Cell-Cell Contact-Mediated Hepatitis C Virus (HCV) Transfer, Productive Infection, and Replication and Their Requirement for HCV Receptors

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Hepatitis C virus (HCV) infection is believed to begin with interactions between cell-free HCV and cell receptors that include CD81, scavenger receptor B1 (SR-B1), claudin-1 (CLDN1), and occludin (OCLN). In this study, we have demonstrated that HCV spreading from infected hepatocytes to uninfected hepatocytes leads to the transfer of HCV and the formation of infection foci and is cell density dependent. This cell-cell contact-mediated (CCCM) HCV transfer occurs readily and requires all these known HCV receptors and an intact actin cytoskeleton. With a fluorescently labeled replication-competent HCV system, the CCCM transfer process was further dissected by live-cell imaging into four steps: donor cell-target cell contact, formation of viral puncta-target cell conjugation, transfer of viral puncta, and postransfer. Importantly, the CCCM HCV transfer leads to productive infection of target cells. Taken together, these results show that CCCM HCV transfer constitutes an important and effective route for HCV infection and dissemination. These findings will aid in the development of new and novel strategies for preventing and treating HCV infection.

Approximately 170 million people, 3% of the world’s population, are currently infected with hepatitis C virus (HCV) (1). The infection frequently leads to hepatitis and liver steatosis and is considered a leading cause of life-threatening chronic liver diseases, such as liver fibrosis, cirrhosis, and hepatocellular carcinoma (2). In the United States and Europe, HCV infection has become the primary cause for liver transplantation (3). Despite intensive research efforts during the last 2 decades, no HCV vaccines have become available (4, 5). The first two HCV-specific antivirals, the HCV protease NS3/NS4 inhibitors telaprevir and boceprevir, were approved by the FDA in 2011, yet combinatorial treatment with these inhibitors and pegylated alpha interferon became the primary cause for liver transplantation (3). Despite intensive research efforts during the last 2 decades, no HCV vaccines have become available (4, 5). 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lular virus transfer (21, 26), while HSV exploits the tight junctions among epithelial cells for viral spreading (22).

In this study, we established a coculture assay, demonstrated CCCM HCV infection in hepatocytes, including primary human hepatocytes (PHHs), and characterized the roles of known HCV receptors and cytoskeletal structures in this process. In addition, we adapted the tetracycline (Tc)-biarsenic dye labeling strategy in combination with three-dimensional (3D) live-cell fluorescence microscopy imaging and dissected the spatial and temporal details of the transfer process. We confirmed the transfer of micrometer-sized HCV core protein puncta into target cells in real time and illustrated the four steps in CCCM HCV transfer: donor cell (D)-target cell (T) contact, formation of viral puncta-target cell interaction (the TC tag is shown in bold). The region from the 5' UTR to NS2 containing the TC-core protein (TCCore) was then cloned between the pIFH1 plasmid using EcoRI and NotI to obtain pIFH1-TCore. All recombinant plasmids were verified by sequencing. SIGENOME SMARTPool small interfering RNAs (siRNAs) against human CD81 (siCD81), SR-B1 (siSR-B1), CLDN1 (siCLDN1), and OCLN (siOCLN) and scrambled control siRNA (siCtrl) were purchased from Dharmacon (Lafayette, CO).

**Viral RNA preparation and siRNA/plasmid/viral RNA transfections.** pIFH1 and pIFH1-TCore were linearized by XbaI and used as the templates to transcribe viral RNAs in vitro using a MEGAscript kit (Ambion, CA). The synthesized viral RNA was purified by acidic phenol-chloroform extraction. Viral RNA transfection into HuH7.5.1 cells, siRNA transfection of HuH7.5.1 cells, pcDNA3-CD81 transfection of HepG2 cells, and pcMV-CLDN1 transfection of CYNK10 cells were performed with the Lipofectamine 2000 reagent (Invitrogen) according to the manufacturer’s instructions. Media were replaced at 6 h posttransfection. 293T cells were transfected with pcDNA3-CD81 and pcDNA3-SR-B1 using the standard calcium phosphate transfection method, and media were replaced after an overnight incubation. Transfection efficiencies of plasmid DNA were monitored by pEGFP-N3 transfection and were 80 to 90% for 293T cells, 40 to 50% for HepG2 cells, and 50 to 60% for NKNT3 and CYNK10 cells.

**Virus production and titration.** HuH7.5.1 cells were either transfected with viral RNAs (JFH1 or JFH1-TCcore) as described above or inoculated with JFH1 viral stock (a generous gift from Wenzhe Ho of Temple University, Philadelphia, PA) at a multiplicity of infection (MOI) of 0.2 to 1. Supernatants containing HCV virions were collected at 30 to 40 days posttransfection or 5 days postinfection, cell debris was removed by low-speed centrifugation (1,000 x g, 4°C, 10 min), and the supernatants were passed through a 0.22-μm-pore-size filter and stored at −70°C. The filtered culture supernatant was concentrated by the addition of a 1/5 volume of 40% PEG 8000 to make a final concentration of 8%, and virus was precipitated at 4°C overnight, followed by centrifugation at 3,500 x g and 4°C for 30 min. The precipitated virus was suspended, titrated as described below, and stored frozen at −70°C. To titrate the virus stock, 10-fold serially diluted stock was added to HuH7.5.1 cells, and medium was changed after 2 h. At 72 h postinfection, immunostaining against HCV core protein was performed. The number of foci formed at the highest dilution was used to calculate the virus titer, which was expressed as the number of focus-forming units per milliliter of supernatant (FFU/ml). The titers of our JFH1 viral stock were usually in the range of 10^4 to 10^6 FFU/ml.

**Coculture assay and transwell assay.** In the coculture assay, HuH7.5.1 cells were infected with HCV for 3 days and used as the donor cells. The donor cells or the target cells were first labeled with PKH26 or CMFDA according to the manufacturer’s instructions. Briefly, cells were trypsinized and labeled with 2 μM PKH26 for 5 min at room temperature, followed by incubation with 1% FBS–phosphate-buffered saline (PBS) to stop the labeling reaction and four extensive washes to remove any residual dye. Alternatively, media containing 1.25 μM (for flow cytometry) or 5 μM (for confocal imaging) CMFDA were directly added to cells growing in culture dishes. The cells were labeled at 37°C and 5% CO2 for 30 min, followed by a medium change to normal growth medium and an extra 30 min of incubation. The labeled donor cells/target cells were counted (or trypsinized and counted for CMFDA-labeled cells) and mixed with unlabeled target/donor cells and seeded to a 24- or 12-well plate at >90% confluence. Unless stated otherwise, the donor cells/target cells were allowed to coincubate for approximately 20 h, followed by collection for immunostaining and flow cytometry or confocal analysis. The transwell...
assay was carried out in a similar fashion, except that neither donor nor target cells were labeled and the donor and target cells were seeded into the upper and lower chambers of the transwell (Corning, Lowell, MA), respectively.

Flow cytometry. Cells from the coculture assay or the target cells from the transwell assay were trypsinized and fixed with 4% paraformaldehyde at room temperature for 10 min, followed by permeabilization with 50 μg/ml saponin at room temperature for 15 min. Staining was performed with mouse anti-core protein primary antibody (2 μg/ml) at room temperature for 60 min, followed by goat anti-mouse IgG-PE (4 μg/ml, for the CMFDA-labeled or Huh7.5.1-GFP cell coculture) or goat anti-mouse IgG-Alexa 488 (2 μg/ml, for the PKH26-labeled coculture) secondary antibody at room temperature for 60 min. Cells were then suspended in PBS and subjected to flow cytometry analysis (with a BD FACSCalibur or BD Accuri C6 flow cytometer). For the detection of CD81 surface expression, siRNA-transfected Huh7.5.1 cells or pcDNA3/CD81-transfected HepG2 cells were removed from culture dishes with 0.5 mM EDTA-PBS and stained with mouse normal IgG (2 μg/ml) or mouse anti-CD81 (2 μg/ml), followed by goat anti-mouse IgG-PE or goat anti-mouse IgG-Alexa 488. Cells were suspended in PBS and immediately analyzed by flow cytometry. Between each step, the cells were extensively washed with PBS.

Confocal microscopy. Cells from the coculture assay were fixed with 4% paraformaldehyde on coverslips, permeabilized with saponin, and blocked in 1% PBS-PBS at room temperature for 30 min. Cells were stained with mouse anti-core protein primary antibody at room temperature for 60 min, followed by goat anti-mouse IgG-Alexa 488 or -Alexa 555 (2 μg/ml) secondary antibody at room temperature for 60 min. The nuclei were stained with 0.25 μg/ml DAPI in PBS at room temperature for 15 min. The coverslips were mounted with Fluorosphore Mountant (Southern Biotech, Birmingham, AL), and microscopic images were taken using a Zeiss Axiovert 200 microscope for epifluorescence images or a Zeiss LSM510 microscope for confocal images. For the detection of OCLN, siCtrl- or siOCLN-transfected Huh7.5.1 cells were fixed, permeabilized, and stained with mouse normal IgG (2.5 μg/ml) or mouse anti-OCLN (2.5 μg/ml), followed by goat anti-mouse IgG-Alexa 555 secondary antibody. For F-actin staining, cells from the coculture assay were fixed and permeabilized as described above and then stained with 1 μg/ml phalloidin-TRITC in PBS at room temperature for 30 min. For the visualization of microtubules, cells from the coculture assay were fixed with −20°C methanol for 3 min and then permeabilized and stained with mouse antitubulin (2 μg/ml), followed by Alexa 555-conjugated secondary antibody. Cells were washed three times with PBS between each step.

Western blotting. Cell lysates were prepared at 48 h posttransfection or 72 h postinfection using a standard radioimmunoprecipitation assay buffer (50 mM Tris-HCl, pH 8.0, 0.5% NP-40, 2 mM EDTA, 137 mM NaCl, 10% glycerol) and separated on a 12% (for HCV core protein or CLDN) or 8% (for SR-B1 or OCLN) polyacrylamide-SDS gel. The proteins were transferred onto nitrocellulose membranes, probed with appropriate primary antibodies (1 μg/ml for mouse anti-HCV core protein and rabbit anti-CLDN1 antibodies, 2 μg/ml for mouse anti-SR-B1 and rabbit anti-OCLN antibodies, 0.5 μg/ml for anti-β-actin antibody) and HRP-conjugated secondary antibodies (0.5 μg/ml), and visualized by the use of homemade enhanced chemiluminescence reagents.

RNA extraction and qRT-PCR for HCV RNA. RNA was isolated from cells or cell debris-free culture supernatants (50 to 200 μl) using the TRizol or TRizol LS reagent, respectively, according to the manufacturer’s instructions and used for quantitative real-time reverse transcription-PCR (qRT-PCR) amplification with a TaqMan one-step RT-PCR kit (Applied Biosystems, Foster City, CA) and HCV-specific primers and probe, which were described previously (29). qRT-PCR was performed on a C1000 Touch thermal cycler (Bio-Rad, CA), and signal detection was with a CFX96 real-time system (Bio-Rad). The PCR program was as follows: 48°C for 30 min, 95°C for 10 min, and then 40 cycles at 95°C for 15 s and 60°C for 1 min. Serially diluted in vitro-transcribed JFH1 RNA was included as a standard in parallel and used to calculate the absolute level of HCV RNA.

ReAsH labeling of JFH1-TCcore and live-cell imaging. Culture supernatants were collected from JFH1-TCcore-transfected Huh7.5.1 cells at day 42 posttransfection, concentrated 30-fold by using PEG 8000 precipitation, and titrated as described above. Huh7.5.1 cells were infected with the concentrated JFH1-TCcore virus (MOI = 0.15) and labeled with ReAsH according to the instructions provided in the TC tag detection kit (Invitrogen). These labeled cells were then used as donor cells. Specifically, at day 3 postinfection, cells were washed once with Opti-MEM medium (Invitrogen) and labeled with the ReAsH dye (final concentration, 1 μM) in Opti-MEM medium. Cells were incubated at 37°C for 30 min and then washed twice with PBS, trypsinized, and mixed at 1:1 ratio with the Huh7.5.1-GFP target cells. The mixed cells were seeded onto polylysine-coated 35-mm glass-bottom dishes to 90% confluence, and the media were replaced with DMEM without phenol red at 2 h postseeding. Live-cell imaging was set up after the medium change. Images were taken starting at 9 h postseeding and continued for 18 h. All images were acquired with a Zeiss LSM 510 confocal microscope fitted with a ×20 air objective and a thermostatic stage incubator set at 37°C in 5% CO2. ReAsH-labeled TCcore was visualized with filter sets with excitation at 488 nm and emission at 528/45 nm. A 13-slice by 0.9-μm Z stack was taken at each selected field every 18 min using LSM 510 LVO software. Images were later processed and analyzed with the ImageJ 1.45s program. Unless otherwise specified, all red/green fluorescence images were Z projections (maximum intensity projection) from the original stack, and all differential interference contrast (DIC) images were the most focused slice from the Z stack (the 7th slice).

Data analysis. Where appropriate, values are expressed as the mean ± standard deviation of triplicate experiments. All statistical analyses were done by one-way and two-way analyses of variance followed by post hoc tests (with the Bonferroni correction or Dunnett’s test). A P value of <0.05 was considered statistically significant, a P value of <0.01 was considered highly significant, and a P value of <0.001 was considered strongly significant. All data are representative of multiple repeated experiments.

RESULTS

Formation of HCV infection foci in a cell density-dependent manner. We repeatedly noticed that during the production and titration of HCV JFH1 virus in Huh7.5.1 cells, more distinct infection foci (clustered HCV-infected cells, usually with 10 to 50 cells) were present at the lower MOI (Fig. 1a, right) and fewer foci were present at the higher MOI and gave rise to approximately 80 to 90% HCV core protein-positive cells (Fig. 1a, middle). Formation of these infection foci at the lower MOI could result from division and proliferation of HCV-infected cells. Alternatively, it could be due to the close proximity of uninfected cells to HCV-infected cells and subsequent CCCM HCV transfer to the neighboring uninfected cells, resulting in infection. To distinguish these two possibilities, we infected Huh7.5.1 cells with JFH1 virus at a lower MOI, removed the excess virus, and then replated the cells at different cell densities. Following 3 days of continued culturing, the cells were processed for analysis of expression of the HCV core protein, a marker widely used for HCV infection. We reasoned that if infection focus formation resulted from proliferation of HCV-infected cells, there would be no correlation between the number of infection foci and cell density; if infection focus formation resulted from CCCM HCV transfer, there would be a positive correlation between the number of foci formed and cell density.
To ensure the validity of the data, we assessed HCV core protein-positive cells by Western blotting for HCV core protein, immunofluorescence staining for HCV core protein followed by immunofluorescence microscopic imaging, manual counting of HCV core protein-positive cells under an immunofluorescence microscope, or flow cytometry analysis. Western blotting showed that the level of HCV core protein expression exhibited a gradual increase with a higher cell density (Fig. 1b). Immunostaining of these cells for HCV core protein followed by immunofluorescence microscopic imaging showed an apparent increase in the number and intensity of core protein-positive cells with increased cell density (Fig. 1c). The formation of core protein-positive cell foci appeared to occur only at higher cell densities. Manual quantitation of core protein-positive cells showed a positive correlation between the number of core protein-positive cells and cell density (Fig. 1d). The positive correlation was further quantified by flow cytometry analysis of core protein-positive cells (Fig. 1e), and a two-variable linear regression analysis gave rise to a correlation coefficient of $R^2 = 0.9883$, which was derived from a standard two-variable mathematical regression analysis: cell confluence versus percentage of core-positive cells (e). REL, relative values (core expression in relation to the loading control β-actin and the value of HCV-infected cells at the 100% confluence was set as 1).

**CCCM HCV transfer to Huh7.5.1 and PHHs.** To characterize CCCM HCV infection, we devised a coculture assay that involved incubation of HCV-infected Huh7.5.1 donor cells with uninfected Huh7.5.1 cells for a certain period of time and determination of the number of core protein-positive target cells from CCCM HCV transfer by immunostaining. CCCM viral transmission requires different lengths of coincubation for different viruses and is often dependent on the ratio of donor cells (D) to target cells (T) (18, 30). Therefore, we started the coculture assay with a D/T ratio of 1:1 and a coincubation time of 20 h for HCV. To distinguish target cells from donor cells, several cell-labeling strategies were exploited. First, we established a GFP-expressing Huh7.5.1 stable cell line with more than 95% GFP-positive cells, as determined by GFP analysis by flow cytometry (data reviewed but not shown), and used cells of this cell line as the target cells in the coculture assay. Incubation of these cells with HCV-infected Huh7.5.1 cells gave rise to core protein-positive and GFP-positive target cells (Fig. 2a, panels i, v, and ix, arrowheads). We also labeled naive Huh7.5.1 cells with a thiol-reactive cell-tracking dye, CMFDA (31), and used these cells as the target cells in the coculture assay. Incubation of CMFDA-labeled cells with HCV-infected Huh7.5.1 cells showed core protein-positive and CMFDA-positive target cells (Fig. 2a, panels ii, vi, and x, arrowheads). Similar results were obtained using a membrane-labeling dye, PKH26, to label the target cells (Fig. 2a, panels iii, vii, and x, arrowheads). We also labeled HCV-infected cells with CMFDA and used them as the donor cells and naive Huh7.5.1 cells as the target cells for the coculture assay. The results showed core protein-positive and CMFDA-negative target cells (Fig. 2a, panels iv, viii, and xii, arrowheads). Quantitation analysis by flow cytometry showed a comparable level of CCCM HCV transfer, i.e., about 20% of the target cells, among the different labeling strategies (Fig. 2b). Noticeably, in both the GFP and CMFDA labeling strategies, the target cells exhibited less intense core protein staining than the donor cells.
donor cells (Fig. 2a), potentially explained by the directionality of HCV transfer. To ascertain that CCCM HCV transfer occurred in PHHs, we performed the coculture assay with CMFDA-labeled HCV-infected Huh7.5.1 cells as the donor cells and PHHs as the target cells. The results showed core protein-positive and CMFDA-negative PHH targets (Fig. 3a, arrowheads), which were equivalent to approximately 20% of the total target cell population, as determined by flow cytometry (Fig. 3b). Taken together, these results confirmed CCCM HCV transfer in both human hepatoma cells and primary hepatocytes.

**CCCM HCV transfer versus cell-free HCV infection.** The core protein-positive target cells that were detected following 20 h of coculturing (Fig. 2 and 3) could also result from infection of target cells with cell-free HCV from HCV-infected donor cells. To address this possibility, we determined the kinetics of CCCM HCV transfer and compared it to that of cell-free HCV infection. We labeled uninfected Huh7.5.1 target cells with PKH26 and performed the coculture assay at a D/T ratio of 1:1 for 0, 3, 6, 9, and 20 h. Flow cytometry analysis showed that core protein-positive and PKH26-positive target cells began to emerge within 3 h of cocultivation and gradually increased in number up to 20 h (Fig. 4a). In parallel, we set up a transwell assay (Fig. 4b), in which HCV-

![FIG 3](image-url) CCCM HCV transfer between Huh7.5.1 cells and PHHs. Human PHHs were cocultured with JFH1-infected CMFDA-labeled Huh7.5.1 cells at a 1:1 ratio for 20 h. The mixed culture was immunostained for HCV core protein and subjected to confocal imaging (a) or flow cytometry (b). The human primary hepatocytes with CCCM HCV transfer are marked with an arrowhead in panel a. *, P < 0.05.

![FIG 4](image-url) CCCM HCV transfer versus cell-free virus infection. (a) JFH1-infected Huh7.5.1 donor cells were cocultured with PKH26-labeled Huh7.5.1 target cells at a 1:1 ratio for 0, 3, 6, 9, or 20 h, followed by immunostaining for HCV core protein and flow cytometry analysis. The HCV core protein-positive and PKH26-positive cells in the upper right quadrant represent cells with CCCM HCV transfer; they were expressed as the percentage of the total number of cells and shown in the upper right quadrant in each dot plot. (b) JFH1-infected Huh7.5.1 donor cells were seeded on top of a 0.4-μm virus-permeable membrane in a transwell insert, and Huh7.5.1 target cells were seeded in the lower chamber of the transwell. The cells were cultured for 20 h with the same total number of cells used in the coculturing assay. The cells in the lower chambers were collected, immunostained for core protein, and analyzed by flow cytometry. (c) CCCM HCV transfer kinetics in coculturing and transwell assays at the same 1:2 ratio of D/T. (d) Similar transwell experiments were performed as described for panel b, except that extended culturing times were used. The HCV RNA in the upper and lower chambers was determined by qRT-PCR (left y axis, bar graph), whereas the percentage of core protein-positive cells in the lower chamber was determined by immunostaining for core protein, followed by flow cytometry (right y axis, dotted line). vge, viral genome equivalents. (e) Cell-free HCV infection kinetics. Huh7.5.1 cells were infected with HCV JFH1 (MOI = 1); cells were harvested at the indicated times, and core protein-positive cells were determined by immunostaining for core protein, followed by flow cytometry. (f) Similar coculturing experiments were performed as described for panel a with different ratios of donor cells/target cells. ***, P < 0.001; NS, not significant.
infected Huh7.5.1 donor cells were seeded on a 0.4-μm-pore-size permeable membrane on the transwell insert in the upper chamber and uninfected Huh7.5.1 cells were seeded in the lower chamber of the transwell. The 0.4-μm-pore-size permeable membrane functions as a physical barrier to separate the donor cells from the target cells and therefore allows only cell-free HCV to diffuse from the upper chamber of the transwell insert into the lower chamber of the transwell to infect cells and at the same time excludes CCCM HCV infection. As the surface area of the insert is only half of that of the bottom of the lower chamber, a D/T ratio of 1:2 was used to achieve a comparable cell confluence between the upper and lower chambers. The cells in the lower chamber were harvested at the same time points as those in the coculture assay, i.e., 0, 3, 6, and 20 h, and stained for HCV core protein expression. Compared to the results of the coculture assay, the transwell assay gave rise to very few core protein-positive target cells in the lower chamber of the transwell during the same time points (Fig. 4c). Therefore, the core protein-positive target cells detected within 20 h of coculturing (Fig. 2 and 3) likely resulted from CCCM HCV infection and not from cell-free infection. To ascertain that cell-free HCV can freely diffuse through the membrane on the insert, the transwell assay was extended to 48 and 72 h. HCV-infected cells in the lower chamber were similarly determined using core protein immunostaining. Core protein-positive cells in the lower chamber began to emerge only at 24 h and increased at 48 h and 72 h (Fig. 4d, line graph); the kinetics was very similar to that of the cell-free HCV infection of Huh7.5.1 cells by direct inoculation (Fig. 4e), which was routinely performed in the laboratory. These results not only support the notion that cell-free HCV infection takes a longer time than CCCM HCV infection but also confirm that cell-free HCV is capable of passing through the membrane and infecting target cells in the lower chamber. In addition, qRT-PCR was performed to compare the HCV RNA levels between the upper and lower chambers at 24, 48, and 72 h. No differences of HCV RNA levels were found (Fig. 4d, bar graph), further confirming that HCV can diffuse through the 0.4-μm-pore-size permeable membrane on the insert and that CCCM infection and not cell-free infection is solely responsible for the newly infected target cells in the 20-h coculture assay. Next, we determined whether CCCM HCV transfer was dependent on the D/T ratio. We performed the coculture assay with the same coincubation time (20 h) and the same total number of donor and target cells but with different D/T ratios. The results showed that the percentage of core protein-positive target cells increased with increasing D/T ratios (Fig. 4f), suggesting that a higher D/T ratio likely provided more opportunities for the target cells to be in contact with the HCV-infected donor cells. The increase in HCV CCCM transfer over the D/T ratio appeared to be modest, likely due to an already higher number (density) of starting cells in the coculturing assay. In addition, our subsequent imaging data showed that CCCM HCV transfer did not necessarily occur between one donor cell and one target cell. Furthermore, in this coculturing experiment, the same total number of cells was maintained when the D/T ratio was increased. Thus, an increase in the D/T ratio does not necessarily translate to a proportional increase in the contact between donor cells and target cells. In other words, there would be more cell-cell contact and CCCM transfer among the donor cells themselves at a higher D/T ratio. However, this type of CCCM transfer was not accounted for in our experimental setting. Taken together, these results showed that CCCM HCV infection occurred more readily than cell-free infection and was dependent on direct cell–cell contact.

**Indispensable roles of CD81, SR-B1, CLDN1, and OCLN in CCCM HCV transfer.** Huh7.5.1 cells were transfected with 100 nM siRNAs specific for CD81, SR-B1, CLDN1, and OCLN. Preexperiments were performed to determine which siRNA concentration allowed maximal knockdown. At 48 h posttransfection, cells were collected for analysis of CD81 expression by flow cytometry (a), SR-B1, CLDN1, and OCLN expression by Western blotting (WB) (b), and OCLN expression by immunofluorescence staining and imaging (c). At 48 h posttransfection, the siRNA-transfected cells were labeled with PKH26 and then cocultured with JFH1-infected Huh7.5.1 cells at a 1:1 ratio for 20 h. The mixed cells were then immunostained for HCV core protein and analyzed by flow cytometry for target cells with CCCM HCV transfer (d). (e) Huh7.5.1 cells that were transfected with all four siRNAs for CD81, SR-B1, CLDN1, and OCLN (50 nM each) were evaluated for CCCM HCV transfer. siRNA control and isotype staining controls were included in each set of experiments, as indicated. ***, P < 0.001; ***, P < 0.001.
Expression of HCV receptors in different cell lines and their permissiveness to cell-free HCV infection

| Cell   | CD81 | SR-B1 | CLDN1 | OCLN | Cell-free HCV infection
|--------|------|-------|-------|------|-------------------------
| Huh7.5.1 | ++   | ++++  | ++++  | ++++  | ++          |
| 293T   | ++++ | +     | ++++  | +    | --          |
| HepG2  | --   | ++++  | ++++  | +    | --          |
| CYNK10 | ++++ | ++    | ++    | +    | --          |
| NKNT3  | ++++ | ++    | ++    | +    | --          |

* a, no expression or infection; +, low expression or infection; ++, medium expression or infection; ++++, high expression or infection.

Roles of intact actin but not the microtubule cytoskeleton in CCCM HCV transfer. To determine the roles of cytoskeleton in CCCM HCV transfer, we performed the coculturing experiments in the presence of the actin polymerization inhibitor cytochalasin D (44, 45). The effects of cytochalasin D on actin polymerization were confirmed by immunofluorescence staining of F actin. As expected, in the absence of cytochalasin D treatment, all of the F-actin filaments were along the boundaries of adjacent cells, where they made contact with each other (Fig. 7a). When the cells were treated with 0.625 μM cytochalasin D, F-actin filaments were polymerized on the tips of filopodia instead of being expressed at cell-cell contacts. When the cytochalasin D concentration was further increased to 2.5 μM and 10 μM, F-actin filaments formed aggregates in the cytoplasm and filopodia were not present. In parallel experiments, a significant decrease in CCCM HCV transfer occurred in the coculture treated with 0.625 μM cytochalasin D and a complete abrogation of CCCM HCV transfer was apparent in the cocultures treated with 2.5 μM and 10 μM cytochalasin D.

To further analyze the roles of these receptors in CCCM HCV transfer, we took advantage of a panel of human cell lines that constitutively express varied levels of HCV receptors and determined the relationships between HCV receptor expression and cell-free HCV infection or CCCM HCV transfer. The human hepatoma cell lines HepG2, NKNT3, and CYNK10 were used. HepG2 cells lack CD81 expression but express medium to high levels of the other three major HCV receptors (38–41). NKNT3 and CYNK10 cells both express very low levels of CLDN1 but have medium to high levels of the other HCV receptors (42, 43). We also included the HCV-susceptible Huh7.5.1 cell line and the nonhepatoma fibroblast cell line 293T as controls. Huh7.5.1 cells expressed all four HCV receptors at a moderate to high level, while 293T cells expressed very high levels of CD81 and OCLN but a low level of SR-B1 and no CLDN1. Expression of all four receptors in these cells was confirmed (Table 1 and data reviewed but not shown). Of all cell lines, only Huh7.5.1 cells were susceptible to cell-free HCV infection (Table 1 and data reviewed but not shown). When each cell line was cocultured with HCV-infected Huh7.5.1 cells, no CCCM HCV transfer was detected in cells other than Huh7.5.1 cells at 24 h after coculturing (Table 1 and data reviewed but not shown). These results confirmed the important roles of all four HCV receptors in both cell-free and CCCM HCV infection. We then introduced into the cells the receptor(s) that was not expressed and/or expressed at a lower level by ectopic expression and assessed the possibility of CCCM HCV susceptibility in those cells. Ectopic expression of CD81 in HepG2 cells (Fig. 6a) gave rise to little CCCM HCV transfer from HCV-infected Huh7.5.1 cells (Fig. 6b). Similarly, ectopic expression of CLDN1 in NKNT3 cells (Fig. 6c) gave rise to little CCCM HCV transfer from HCV-infected Huh7.5.1 cells (Fig. 6d); ectopic expression of CLDN1 in CYNK10 cells gave rise to little CCCM HCV transfer from HCV-infected Huh7.5.1 cells (data reviewed but not shown); ectopic expression of both SR-B1 and CLDN1 in 293T cells (Fig. 6e) gave rise to little CCCM HCV transfer from HCV-infected Huh7.5.1 cells (Fig. 6f). Taken together, these results suggest that expression of all four major HCV receptors, CD81, SR-B1, CLDN1, and OCLN, is essential but not sufficient for CCCM HCV infection.

FIG 6 No CCCM HCV transfer from Huh7.5.1 cells to cells of other hepatoma and nonhepatoma cell lines. HepG2 cells were transfected with pcDNA3-CD81, NKNT3 cells were transfected with pCMV-CLDN1, and 293T cells were transfected simultaneously with both pcDNA3-SR-B1 and pCMV-CLDN. At 48 h posttransfection, the transfected cells were collected for analysis of CD81 expression in HepG2 cells by flow cytometry (a), CLDN1 expression in NKNT3 cells by Western blotting (c), or SR-B1 and CLDN1 expression in 293T cells by Western blotting (e). Simultaneously, these receptor-expressing HepG2, NKNT3, or 293T cells were labeled with PKH26 or CMFDA and cocultured with JFH1-infected Huh7.5.1 cells at a 1:1 ratio for 20 h. The mixed cell aggregates in the cytoplasm and filopodia were not present. In parallel experiments, a significant decrease in CCCM HCV transfer occurred in the coculture treated with 0.625 μM cytochalasin D and a complete abrogation of CCCM HCV transfer was apparent in the cocultures treated with 2.5 μM and 10 μM cytochalasin D.

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D (Fig. 7b). We also performed similar experiments in the presence of the microtubule-depolymerizing agent nocodazole (46, 47) and determined its effects on CCCM HCV transfer. As expected, nocodazole treatment disrupted the microtubule cytoskeleton in a dose-dependent manner, as determined by antitubulin immunofluorescence staining, beginning with a gradual loss of the microtubule network, followed by more diffuse tubulin staining patterns in the cytoplasm (Fig. 7c). However, nocodazole-induced disruption of microtubules did not lead to significant changes in the level of CCCM HCV transfer (Fig. 7d). The inhibition of CCCM HCV transfer by cytochalasin D treatment (Fig. 7b) was not due to the cytotoxicity of cytochalasin D (48; unpublished data) or any adverse effects of cytochalasin D on cell-free HCV infection (data reviewed but not shown). Therefore, these results suggest that an intact actin network but not the microtubule cytoskeleton may be required for CCCM HCV transfer.

Live-cell imaging of the CCCM HCV transfer process in Huh7.5.1 cells from conjugate formation to transfer. To directly dissect the spatial and temporal details of CCCM HCV transfer, we attempted to construct a fluorescein-labeled HCV clone to characterize viral transfer in live cells. We first attempted to generate GFP-labeled HCV by introducing GFP in frame with the HCV core protein gene in JFH1 but were unsuccessful (data reviewed but not shown). This was likely due to the large size of the foreign GFP protein and subsequent alterations in HCV assembly. We therefore turned to the TC tag-biarsenical dye labeling system to track HCV core protein. Huh7.5.1 cells were transfected with in vitro-transcribed full-length JFH1 RNA or JFH1-TCcore RNA and monitored HCV replication and production using qRT-PCR. Compared to JFH1, JFH1-TCcore showed a slightly delayed viral replication (Fig. 8b) and virus production (Fig. 8c) and a 2- to 4-fold lower maximal level of virus production (Fig. 8c). Nevertheless, labeling of JFH1-TCcore-transfected and -infected cells with the biarsenical compound did not alter the HCV-TCcore infectivity (data reviewed but not shown) (50). We then determined the labeling specificity of the biarsenical compound (ReAsH) and the subcellular localization of the TC-tagged core protein. Huh7.5.1 cells were transfected with in vitro-transcribed full-length JFH1 or JFH1-TCcore RNA and stained with ReAsH and then HCV core protein using an anti-HCV core protein antibody. ReAsH labeled only JFH1-TCcore-transfected cells and not JFH1-transfected cells (Fig. 8d). There appeared to be a complete overlap of ReAsH labeling with HCV core protein staining, confirming that ReAsH specifically bound to the TC tag and that the tag did not alter the subcellular localization of HCV core protein. Taken together, these results demonstrate the feasibility of using the JFH1-TCcore and ReAsH labeling system to track HCV core protein in live cells.

We performed live-cell imaging to track CCCM HCV transfer with the JFH1-TCcore system. We infected Huh7.5.1 cells with JFH1-TCcore for 3 days and then labeled the cells with ReAsH. We used ReAsH-labeled cells as donor cells and Huh7.5.1 cells stably expressing GFP (Huh7.5.1-GFP cells) as target cells and performed the coculturing assay under live confocal imaging. Red (ReAsH)/green (GFP) fluorescence and DIC images in the se-
lected fields were taken every 18 min throughout the 18-h coculturing period. A single HCV-infected ReAsH-labeled Huh7.5.1 donor cell (Fig. 9a, labeled D) had three recorded sequential transfer events (yellow arrowheads) to three contacted target Huh7.5.1-GFP cells (Fig. 9a, labeled T1, T2, and T3) during this time period (see Movie S1 in the supplemental material). For each transfer event, the donor cell either accumulated large numbers of viral puncta on the contact surface between the

FIG 8 Characterization of recombinant JFH1-TCcore HCV. (a) JFH1-TCcore recombinant HCV clone. A TC tag-containing 12-amino-acid peptide was inserted after the third amino acid of the HCV JFH1 core protein. (b and c) Huh7.5.1 cells were transfected with an equal amount of *in vitro*-transcribed full-length HCV JFH1 RNA or HCV JFH1-TCcore RNA. Transfection medium was replaced with fresh culture medium, and the cells were extensively washed 6 h after transfection. RNA was isolated from the cells and the culture supernatants at the indicated time points and was subjected to qRT-PCR to monitor HCV replication in the cells (b) and HCV production in the culture supernatants (c), respectively. (d) ReAsH labeling specificity of TCcore in JFH1-TCcore RNA-transfected Huh7.5.1 cells. Huh7.5.1 cells were transfected with JFH1 or JFH1-TCcore RNA. At 3 days posttransfection, the cells were first labeled with ReAsH and then immunostained for detection of the HCV core protein and stained with DAPI for detection of the nucleus.

FIG 9 Live-cell imaging of CCCM HCV transfer. JFH1-TCcore-infected ReAsH-labeled Huh7.5.1 donor cells (red) and Huh7.5.1-GFP target cells (green) were cocultured and imaged live for 18 h. (a) CCCM HCV transfer from one ReAsH-positive TCcore-positive donor cell (labeled D) to three GFP-positive target cells (labeled T1, T2, and T3 and outlined) sequentially over 18 h of coculturing. The ReAsH, GFP, and DIC overlay is shown in the upper panels, from which the boxed area was magnified and examined in the middle (ReAsH only) and the lower (ReAsH and GFP overlay) panels. HCV transfer from the donor cell (D) to target cells T1 (i and ii) and T2 (iii), and T3 (vi) is indicated by yellow arrowheads. The yellow arrow (v) indicates T3’s filopodia toward D after viral transfer. Times are shown in hours:minutes. (b) Concurrent HCV transfer from one donor cell (red) to two target cells (green). Orthogonal views are shown, and the positions of the perpendicular planes are indicated as notches. (c) 3D reconstruction of a transfer event between a donor cell and one target cell observed in panel b. This transfer event occurred at a time later than that shown in panel b. (d) Transfer of a viral punctum (circled in the lower panels) into a target cell: (i) donor cell-target cell in contact before conjugate formation; (ii) formation of viral puncta-target cell conjugation; (iii) transfer of viral puncta; and (iv) postransfer. The lower panels are 3D reconstructions of the boxed area in the corresponding upper panels. (e) Tracking of transfer events in 13 donor cells in one imaging field; each line represents one donor cell-target cell pair.
donor cell and the target cell (Fig. 9a, D and T1 above panels i and ii) or produced filopodia containing viral puncta at the contact site (Fig. 9a, D and T2 above panel III and D and T3 above panel iv) prior to the transfer. Simultaneous transfer of HCV from one single donor cell to multiple target cells was also recorded (Fig. 9b; see Movies S2 and S3 in the supplemental material). The orthogonal view (Fig. 9b) and 3D reconstruction of the transfer process (Fig. 9c; see Movies S3 and S4 in the supplemental material) confirmed that the transferred viral puncta were located inside the target cell, as the viral puncta were visualized as yellow rather than red. The sizes of the pretransfer viral puncta were estimated to be approximately 0.5 μm to 2 μm, suggesting that the puncta contained more than a single virus. The number of core protein puncta that were transferred from one donor cell to one target cell could reach up to 100 during the 18-h coculturing period. Tracking the transfer of a single viral puncta revealed four distinct steps in the CCCM HCV transfer process: donor cell–target cell contact, viral puncta–target cell conjugate formation, transfer of viral puncta, and posttransfer (Fig. 9d). The transfer began with contact initiated between the donor and target cells (Fig. 9d, panel i). Viral puncta were formed within the donor cell and transitioned to the contact site between the donor and the target cells (Fig. 9d, panel i). This was followed by formation of viral puncta–target cell conjugates (Fig. 9d, panel ii). The transfer took place when the viral puncta were localized in the target cell (Fig. 9d, panel iii). The viral puncta likely dissipated into a few smaller puncta in the target cell (Fig. 9d, panel iv) and presumably disappeared as viral uncoating took place. A single transfer event was estimated to take approximately 18 min to complete. To further understand the transfer process, we analyzed the transfer events in the entire tracked field (450 μm by 450 μm). There were a total of 33 ReAssH-positive donor cells and 160 Huh7.5.1-GFP target cells in the field at the beginning of tracking (time zero). During the 18 h of tracking, 13 cells (13/33 = 39%) formed conjugates with target cells, 7 donor cells showed transfer (7/33 = 21%), and a total of 8 transfer events occurred (Fig. 9e). Each of these processes was confirmed with 3D reconstruction, as shown in Fig. 9c and d. The average duration of viral puncta–target cell conjugation between a donor cell and a target cell was calculated to be 408 min, and the average time for the actual transfer was calculated to be 171 min. The latter was much longer than the 18 min for a single transfer event, as estimated above; this is because more than one viral punctum was transferred between one donor cell and one target cell. Taken together, these results provide the temporal and spatial details of CCCM HCV transfer and demonstrate the high efficiency of this transmission route.

**CCCM HCV transfer and productive HCV infection.** To determine whether CCCM HCV transfer leads to productive HCV infection, we took advantage of a neutralizing HCV antibody (CBH-5) that has been shown to be very effective in blocking free HCV infection by over 99% at a concentration of 5 μg/ml (data reviewed but not shown). We then performed the above-mentioned coculture HCV transfer assay in the presence of 5 μg/ml CBH-5 or a control isotope-matched antibody (R04) (51, 52) and determined the percentage of HCV core protein-positive target cells at 24, 48, and 72 h of coculturing. The percentage of HCV core protein-positive target cells increased over time (Fig. 10a and b). Compared to the results obtained with the R04 control, CBH-5 treatment showed a slightly lower but still significant percentage of HCV core protein-positive cells and significant increases in the percentage of HCV core protein-positive cells over time. In addition, the mean fluorescence intensity of the core protein-positive cells exhibited increases over time in both R04- and CBH-5-treated cocultures (Fig. 10c). The increases in the percentage and the mean fluorescence intensity of HCV core protein-positive target cells over an extended period of time in the presence of the neutralizing HCV antibody CBH-5 suggest that CCCM HCV transfer leads to productive HCV infection and replication. Culture supernatants were collected from cocultures that were treated with each of the antibodies and tested for their infectivity. Very few HCV core protein-positive cells were detected with CBH-5–treated culture supernatants at each time point (Fig. 10d), confirming the HCV-neutralizing activity of the CBH-5 antibody.

To further ascertain the productive nature of CCCM HCV transfer, we set up CCCM HCV transfer using stable GFP-expressing Huh7.5.1 as the target cells in the coculture experiments. Following the initial 20-h coculturing in the presence of 5 μg/ml CBH-5 antibody, we used fluorescence-activated cell sorting (FACS) to sort out the GFP-positive target cells and then monitored HCV replication in these sorted cells. FACS yielded more than 97% GFP-positive cells, of which about 2% were HCV core protein-positive cells (Fig. 10e and f). Similarly, the percentage of HCV core protein-positive and GFP-positive cells showed increases over an extended period of time up to 7 days. In parallel, there was a significant increase of HCV RNA in these cells between day 4 and day 7 (Fig. 10g) and the infectivity of the culture supernatants between day 4 and day 7 (Fig. 10h). Similar results were also obtained by selective elimination of the HCV-infected Huh7.5.1 donor cells from the coculturing and monitoring of HCV infection and replication in the remaining target cells (data reviewed but not shown). Taken together, these results support the notion that CCCM HCV transfer leads to productive infection and replication.

**DISCUSSION**

Despite significant advances since the establishment of the JFH1-based HCV cell culture system in 2005 (53–55), the process of HCV transmission and spreading in the liver following infection remains poorly understood. The liver is one of the organs with an extremely high cell density (2 × 10^9 to 3.0 × 10^10 hepatocytes/cm^2 [56]), which provides HCV with numerous cell-cell contact sites. In chronically HCV-infected liver, viral replication and the intrahepatic HCV RNA level are very low (7 to 64 genomic equivalents per cell) (57, 58), and nAbs and other immunological responses are often present (59). However, even with the presence of those nAbs that are capable of neutralizing cell-free infectivity in vitro (15), the prevalence of infected hepatocytes in the livers of chronically infected patients is normally high (57, 60). Our data show that cell-cell contact facilitates HCV spreading among hepatocytes (Fig. 1 and 2), including PHHs (Fig. 3). We further show that CCCM HCV infection is a rapid process compared to cell-free HCV infection (Fig. 4). Previous studies have shown that discrete, localized infectious foci and a gradient dispersion of the viral genome around the center of the foci are observed in patient liver biopsy specimens (58, 60, 61) and that CCCM HCV infection is relatively less sensitive to nAbs and neutralizing patient sera than cell-free HCV infection (62). Taken together, it is highly conceivable that CCCM HCV occurs in vivo, and it could even be more
favorable than cell-free infection, given the compact nature of the liver. Thus, CCCM HCV transmission should be considered in designing any future anti-HCV vaccines and therapeutics.

CD81, the first discovered HCV receptor, was identified by screening for HCV E2 binding (33) and is believed to mediate HCV cell-free entry in both an early E2-binding step and a late postbinding step in association with CLDN1 (8,12, 13). SR-B1, another E2-binding HCV receptor, also mediates cell-free HCV attachment and entry in an early step in this multistep entry process (63–66). The other two, CLDN1 and OCLN, are both tight-junction proteins recently identified to be HCV coreceptors by screening for cellular determinants able to confer HCV pseu-

dotyped particle (HCVpp) entry into nonsusceptible cells (32, 34). To determine the role of each of these four HCV receptors in CCCM HCV infection, we first used an siRNA strategy to knock down the expression of each of them individually in Huh7.5.1 target cells and found between 46% and 72% decreases in CCCM HCV infection, suggesting a fundamental role of each of them in this process (Fig. 5). Furthermore, simultaneous knockdown of all of them almost completely abolished CCCM HCV infection, suggesting that they mediate CCCM HCV infection in a combinato-

rial mechanism. To further understand the roles of the four receptors in CCCM infection, we performed the coculture assay using CD81-negative HepG2 cells, NKN73 and CYNK10 cells expressing low levels of CLDN1, and 293T cells expressing low levels of SR-B1 and no CLDN1 (Table 1 and data reviewed but not shown). All these cell lines are refractory to both cell-free and CCCM HCV infection (Table 1 and data reviewed but not shown). However, ectopic expression of the receptor(s) that was missing or expressed at a low level did not render these cells susceptible to CCCM HCV infection (Fig. 6). These results together suggest that all four HCV receptors are important for CCCM HCV infection. Consistent with our findings, previous studies have shown that SR-B1 antag-

onists and an anti-CLDN1 serum inhibit CCCM HCV infection (67). However, the roles of CD81 in CCCM HCV infection appear to be controversial in the literature. Using soluble CD81 and anti-

CD81 antibody, one group has shown that CCCM HCV infection can be either CD81 dependent or CD81 independent (62). The same group later used a different HCV strain and concluded that CD81 is not necessary for CCCM HCV infection (68). It is clear that the roles of CD81 in CCCM HCV infection merit further investigation.

CCCM infection has been shown in many different animal models, and its role in immune evasion and rapid viral dissemination has recently gained more attention (18). CCCM viral trans-

mission in animal cells could occur in a variety of ways. Among the known ways are through virological synapses (HIV-1 and HTLV-

1), nanotubules (HIV-1), viral transfer across tight junctions

FIG 10 CCCM HCV transfer leads to productive HCV infection. (a to d) Infected Huh7.5.1 cells were cocultured with GFP-expressing Huh7.5.1 cells at a 1:4 ratio in the presence of 5 μg/ml neutralizing antibody CBH-5 or a control antibody, R04. The coculture was replated every 24 h, and fresh antibodies were added. Meanwhile, aliquots of cells were collected for core protein staining, followed by flow cytometry analysis for GFP-positive/core protein-positive cells (a to c); the supernatants were also collected and used to infect uninfected Huh7.5.1 cells, and their infectivity was determined by immunostaining and flow cytometry analysis for HCV core protein-positive cells at 3 days postinfection (d). (a) Representative kinetics of GFP-positive/core protein-positive cells; (b) data from multiple repeats; (c) difference in mean fluorescence intensity (dMFI) of core protein staining of GFP-positive cells from multiple repeats. (e to h) Infected Huh7.5.1 cells were cocultured with GFP-expressing Huh7.5.1 cells at a 1:4 ratio in the presence of 5 μg/ml neutralizing antibody CBH-5. GFP-expressing target cells were sorted out from the cocultures at day 1 (D1) and further cultivated for an extended period of time. Core protein-positive cells in the purified GFP-positive cell population were determined at day 4 (D4) and day 7 (D7) by immunostaining for the core protein. (e) Representative kinetics of GFP-positive/core protein-positive cells; (f) data from multiple repeats. D1#, cocultured cells prior to sorting. In addition, cells and their culture supernatants were collected at day 4 and day 7 and assayed for HCV RNA in the cell by qRT-PCR (g) or for the infectivity of the supernatant (h), respectively, as described for panel d. **, P < 0.01; ***, P < 0.001; NS, not significant.
of HCV-loaded vesicles with the plasma membrane (the actual release process) has never been captured with fluorescence or electron microscopy. Our live-cell imaging studies showed that during CCCM HCV transmission, large amounts of mobile viral puncta moved to the contacted surfaces between the donor and target cells and formed conjugates with the target cell, followed by transfer of the entire viral puncta (Fig. 9). These core protein puncta are micrometer sized and are unlikely to be single viral particles but are more likely to be vesicles loaded with virus particles (100 to 1,000 virions/punctum, on the basis of size) (28, 50). The transfer of a single core protein punctum took 18 min (Fig. 9d), and the transfer process lasted 171 min, on average (Fig. 9e). Moreover, the transfer events could occur at multiple donor cell-target cell contact sites (Fig. 9a, panel ii) or concurrently between one donor and multiple targets (Fig. 9b). Therefore, during the entire CCCM transfer process, a great number of viral puncta, which correspond to an even greater number of viral particles, can be delivered into the target cell, in contrast to the cell-free infection, in which one virion enters one target cell at a time. It is not guaranteed that each of the viral particles in each of the transferred puncta proceeds through the downstream viral life cycle in the target cell, yet the high input of CCCM viral transfer certainly leads to productive infection of the target cell (Fig. 10).

In summary, we showed that HCV infection occurs through CCCM transmission, resulting in efficient and productive infection and replication. In addition, we demonstrated the requirements for HCV receptors in CCCM infection and the involvement of the actin cytoskeleton in this process. Furthermore, the three-dimensional live-cell microscopic imaging showed the kinetics and process of transfer of intact core protein puncta from donor cells to target cells. These findings will add to our understanding of HCV infection and transmission and provide new avenues for the development of novel anti-HCV therapeutics.

ACKNOWLEDGMENTS

We thank Charles Rice of Rockefeller University, New York, NY, for HuH7.5.1 cells, Ira Fox of the University of Pittsburgh, Pittsburgh, PA, for NKNT3 and CYNK10 cells, Wenze He of Temple University, Philadelphia, PA, for the JFH1 virus stock, Takaji Wakita of the National Institute of Infectious Diseases, Japan, for JFH1 recombinant plasmid DNA, and Steve Foung of Stanford University, Stanford, CA, for antibodies CBH-5 and R04. We also thank Ignacy Gryczynski, Rafal Fudala, and I-Fen Chang for their assistance on confocal imaging, Larry Oakford for his assistance on confocal data processing, and Andy (Qigui) Yu, Randy Brutkiewicz, Cheng Kao, and William Sullivan for their advice and input throughout the study.

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