

FIG 2 Long-term culture of MA/JFH-1.1 and MA/JFH-1.2 RNA-transfected cells. Ten micrograms of HCV RNA was transfected into Huh7.5.1 cells, and cells were passaged every 2 to 5 days, depending on cell status. Culture medium was collected after every passage, and HCV core protein levels were measured. Transfection was performed twice for each chimeric RNA (1 and 2 for each construct). (A) HCV core protein levels in culture medium from MA/JFH-1.1 and MA/JFH-1.2 RNA-transfected cells. (B) Immunostained cells at 3 days after transfection (a to d), at 21 days after transfection (e to h), and at the time

TABLE 1 HCV core protein levels and infectivity in culture medium immediately after RNA transfection (day 3) and after long-term culture (days 35 to 49)

Sample period and virus	Sample no.	Day no. ^a	HCV core (fmol/liter)	Infectivity (FFU/ml)
After transfection				
MA/JFH-1.1	1	3	1.06×10^3	5.00×10^1
	2	3	1.14×10^3	5.70×10^1
MA/JFH-1.2	1	3	2.14×10^3	7.30×10^1
	2	3	2.15×10^3	9.30×10^1
After long-term culture				
MA/JFH-1.1	1	42	3.38×10^5	1.62×10^5
	2	42	4.70×10^5	3.23×10^5
MA/JFH-1.2	1	35	2.27×10^5	1.61×10^5
	2	49	4.93×10^5	3.27×10^5

^a For the long-term culture, the days are those of peak core protein levels.

as a 5' UTR replacement from J6CF (genotype 2a) to JFH-1 enhanced virus production of chimeric J6CF virus harboring the region of NS2 to 3' X of JFH-1 (J6/JFH-1) (A. Murayama et al., unpublished data). The core protein accumulation levels with MA/JFH-1.2 RNA-transfected cells were higher than those with MA/JFH-1.1 ($P < 0.05$) (Fig. 1B). Similarly, core protein and HCV RNA levels in the medium of MA/JFH-1.2 RNA-transfected cells were higher than those of MA/JFH-1.1 ($P < 0.05$) (Fig. 1C and D). Infectivity on day 3 was also higher than with MA/JFH-1.1 ($P < 0.05$) (Fig. 1E), indicating that the 5' UTR of JFH-1 enhanced virus production. However, infectivity of medium from MA/JFH-1.2 RNA-transfected cells on day 3 remained 6.4-fold lower than that of JFH-1 although HCV RNA levels in the medium were similar to those of JFH-1 (Fig. 1D and E).

These results indicate that 2b/2a chimeric RNA is able to replicate autonomously in Huh7.5.1 cells and produce infectious virus although infectivity remains lower than that of JFH-1.

Assembly-enhancing mutation in core region introduced during long-term culture. Because MA/JFH-1.1 and MA/JFH-1.2 replicated efficiently but produced small amounts of infectious virus, we performed long-term culture of these RNA-transfected cells in order to examine whether these chimeric RNAs would continue replicating and producing infectious virus over the long term. We prepared two RNA-transfected cell lines for each construct (MA/JFH-1.1 and MA/JFH-1.2) as both of these replicated and produced infectious virus at different levels.

Immediately after transfection, core protein levels and infectivity in culture medium were low (1.06×10^3 to 2.15×10^3 fmol/liter and 5.00×10^1 to 9.30×10^1 FFU/ml, respectively) (Fig. 2A and Table 1) although a considerable number of core protein-positive cells were observed by immunostaining (Fig. 2B, frames a to d). Subsequently, core protein levels in the culture medium decreased gradually (Fig. 2A), and core protein-positive cells were rare (Fig. 2B, frames e to h). However, at 30 to 40 days

of peak core levels (days 42 to 49). Infected cells were visualized with anti-core protein antibody (green), and nuclei were visualized with DAPI (blue). (C) Infection of naive cells by culture medium at an MOI of 0.001. (D) Immunostained cells at 15 days after infection with medium at peak core protein levels (Fig. 2A) at an MOI of 0.001. Infected cells were visualized with anti-core antibody (green), and nuclei were visualized with DAPI (blue).

after transfection, core protein levels in the supernatants of all chimeric RNA-transfected cells increased and reached 2.27×10^5 to 4.93×10^5 fmol/liter (Fig. 2A and Table 1). Infectivity in the culture medium also increased (1.61×10^5 to 3.27×10^5 FFU/ml) (Table 1), and at this point, most of the cells were core protein positive (Fig. 2B, frame i to l).

As the infectivity of culture supernatant of MA/JFH-1 RNA-transfected cells appeared to increase after long-term culture, we compared viral spread by infection with these supernatants on day 3 (immediately after transfection) and for each peak in core protein levels (after long-term culture). When naïve Huh7.5.1 cells were infected with supernatant on days corresponding to a peak in core protein levels at a multiplicity of infection (MOI) of 0.001, core protein levels in the medium increased rapidly and reached 0.64×10^6 to 1.13×10^6 fmol/liter by day 15 after infection (Fig. 2C). Immunostained images showed that most cells were HCV core protein positive on day 15 (Fig. 2D). When naïve Huh7.5.1 cells were infected with supernatant from day 3 at an MOI of 0.001, core protein levels in the medium did not increase under these conditions (Fig. 2C). These results indicate that both MA/JFH-1 chimeric viruses (MA/JFH-1.1 and MA/JFH-1.2) acquired the ability to spread rapidly after long-term culture.

As the characteristics of the MA/JFH-1 virus changed in long-term culture, we analyzed the possible mutations in the viral genome from the supernatant at each peak in core protein levels (Table 1, days at peak core levels). Nine- to 12-nucleotide mutations were found in the viral genome from each supernatant, and the detected mutations were distributed along the entire genome. Among these mutations, a common nonsynonymous mutation was found in the core region (Arg to Gly at amino acid [aa]167, R167G).

In order to test the effects of R167G on virus production, an R167G substitution was introduced into MA/JFH-1.2 as MA/JFH-1.2 replicated and produced infectious virus more efficiently than MA/JFH-1.1. HCV core protein levels in cells and medium of MA/JFH-1.2 with R167G (MA/JFH-1.2/R167G) were higher than with MA/JFH-1.2 ($P < 0.05$) (Fig. 3A and B). HCV RNA levels in the medium of MA/JFH-1.2/R167G RNA-transfected cells were also higher than with MA/JFH-1.2 ($P < 0.05$) (Fig. 3C). Infectious virus production was also increased by the R167G mutation ($P < 0.05$) (Fig. 3D) and was 8.7-fold higher than that of JFH-1 RNA-transfected cells on day 3 ($P < 0.05$) (Fig. 3D).

We then tested whether R167G was responsible for the rapid spread observed in culture supernatant after long-term culture by monitoring virus spread after infection of naïve Huh7.5.1 with culture medium taken 3 days after RNA transfection of MA/JFH-1.2 and MA/JFH-1.2/R167G at an MOI of 0.005. Core protein levels in medium from MA/JFH-1.2/R167G-infected cells increased with the same kinetics as levels of JFH-1 (Fig. 3E), and the population of core protein-positive cells was almost the same as with JFH-1-infected cells (Fig. 3F), indicating that MA/JFH-1.2/R167G virus spread as rapidly as JFH-1 virus. In contrast, we observed no infectious foci in the MA/JFH-1.2 virus-inoculated cells (Fig. 3F). These data suggest that the R167G mutation in the core region was a cell culture-adaptive mutation and that it enhanced infectious MA/JFH-1.2 virus production.

In order to determine whether R167G enhances RNA replication or other steps in the viral life cycle, we performed a single-cycle virus production assay (11) using Huh7-25 cells, a HuH-7-derived cell line lacking CD81 expression on the cell surface (1).

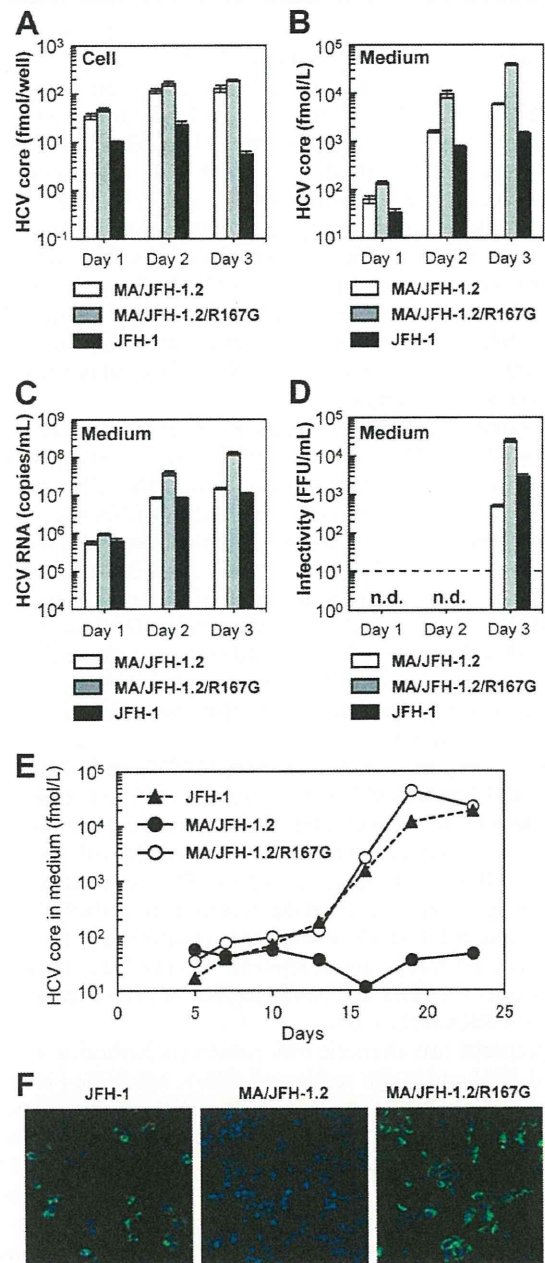


FIG 3 Effects of R167G on replication and virus production of MA/JFH-1.2 in Huh7.5.1 cells. Ten micrograms of HCV RNA was transfected into Huh7.5.1 cells, and cells and medium were harvested on days 1, 2, and 3. HCV core protein levels in the cells (A) and culture medium (B) and HCV RNA levels in the medium (C) and the infectivity of culture medium (D) from HCV RNA-transfected Huh7.5.1 cells are shown. n.d., not determined. Dashed line indicates the detection limit. Assays were performed three times independently, and data are presented as means \pm standard deviation. (E) HCV core protein levels in culture medium from cells infected with medium at 3 days posttransfection at an MOI of 0.005. (F) Immunostained cells at 19 days postinfection. Infected cells were visualized with anti-core antibody (green), and nuclei were visualized with DAPI (blue).

This cell line can support replication and infectious virus production upon transfection of HCV genomic RNA but cannot be reinfectious by progeny virus, thereby allowing observation of a single cycle of infectious virus production without the confounding ef-

fects of reinfection. R167G did not affect HCV core protein levels in the chimeric RNA-transfected Huh7-25 cells (Fig. 4A), demonstrating that R167G did not enhance RNA replication. Nevertheless, R167G increased HCV core protein levels in the medium ($P < 0.05$ on days 2 and 3) and infectivity (Fig. 4B and C). These results suggest that R167G did not affect RNA replication but affected other steps such as virus assembly and/or virus secretion.

Virus particle assembly efficiency was then assessed by determining intracellular-specific infectivity from infectivity and RNA titer in the cells, as reported previously (11). As shown in Fig. 4G, R167G enhanced intracellular-specific infectivity of MA/JFH-1.2 virus 10.2-fold. Virus secretion efficiency was also calculated from the amount of intracellular and extracellular infectious virus, but R167G had no effect (Fig. 4G).

To confirm the effects of Arg167 in other HCV strains, we tested its effects on JFH-1. As aa 167 of JFH-1 is Gly, we replaced it with Arg (G167R). HCV core protein levels in the cells were not affected by G167R (Fig. 4D), and no effects on RNA replication were confirmed. HCV core protein levels in the medium and infectivity decreased after G167R mutation (Fig. 4E and F). As the G167R mutation decreased intracellular infectious virus production of JFH-1 to undetectable levels, we were unable to determine the intracellular-specific infectivity and virus secretion efficiency of JFH-1 G167R (Fig. 4G). These results indicate that Gly is favored over Arg at core position 167 for infectious virus assembly in multiple HCV strains.

MA harboring the R167G mutation, 5' UTR, and N3H (NS3 helicase) and N5BX (NS5B to 3' X) regions of JFH-1 replicated and produced infectious chimeric virus. In order to establish a genotype 2b cell culture system with the MA strain with minimal regions of JFH-1, we attempted to reduce JFH-1 content in MA/JFH-1.2. We previously reported that replacement of the N3H and N5BX regions of JFH-1 allowed efficient replication of the J6CF strain, which normally cannot replicate in cells (21). Thus, we tested whether the N3H and N5BX regions of JFH-1 could also support MA RNA replication.

We prepared two chimeric MA constructs harboring the 5' UTR and N3H and N5BX regions of JFH-1, MA/N3H+N5BX-JFH1 (Fig. 5A) and MA/N3H+N5BX-JFH1/R167G. After *in vitro* transcribed RNA was transfected into Huh7.5.1 cells, intracellular core protein levels of MA/N3H+N5BX-JFH1 and MA/N3H+N5BX-JFH1/R167G RNA-transfected cells increased in a time-dependent manner and reached almost the same levels as with MA/JFH-1.2 RNA-transfected cells on day 5 (Fig. 5B). Extracellular core protein and HCV RNA levels of MA/N3H+N5BX-JFH1 and MA/N3H+N5BX-JFH1/R167G RNA-transfected cells also increased in a time-dependent manner (Fig. 5C and D). However, they were more than 10 times lower than with MA/JFH-1.2 RNA-transfected cells although intracellular core levels were comparable on day 5 (Fig. 5B to D).

We then tested whether the medium from MA/N3H+N5BX-JFH1 and MA/N3H+N5BX-JFH1/R167G RNA-transfected cells was infectious. Infectivity of the medium from MA/N3H+N5BX-JFH1 RNA-transfected cells was below the detection limit, and that of MA/N3H+N5BX-JFH1/R167G RNA-transfected cells on day 5 was very low ($3.3 \times 10^1 \pm 2.1 \times 10^1$ FFU/ml) (Fig. 5E). To confirm infectivity, the culture media were concentrated, and their infectivity was determined. Infected foci were observed after infection with concentrated medium in MA/N3H+N5BX-JFH1/R167G RNA-transfected cells (Fig. 5F), and infectivity was found

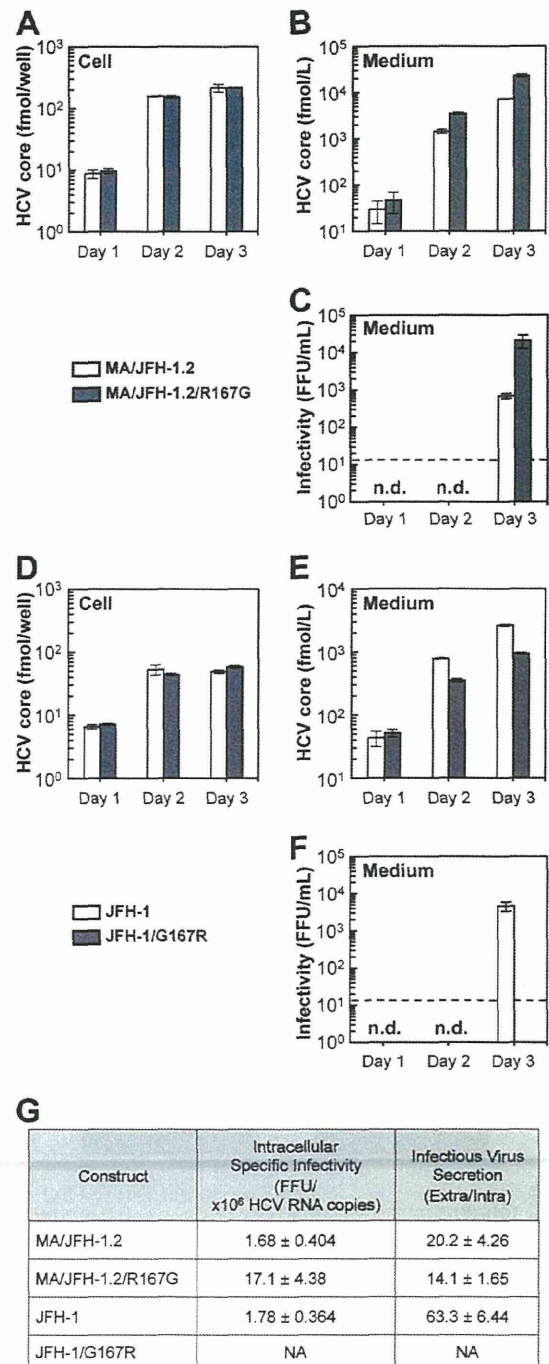


FIG 4 Effects of R167G on replication and virus production of MA/JFH-1.2 and JFH-1 in Huh7-25 cells. Ten micrograms of HCV RNA was transfected into Huh7-25 cells, and cells and medium were harvested on days 1, 2, and 3. HCV core protein levels in cells (A and D) and in medium (B and E) were measured, and infectivity of medium (C and F) was determined. n.d., not determined. Dashed line indicates the detection limit. (G) Intracellular specific infectivity and virus secretion efficiency of chimeric HCV RNA-transfected cells. Intracellular and extracellular infectivity of day 3 samples was determined, and specific infectivity and virus secretion rate were calculated. Assays were performed three times independently, and data are presented as means \pm standard deviation. NA, not available.

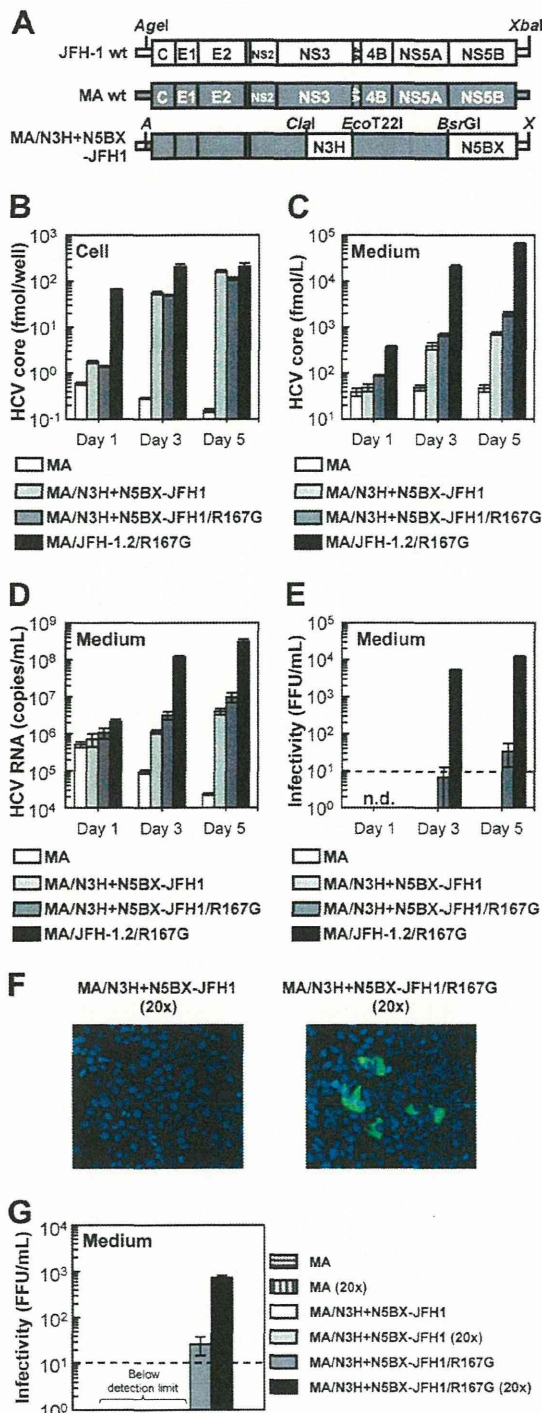


FIG 5 Replication and virus production of MA/N3H+N5BX-JFH1/R167G in Huh7.5.1 cells. (A) Schematic structures of JFH-1, MA, and MA/N3H+N5BX-JFH1. The junction of JFH-1 and MA in the 5' UTR is an AgeI site; the junctions of MA and JFH-1 in the NS3 regions are ClaI and EcoT22I sites, and the junction in the NS5B region is a BsrGI site. A, AgeI; X, XbaI. (B to G) Chimeric HCV RNA replication in Huh7.5.1 cells. Ten micrograms of HCV RNA was transfected into Huh7.5.1 cells, and cells and medium were harvested on days 1, 3, and 5. HCV core protein levels in cells (B) and in medium (C) and HCV RNA levels in medium (D) were measured, and infectivity of medium (E) was determined. Assays were performed three times independently, and data are presented as means \pm standard deviation. n.d., not determined. Dashed line indicates the detection limit. (F) Immunostained cells. Huh7.5.1

to be $7.27 \times 10^2 \pm 7.57 \times 10^1$ FFU/ml (Fig. 5G). No infected foci were observed after infection of MA/N3H+N5BX-JFH1 RNA-transfected cells, even when medium was concentrated (Fig. 5F), although intracellular and extracellular core protein levels were comparable to those with MA/N3H+N5BX-JFH1/R167G RNA-transfected cells (Fig. 5B and C). These results indicate that replacement of the 5' UTR and N3H and N5BX regions in JFH-1 were necessary to rescue autonomous replication in the replication-incompetent MA strain and for secretion of infectious chimeric virus. However, the secretion and infection efficiencies of the virus were low.

Cell culture-adaptive mutations enhanced infectious virus production of MA/N3H+N5BX-JFH1/R167G. Because MA/N3H+N5BX-JFH1/R167G replicated efficiently but produced very small amounts of infectious virus, we performed a long-term culture of the RNA-transfected cells in order to induce cell culture-adaptive mutations that could enhance infectious virus production. We prepared RNA-transfected cells using two constructs, MA/N3H+N5BX-JFH1 and MA/N3H+N5BX-JFH1/R167G; both of these replicated efficiently, and MA/N3H+N5BX-JFH1/R167G produced infectious virus at low levels while MA/N3H+N5BX-JFH1 did not. Immediately after transfection, the HCV core protein levels in the medium of each RNA-transfected cell culture peaked at 3.0×10^3 fmol/liter and declined thereafter. However, the core protein level in the medium with MA/N3H+N5BX-JFH1/R167G RNA-transfected cells continued to increase and reached a peak of 2.7×10^5 fmol/liter 54 days after transfection, at which point most cells were core protein positive (Fig. 6B). The core protein level in the medium with MA/N3H+N5BX-JFH1 RNA-transfected cells did not increase and core-positive cells were scarce on day 54 (Fig. 6B). We analyzed the viral genome in the culture supernatants from day 54 for possible mutations and identified four nonsynonymous mutations in the MA/N3H+N5BX-JFH1/R167G genome: L814S (NS2), R1012G (NS2), T1106A (NS3), and V1951A (NS4B). In order to test whether these amino acid substitutions enhance infectious virus production, L814S, R1012G, T1106A, and V1951A were introduced into MA/N3H+N5BX-JFH1/R167G, and the product was designated MA/N3H+N5BX-JFH1/5am (where am indicates adaptive mutation). On day 1, although HCV core protein levels in the MA/N3H+N5BX-JFH1/5am RNA-transfected cells were higher than those of MA/N3H+N5BX-JFH1/R167G RNA-transfected cells, they were still lower than those of MA/JFH-1.2/R167G RNA-transfected cells; however, on days 3 and 5, they reached a level comparable to that of MA/JFH-1.2/R167G RNA-transfected cells (Fig. 6C). HCV core protein and HCV RNA levels in the medium of MA/N3H+N5BX-JFH1/5am RNA-transfected cells were higher than those of MA/JFH-1.2/R167G RNA-transfected cells ($P < 0.05$, Fig. 6D and 6E, respectively). MA/N3H+N5BX-JFH1/5am, containing the four additional adaptive mutations, produced infectious virus at the same level as MA/JFH-1.2/R167G on day 5 (Fig. 6F). These results indicate that the

cells were infected with concentrated medium from RNA-transfected cells on day 5. Infected cells were visualized with anti-core antibody (green), and nuclei were visualized with DAPI (blue). (G) Infectivity of concentrated culture medium from HCV RNA-transfected cells. Culture medium was concentrated by 20 times. Infectivities of original and concentrated culture media were determined. Dashed line indicates detection limit.

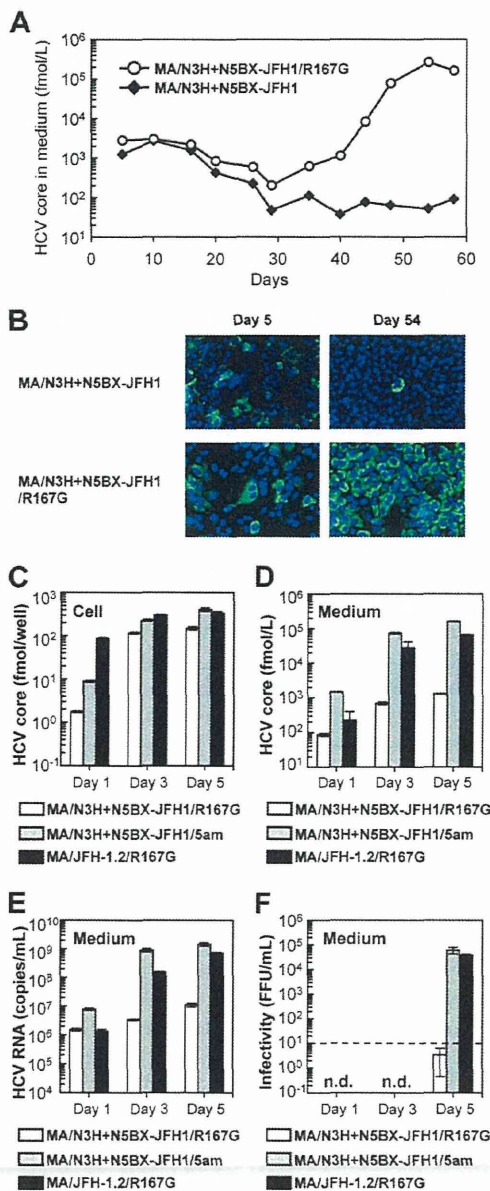


FIG 6 Cell culture-adaptive mutations enhanced infectious virus production of MA/N3H+N5BX-JFH1/R167G. (A) Long-term culture of MA/N3H+N5BX-JFH1 and MA/N3H+N5BX-JFH1/R167G RNA-transfected cells. Ten micrograms of HCV RNA was transfected into Huh7.5.1 cells, and cells were passaged every 2 to 5 days, depending on cell status. Culture medium was collected after every passage, and HCV core protein levels were measured. HCV core protein levels in culture medium from MA/N3H+N5BX-JFH1 and MA/N3H+N5BX-JFH1/R167G RNA-transfected cells are presented. (B) Immunostained cells on days 5 and 54 after transfection. Infected cells were visualized with anti-core antibody (green), and nuclei were visualized with DAPI (blue). (C to F) Effect of four additional cell culture-adaptive mutations on virus production. Ten micrograms of HCV RNA was transfected into Huh7.5.1 cells, and cells and medium were harvested on days 1, 3, and 5. HCV core levels in cells (C) and in medium (D) and HCV RNA levels in medium (E) were measured, and infectivity of medium (F) was determined. Assays were performed three times independently, and data are presented as means \pm standard deviation. n.d., not determined. Dashed line indicates the detection limit.

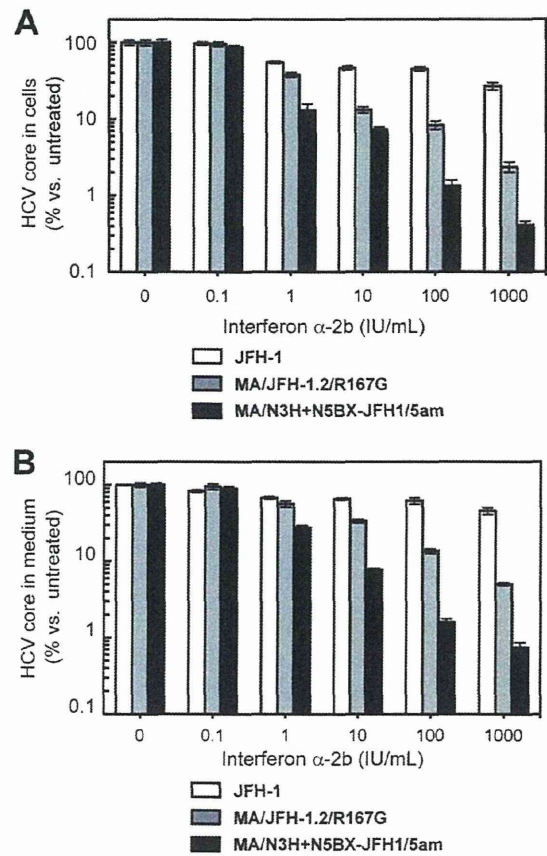


FIG 7 Comparisons of interferon sensitivity between JFH-1, MA/JFH-1.2/R167G and MA/N3H+N5BX-JFH1/5am. Two micrograms of HCV RNA was transfected into Huh7.5.1 cells, and interferon was added at the indicated concentrations at 4 h after transfection. HCV core protein levels in cells (A) and in medium (B) on day 3 were measured, and data are expressed as percent versus untreated cells (0 IU/ml). Assays were performed three times independently, and data are presented as means \pm standard deviation.

four additional adaptive mutations enhance infectious virus production and that MA/N3H+N5BX-JFH1/5am RNA-transfected cells replicate and produce infectious virus as efficiently as MA/JFH-1.2/R167G RNA-transfected cells.

Comparison of interferon sensitivity between JFH-1, MA/JFH-1.2/R167G, and MA/N3H+N5BX-JFH1/R167G. Using the newly established genotype 2b infectious chimeric virus, we compared interferon sensitivity between the JFH-1, MA/JFH-1.2/R167G, and MA/N3H+N5BX-JFH1/5am viruses. JFH-1 or MA chimeric viral RNA-transfected Huh7.5.1 cells were treated with 0.1, 1, 10, 100, or 1,000 IU/ml interferon α -2b, and HCV core protein levels in the cells and in culture media were compared. Interferon decreased HCV core protein levels in the JFH-1 RNA-transfected cells and in the medium in a dose-dependent manner, and production was inhibited to 26.8% \pm 3.0% and 45.6% \pm 4.7%, respectively, of control levels (Fig. 7A and B, respectively). In contrast, HCV core protein levels in cells and medium of MA/JFH-1.2/R167G and MA/N3H+N5BX-JFH1/5am RNA-transfected cells decreased more pronouncedly in a dose-dependent manner (Fig. 7A and B, respectively). HCV core protein levels in cells and medium from MA/N3H+N5BX-JFH1/5am RNA-transfected cells were lower than those from MA/JFH-1.2/

R167G RNA-transfected cells (Fig. 7A and B, respectively) ($P < 0.05$ at 1, 10, 100, and 1,000 IU/ml), indicating that the MA/N3H+N5BX-JFH1/5am virus was more sensitive to interferon than the MA/JFH1.2/R167G virus, which contained more regions from JFH-1.

DISCUSSION

In this study, we developed a novel infectious HCV production system using a genotype 2b chimeric virus. To improve infectious virus production, we introduced two modifications into the chimeric genome.

First, we replaced the 5' UTR from MA with that of JFH-1. Similarly to J6/JFH-1, replacement of the 5' UTR increased core protein accumulation in both the cells and medium when these RNAs were transfected into Huh7.5.1 cells (Fig. 1). The same trend was observed when these RNAs were transfected into Huh7-25 cells (data not shown), indicating that the 5' UTR of JFH-1 enhanced RNA replication. There are two genetic variations in J6CF and seven in MA in the region we replaced (nt 1 to 154 for J6CF and nt 1 to 155 for MA), and some of these mutations may affect RNA replication by changing the RNA secondary structure, RNA-RNA interactions, or binding of host or viral proteins.

Second, we introduced a cell culture-adaptive mutation (R167G) in the core region. This mutation was induced by long-term culture of MA/JFH-1 RNA-transfected cells (Fig. 2). MA/JFH-1 chimeric RNA (MA/JFH-1.1 and MA/JFH-1.2) replicated when synthesized RNA was transfected into the cells. However, infectious virus production was low, and virus infection did not spread over the short term. In early stages of long-term culture, the number of core protein-positive cells gradually decreased, and core protein-positive cells were scarcely detectable. Subsequently, the population of core protein-positive cells increased, reaching almost 100%. At this time point, we identified a common mutation in the core region (R167G) of the viral genome as a cell culture-adaptive mutation and found that it enhanced infectious virus production (Fig. 3). Several nonsynonymous mutations other than R167G were identified in the viral genome from each supernatant, and these mutations may enhance infectious virus production. However, there was a discrepancy between RNA levels and the infectivity of the culture media of MA/JFH-1.2 and MA/JFH-1.2/R167G RNA-transfected cells (Fig. 3C and D). The MA/JFH-1.2/R167G mutant had a 2-log increase in viral infectivity compared to that of MA/JFH-1.2 but only a 1-log increase in secreted RNA. The replication efficiency of MA/JFH-1.2 RNA-transfected cells was comparable to that of MA/JFH-1.2/R167G RNA-transfected cells, but the efficiency of infectious virus assembly within the cells was low, indicating that mainly noninfectious virus may be produced.

Infection of MA/JFH-1.2/R167G virus spreads rapidly, similarly to that of the JFH-1 virus, when it is inoculated into naïve Huh7.5.1 cells. On a single-cycle virus production assay, we found that the R167G mutation did not affect RNA replication or virus secretion but enhanced infectious virus assembly within the cells (Fig. 4). Efficient infectious virus assembly within the cells was mainly responsible for the rapid spread and high virus production of MA/JFH-1.2/R167G.

The amino acid at 167 (aa 167) is located in domain 2 of the core region, which is important for localization of the core

protein (3, 8). Lipid droplet localization of the core protein and/or NS5A is important for infectious virus production (4, 18, 26). The interaction between the core protein and NS5A is also important for infectious virus production (16). Thus, aa 167 affects infectious virus production possibly by altering subcellular localization of the core protein or interaction between the core protein and NS5A. We examined the amino acid sequence of the core protein in 2,078 strains in the Hepatitis Virus Database (<http://s2as02.genes.nig.ac.jp/>) and found that aa 167 is Gly in all other strains. These data strongly suggest that Gly at aa 167 is important for the HCV life cycle. As the MA strain was cloned from the serum of a patient with chronic hepatitis C, the low virus production by this Gly at aa 167 may be important for persistent infection.

We then attempted to reduce the contents of JFH-1 from MA/JFH-1.2/R167G. We previously reported that the N3H and N5BX regions of JFH-1 were sufficient for replication of the J6CF strain (21). We also reported that this effect was observed only in genotype 2a strains (J6CF, JCH-1, and JCH-4). In this study, we tested whether the N3H and N5BX regions of JFH-1 could also support replication of a genotype 2b strain, MA. We constructed an MA chimeric virus harboring the N3H and N5BX regions of JFH-1 and combined this with the 5' UTR of JFH-1 and the R167G mutation (MA/N3H+N5BX-JFH1/R167G). This chimeric RNA was able to replicate in the cells and produce infectious chimeric virus in culture medium although infectious virus production levels were low (Fig. 5).

We showed in this paper that the N3H and N5BX regions of JFH-1 were able to support RNA replication by both genotype 2a clones and genotype 2b clones, but the nucleotide sequence similarity between JFH-1 and MA was lower than that between JFH-1 and J6CF (77% versus 89%, respectively). Compared to MA/JFH-1.2/R167G, MA/N3H+N5BX-JFH1/R167G RNA showed the same levels of RNA replication and low levels of infectious virus production. To clarify whether there were any differences in the characteristics of the secreted virus, we performed density gradient ultracentrifugation with the MA/JFH-1.2/R167G and MA/N3H+N5BX-JFH1/R167G viruses. The distributions of the HCV core protein and infectivity showed similar profiles (data not shown).

The differences between MA/JFH-1.2/R167G and MA/N3H+N5BX-JFH1/R167G are the NS2, NS3 protease domain (N3P), and NS4A to NS5A regions. Nucleotide variation(s) other than aa 167 in these regions of the MA strain may be associated with reduced virus assembly. We identified four additional cell culture-adaptive mutations, L814S (NS2), R1012G (NS2), T1106A (NS3), and V1951A (NS4B), which resulted from long-term culture of MA/N3H+N5BX-JFH1/R167G RNA-transfected cells. Consequently, cells transfected with MA/N3H+N5BX-JFH1/5am constructed by insertion of these four adaptive mutations into MA/N3H+N5BX-JFH1/R167G replicated and produced infectious virus as efficiently as MA/JFH-1.2/R167G RNA-transfected cells (Fig. 6).

This system is able to contribute to studies into the development of antiviral strategies. It has been reported that HCV genotype 2a was more sensitive to interferon therapy than HCV genotype 2b in a clinical study (20). To assess the interferon resistance of genotype 2b, a cell culture system with multiple genotype 2b strains is necessary. The previously reported replicable genotype 2b chimeric virus harbored only structural

regions of 2b strains (6, 27). The 2b/JFH-1 chimeric virus containing the region of the core protein to NS2 from the J8 strain (genotype 2b) and the region of NS3 to 3' X of JFH-1 was able to replicate and showed that there were no differences in interferon sensitivity among the JFH-1 chimeric viruses of other genotypes (6, 27). Another 2b/JFH-1 chimeric virus containing the regions of the core protein to NS2 (nt 342 to 2867) of a genotype 2b strain and of NS2 to 3' UTR (nt 2868) of JFH-1 has been reported (6, 27). The authors reported that their 2b/JFH-1 chimeric virus was more sensitive to interferon than JFH-1 (6, 27). We developed the genotype 2b HCV cell culture system with another HCV genotype 2b strain (MA). We identified a virus assembly-enhancing mutation in the core region, the minimal JFH-1 regions necessary for replication, and four additional adaptive mutations that enhance infectious virus production and demonstrated that MA harboring the five adaptive mutations and the 5' UTR and N3H and N5BX regions of JFH-1 (MA/N3H+N5BX-JFH1/5am) could replicate and produce infectious virus efficiently.

Using these novel genotype 2b chimeric viruses, we assessed interferon sensitivity. We found that MA/JFH-1.2/R167G chimeric virus and MA/N3H+N5BX-JFH1/5am virus were more sensitive to interferon than the JFH-1 virus (Fig. 7). Furthermore, we found that MA/N3H+N5BX-JFH1/5am was more sensitive to interferon than MA/JFH-1.2/R167G, indicating that the genetic variation(s) in the NS2, N3P, and NS4A to NS5A regions affect interferon sensitivity. Although genotype 2a viruses are more sensitive to interferon than genotype 2b viruses in clinical studies, JFH-1 displayed interferon resistance in our study.

These results suggest that the JFH-1 regions in the 2b/JFH-1 virus affect the interferon sensitivity of the chimeric virus. Moreover, it was reported that amino acid variations in E2, p7, NS2, and NS5A were associated with the response to peginterferon and ribavirin therapy in genotype 2b HCV infection (10). Therefore, our MA/JFH-1 chimeric virus harboring minimal regions from JFH-1 (MA/N3H+N5BX-JFH1/5am) is more suitable for assessing the characteristics of the MA strain than the MA/JFH-1 chimeric virus, which includes a nonstructural region from JFH-1 (MA/JFH-1.2/R167G). We showed here that replacement of the 5' UTR and N3H and N5BX regions in MA with those from JFH-1 is able to convert MA into a replicable virus. Using the same strategy, numerous HCV cell culture systems with various genotype 2b strains, as well as genotype 2a strains, may be available.

In conclusion, we established a novel HCV genotype 2b cell culture system using a chimeric genome in MA harboring minimal regions from JFH-1. This cell culture system using the chimeric genotype 2b virus will be useful for characterization of genotype 2b viruses and the development of antiviral strategies.

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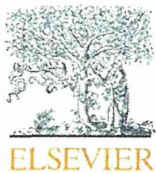
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Potent and selective inhibition of hepatitis C virus replication by novel phenanthridinone derivatives

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ABSTRACT

A number of novel phenanthridinone derivatives were examined for their inhibitory effect on hepatitis C virus (HCV) replication in Huh-7 cells harboring self-replicating subgenomic viral RNA replicons with a luciferase reporter (LucNeo#2). The activity of compounds was further confirmed by inhibition of viral RNA copy number in different subgenomic and full-genomic replicon cells using real-time reverse transcription polymerase chain reaction. Among the compounds, 4-butyl-11-(1,1,1,3,3,3-hexafluoro-2-hydroxypropan-2-yl)-7-methoxy-[1,3]dioxolo[4,5-c]phenanthridin-5(4H)-one (HA-719) was found to be the most active with a 50% effective concentration of $0.063 \pm 0.010 \mu\text{M}$ in LucNeo#2 cells. The compound did not show apparent cytotoxicity to the host cells at concentrations up to $40 \mu\text{M}$. Western blot analysis demonstrated that HA-719 reduced the levels of NS3 and NS5A proteins in a dose-dependent fashion in the replicon cells. Interestingly, the phenanthridinone derivatives including HA-719 were less potent inhibitors of JFH1 strain (genotype 2a HCV) in cell-free virus infection assay. Although biochemical assays revealed that HA-719 proved not to inhibit NS3 protease or NS5B RNA polymerase activity at the concentrations capable of inhibiting viral replication, their molecular target (mechanism of inhibition) remains unknown. Considering the fact that most of the anti-HCV agents currently approved or under clinical trials are protease and polymerase inhibitors, the phenanthridinone derivatives are worth pursuing for their mechanism of action and potential as novel anti-HCV agents.

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1. Introduction

Hepatitis C virus (HCV) infection is a worldwide problem. More than 130 million individuals are infected with this virus, and 3–4 million are newly infected every year. In general, HCV infection proceeds to chronic infection [1], which often induces liver cirrhosis and hepatocellular carcinoma [2] liver transplantation is the only way to rescue patients with the end-stage liver disorders caused by HCV infection [3]. Protective vaccines are not available so far, and pegylated interferon (PEG-IFN) and the nucleoside analog ribavirin are the standard treatment for HCV infection [4–6]. However, many patients cannot tolerate the serious side effects of PEG-IFN and ribavirin. Therefore, the development of novel agents with better efficacy and tolerability is still mandatory.

HCV is an enveloped virus belonging to the hepacivirus genus of the family *Flaviviridae* [7,8]. The viral genome consists of positive sense single RNA coding a polyprotein cleaved by viral and host proteases into four structural and six non-structural proteins. Non-structural proteins are involved in the replication of HCV genome [9]. The discovery of effective anti-HCV agents was greatly hampered by the lack of cell culture systems that allowed robust propagation of HCV in laboratories. However, the development of HCV RNA replicon systems [10] and recent success in propagating infectious virus particles in vitro have provided efficient tools for screening new antiviral agents against HCV replication [11,12]. Furthermore, replicons containing a reporter gene, such as luciferase and green fluorescence protein, have provided fast and reproducible screening of a large number of compounds for their antiviral activity [13–15].

Currently, two NS3 protease inhibitors, terapeutic and bocoprevir, have been licensed and a considerable number of novel anti-HCV agents are under clinical trials [16,17]. Most of them are directly acting inhibitors of NS3 protease or NS5B polymerase. However, the

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emergence of HCV mutants resistant to most of these agents has also been reported [18]. To circumvent the drug-resistance, it seems necessary to use more than two directly acting drugs targeting different molecules for inhibition of viral replication [19]. Thus, in addition to the protease and polymerase inhibitors, novel compounds with a unique mechanism of action are highly desired.

We have recently identified some compounds with a novel phenanthridinone structure as moderate inhibitors of HCV replication [20]. This prompted us to synthesize a number of phenanthridinone derivatives and investigate their anti-HCV activity. After optimization of chemical structures, we have obtained the compounds that exert anti-HCV activity in the nanomolar range. Interestingly, these compounds did not inhibit the enzymatic activity of NS3 protease or NS5B RNA polymerase at the concentrations capable of inhibiting HCV replication in replicon cells.

2. Materials and methods

2.1. Compounds

More than 100 phenanthridinone derivatives were synthesized and used in this study. The synthesis of these compounds has been described previously [20,21]. Cyclosporin A (CsA) was purchased from Sigma–Aldrich. All compounds were dissolved in dimethyl sulfoxide (DMSO) (Nacalai Tesque) at a concentration of 20 mM or higher to exclude the cytotoxicity of DMSO and stored at -20°C until use.

2.2. Cells

Huh-7 cells were grown and cultured in Dulbecco's modified Eagle medium with high glucose (Gibco/BRL) supplemented with 10% heat-inactivated fetal bovine serum (Gibco/BRL), 100 U/ml penicillin G, and 100 $\mu\text{g}/\text{ml}$ streptomycin. Huh-7 cells containing self-replicating subgenomic HCV replicons with a luciferase reporter, LucNeo#2 [22], were maintained in culture medium containing 1 mg/ml G418 (Nakarai Tesque). The subgenomic replicon cells without reporter #50-1 and the full-genomic replicon cells NNC#2 [23] were kindly provided by Dr. Hijikata (Kyoto University, Kyoto, Japan). These cells were also maintained in culture medium containing 1 mg/ml G418.

2.3. Anti-HCV assays

The anti-HCV activity of the test compounds was determined in LucNeo#2 cells by the previously described method with some modifications [24]. Briefly, the cells (5×10^3 cells/well) were cultured in a 96-well plate in the absence of G418 and in the presence of various concentrations of the compounds. After incubation at 37°C for 3 days, the culture medium was removed, and the cells were washed twice with phosphate-buffered saline (PBS). Lysis buffer was added to each well, and the lysate was transferred to the corresponding well of a non-transparent 96-well plate. The luciferase activity was measured by addition of the luciferase reagent in a luciferase assay system kit (Promega) using a luminometer with automatic injectors (Berthold Technologies).

The activity of the test compounds was also determined by the inhibition of HCV RNA synthesis in LucNeo#2, #50-1, and NNC#2 cells [23,25]. The cells (5×10^3 cells/well) were cultured in a 96-well plate in the absence of G418 and in the presence of various concentrations of the compounds. After incubation at 37°C for 3 days, the cells were washed with PBS, treated with lysis buffer in TaqMan[®] Gene Expression Cell-to-CT[™] kit (Applied Biosystems), and the lysate was subjected to real-time reverse transcription polymerase chain reaction (RT-PCR), according to the

manufacturer's instructions. The 5'-untranslated region of HCV RNA was quantified using the sense primer 5'-CGGGAGAGCCA-TAGTGG-3', the antisense primer 5'-AGTACCACAAGGCCTTTCG-3', and the fluorescence probe 5'-CTGCGGAACCGGTGAGTACAC-3' (Applied Biosystems).

The inhibitory effect of the test compounds on the replication of a genotype 2a strain was evaluated by the infection of Huh-7.5.1 cells, kindly provided by Dr. Chisari at Scripps Institute, with cell-free JFH-1 virus, as previously described [11]. At 48 h after virus infection, the cells were treated with SideStep Lysis and Stabilization Buffer (Agilent Technologies), and the lysate was subjected to real-time RT-PCR for quantification of HCV RNA [25].

2.4. Cytotoxicity assay

Huh-7 cells (5×10^3 cells/well) were cultured in a 96-well plate in the presence of various concentrations of the test compounds. After incubation at 37°C for 3 days, the number of viable cells was determined by a dye method using the water soluble tetrazolium Tetracolor One[®] (Seikagaku Corporation), according to the manufacturer's instructions. The cytotoxicity of the compounds was also evaluated by the inhibition of host cellular mRNA synthesis. The cells were treated with lysis buffer in the kit, as described above, and the cell lysate was subjected to real-time RT-PCR for amplification of a part of glyceraldehyde-3-phosphate dehydrogenase (GAPDH) RNA using a TaqMan[®] RNA control reagent (Applied Biosystems).

2.5. Immunoblotting

LucNeo#2 cells (5×10^3 cells/well) were cultured in a 96-well plate in the presence of various concentrations of the test compounds. After incubation at 37°C for 4 days, the culture medium was removed, and the cells were washed with PBS and treated with lysis buffer (RIPA Buffer[®], Funakoshi). The protein concentration of the lysate was measured by Bradford protein assay method (Bio-Rad). Then, the lysate was subjected to sodium dodecyl sulfate polyacrylamide gel electrophoresis (SDS-PAGE). The primary antibodies used for protein detection were anti-NS3 (Thermo Scientific), anti-NSSA (Acris Antibodies), and anti-GAPDH (Santa Cruz Biotechnology) mouse monoclonal antibodies.

2.6. Protease and polymerase inhibition assays

The effect of the test compounds on NS3 protease activity was determined by a fluorescence resonance energy transfer-based assay using SensoLyte[®] 520 HCV Assay Kit (AnaSpec), according to the manufacturer's instructions. The inhibition assay for NS5B polymerase was performed at 37°C for 60 min in a 384-well plate. A reaction mixture (30 $\mu\text{l}/\text{well}$) contains 20 mM Tris-HCl (pH 7.6), 10 mM MgCl_2 , 20 mM NaCl, 1 mM dithiothreitol, 0.05% Tween 20, 0.05% pluronic F127, 1 μM [³H]GTP (0.1 $\mu\text{Ci}/\text{well}$) plus cold GTP, 5 nM poly(rC), 62.5 nM biotinylated dG₁₂, 45 nM recombinant NS3 protease, and various concentrations of the compounds. The reaction was stopped by streptavidin scintillation proximity assay beads in 0.5 M ethylenediaminetetraacetic acid. The plate was counted with a microbeta reader on the following day.

3. Results

When a number of phenanthridinone derivatives were examined for their antiviral activity in LucNeo#2 cells, three phenanthridinone derivatives, 5-butyl-2-(1,1,1,3,3,3-hexafluoro-2-hydroxypropan-2-yl)-3,8-dimethoxyphenanthridin-6(5H)-one (KZ-16), 4-butyl-11-(1,1,1,3,3,3-hexafluoro-2-hydroxypropan-2-yl)-[1,3]dioxolo[4,5-c]