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A DISULFIDE-BONDED DIMER OF THE CORE PROTEIN OF HEPATITIS C VIRUS IS IMPORTANT FOR VIRUS-LIKE PARTICLE PRODUCTION

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Hepatitis C virus (HCV) core protein forms the nucleocapsid of the HCV particle. Although many functions of core protein have been reported, how the HCV particle is assembled is not well understood. Here we show that the nucleocapsid-like particle of HCV is composed of a disulfide-bonded core complex (dbc-complex). We also found that the disulfide-bonded dimer of the core (dbd-core) is formed at the endoplasmic reticulum (ER) where the core protein is initially produced and processed. Mutational analysis revealed that the cysteine residue at amino-acid position 128 (Cys128) of the core, a highly conserved residues among almost all reported isolates, is responsible for dbd-core formation and virus-like particle production with no effect on the replication of HCV RNA genome and the several known functions of the core, including RNA binding ability and localization to the lipid droplet. The Cys128 mutant core showed a dominant-negative effect in terms of HCV-like particle production. These results suggest that this disulfide bond is critical for the HCV virion. We also obtained the results that the dbc-complex in the nucleocapsid-like structure was sensitive against proteinase K but not trypsin digestion, suggesting that the capsid is built up of a tightly packed structure of the core with its amino (N)-terminal arginine-rich region concealed inside.
Hepatitis C virus (HCV) infection is a major cause of chronic hepatitis, liver cirrhosis, and hepatocellular carcinoma, affecting approximately 200 million people worldwide (13, 29, 44). Current treatment strategies, including interferon coupled with ribavirin, are not effective for all patients infected with HCV. An error-prone replication strategy allows HCV to undergo rapid mutational evolution in response to immune pressure, and thus evade adaptive immune responses (10). New approaches to HCV therapy include the development of specifically targeted antiviral therapies for hepatitis C (STAT-Cs), which target such HCV proteins as NS3/4A, serine protease, and the RNA-dependent RNA polymerase NS5B (3). Despite potent antiviral activity for some of these approaches, many resistant HCV strains have been reported after treatment with existing STAT-Cs (23, 48, 51). Therefore, identification of new targets that are common to all HCV strains and are associated with low mutation rates is an area of active research.

HCV has a 9.6-kb, plus-strand RNA genome composed of a 5’-untranslated region (UTR), an open reading frame that encodes a single polyprotein of about 3000 amino acids, and a 3’-UTR. The polyprotein is processed by host and viral proteases to produce three structural proteins (core, E1, and E2) and seven nonstructural proteins (p7,
NS2, NS3, NS4A, NS4B, NS5A, and NS5B (14, 16, 17, 22, 49). HCV core protein is produced co-translationally via carboxyl (C)-terminal cleavage to generate an immature core protein, 191 amino acids in length, on the endoplasmic reticulum (ER) (16). This protein consists of three predicted domains: the N-terminal hydrophilic domain (D1), the C-terminal hydrophobic domain (D2), and the tail domain (33), which serves as a signal peptide for the E1 envelope protein. The D1 includes a number of positively charged amino acids responsible for viral RNA binding (amino acids 1-75) (43) and the region involved in multimerization of core via homotypic interactions (amino acids 36-91 and 82-102) (32, 40) (Supplementary Fig. 1). The hydrophobic D2 includes the region responsible for core association with lipid droplets (LDs) (amino acids 125-144) (7, 18, 37), which accumulate in response to core production (1, 6).

Many functions of core protein have been reported (13, 38, 50). Yet because infectious HCV particles cannot be appropriately produced in currently available experimental systems, HCV particle assembly has not been elucidated to date. A cell culture system that reproduces the complete lifecycle of HCV in vitro was developed by Wakita et al. using a cloned HCV genome (JFH1) (53). Using this system, the assembly of infectious HCV particles was found to occur near LDs and ER-derived LD-associated membranes (36, 47). Neither the structures nor functions of the virus proteins involved
in virus particle assembly are known, however. To elucidate this point, we have
analyzed the biochemical characteristics of the proteins within the fraction containing
the HCV particle, and found a disulfide-bonded core protein complex. We revealed that
the disulfide-bonded dimer of core (dbd-core) was formed by a single cysteine residue
at amino-acid position 128 on the ER. The roles of the disulfide bond of the core in the
virus-like particle formation are discussed in this paper.
MATERIALS AND METHODS

Cell culture: The HuH-7 and HuH-7.5 human hepatoma cell lines were grown in Dulbecco’s modified Eagle’s medium (Nacalai Tesque, Kyoto, Japan) supplemented with 10% fetal bovine serum, 100 U/ml nonessential amino acids (Invitrogen, Carlsbad, CA), and 100 μg/ml each penicillin and streptomycin sulfate (Invitrogen).

Antibodies: The antibodies used for immunoblotting and indirect immunofluorescence analysis were specific for core (#32-1), FLAG M2 (Sigma-Aldrich, St Louis, MO), c-myc (Sigma-Aldrich), NS5A (CL1), ADRP (StressGen, Victoria, Canada), Calnexin-NT (StressGen), and GAPDH (Chemicon, Temecula, CA). Antibodies specific for core (#32-1) were a gift from Dr Kohara (The Tokyo Metropolitan Institute of Medical Science, Japan). Rabbit polyclonal anti-NS5A CL1 antibodies have been described previously (36).

Plasmid construction: All plasmids were generated by inserting PCR-amplified fragments into expression plasmids. The plasmids, primer sequences, templates for the PCRs, and restriction enzyme sites used to construct the plasmids are listed in Supplementary Table. Plasmids pJFH1^{E2FL} (full-length HCV genome with FLAG
epitope in E2 HVR), pJFH1^{AAA99} (encoding a NS5A mutant of JFH1^{E2FL}, resulting in non-infectious HCV particles), pJFH1^{PP/AA} (encoding a core mutant of JFH1^{E2FL}, which allows replication in cells but prevents HCV particle production), and pcDNA3-core^{WT} (expression plasmid encoding full-length JFH1 core) have been previously described (36). Plasmid pJ6/JFH1, which contains the full-length HCV genome encoding structural proteins from the J6 strain and nonstructural proteins from the JFH1 strain, was kindly provided by Charles M. Rice (The Rockefeller University, New York, USA).

**In vitro transcription:** RNA for transfection was synthesized as described previously (36). In brief, plasmids carrying the HCV RNA sequence were linearized with XbaI and used as templates for *in vitro* transcription with MEGAscript T7 (Ambion, Austin, TX).

**Transfection:** Ten micrograms of JFH1^{E2FL}, JFH1^{C128A}, JFH1^{C184A}, JFH1^{C128/184A}, or JFH^{AAA99} and J6/JFH1 or J6/JFH1^{AAA99} RNA were transfected into HuH-7 and HuH-7.5 cells (1.0 × 10^7 cells) by electroporation (260 V, 0.95 µF) using a GENE PULSER II system (BioRad, Hercules, CA). Core expression plasmids were transfected into HuH-7 cells using Lipofectamine LTX (Invitrogen) according to the manufacturer’s protocol.
**HCV particle precipitation:** Culture medium from HCV RNA–transfected cells were concentrated using Amicon Ultra-15 centrifugal filters with Ultracell-100 membranes (Millipore, Billerica, MA) and mixed with sucrose solution in PBS to a final sucrose concentration of 2%. This mixture was ultracentrifuged (100,000 × g; 4°C for 2 h) and the HCV particles were obtained as a pellet. The pellet was then suspended in culture medium for infection experiments or PBS for immunoblot analysis.

**Indirect immunofluorescence analysis:** Indirect immunofluorescence analyses of HCV infection and the cellular localization of HCV proteins were performed as described previously (36).

**Protease protection assay:** Concentrated culture medium from JFH1E2FL RNA–transfected HuH-7 cells was fractionated using 20–50% sucrose density gradients and the HCV RNA titer was measured in quantitative RT-PCRs as described below. Fractions with high HCV RNA titers were collected and JFH1E2FL particles were obtained as a pellet after ultracentrifugation (100,000 × g; 4°C for 2 h). The pellet was suspended in PBS and treated with 10 µg/ml trypsin or 5 µg/ml proteinase K in the presence or absence of 1% NP-40 at 37°C for 15 min, respectively, unless otherwise
The reaction was quenched by the addition of protease inhibitor cocktail (Nacalai Tesque) followed by SDS-PAGE under non-reducing conditions and immunoblotting specific for core protein.

**Immunoblot analysis:** Samples were subjected to SDS-PAGE in sample buffer (62.5 mM Tris-HCl [pH 7.8], 1% SDS, and 10% glycerol) with or without 5% β-ME or 50 mM DTT for reducing or non-reducing conditions, respectively. N-ethylmaleimide (NEM) (Nacalai Tesque) was added to the sample buffer to final concentration of 5 mM in indicated samples. Proteins were transferred to polyvinylidene difluoride membrane, and blocked in blocking buffer for 1 h at room temperature with gentle agitation. After incubation with primary antibodies overnight at 4°C, the membrane was washed three times for 5 min in washing buffer at RT with gentle agitation. Then, the membrane was incubated with HRP-conjugated secondary antibodies for 1 h at RT. After three washes in washing buffer, proteins were detected using Western Lightning (PerkinElmer, Waltham, MA) or ECL Advance (GE Healthcare, Buckinghamshire, England) and Kodak MXJB plus medical X-ray film (Kodak, Rochester, NY) or an LAS-4000 system (Fujifilm, Tokyo, Japan)
Preparation of LDs: LDs were prepared as described previously (36).

Preparation of MMFs: MMFs were collected as previously described (15) with some modifications. In brief, cells were collected in homogenization buffer (20 mM Tris-HCl [pH 7.8], 250 mM sucrose, and 0.1% ethanol supplemented with protease inhibitor cocktail) and homogenized on ice using 40 strokes of a dounce homogenizer. The samples were then centrifuged at 1000 × g for 10 min at 4°C. Supernatant was collected in a new tube and centrifuged again at 16,000 × g for 20 min at 4°C. Supernatant was further centrifuged at 100,000 × g for 60 min at 4°C. The MMF precipitate was homogenized in lysis buffer (1% NP-40, 0.1% SDS, 20 mM Tris-HCl [pH 8.0], 150 mM NaCl, 1 mM EDTA, and 10% glycerol supplemented with protease inhibitor cocktail) using a dounce homogenizer.

Quantitative reverse transcription (qRT)-PCR analysis: qRT-PCR analysis for the HCV RNA titer was performed as described previously (36).

Enzyme-linked immunosorbent assay (ELISA) specific for core: Core in culture medium was quantified using an ELISA according to the manufacturer’s protocol (HCV
antigen ELISA test; Ortho-Clinical Diagnostics, Raritan, NJ).

RNA-protein binding precipitation assay: Core\textsuperscript{WT} or core\textsuperscript{C128A} were translated \textit{in vitro} from pcDNA3-core\textsuperscript{WT} or pcDNA3-core\textsuperscript{C128A}, respectively, using the TNT Coupled Rabbit Reticulocyte Lysate system (Promega, Madison, WI) according to the manufacturer’s protocol. These proteins were incubated with poly-U agarose (Sigma) in 50 mM HEPES (pH 7.4), 100 mM NaCl, 0.1% NP-40, and 20 U RNase inhibitor at 4°C for 2 h with or without RNase A. After five washes, resin-bound core proteins were immunoblotted.
RESULTS

The HCV particle contains core complex formed by a disulfide bond

To analyze the core protein of the HCV particle, we first subjected the concentrated culture medium of HuH-7 cells transfected with \textit{in vitro} transcribed JFH1$^{\text{E2FL}}$ RNA to ultracentrifugation. After the resulting pellet was resuspended in culture medium, we confirmed the presence of infectious HCV particles based on infectivity against HuH-7.5 cells (Fig. 1a). The infectious JFH1$^{\text{E2FL}}$ particle–containing pellet was separated by SDS-PAGE under non-reducing conditions, and immunoblot analysis showed the presence of a core antibody–reactive protein that was approximately twice the size of core (38 kDa), in addition to the expected 19-kDa core protein (Fig. 1b, lane 1). Because treatment with dithiothreitol (DTT) eliminated the larger core antibody–reactive band while levels of core monomer increased (Fig. 1b, lanes 2-6), the larger protein likely represented a core-containing complex formed by disulfide bonds. This complex was also found in J6/JFH1-derived particles (Supplementary Fig. 2), indicating that the complex was not specific for JFH1$^{\text{E2FL}}$.

To determine whether the core complex is a component of the HCV particle, a protease protection assay was performed using RNase-resistant HCV particles fractionated based on buoyant density. Concentrated culture medium from HuH-7 cells
transfected with *in vitro* transcribed JFH1<sup>E2FL</sup> RNA were fractionated using a 20-50% sucrose density gradient and JFH1<sup>E2FL</sup> particles which were presumed to contain both infectious and non-infectious particles, were collected from fractions with high HCV RNA titers using ultracentrifugation (**Fig. 2a**, fraction #8 to #13). The core protein from the collected fractions was analyzed by immunoblotting after SDS-PAGE under the non-reducing condition, showing only the core complex (**Fig. 1c**, right panel).

To examine whether the complex contributes to the infectivity of the particles, we analyzed the core complex in the fractions containing infectious and non-infectious HCV particles (**Fig. 2a**, filled and open arrowheads, respectively). Both infectious and non-infectious HCV particle containing fraction consists the core complex (**Fig. 2b**). To confirm this further, a pellet containing mutant JFH1<sup>AAA99</sup> particles—a mutant of JFH1<sup>E2FL</sup> that produces primarily non-infectious particles (36)—was analyzed in a similar manner. These core complexes were found in both pelleted particles of JFH1<sup>AAA99</sup> and J6/JFH1<sup>AAA99</sup> which was a mutant J6/JFH1 with similar substitution to JFH1<sup>AAA99</sup> (**Supplementary Fig. 2**). These results indicated that the core complex was present in both the infectious and non-infectious HCV-like particles.

The core monomer observed in the pellet samples (**Fig. 1b**) may be from the
secreted core or the debris of apoptotic cells, because the core is known to be secreted from cells expressing this protein under particular conditions (42), and JFH1 strain is known to cause apoptosis (45). The core complex–specific signals in the HCV particles seem to be increased with the NP-40 treated samples for some unknown reason (Fig. 1c; lanes 1 and 2). Although the intermolecular disulfide bond is known to be artificially formed in denaturing SDS-PAGE in the absence of reducing agents, the core complex was still observed even in the presence of NEM, which is alkylating agent for free sulphydryls, during sample preparation (Fig. 2c), indicating that the core complex was naturally present in the virus-like particles.

The HCV nucleocapsid is covered with lipid membranes and E1 and E2 envelope proteins, making it resistant to proteases. As expected, in the absence of NP-40, the core complex was resistant to proteinase K (Fig. 1c, lane 3), whereas proteinase K was able to digest core protein in whole-cell lysates collected from JFH1 E2FL-transfected HuH-7 cells (Fig. 1c, left panel). Disrupting the envelope structure with NP-40 made the core complex susceptible to proteinase K treatment (Fig. 1c, lane 4), indicating that the core complex was indeed a component of the HCV particle.

The disulfide-bonded core complex (dbc-complex) forms on the ER
To investigate the subcellular site at which the dbc-complex forms, LD and microsomal membrane fractions (MMFs) from JFH1\textsuperscript{E2FL} replicating HuH-7 cells were analyzed by immunoblotting. We first analyzed the dbc-complex in LDs, because the LD is involved in infectious HCV particle formation (36, 47). The purity of the LD fraction was determined using immunoblot analysis of calnexin and adipocyte differentiation-related protein (ADRP), an ER and an LD marker protein, respectively (Fig. 3a, upper panel). The core protein was then analyzed in the LD fraction. As shown in Figure 3a (lower panel), the dbc-complex was observed in the LD fraction from JFH1\textsuperscript{E2FL} RNA–transfected HuH-7 cells. We next analyzed the core protein in the ER-containing MMF, because the core protein is first translated and processed on the ER (16). As shown in Figure 3b, the dbc-complex was observed in the MMF from JFH1\textsuperscript{E2FL} RNA–transfected HuH-7 cells. These results suggest that the dbc-complex is first formed at the ER. To assess the possibility that dbc-complex-containing HCV particles were also assembled on the ER, the sensitivity of the dbc-complex to protease treatment was analyzed. The dbc-complex in the MMF was susceptible to protease treatment in the absence of NP-40, indicating that the dbc-complex on the ER was not part of a HCV particle (data not shown).
dbc-Complex is most likely a disulfide-bonded dimer form of the core

In order to examine whether the core itself have a potential to form
dbc-complex, we analyzed dbc-complex formation of full length wild-type core
(core\textsuperscript{WT}) expressed from pcDNA3-core\textsuperscript{WT} (36), the expression plasmid encoding 191
amino acid full length core of JFH1 strain. We used this expression plasmid because the
core from this plasmid was likely to be processed correctly enough to produce
infectious HCV particles when co-transfected with JFH1\textsuperscript{dc3} RNA, which is a core
deletion mutant of JFH1 (36). As results, the dbc-complex formation was observed from
the MMF of core\textsuperscript{WT} expressing cells both in the absence and the presence of NEM (Fig.
4b; lane 2 and data not shown, respectively). We next investigated the effect of the
amino acid region of E1 on the production of dbc-complex, because it has been reported
that the efficient processing of core protein is dependent on the presence of some E1
sequence to ensure the insertion of the signal sequence for E1 in the
translocon/membrane machinery (34). Then the dbc-complex was also observed when
the core was expressed from a pcDNA3-C-E1/25, which encodes the full length core
followed by the N-terminal 25 amino acid sequence of E1 to ensure that the core is
processed properly (Supplementary Fig. 3a). These data showed that the dbc-complex
was formed by expression of the core protein only in the cells.
Next, we examined the structural components of the dbc-complex. Because the dbc-complex was twice the size of a core monomer, it likely was disulfide-bonded dimer form of the core (dbd-core). So, we investigated whether the core molecules with different tags were able to form the dbd-core. We first generated expression plasmids encoding core with the N-terminal FLAG and Myc tags (pcDNA3-FLAG-core and pcDNA3-Myc-core, respectively; Fig. 4a). The tagged core proteins were expressed or co-expressed with core\textsuperscript{WT} in HuH-7 cells and the MMF was analyzed by SDS-PAGE. The FLAG or Myc tag shifted the positions of the monomer and the complex bands (Fig. 4b; lanes 3 and 4), compared with the core\textsuperscript{WT} (Fig. 4b; lane 2). When the core\textsuperscript{WT} was co-expressed with FLAG-core or Myc-core, the core complex with an intermediately size was observed in addition to the bands obtained when the constructs were independently expressed (Fig. 4b; lanes 5 and 6, filled arrows); the intermediate band disappeared after treatment with β-mercaptoethanol (β-ME) (Supplementary Fig. 3b; lanes 11 and 12, filled arrows), indicating that core\textsuperscript{WT} and tagged core formed a heteromeric disulfide-bonded dimer. These results demonstrated that the dbc-complex on the ER is a dbd-core. Although we tried to detect the hetero-/homo-dimer consisting the tagged-core by using anti-FLAG or anti-Myc antibodies, these dimers but the monomeric forms of the tagged-core were not detected, possibly because of the less
sensitivity and specificity of the antibodies compared to the anti-core antibody we used especially against epitopes in the dbd-core. Above results coupled with the similarity of the molecular size and sensitivity against β-ME and DTT, suggested the dbc-complex in the HCV particle is most likely a dbd-core.

Core cysteine residue 128 (Cys128) mediates dbd-core formation

Our results showed that core from JFH1E2FL forms a disulfide-bonded dimer on the ER. A search for cysteine residues in JFH1E2FL core identified amino-acid positions 128 (Cys128) and 184 (Cys184) (Supplementary Fig. 1). These residues are highly conserved in core proteins from the approximately 2000 reported HCV strains (HCVdb, http://www.hcvdb.org/; Hepatitis Virus Database; http://s2as02.genes.nig.ac.jp/). To determine which cysteine residue mediated disulfide bond formation, we generated point mutations in JFH1E2FL that substituted Cys128 and/or Cys184 with Alanine (Ala)

(C128A, C184A and C128/184A in JFH1C128A, JFH1C184A and JFH1C128/184A, respectively; Fig. 5a). As shown in Figure 5b, core protein from JFH1C128A and JFH1C128/184A failed to form a dbd-core under non-reducing condition, whereas core protein from JFH1C184A formed the dimer, indicating that Cys128 was the responsible residue. Similar results were observed when Cys was substituted to Serine (Ser) instead of Ala
Recently, Majeau et al. reported that the core protein of J6/JFH1 strain with Cys128 substitutions to Ala or Ser were unstable in both *Pichia pastoris* and human hepatoma cell line HuH-7.5 (31), although we did not detect any noticeable degradation of the mutant cores of JFH1 strain (Fig. 5b and Supplementary Fig. 5c). This difference may result from differences in sample preparation as we used full length genome of JFH1<sup>E2FL</sup> strain, bearing JFH1 strain core, and HuH-7 cells instead of core expressing plasmid for J6 strain and HuH-7.5.

To exclude the possibility that mutation of Cys128 inhibited dbd-core formation by creating a conformational change, T127A and G129A core mutants (JFH1<sup>T127A</sup> and JFH1<sup>G129A</sup>, respectively) were created and examined for the effects on dbd-core formation and infectious virus particle production. These mutants formed dbd-core and infectious HCV particles were detected in the culture medium (Supplementary Fig. 4a-c), supporting an essential role for Cys128 in dbd-core and particle formation.

**dbd-Core contributes to HCV particle production**

To examine the functional roles of dbd-core, infectious HCV particle production, HCV replication efficiency, co-localization of core and the LD, and
RNA-binding of mutant and wild-type (JFH1E2FL) core were evaluated. Culture medium from HuH-7 cells transfected with JFH1C128A or JFH1C128/184A RNA contained significantly fewer infectious HCV particles compared with results obtained with JFH1E2FL or JFH1C184A RNA (Fig. 5c). We also found significant decreases in the levels of HCV RNA and the core protein in the culture medium of HuH-7 cells transfected with JFH1C128A or JFH1C128/184A RNA (Fig. 5d, e). Similar results were observed with J6/JFH1 C128A or C128/184A mutant strain (data not shown). To investigate whether these results were due to suppressed HCV replication, HCV RNA and protein levels in cells transfected with mutant RNA were analyzed using qRT-PCR and immunoblot analyses, respectively. Compared with results obtained with JFH1E2FL, no significant changes were observed in the cellular HCV RNA titer at days 1, 3 and 5 post-transfection or in the expression of the HCV nonstructural protein NS5A (Fig. 6a, b). This indicated that substitution of Cys128 did not significantly affect HCV RNA genome replication or viral protein production, demonstrating that the dbd-core functions during HCV particle production rather than HCV genome replication. Similar results were observed using RNA of JFH1 mutant strain which Cysteine of position 128 were substituted to Ser instead of Ala; JFH1C128S (Supplementary Fig. 5a, b, d).

The subcellular localizations of core and NS5A in HuH-7 cells transfected with
HCV RNA were investigated using indirect immunofluorescence and confocal microscopy, because recruiting HCV proteins to the LD is an important step in infectious HCV particle production (36, 47) and core trafficking to the LD is dependent on SPP-mediated cleavage of the tail region (34, 41). JFH1\textsuperscript{C128A} mutant core and NS5A were efficiently trafficked to the LD, as was observed with wild-type JHF1\textsuperscript{E2FL} (Fig. 6c), suggesting that SPP cleavage and core maturation were not affected by the C128A mutation. Similar results were obtained with core derived from JFH1\textsuperscript{C184A} and JFH1\textsuperscript{C128/184A} (Supplementary Fig. 6), and also, Ser mutant JFH1\textsuperscript{C128S} (Supplementary Fig. 5e).

Because HCV core protein can bind RNA, including the HCV genome during viral particle assembly (43), we analyzed RNA binding by core using \textit{in vitro} translated core\textsuperscript{C128A}, core\textsuperscript{WT}, and poly-uridine (U) agarose resin. Core\textsuperscript{C128A} and core\textsuperscript{WT} similarly bound with poly-U resin (Fig. 6d).

\textbf{dbd-Core is important for HCV particle assembly}

The mutational analysis of core indicated that core\textsuperscript{C128A} and core\textsuperscript{WT} similarly localize to LDs, recruit NS proteins to the LD, and bind to RNA. Moreover, this mutation did not markedly affect HCV genome replication. How does core\textsuperscript{C128A} affect
the production of HCV particles? An important function of core protein is multimerization, which is followed by capsid construction and packaging of the RNA genome in the viral particles. We therefore determined whether core\(^{C128A}\) had a dominant-negative effect on virus-like particle production. Wild-type JFH1\(^{E2FL}\) RNA and different amounts of JFH1\(^{C128A}\) RNA were co-transfected into HuH-7 cells and the HCV RNA titer and infectivity of the virus-like particles in the culture medium were analyzed. As expected, the HCV RNA titer in the cells increased with higher levels of transfected RNA (Supplementary Fig. 7a). In contrast, the HCV RNA titer and infectivity in the culture medium decreased in a JFH1\(^{C128A}\) RNA dose–dependent manner when this mutant RNA was co-transfected with wild-type RNA (Fig. 7a, b). This suppressive effect was not observed when either wild-type RNA or core deletion mutant JFH1\(^{dc3}\) RNA was used instead of mutant RNA in a similar experiment (Supplementary Fig. 7b-e), indicating that higher levels of HCV RNA alone did not inhibit HCV particle production. Thus, core\(^{C128A}\) had a dominant-negative effect on HCV particle production. Together, these results suggest that dbd-core is involved in the assembly of HCV particles.

The nucleocapsid-like particle of HCV was resistant to trypsin treatment
To further investigate the structure of the HCV nucleocapsid-like particle most likely formed by dbd-core, we examined the sensitivity of the particle to trypsin, which cleaves polypeptides at the C-terminal end of basic residues. Whereas trypsin digested core in the whole-cell lysates (Fig. 8a, left panel), dbd-core from buoyant density-fractionated JFH1\textsuperscript{E2FL} particles was resistant to digestion despite NP-40 treatment (Fig. 8a, right panel), although it was sensitive to proteinase K which have a broad specificity (Fig. 1c). The N-terminal hydrophilic domain of the core protein (from residues 6-121) contains a number of trypsin cleavage sites (25 sites, in JHF1 strain) (Supplementary Fig. 1), suggesting that the N-terminal domain faces inward and/or the conformation prevents protease access. To address this idea, buoyant density-fractionated JFH1\textsuperscript{E2FL} particles were treated with trypsin under more strict conditions in the presence of NP-40. Cleavage of dbd-core by various levels of trypsin correlated with the appearance of a shorter molecule (Fig. 8b, white arrowhead). The shorter molecule was presumed to be partially digested dbd-core with an intact N-terminal region because it was recognized by anti-core antibodies, which bind an epitope located in amino-acids 20-40 of core (personal communication from Dr. M. Kohara, The Tokyo Metropolitan Institute of Medical Science, Japan). These results suggest that dbd-core is assembled into the nucleocapsid-like particle such that most of
the N-terminal domain is inside.
DISCUSSION

In this study, we have shown that the nucleocapsid-like particle of HCV contains most likely a dimer of core protein that is stabilized by a disulfide bond. Mutational analysis revealed that Cys128 forms the disulfide bond between core monomers. Several reports have shown that disulfide bonds in the capsid proteins of some viruses are involved in virus particle assembly and stabilization of the viral capsid structure (4, 21, 27, 28, 57); these viruses are characterized by icosahedral nucleocapsids. Because, like these viruses, the HCV virion is spherical (2, 20), it has been suggested that HCV may contain a nucleocapsid with a similar structure (20). We found the dbc-complex which is most likely to be the dbd-core in JFH1\textsuperscript{E2FL} virus-like particles (Figs. 1c and 8a). The dbd-core in the capsid structure was digested by proteinase K but not trypsin in the presence of NP-40 (Figs. 1c and 8a, lane 4). The resistance to trypsin suggested a tight conformation for dbd-core in the capsid with no exposed trypsin cleavage sites. The truncated form of dbd-core that was observed under certain trypsin treatment conditions likely resulted from cleavage in the C-terminal portion of the protein (e.g., arginine residues at positions 149 and 156) (Supplementary Fig. 1), although it is possible that the truncation of dbd-core was due to non-specific cleavage by trypsin. These results imply that dbd-core is configured such that the N-
and C-terminal ends are located at the inner and outer surface of the capsid, respectively.

Because the N-terminal region of core includes the RNA binding domain (43), the HCV RNA genome likely interacts with core as it is packed in the nucleocapsid. On the other hand, the C-terminal hydrophobic domain probably faces the lipid membranes to form the envelope structure. Only part of the N-terminal hydrophilic region of the core protein has been structurally examined using X-ray crystal structural analysis (35), and using structural bioinformatics and nuclear magnetic resonance analysis (11). Although the C-terminal half of core has been structurally investigated by bioinformatics (8), the 3D structure containing the Cys128 residue is unknown. Thus, determination of the structure of the core in the nucleocapsid containing Cys128 residue should be required for understanding the whole structure of this protein in the virus particles.

Because co-transfection of JFH1\textsuperscript{C128A} RNA with wild-type JFH1\textsuperscript{E2FL} RNA inhibited particle production in a mutant RNA dose-dependent manner (Fig. 7a, b), the C128A core variant clearly inhibited HCV particle formation by wild-type core. Cys128 was also reported previously to be a residue included in the region important for the production of infectious HCV (39). This residue is located near the N-terminal end of the hydrophobic region of the core (amino acids 122-177) and belongs to the hydrophilic side of an amphipathic helix expected to interact in-plane of the membrane.
interface (7). Therefore, it is possible to think that the dbd-core formation via Cys128 can stabilize the interaction between core and the membranes. The N-terminal half of core (amino acids 1-124) reportedly assembles into nucleocapsid-like particles in the presence of 5’-UTR from HCV RNA (24), suggesting that some nucleocapsid-like particles may assemble via only homotypic interactions from the core protein. In addition to weak homotypic interactions, the HCV core protein forms a disulfide bond to stabilize the capsid structure, thus making dbd-core dispensable in the stable virus-like particle. We observed that culture medium from JFH1^{C128A} or JFH1^{C128S}-transfected cells included slight infectivity (Fig. 5c or Supplementary Fig. 5d). This made us speculate that this mutant may produce some infective particle-like structure formed by homotypic interaction of the core. Such a slight infectivity may have reflected the optimized in vitro culture conditions compared with in vivo conditions, allowing relatively unstable virus particles to survive.

A nucleocapsid must be resistant to environmental degradation, yet still be able to disassemble after infection. Disulfide bonds could help with these process by switching between a stable and unstable virus capsid based on different intracellular and extracellular oxidation conditions (12, 30). During the virus life cycle, the disulfide bond strengthens the viral capsid structure and protects the viral genome from oxidative...
conditions and cellular nucleases when virus particles are formed. Upon infection, the
disulfide bond may be cleaved under cytoplasmic reducing conditions, thereby releasing
the viral genome into the cell for replication. HCV may utilize the core protein disulfide
bond in this way as HCV enters the host cell via clathrin-mediated endocytosis (5) into
a low-pH, endosomal compartment (25, 52); this is presumably followed by endosomal
membrane fusion and release of the viral capsid into the cytoplasm (38).

Treatment of HCV with pegylated interferon in combination with ribavirin is
not effective for all patients. Recently, drugs targeting the viral proteins NS3/4A and
NS5B have been examined in clinical trials. Although these drugs are relatively specific,
resulting in fewer side effects and potent antiviral activity, monotherapy can be
complicated by rapidly emerging resistant variants, carrying mutations that reduce drug
efficacy, perhaps due to conformational changes in the target (23, 48, 51). Therefore,
viral proteins that are highly conserved among strains and those characterized by low
mutation rates may be better targets for drug development. Because the core protein is
the most conserved HCV protein and Cys128 is conserved among almost all examined
HCV strains, drugs that interact with Cys128 and/or region around or near this residue
will likely show broad spectrum efficacy to block the stable infectious particle
formation. Structural analysis of dbd-core should aid the development of new STAT-Cs
that target Cys128 by direct interaction with the sulfide group and/or region around this
residue. Until now and still, the mechanism of disulfide bond formation of core on the
ER is unknown. Dimerization of capsid protein by disulfide bond has been reported in
some enveloped viruses (9, 19, 54, 56), although some were shown not to be important
for virus particle formation (26, 55). Unlike the vaccinia virus (46), no redox system of
its own has been reported for these viruses. Therefore, further investigations addressing
the mechanisms underlying dbd-core formation on the ER may reveal new mechanism
for disulfide bond formation of viral proteins in infected cells.
ACKNOWLEDGMENTS

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FIGURE LEGENDS

Figure 1. The HCV-like particle consists of a core complex formed by a disulfide bond. (a) The infectivity of the pellet fraction collected from concentrated culture medium from JFH1\textsuperscript{E2FL} RNA–transfected HuH-7 cells was analyzed as described in the Materials and Methods. “input” represents the same volume of concentrated culture medium used to pellet the virus-like particles. (b) Immunoblot analysis of the core in pellets containing JFH1\textsuperscript{E2FL} virus particles treated with various levels of DTT (lanes 1, 2, 3, 4, 5 and 6 represent 0, 1.56, 3.13, 6.25, 12.5 and 25 mM, respectively). (c) Immunoblot analysis of core in JFH1\textsuperscript{E2FL} particles collected from sucrose density gradient fractions with high HCV RNA titers (particle) (Fig. 2a, fraction #8 to #13) and treated with 5 µg/ml proteinase K at 3°C for 15 min in the presence or absence of 1% NP-40 (right panel). As a positive control, whole-cell lysate (WCL) prepared from JFH1\textsuperscript{E2FL} RNA–transfected HuH-7 cells in lysis buffer was treated with 5 µg/ml proteinase K at 37°C for 15 min (left panel). Data are representative of three independent experiments.

Figure 2. HCV nucleocapsid-like particle consists of core complex. (a) HCV RNA titer in culture medium separated on a 20-50% sucrose density gradient. Concentrated
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**Figure 3.** The core complex is formed at the LD and ER. (a) The LD fraction and whole-cell lysate (WCL) were collected from JFH1\textsuperscript{E2FL} RNA–transfected HuH-7 cells
on day 5 post-transfection. Immunoblot analysis of the LD marker adipose
differentiation-related protein (ADRP) and the ER marker calnexin in the LD fraction
(upper panel). Immunoblot analysis of core in the LD fraction treated with or without 50
mM DTT (lower panel). (b) Immunoblot analysis of core protein in the MMF and WCL
collected from JFH1E2FL-producing HuH-7 cells on day 5 post-transfection in the
presence or absence of 5% β-mercaptoethanol (β-ME). Data are representative of three
independent experiments.

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FLAG-tagged (FLAG-core), and Myc-tagged (Myc-core) cores. (b) Immunoblot
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**Figure 6.** Site-directed mutagenesis has no effect on HCV replication. (a) Real-time qRT-PCR analysis of the HCV RNA titer using total cellular RNA collected at the indicated time points from cells transfected with JFH1\textsuperscript{E2FL} (WT) (open circles),
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Infectivity of culture medium collected from HuH-7 cells that had been transfected with the indicated ratio of WT and C128A RNA was analyzed as described in the Materials and Methods. Data represent the means ± s.d. from three independent experiments (a) or are representative of three independent experiments (b).

**Figure 8.** The nucleocapsid-like particle of JFH1\(^{E2FL}\) is assembled with the C-terminal region of core on the outer surface. (a) Immunoblot analysis of core in JFH1\(^{E2FL}\) particles collected from sucrose density gradient fractions with high HCV RNA titers (particle) (Fig. 2a, fraction #8 to #13). Fractions were treated with 10 µg/ml trypsin at 37°C for 15 min in the presence or absence of 1% NP-40 (right panel). As a positive control, whole cell lysate (WCL) prepared from JFH1\(^{E2FL}\) RNA–transfected HuH-7 cells in lysis buffer was treated with 10 µg/ml trypsin at 37°C for 15 min (left panel). (b) Immunoblot analysis of core in JFH1\(^{E2FL}\) particles collected from sucrose density gradient fractions with high HCV RNA titers. Fractions were treated with the indicated concentrations of trypsin at 37°C for 10 min in the presence of 1% NP-40. Open and filled arrows indicate the positions of dbd-core and the trypsin-digested fragment, respectively. Data are representative of three independent experiments.
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HuH-7 cells transfected with pcDNA3 (vector) and/or core expression plasmids (core191, FLAG-core, and Myc-core) as indicated. Samples were treated with or without 5% β-mercaptoethanol (β-ME). Filled arrowheads indicate the positions of the intermediate core complexes formed by core\textsuperscript{WT} and tagged core. Data are representative of two (a) or three (b) independent experiments.

**Supplementary Figure 4.** Site-directed mutagenesis of amino-acid position 127 or 129 had no effect on HCV replication or the production of HCV particles. (a) Immunoblot analysis of core in microsomal membrane fractions collected on day 3 post-transfection from cells transfected with JFH1\textsuperscript{E2FL} (WT), JFH1\textsuperscript{T127A} (T127A), or JFH1\textsuperscript{G129A} RNA. Samples were treated with or without 5% β-mercaptoethanol (β-ME). (b, c) Real-time qRT-PCR analysis of HCV RNA titers in total cellular RNA (b) or culture medium (c) collected on day 5 post-transfection. Data are representative of three independent experiments (a) or are the means ± s.d. from three independent experiments (b, c).

**Supplementary Figure 5.** Analysis of core C128S mutant. (a) Real-time qRT-PCR analysis of HCV RNA titers in culture medium collected at the indicated time points from HuH-7 cells transfected with JFH1\textsuperscript{E2FL} (WT, open circles) or JFH1\textsuperscript{C128S} (C128S, filled circles) RNA. (b) Real-time qRT-PCR analysis of the HCV RNA titer using total
cellular RNA collected at the indicated time points from cells transfected with WT (open circles) or (C128S) (filled circles). (c) Immunoblot analysis of core in microsomal membrane fraction collected on day 3 post-transfection from cells transfected with JFH1E2FL (WT) or JFH1C128S RNA (C128S). (d) Infectivity of culture medium collected and concentrated on day 5 post-transfection from HuH-7 cells transfected with WT or C128S RNA. (e) Confocal microscopy of the subcellular localization of the LD (green), core (blue), NS5A (red), and nucleus (DAPI) (grey) in cells transfected with JFH1E2FL (WT) or JFH1C128S RNA (C128S) on day 3 post-transfection. Data are the means ± s.d. from three independent experiments (c, b) or are representative of three independent experiments (c, d, e).

**Supplementary Figure 6.** Subcellular localization of HCV proteins. Confocal microscopy of the subcellular localizations of the lipid droplet (LD), core, NS5A, and the nucleus (DAPI) three days post-transfection with JFH1C184A (C184A) or JFH1C128/184A (C128/184A). Scale bar indicates 10 µm. Data are representative of three independent experiments.

**Supplementary Figure 7.** Transfection of various amounts of HCV RNA had no
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Figures and Legends for:

A DISULFIDE-BONDED DIMER OF THE CORE PROTEIN OF HEPATITIS C VIRUS IS IMPORTANT FOR VIRUS-LIKE PARTICLE PRODUCTION

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Figure 1. The HCV-like particle consists of a core complex formed by a disulfide bond. (a) The infectivity of the pellet fraction collected from concentrated culture medium from JFH1 E2FL RNA–transfected HuH-7 cells was analyzed as described in the Materials and Methods. “input” represents the same volume of concentrated culture medium used to pellet the virus-like particles. (b) Immunoblot analysis of the core in pellets containing JFH1 E2FL virus particles treated with various levels of DTT (lanes 1, 2, 3, 4, 5 and 6 represent 0, 1.56, 3.13, 6.25, 12.5 and 25 mM, respectively). (c) Immunoblot analysis of core in JFH1 E2FL particles collected from sucrose density gradient fractions with high HCV RNA titers (particle) (Fig. 2a, fraction #8 to #13) and treated with 5 μg/ml proteinase K at 3°C for 15 min in the presence or absence of 1% NP-40 (right panel). As a positive control, whole-cell lysate (WCL) prepared from JFH1 E2FL RNA–transfected HuH-7 cells in lysis buffer was treated with 5 μg/ml proteinase K at 37°C for 15 min (left panel). Data are representative of three independent experiments.
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Figure 7 (Hijikata)

**a**

[Graph showing HCV RNA titer in culture medium for different transfected RNA ratios (WT : C128A).]

**b**

[Images showing culture medium collected from transfected HuH-7 cells for different RNA ratios (WT : C128A).]

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Supplementary information for:

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<table>
<thead>
<tr>
<th>Plasmid name</th>
<th>Primer sequences (5’-3’)</th>
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