel and plated.

For infection experiments, 3D grown enteroids were dissociated for 4 min at  $37^{\circ}\text{C}$  in a water bath with Trypsin-EDTA 0.05% (Invitrogen) and mixed by repeated pipetting in order to obtain a single cell suspension. A minimum of  $1 \times 10^5$  cells per well were plated in 96-wells plates (Nunclon, ThermoScientific) coated with 33  $\mu$ g/mL human collagen type IV (Sigma-Aldrich) in molecular-grade water. After 24 to 48 h of culture at  $37^{\circ}\text{C}$  and 5%  $\text{CO}_2$  with OGM supplemented with Y-27632 (10  $\mu$ M, Sigma-Aldrich), medium was replaced with differentiation medium made of 50% Intestinal Organoid Growth Medium basal component, and 50% complete medium without growth factor (CMGF-), which is advanced DMEM/F12 medium (ThermoFisher) supplemented with 2 mM GlutaMax (Life technologies) and 10 mM HEPES, for 4 days to induce cellular differentiation.

**Infection experiments.** Infections were performed on confluent monolayers 4 days after initiating differentiation. When not specified, the viral inoculum concentration used per well was as described in Table 3. Viral inocula were prepared with CMGF- supplemented with various concentrations of glycochenodeoxycholic acid (GCDCA, Sigma-Aldrich), ranging from 5  $\mu$ M to 1 mM, and 500  $\mu$ M when not specified. Cells were incubated with viral inoculum for 1 to 2 h. Inocula were then removed, and cells were washed 3 times with CMGF-. 100  $\mu$ L of differentiation medium with GCDCA were then added to each well, and for each set of infections, one plate was immediately frozen at  $-20^{\circ}\text{C}$ . The other plate was incubated at  $37^{\circ}\text{C}$  under 5%  $\text{CO}_2$  for 3 to 7 days, then frozen at  $-20^{\circ}\text{C}$  until undergoing RNA extraction.

**HuTu80 culture and infection.** The HuTu80 cell line (ATCC, number HTB-40) was cultured in DMEM supplemented with 2 mM Glutamax, 0.3% PenStrep and 5% FBS at  $37^{\circ}\text{C}$  under 5%  $\text{CO}_2$ . Monolayers were made by dissociating cells from a confluent flask and seeding 24-well plates with  $5 \times 10^5$  cells in 0.5 mL of culture medium. Before infection, the medium was replaced with 0.5 mL of virus growth

**TABLE 3** Viral concentrations of inocula used for infection experiments

Virus	Strain	Concn / well	Genotype
HuNoV	TCH11-64	$1 \times 10^5$	GII.4
	TCH04-577	$1 \times 10^6$	GII.3
HuSaV	S611	$1 \times 10^8$	GI.1
	S513	$1 \times 10^7$	GI.2

media, which is a 3% FBS culture media supplemented with the bile acid GCDCA at concentrations varying from 250  $\mu$ M to 1 mM. The viral inoculum was then added to each well, and cells were washed twice with 2% FBS DMEM 1 day after inoculation. One mL of virus growth medium was then added to each well, and 100  $\mu$ L of supernatant was harvested at each time point and frozen until RNA extraction. The protocol was adapted from (19).

**RNA extraction from cell cultures.** RNA was extracted from cell cultures in 96-well plates with the NucliSens kit (bioMérieux) using the EasyMag automated system. The plates were left at room temperatures to thaw, and then 100  $\mu$ L of lysis buffer was added to each well and incubated for 10 min. The 200  $\mu$ L-content of each well was then homogenized and transferred to an EasyMag well containing 2 mL of lysis buffer, and 50  $\mu$ L of magnetic silica was added. The manufacturer's program was then used for nucleic acid extraction with 100  $\mu$ L elution buffer.

**Viral quantification by qRT-PCR.** The genomic viral titers were measured by qRT-PCR, using the RNA UltraSense One-Step Quantitative RT-PCR kit. For HuSaV, the forward primers were SaV1F, 5'-TTG GCC CTC GCC ACC TAC-3' and SaV124F, 5'-GAY CAS GCT CTC GCY ACC TAC-3', the reverse primer was SaV1245R, 5'-CCC TCC ATY TCA AAC ACT A-3' and the probe was SaV124TP, 5'-FAM-CCR CCT ATR AAC CA-MGBNFQ (54). For HuNoV, the primers were used as previously described (55). Initial denaturation was carried out at 55°C for 15 min followed by 5 min at 95°C, and then 45 cycles consisting of 15 s at 95°C followed by 30 s at 60°C and 30 s at 65°C were performed.

Quantification of viral copy numbers was calculated with a  $2 \times 6$  points standard curve made by 10-fold serial dilutions of synthetic DNA (Integrated DNA Technologies), with the target sequence between nucleotide 4626 and 5683 of HuSaV strain Mc114 (GenBank [AY237422.3](#)), or the sequence between nucleotide 4191 and 5863 of HuNoV GII.4 Houston virus (56) (GenBank [EU310927](#)).

**Gene expression assay by fluidic qPCR.** To assess the differentiation of intestinal cells at the time of infection, the expression of 6 genes was measured in RNA extracted 2 h postinfection (hpi) using a microfluidic qPCR system (Standard Biotoools). Primer and probes for gene expression assays (GEA, Applied Biosystems) were chosen to target a reference gene (GAPDH), markers of proliferating (Ki67) and stem cells (LGR5, CD44) and two differentiation markers, Sucrase-Isomaltase (SI) for enterocytes and Mucine 2 (Muc2) for goblet cells. Briefly, 7  $\mu$ L of RNA was reverse-transcribed using the SuperScript II kit (Thermo Scientific), as per the manufacturer's instructions, in 20  $\mu$ L reaction mixtures with 2.5  $\mu$ M random hexamers and nonamers (Thermo Scientific). Complementary DNA (1.25  $\mu$ L per sample) was pre-amplified in 5  $\mu$ L reaction mixtures using 1  $\mu$ L of Preamp Master Mix (Standard Biotoools) with 1.25  $\mu$ L of 0.2 $\times$  primers-probe assays pooled, through 2 min denaturation at 95°C followed by 14 cycles of 15s denaturation at 95°C and 4 min. hybridation-elongation at 60°C. Immediately after the preamplification, PCR products were diluted (1:5) by adding 20  $\mu$ L of TE and stored at 4°C prior to use in qPCR. For the qPCR assay, a Flex Six integrated fluidic circuit (Standard Biotoools) was loaded on one side with 12  $\times$  3  $\mu$ L reaction mixtures based on the UltraSense One-Step Quantitative RT-PCR kit (Thermo Scientific) in a BioMark device (Standard Biotoools), together with 1 $\times$  GE Sample Loading Reagent (Standard Biotoools) and 1.35  $\mu$ L of preamplified DNA, and on the other side with 3  $\mu$ L of assay mix containing 1.5  $\mu$ L of one gene expression assay (10 $\times$  final) and 1.5  $\mu$ L assay loading reagent (Standard Biotoools), in duplicate for each of the six assays. The qPCR program included a thermal mix with 30 min at 25°C and 1h at 70°C, 5 min. denaturation at 95°C and 45 cycles of 15s denaturation at 95°C, 30s hybridation at 60°C and 30s elongation at 65°C with fluorescence acquisition. It was run in a BioMark device (Standard Biotoools). Results were analyzed using the BioMark analysis software (Standard Biotoools).

**Evaluation of cell viability during HuSaV infection.** Infected HIE monolayers were incubated with 25  $\mu$ L of Blue Cell Titer (Promega) diluted in 75  $\mu$ L of differentiation medium, for 3 h. Fluorescence was then measured with a TECAN Infinite 200 Pro (Tecan Group Ltd.) at an excitation wavelength of 560 nm and emission wavelength of 590 nm. Next, the wells were washed twice with CMGF- and the differentiation medium was replaced. Wells without cells were used as baseline absorbance values. The percentage of viable cells was calculated by comparing the absorbance of the measured wells for each condition (inoculated with live or heat-treated S611) to the corresponding control wells, which contained cells that were not infected and were considered to display 100% viability value at each time point.

**Controls and statistical analysis.** Viral replication was evaluated by calculating the geometric mean fold change for each set of experiments, as the mean of triplicate  $2^{\Delta\Delta Ct}$  measured between a late time point (usually 72 hpi) and the first time point (1 to 2h pi) for each replicate. Hence, a geometrical mean fold change above 1 corresponds to an increase in viral genome quantities over time, and below 1, to a decrease. The virus was considered to replicate when the geometrical mean fold change was equal or greater than 3 ( $\sim 0.5 \log_{10}$ ). For most experiments with HuSaV, HIE monolayers were inoculated with HuNoV GII.4 (TCH11-64) in parallel as a positive control. Three experiments were excluded from analysis

when HuNoV unexpectedly did not replicate (fold increase 0.5 log<sub>10</sub> or less), indicating that the culture conditions were suboptimal. Statistical analyses were run using GraphPad Prism version 9.4.0.

## SUPPLEMENTAL MATERIAL

Supplemental material is available online only.

**SUPPLEMENTAL FILE 1**, TIF file, 0.1 MB.

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