Lipid Droplet-Binding Protein TIP47 Regulates Hepatitis C Virus RNA Replication through Interaction with the Viral NS5A Protein

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Abstract
The nonstructural protein NS5A has emerged as a new drug target in antiviral therapies for Hepatitis C Virus (HCV) infection. NS5A is critically involved in viral RNA replication that takes place at newly formed membranes within the endoplasmic reticulum (membranous web) and assists viral assembly in the close vicinity of lipid droplets (LDs). To identify host proteins that interact with NS5A, we performed a yeast two-hybrid screen with the N-terminus of NS5A (amino acids 1–31), a well-studied α-helical domain important for the membrane tethering of NS5A. Our studies identified the LD-associated host protein, Tail-Interacting Protein 47 (TIP47) as a novel NS5A interaction partner. Coimmunoprecipitation experiments in HuH7 hepatoma cells confirmed the interaction of TIP47 with full-length NS5A. shRNA-mediated knockdown of TIP47 caused a more than 10-fold decrease in the propagation of full-length infectious HCV in HuH7.5 hepatoma cells. A similar reduction was observed when TIP47 was knocked down in cells harboring an autonomously replicating HCV RNA (subgenomic replicon), indicating that TIP47 is required for efficient HCV RNA replication. A single point mutation (W9A) in NS5A that disrupts the interaction with TIP47 but preserves proper subcellular localization severely decreased HCV RNA replication. In biochemical membrane flotation assays, TIP47 cofractionated with HCV NS3, NS5A, NS5B proteins, and viral RNA, and together with nonstructural viral proteins was uniquely distributed to lower-density LD-rich membrane fractions in cells actively replicating HCV RNA. Collectively, our data support a model where TIP47—via its interaction with NS5A—serves as a novel cofactor for HCV infection possibly by integrating LD membranes into the membranous web.

Introduction
Plus-strand RNA viruses induce a highly regulated process of membrane rearrangements and novel vesicle formation in infected cells, in order to create a suitable environment for viral RNA replication as well as assembly and release of new virions. These viruses replicate their genome in membrane-associated complexes; however, the origin of the membranes used for replication varies from virus to virus. Infection with Hepatitis C Virus (HCV), a single- plus-stranded RNA virus within the Flaviviridae family, triggers rearrangements of intracellular membranes, resulting in membranous vesicles of heterogeneous size and morphology [1]. This so-called “membranous web” is known to include ER membranes [2] and is considered to be the site of HCV RNA replication [3]. Expression of the non-structural viral protein NS4B in mammalian cells leads to membrane alterations resembling the membranous web [1]. However, molecular details of how the membranous web is formed in HCV-infected cells are still lacking.

The viral non-structural protein NS5A is a proline-rich, hydrophilic phosphoprotein that functions as a key regulator of HCV RNA replication and assembly [4]. NS5A has recently emerged as a major drug target in HCV infection, although the precise mode-of-action of NS5A-targeting drugs is not fully understood [5]. NS5A has no intrinsic enzymatic activity and is thought to function mainly as an adaptor protein for a wide variety of host proteins, including factors involved in host signaling, membrane trafficking, and lipoprotein synthesis pathways [6–8].

At the ER, NS5A is a component of the HCV replication complex that includes the two viral proteases NS2 and NS3/4A, the virally encoded RNA-dependent RNA polymerase NS5B, and the NS4B protein [1]. The N-terminus of NS5A forms a 30-amino-acid amphipathic α-helix that is highly conserved among HCV isolates [9]. Disruption of the amphipathic nature of the α-
Hepatitis C Virus (HCV) belongs to the Flaviviridae family, and is an enveloped, positive, single-stranded RNA virus containing a 9.6 kb genome. Plus-strand RNA viruses induce a highly regulated process of membrane rearrangements and novel vesicle formation in infected cells (the “membranous web” for HCV) to create a suitable environment for RNA replication as well as for the assembly and release of new virions. HCV assembly occurs close to lipid droplets (LDs), cell organelles involved in fat storage and utilization. The viral non-structural protein NS5A plays a critical role in both viral RNA replication that occurs within the membranous web and viral assembly at LDs. We identified the host protein TIP47 as a novel host interaction partner for NS5A. TIP47 is a LD-binding protein also known to function as cargo in late-endosome-to-Golgi vesicular transport. Our data support a model where the recruitment of TIP47 by NS5A is required for viral RNA replication, indicating that LDs play a previously unrecognized role not only in viral assembly but also in RNA replication.

Results

Interaction of the N-terminal amphipathic α-helix of NS5A with TIP47

To identify host factors that interact with HCV NS5A, a Y2H assay was performed using the N-terminal amphipathic α-helix as bait (Figure 1A,B). The screen resulted in 22 confirmed positive clones. 12 of the 22 clones were found to be TIP47 (by sequencing and 8 by restriction digest), thus identifying TIP47 as a novel interaction partner with this domain in NS5A. Using a quantitative Y2H assay, the binding site of NS5A to TIP47 was mapped to a minimal domain spanning amino acids 1–14 in NS5A (Figure 1C). A tryptophan-to-alanine mutation at position nine (W9A) decreased protein binding by 6-fold compared to the wild-type sequence (Figure 1C), but did not interfere with the subcellular localization of NS5A when introduced into the full-length NS5A protein (Figure S1). These results are consistent with previous data indicating that the N-terminus of NS5A is involved in protein-protein interactions essential for HCV propagation independently from its reported role in membrane localization [11].

The TIP47-NS5A interaction was confirmed in coimmunoprecipitation experiments in human HuH7 hepatoma cells. Myc-tagged TIP47 was overexpressed together with FLAG-tagged NS5A, or a deletion mutant in which the N-terminal 31 amino acids were missing (Δ31-NS5A-Flag). TIP47 coimmunoprecipitated with full-length but not N-terminally truncated NS5A, confirming that these proteins interact in a manner dependent on the amphipathic α-helix of NS5A (Figure 1D).

Furthermore, the TIP47-NS5A interaction was confirmed in the context of active HCV RNA replication in HuH7.5 cells harboring a subgenomic HCV replicon expressing only the non-structural viral proteins NS3–5B (Figure 1E). This replicon is sufficient to autonomously replicate HCV RNA but cannot support infectious particle production since critical structural proteins as well as p7 and NS2 are lacking (Figure 2A, HCV Replicon, genotype 1b) [25]). Collectively, these results identify TIP47 as a novel host factor binding to the N-terminus of NS5A.

TIP47 is required for HCV RNA replication

To study TIP47’s role in HCV infection, we introduced shRNAs directed against TIP47 or non-targeting control shRNAs into HuH7.5 cells via lentiviral transduction. No toxicity or significant alterations in LD content were observed in cells after TIP47 knockdown (Figure S2). Transduced cells were then transfected with replicon RNA (genotype 1b) and selected with G418. The replicon RNA encodes the neomycin phosphotransferase selection gene thus only cells actively replicating HCV RNA survive (Figure 2A). After four weeks, surviving clone cells were fixed, stained with crystal violet and counted. We found that down regulation of TIP47 expression resulted in a more than 10-fold decrease in colony formation as compared to control cells, indicating that TIP47 plays an important role in HCV RNA replication (Figure 2B, C). The loss of TIP47 expression was verified by western blotting (Figure 2D). These studies reveal that TIP47 functions as a new cellular regulator of HCV RNA replication.

We confirmed these results in the context of fully infectious HCV. Here, shRNA-transduced cells were transfected with infectious RNA of HCV-JFH1 (genotype 2a), carrying firefly luciferase as a reporter gene (Figure 2A, Luc-JFH1). Cells were harvested at different time points after transfection and subjected to luciferase assays. All values were expressed as the fold increase of luciferase.
per well over an early 4-hour time point before RNA replication occurs [26,27]. We observed a 10-fold decrease in luciferase ratios at 72 hours post-transfection when TIP47 expression was down regulated compared to control cells (Figure 2E,F). When we reconstituted TIP47 expression in shRNA-containing cells via overexpression of an shRNA-resistant allele, HCV infection was restored underscoring the specific effect TIP47 plays in HCV infection (Figure 2E). In accordance with these observations, overexpression of Myc-TIP47 in cells treated with control shRNAs enhanced HCV replication, pointing to a limiting function of TIP47 expression for HCV infection in cells (Figure 2E).

TIP47 regulates HCV RNA replication via interaction with NS5A

To examine whether the interaction with NS5A is important for TIP47’s role in HCV RNA replication, we introduced the W9A mutation that decreased the NS5A-TIP47 interaction into the replicon cells, and transfected the mutated version into Huh7.5 cells as described above (Figure 3A, HCV-Replicon). The W9A mutation almost completely abolished colony formation in transfected Huh7.5 cells (Figure 3B, compare WT to W9A). To test for revertant or compensatory mutations, we sequenced the NS5A region in the few surviving colonies, and found that all sequenced clones maintained the W9A mutation, but discovered that all clones carried a second isoleucine-to-valine mutation at position 12 (I12V). To test whether the I12V mutation suppresses the RNA replication defect induced by the W9A mutation, the I12V mutation alone as well as the combined W9A-I12V mutation was cloned back into the wild-type replicon RNA and studied in HCV replicon assays. The I12V mutation alone resulted in similar colony numbers as the wild-type replicon RNA (Figure 3B, compare W9-V12 to WT). However, when combined with the
Figure 2. TIP47 is required for HCV RNA replication. A) Schematic representation of HCV Replicon (Con1, genotype 1b), a subgenomic bicistronic HCV replicon with neomycin as a reporter gene, and Luc-JFH1 (genotype 2a), a full-length bicistronic HCV infectious clone with firefly luciferase as a reporter gene. B) Crystal violet-stained Huh7.5 cells transfected with control shRNA or shRNA against TIP47 and HCV replicon RNA (A). G418 was added to the medium (1 mg/ml). Cells were kept under selective pressure for 4 weeks. Only cells actively replicating HCV RNA, and thus expressing the neomycin phosphotransferase survive. C) Quantification of colony numbers/plate in (B) (mean ± standard deviation; n = 3, *P < 0.05). D) Western blot analysis of TIP47 expression in shRNA-expressing Huh7.5 cells at the time of transfection in (B). E) Transfection of shRNA-expressing Huh7.5 cells with Luc-JFH1 RNA (A) and Myc-TIP47 expression plasmid (where indicated). HCV replication was measured by determining the fold increase of luciferase (RLU) per well over a 4 h time point (mean ± standard deviation; n = 3; *P < 0.05, **P < 0.01). F) Western blot analysis of TIP47 expression in shRNA-expressing Huh7.5 cells at time of transfection in (E).

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Figure 3. TIP47 is involved in HCV RNA replication via interaction with NS5A. A) Schematic representation of the HCV Replicon (genotype 1b) and Luc-JFH1 (genotype 2a), with mutations at position 9 and 12 in the NS5A N-terminus. B) Crystal violet-stained Huh7.5 cells transfected with wild-type or mutant HCV replicon RNA (genotype 1b). 24 h post-transfection, G418 was added to the medium (1 mg/ml). Cells were kept under selective pressure for 4 weeks before being stained with crystal violet. These results are representative of three independent experiments. C) Luciferase assay of Huh7.5 cells transfected with wild-type Luc-JFH1 RNA (genotype 2a) or Luc-JFH1 RNA containing indicated NS5A mutations. Values are expressed as fold increase of luciferase (RLU) per well, over a 4 h time point (mean ± standard deviation; n = 3, **P < 0.01, ***P < 0.001). The decrease of A9-I12 and W9-I12 is significant in relation to WT (indicated in grey *), whereas the increase in A9-V12 is significant in relation to A9-I12 where indicated (indicated in black *). D) Amino acid sequences of the NS5A N-terminus of HCV replicon (genotype 1b) and HCV-JFH1 (genotype 2a). Positions 9 and 12 are highlighted in bold black and mutations are highlighted in color. Summary of respective replication phenotypes in the HCV replication assay (genotype 2a, Figure 3B, C). Wild-type replication phenotypes were set as equal to (++++) in their respective assays, while reduced replication levels were defined in the Replicon assay as (+++) for ~90% of wild-type colonies, (+) for ~20% colonies, or (−) for no colonies. In the Luciferase assay, luciferase values were described as (+++) for values ~19% of wild-type, (+) for ~2%, and (−) for values below 1%.
W9A mutation, we found that the I12V mutation partially restored colony formation, suggesting that this mutation evolved to overcome the replication defect induced by the loss of the TIP47 binding (Figure 3B, compare A9-V12 to W9A). Using full-length Luc-JFH1 constructs (Figure 3A, Luc-JFH1), we obtained similar results as with the HCV replicon: a near complete defect in RNA replication with the A9-V12 mutation and a partial rescue of this defect with the A9-I12 mutation (Figure 3C,D). However, it must be noted that the HCV genotype 2a NS5A protein contains valine rather than isoleucine at position 12 in its wild-type sequence (Figure 3D). Surprisingly, a valine to isoleucine mutation at position 12 alone was lethal in genotype 2a (Figure 3D). Similar to the HCV replicon, we obtained similar results as with the HCV replicon: a near complete defect in RNA replication with the A9-I12 mutation and a partial rescue of this defect with the A9-V12 mutation. After immunoprecipitation with Flag antibodies, TIP47 was detected by western blotting with antibodies specific for Myc.

Next, we tested the effect of these mutations on TIP47 binding to NS5A. Single and double mutations were cloned into the FLAG-tagged NS5A expression vector, transfected along with Myc-tagged TIP47 in Huh7 hepatoma cells, and subjected to immunoprecipitation and western blot assays to determine the extent of the interaction. While cells singly expressing either wild-type or I12V-mutant NS5A co-immunoprecipitated efficiently with TIP47, a significant decrease in binding was observed with the W9A mutant (Figure 3E, compare lane 7 with lane 8), consistent with the results obtained in the Y2H assays (Figure 1C). In contrast, TIP47 binding was restored in cells expressing the double-mutant NS5A construct, indicating that the I12V mutation rescued HCV RNA replication by re-establishing the interaction with TIP47 (Figure 3E, lane 10). These results underline the importance of the NS5A-TIP47 interaction for HCV RNA replication.

Altered membrane association of TIP47 during HCV RNA replication

As HCV RNA replication occurs within the membranous web, we next examined whether TIP47 associates with HCV replicase-containing intracellular membranes. We performed membrane flotation assays in naive or HCV replicon-containing Huh7.5 cells (genotype 1b), in which membrane compartments of crude cell lysates (CL) were separated in a self-forming linear 10–20–30% iodixanol density gradient as described [29]. Twenty-two fractions were collected from top to bottom and assessed by western blotting for TIP47 expression. Cell lysates were also analyzed by western blotting before gradient centrifugation to assess the overall cellular expression of the proteins. No significant differences in expression levels were found for TIP47 in the lysates of HCV replicon–expressing cells compared to Huh7.5 control cells (Figure 4A, CL).

We divided the 22 fractions obtained from the gradient into three areas based on their membrane-associated protein content. The lowest-density membrane fractions (fractions 1–8) were rich in ADRP, a constitutive LD-binding protein [30], but devoid of calreticulin, an integral ER marker, and were therefore labeled “LD-rich” (Figure 4A). Fractions 9–15 contained calreticulin and endosomal Rab proteins (Figure S2A, B) as well as ADRP and were labeled “ER/endosome-rich”, while fractions 16–22 contained soluble and aggregated proteins as previously determined [31] (Figure 4A).

In naïve Huh7.5 cells, TIP47 was mainly found in the ER/endosome-rich and the soluble/aggregated protein fractions, but not within low-density LD-rich membranes (Figure 4A, upper 5 panels). This distribution changed in HCV replicon–containing cells, where TIP47 was found in low-density LD-rich membranes (Figure 4A, lower 5 panels). Overall the fractions of membrane-bound TIP47 increased—especially in LD-rich fractions, with about 7% of total TIP47 in fraction 1 alone—while soluble TIP47 levels decreased (Figure 4B). A slight increase of TIP47 in ER/endosome-rich fractions 9–11 was also observed (Figure 4B). In contrast, the subcellular distributions of LD- and ER-specific markers, ADRP and calreticulin, did not change significantly upon HCV RNA replication, except for a slight increase in calreticulin in fraction 1 (Figure 4A). These results indicate that in cells that actively replicate HCV, TIP47's membrane association is enhanced with enrichment in LD-rich membranes.

Cofractionation of TIP47 with viral RNA and NS5A, NS5B, and NS3 proteins

To test whether TIP47 cofractionated with HCV replication complexes, we blotted the gradient fractions with antibodies against HCV NS3, NS5B, and NS3 proteins. In HCV replicon–containing cells, the viral proteins NS3, NS5B, and NS5A cofractionated with TIP47 in ER/endosome-rich fractions, and like TIP47, were also found in LD-rich fractions, most prominently in fraction 1 (Figure 4A,B). As all three viral proteins are part of the HCV replication complex, this finding supports a model in which TIP47 associates with HCV RNA replication complexes both within the ER and at LD-rich membranes. In support of this model, viral RNA was detected in LD- and ER/endosome-rich fractions with highest concentrations in fractions 1 and 9 underscoring that these fractions harbor actively replicating replicase complexes (Figure 4C).

We also performed similar membrane flotation assays in Huh7.5 cells expressing the NS5–3B polyprotein from a DNA-based expression vector, which allowed us to compare wild-type and W9A mutant NS5A in the context of the HCV replicase independently from RNA replication. Redistribution to LD-rich membranes was reduced for both, TIP47 and NS5A, when the NS5A W9A mutant was tested, indicating that the joint association with these membranes depends, at least in part, on efficient NS5A-TIP47 interactions (Figures 4D, E).

As TIP47 has been implicated as an adaptor protein in late-endosome-to-Golgi trafficking, we also tested the distribution of several endosomal Rab GTPases in the gradient fractions, including the previously described TIP47 interaction partner Rab9 [21,22]. Rab9 was not found in LD-rich fractions in control cells, and showed only a modest (3% of all protein) enhancement in fraction 1 in HCV replicon–expressing cells (Figure S3A, B). In contrast, the early endosomal protein Rab5 was found in LD-rich fractions in uninfected and HCV replicon-containing cells, consistent with previous reports demonstrating its association with the surface of LDs [32] (Figure S3A, B). In HCV replicon–containing cells, a strong redistribution to LD-rich fractions was observed for Rab7, a late endosomal GTPase previously shown to play a critical role in HCV RNA replication [26,33] (Figure S3A, B). Interestingly, we noted a marked (~3-fold) overall increase in Rab9 expression in the cell lysates of HCV replicon–containing cells compared to Huh7.5 control cells, suggesting enhanced late-endosome-to-Golgi trafficking in HCV replicon–expressing cells (Figure S3A).
Figure 4. TIP47 cofractionates with lipid droplets, viral nonstructural proteins, and viral RNA during HCV RNA replication. A) Western blot following the membrane flotation assay in an iodixanol gradient. Briefly, cytoplasmic lysates (CL) of either naïve Huh7.5 cells or Huh7.5 cells harboring a subgenomic HCV replicon (genotype 1b) were mixed 1:1 with 60% iodixanol (in PBS/0.25 M sucrose) for a final 30% iodixanol-.
Association of TIP47 with LDs during HCV infection

To independently study the association of TIP47 with LDs during HCV replication, we isolated highly purified LD fractions free of ER contaminations from naïve Huh7.5 or replicon cells. These studies confirmed that TIP47, together with NS5A, was selectively recruited to purified LD fractions in replicon cells but was not present at LDs in naïve Huh7.5 cells (Figure 5A). As TIP47 is known to associate with LDs during active LD formation, we examined the LD content in replicon cells. Of note, replicon-containing cells do not express the viral protein core, known to induce LD accumulation in infected cells [34,35]. Contrarily, we found that the number of LDs per cell was significantly down regulated in replicon cells in comparison to naïve Huh7.5 cells excluding the possibility that enhanced LD formation triggered the translocation of TIP47 to LDs in replicon cells (Figure 5B).

In indirect immunofluorescence microscopy, more than 60% of NS5A colocalized with TIP47 in replicon cells (genotype 1b) (Figures 5C, E). Interestingly, only ~15% of NS5A colocalized with LDs in these cells, confirming results from membrane flotation assays showing that the majority of NS5A and TIP47 are found at ER/endosome-rich membranes or are soluble (Figures 5C, F). Interestingly, the recruitment to LDs was enhanced in cells expressing full-length JFH1 virus (genotype 2a) with ~50% of NS5A colocalizing with TIP47 and the majority of the protein found at LDs (Figures 5D–F). This is likely due to the steatogenic properties of the viral core protein expressed in the full-length virus but not in the replicon-expressing cells [34,35]. In non-transfected cells, no significant association of TIP47 with LDs was observed, confirming previous reports and our own findings in membrane flotations and LD isolation assays as described above (Figures 4A, B, 5A) [18].

In contrast, no colocalization of Rab9 with NS5A or LDs was observed in cells infected in a single-round-infection with the Jc1 strain of HCV (Figure S3C). In naïve and transfected cells, Rab9 showed the same highly concentrated localization to one perinuclear area, likely late endosomes, that did not overlap with LDs. NS5A colocalized with LDs in perinuclear areas surrounding punctate Rab9-containing subcellular regions, but no actual colocalization between Rab9 and NS5A was observed (Figure S3C). Thus, Rab9 appears not involved in the newly described HCV cofactor role of TIP47 and supports the model that the LD-binding rather than the vesicular transport properties of TIP47 are required to support HCV RNA replication.

Discussion

In the present study, we identified TIP47 as a novel host factor that interacts with the N-terminal amphipathic α-helix of NS5A. Previous studies suggested that this protein domain, in addition to being the membrane anchor for NS5A, is also involved in protein-protein interactions that are essential for HCV RNA replication [11]. In a quantitative Y2H assay, we showed that a single point mutation (W9A) in NS5A—a previously implicated in the protein:protein interactor function of the domain—significantly decreased binding between the amphipathic α-helix and TIP47, supporting the model that this domain is a critical interaction domain with host factors in NS5A and that cellular TIP47 is a bone fide interaction partner. The W9A point mutation is located at the hydrophobic face of the helix, which is predicted to contact the cytoplasmic leaflet of the ER membrane, indicating that the membrane-binding face of the helix, rather than the polar-charged cytosolic side, mediates TIP47 interaction. The interaction was confirmed in coimmunoprecipitation assays in the context of actively replicating HCV.

Two independent sets of experiments support the model that TIP47 plays a critical role in HCV RNA replication. First, shRNA-mediated knockdown of TIP47 markedly decreased viral propagation of full-length infectious clones of HCV as well as viral RNA replication in hepatoma cells containing subgenomic replicons. The fact that a similar degree of suppression was observed in both viral systems is consistent with the notion that RNA replication is the major step in the viral life cycle affected by TIP47. However, we cannot exclude that other steps in the viral life cycle may also depend on functionally intact TIP47. These include particle assembly known to require LDs [12].

The second line of evidence supporting a critical role for the TIP47-NS5A interaction in HCV infection comes from replication studies of mutant HCV replicons. The W9A mutation we used was previously described to leave the amphipathic structure of the N-terminal α-helix in NS5A intact, but was shown to cause significant HCV RNA replication defects [11]. We confirmed and extended these studies by identifying TIP47 as a cellular factor that binds to the NS5A amphipathic α-helix in a W9-dependent manner. We also identified I12V as a compensatory mutation that restored NS5A binding and partially restored the replication deficit when introduced into W9A-mutant NS5A. Since amino acids 9 and 12 are predicted to be “neighbors” in the α-helical structure of the NS5A N-terminus (with 3.6 residues per turn), we hypothesize that both residues are important for TIP47 binding.

In both genotypes (1b and 2a), a combination of A9-I12 in NS5A was detrimental to replication, while the A9-V12 combination partially rescued replication. Why the suppressor mutation at position 12 restores binding to W9A-mutant NS5A to near wild-type levels, but only partially restores the RNA replication defect, remains unclear. A possible explanation is that W9 is involved in NS5A functions other than TIP47 recruitment and that these functions cannot be compensated by the I12V mutation.

A striking finding in our study is that LD membranes are directly linked to viral RNA replication. It has been proposed that the HCV core protein brings ER membranes in close proximity to LDs to create a localized environment with a high concentration of membranes for viral replication and assembly [12,36]. Here, we show that a close LD/ER membrane interface is also established in the absence of viral structural proteins, most notably in the absence of core. Core is known to coat the surface of LDs, a critical step for the recruitment of viral RNA-replication.
Role of TIP47 in Hepatitis C Virus Replication

A

|        | Lysate | LDs |
|--------|--------|-----|
| Huh7.5 | Replicon | Huh7.5 | Replicon |
| ADRP   |         |      |      |
| TIP47  |         |      |      |
| NS5A   |         |      |      |
| Calreticulin |      |      |      |

B

- **Huh7.5**
- **Replicon Cells (1b)**

| LDs per Cell | LD Area per Cell (μm²) |
|--------------|-------------------------|
| 150          | 250                     |
| 200          | 300                     |

C

- **TIP47**
- **NS5A**
- **Lipid Droplets**
- **Merged + DAPI**

D

- **non-transfected**
- **Replicon Cells (1b)**
- **Luc-JFH1 (2a)**

E

- **% NS5A that colocalizes with TIP47**
- **Replicon Cells**
- **Luc-JFH1**

F

- **% NS5A that colocalizes with LDs**
- **Replicon Cells**
- **Luc-JFH1**

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complexes to the vicinity of LDs, and this process is essential for the encapsidation of viral RNA into progeny virions [12–15].

The finding that viral NS3, NS5B, and NS5A proteins together with TIP47 and viral RNA access LD membranes in the absence of core points to LD membranes as critical parts of the membranous web independent from viral assembly (Figure 6). This model is supported by the finding that Rab9, a GTPase previously shown to associate with HCV replication complexes, is also newly distributed to LD membranes in HCV replicon-containing cells, underscoring the relevance of these membranes for HCV RNA replication.

How this redistribution is orchestrated and whether the LD-binding functions of TIP47 or NS5A are important for this process, remain unknown. Our experiments in cells expressing the HCV replicase containing wild-type or W9A mutant NS5A underscores that the TIP47/NS5A interaction is participating in this process. Our observation that LD content is reduced in HCV replicon cells as compared to parental cells is interesting and suggests that LDs are being consumed during HCV replication, as a potential source for membranes or energy. This may explain why the colocalization between NS5A and LDs is relatively low in replicon cells as compared to cells infected with full-length HCV as the overall LD content is reduced. As LDs are overall upregulated in cells infected with full-length HCV, this consumption is likely counterbalanced in the presence of structural genes, most notably core, to provide select assembly “platforms” to enable productive HCV virion formation. A similar “consumption” of LDs during viral infection was recently described for Dengue virus infection and involved autophagy [37].

This is the first demonstration of TIP47’s involvement in viral RNA replication and LD consumption. Previous studies have linked the host factor to assembly roles in other viral life cycles, such as HIV, Dengue, and Vaccinia virus. In HIV, TIP47 functions as a connector protein by binding the cytoplasmic domain of ENV and the matrix (MA) domain of Gag [38–40]. Similarly to the NS5A membrane anchor in our study, the TIP47-binding region in MA overlaps a hydrophobic region implicated in Gag anchoring and binding at the plasma membrane [38–40]. It is therefore possible that TIP47 binding to viral factors serves to compete with membrane anchoring of host binding partners, thus aiding in their release from membrane attachment and facilitating relocation to other membranes and/or interaction with other proteins.

In Dengue virus infection, TIP47 was shown to be required for the Dengue capsid protein to associate with LDs and promote viral assembly [41,42]. In contrast, in Vaccinia virus, the viral p37 protein was shown to associate with TIP47-, Rab9-, and MPR-containing membranes, and mutations in p37 that block association with TIP47 were shown to inhibit plaque formation [43]. These studies indicated that p37 interacts with proteins associated with late endosome-derived transport vesicles to facilitate Vaccinia viral assembly [43].

During HCV infection, Rab9 is only moderately redistributed to LD-rich, TIP47- and NS5A/NS3-containing membranes, and does not colocalize with NS5A in HCV RNA-replicating cells. However, we find that the expression levels of Rab9 are 3-fold up-regulated in HCV replicon-expressing cells, suggesting that HCV infection may overall increase endosome-to-Golgi trafficking independently from TIP47. Future studies will further characterize TIP47’s involvement in HCV infection, and the precise role of LD membranes in HCV RNA replication.

Materials and Methods

Plasmids

The N-terminal 31 amino acid–coding sequence of wild-type (genotype 1b, Con1) or W9A-mutant NS5A was cloned into the bait plasmid pBTM116 [44]. The TIP47 coding sequence was cloned into the prey plasmid pACT2 (Addgene plasmid 11343). The NS5A (genotype 1b, Con1) or W9A-mutant NS5A was cloned into the bait plasmid pBTM116 [44]. Luciferase-Jc1 (genotype 2a), Luciferase-JFH1 (genotype 2a), Jc1 [46], and HCV-Replicon (genotype 1b, Con1) [47] have been described. pGL3.5 (expression vector for firefly luciferase) was purchased from Promega. All NS5A mutant HCV-Replicon plasmids were constructed by site-directed mutagenesis of pNS5A-Flag. All mutant pNS5A-Flag plasmids are constructed by site-directed mutagenesis of pNS5A-Flag using the QuickChange II Site-Directed Mutagenesis Kit (Agilent Technologies). The expression plasmid for Myc-TIP47 was previously described [45]. Luciferase-Jc1 (genotype 2a), Luciferase-JFH1 (genotype 2a), Jc1 [46], and HCV-Replicon (genotype 1b, Con1) [47] have been described. pGL3.5 (expression vector for firefly luciferase) was purchased from Promega. All NS5A mutant HCV-Replicon plasmids were constructed by site-directed mutagenesis of pNS5A-Flag using the QuickChange II XL Site-Directed Mutagenesis Kit (Agilent Technologies). ΔE-Jc1-NS5A-HA was constructed by deleting a portion of the E1/E2 genes from amino acids 313–567 in the Jc1 polyprotein and by inserting a XbaI/Pmel linker, followed by 3xHA and 6xHis tags into domain 3 of NS5A (at amino acid position 361, as described in [48]). NS3-5B and NS3-5B W9A expression plasmids were constructed by cloning the NS3–5B region of the HCV-Replicon (Con1, genotype 1b) containing either wild-type or W9A mutant NS5A into the pEBB expression plasmid. Small hairpin RNAs targeting TIP47 (1445: GGGGCTCATTT-GAAACTTA) or non-targeting small hairpin RNAs (GCGAG-GACGGTGATCTGAAA) were cloned into pSicoR (Addgene plasmid 11579) as described in [49]. pMD2.G (Addgene plasmid 12259) pMDLg/pRRE (Addgene plasmid 12251) and pRSV-Rev (Addgene plasmid 12253) have been described [50].

Cell lines and culture conditions

Huh7 and HEK293T cells were obtained from the American Type Culture Collection, Huh7.5 cells from Charles M. Rice, and Huh7 Lunet cells from Ralf Bartenschlager. All cells were maintained in DMEM supplemented with 10% fetal bovine serum (FBS).
were grown under standard cell culture conditions and transfected with FuGENE6 (Roche) or X-tremeGENE9 (Roche), according to the manufacturer’s protocol. Calcium phosphate-mediated transfection of HEK293T cells was used for the production of lentiviral particles. Huh7 Lunet cells were used for the microscopy studies due to their superior properties for immunofluorescence microscopy [51], if not noted otherwise. Viral RNA replication and infectious virus release properties are similar in both Huh7.5 and Huh7 Lunet cell lines [52].

Antibodies and reagents
The following antibodies were obtained commercially and used as indicated (WB: Western Blot, IP: Immunoprecipitation, IF: Immunofluorescence microscopy): α-NS3 (abcam, #ab65407-100, WB), α-NS5A (Austral, #HCM-131-5, WB, IF), αNS5B (Abcam, #ab100895, WB), α-Calreticulin (Stressgen, SPA-600, WB), α-HA (Roche, 3F10, IF), α-Flag M2 (Sigma, #F-3165, IF, WB, α-Flag-HRP (Sigma, #A8592-2MG, WB), α-Flag (Sigma, #F7425, IF replicon cells), α-TIP47 (Sigma, #HPA006427, IF), α-myc-HRP (Santa Cruz, #sc-70463HRP, WB, α-HA (Roche, #3F10, IF), α-GAPDH (Abcam, #ab8245-100, WB), α-ADRP (Abcam, #ab37516, WB), α-Rab9 (Thermo Scientific, #MA3-067, WB, IF), α-Rab5 (Abcam, #ab18211, WB), α-Rab7 (Abcam, #ab50533, WB), α-mouse Alexa 488 (Invitrogen), α-rabbit Alexa 594 (Invitrogen), α-mouse Alexa 647 (Invitrogen), α-rat Alexa 647 (Invitrogen). Enzymes for molecular cloning were purchased from New England Biolabs; cell culture reagents, and Image IT-Fx-Signal Enhancer from Invitrogen; ProteinG-agarose and ProteinA-agarose from Sigma; and fine chemicals, if not noted otherwise, from Sigma.

Yeast-Two-Hybrid Screen and Quantitative Yeast-Two-Hybrid Assay
The Yeast-2-Hybrid Screen was done using the MATCHMAKER Kit (Clontech) according to the manufacturer’s recommendations. A quantitative β-galactosidase assay was performed as described in the manual provided with the Clontech MATCHMAKER kit, which we used as an indirect assessment of the strength of interaction [53]. Briefly, yeasts were transformed with either bait plasmid, prey plasmid, or both, and plated out onto the appropriate media. Individual clones were inoculated into 5 ml of appropriate liquid media and cultured overnight at 30°C with shaking, after which 2 ml of this overnight culture was used to inoculate 6 ml of YPDA liquid media (1% yeast extract, 2% bactopeptone, 2% dextrose, 2% bacto agar), and further shaken at 30°C for 4 h. Cell extracts were then prepared and assayed for β-galactosidase activity. Units are expressed in terms of ΔA₄₂₀ per minute per equivalent number of yeast cells.

Immunoprecipitation and western blotting
For the immunoprecipitation experiments, Huh7 cells were transfected with plasmid DNA using Fugene6 (Roche) or X-tremeGene9 (Roche) according to the manufacturer’s protocol. For immunoprecipitation experiments, transfected cells were lysed 48 h post-transfection in lysis buffer [150 mM NaCl, 1% NP-40, 1 mM EDTA, 50 mM Tris HCl, pH7.4 and protease inhibitor cocktail (Sigma)] for 30 min and passed 10 times through a G23 needle. Clarified lysates were pre-cleared with protein G agarose (Invitrogen) for 1 h at 4°C and after centrifugation, lysates were immunoprecipitated with protein G agarose (Invitrogen) and Flag antibody for 16 h at 4°C, washed five times in lysis buffer and resuspended in Laemmli buffer for SDS-PAGE. For the immuno-
precipitation experiments with the replicon cell line, replicon cells were harvested and then processed as described above with minor changes. Protein A agarose, and anti-TIP47 antibodies for pull-down and anti-Flag antibodies as control were used. For chemiluminescent detection, we used SuperSignal West Pico or SuperSignal West Femto (Thermo Scientific) and ECL Hyperfilm (Amersham).

**Lentivirus production, transduction and western blotting**

Lentiviral particles were produced as described [54]. Briefly, 293T cells were transfected with the pSicoR constructs, the two HIV-based packaging constructs pMD2.G/pRRE and pRSV-Rev, and a construct expressing the glycoprotein of vesicular stomatitis virus pMD2.G. The culture supernatant containing pseudotyped lentiviral particles was concentrated by ultracentrifugation for 16 h at 20,000 rpm in a SW28 rotor. Infectious titers were determined by transducing cells with serial dilutions of the viral stocks and FACS analysis 2 days post-transduction. Transductions were carried out overnight at 37°C.

For western blot analysis of lentivirus-transduced cells, cells were lysed with RIPA buffer (1% NP-40, 10 mM Tris pH7.5, 150 mM NaCl, 1 mM EDTA) for 30 min at 4°C, and mixed 1:1 with Laemmli buffer followed by SDS-PAGE.

**In vitro transcription of HCV RNA and transfection**

Plasmids encoding HCV reporter viruses were linearized with PvuII (Jc1 and JFH1 viruses) or ScaI (HCV replicon), and purified by phenol-chloroform extraction. In vitro transcription and RNA purification was carried out using the MegaScript T7 kit (Ambion), according to the manufacturer’s protocol. For RNA transfection, Huh7.5 or Huh7.5 Lunet cells were trypsinized, washed once in PBS, and resuspended in Dulbecco’s Modified Eagle’s Medium (DMEM) medium without serum at a concentration of 5 × 10^6 cells/ml. 500 μl of cell suspension was mixed with 100 μl of DMEM, 10 μg HCV RNA and 5 μg of Myc-TIP47 plasmid DNA (where indicated) and pulsed at 220 V and 975 μF with the Gene Pulser II (Biorad).

**Luciferase Assay**

After transfection with HCV RNA, Huh7.5 cells were seeded in 12-well plates. At indicated time points, the cells were treated with Passive Lysis Buffer (Promega). Luciferase activity was measured using Luciferase Assay Systems (Promega) on a MonoLight 2010 luminometer (Pegasus Scientific Inc.).

**HCV-replicon assay and HCV-replicon cell line**

1 μg of HCV replicon RNA and 1 μg of pGL3.5 (Firefly luciferase control plasmid) was transfected into Huh7.5 cells as described above. After transfection, cells were plated in 10 cm dishes for the replicon assay and into 12-well plates for the luciferase assay, and incubated at 37°C. To check for differences in transfection efficiencies, cells for the luciferase assay were lysed and luciferase activity was measured as described above. For the replicon assay, 24 h post-transfection 1 mg/ml G418 was added to the medium. Cells were kept under selective pressure for 3–4 weeks with media changed twice a week. Cells were then washed with PBS and stained with Crystal Violet. To establish an HCV replicon cell line, cells were kept under continuous G418 (800 μg/ml) pressure, and split twice a week.

**Immunofluorescence and LD staining**

Cells grown on coverslips were fixed with 4% paraformaldehyde for 10 min at room temperature, washed with phosphate-buffered saline (PBS), and permeabilized either in 0.1% saponin in PBS (for TIP47) or 0.2% Triton X-100 in PBS (for Rab9) for 10 min. After incubation with Image IT-Fx-Signal Enhancer (Invitrogen) for 30 min, cells were washed with PBS and incubated in blocking solution (DMEM with 5% FBS + 500 μg/ml goat serum) for 20 min. Cells were washed and then incubated with primary antibodies either in PBS with 1% BSA (α-TIP47) or 0.2% Triton X-100/ gelatin (α-Rab9) for 1 h, washed, and then incubated with secondary antibodies for 30 min. For oil-red-O (ORO) staining coverslips were incubated for 5 min in 60% isopropanol and stained with ORO staining solution (stock 0.5 g ORO (Sigma) in 100 ml of isopropanol; the stock was diluted 6:4 (stockwater) and filtered before use) and differentiated in 60% isopropanol for 1 sec. For BODIPY staining, fixed cells were stained for 10 min with 1 μg/ml BODIPY 493/503 (Molecular Probes) in PBS and then washed 3 times in PBS. For LipidToxRed Stain, fixed cells were stained for 30 min with LipidToxRed Neutral Stain (Molecular Probes, 1:200) in PBS, and then washed 3 times in PBS. Coverslips were embedded in Mowiol (Calbiochem) mounting medium or in VECTASHIELD mounting medium with DAPI (Vector Laboratories).

**Epifluorescence microscopy and quantification of images**

Cells were analyzed with Axio observer Z1 microscopy (Zeiss) equipped with EC Plan Neofluar 20x/0.5 PHM27, EC Plan Neofluar 40X/0.75 PH, and Plan Apo 63x/1.4 Oil DIC M27 objectives, filter sets 38HE, 43HE, 45, and 50, Optovar 1.25 and 1.6X magnification, and an AxioCAM MRM REV 3. For quantification of LDs in cells or colocalization of proteins with LDs we used the automatic measurement program of the Volocity software (Perkin Elmer).

**Membrane flotation assay**

Huh7.5 cells or HCV replicon cells (as described above) were trypsinized, washed and counted. 3 × 10^6 cells total for each sample were resuspended in 3.5 ml PBS containing 0.25 M sucrose (PBS/sucrose) plus protease inhibitor cocktail (Sigma). The cells were then lysed with 200 passages in a tight fitting dounce homogenizer (Wheaton) to ensure approximately 90% lysis. The cell lysate was then spun at 2500 × g for 10 min at 4°C to pellet cellular debris and nuclei. The resulting supernatant was referred to as crude lysate. Equal amounts of protein (5 mg) for each cell type was adjusted in volume to 2 ml with PBS/sucrose and mixed with 2 ml of 60% iodoxanol (Sigma) resulting in a 50% iodoxanol concentration. A discontinuous iodoxanol gradient (10%, 20%) was layered on top of the lysate/iodoxanol mixture, and the gradient was spun at 200,000 × g for 16 h at 4°C in a SW41T Rotor. A total of 22 fractions (500 μl each) were collected from top to bottom. The fractions were mixed 1:1 with Laemmli buffer for SDS-PAGE. Quantification of protein expression in western blots was done in ImageJ Software.

**RNA extraction and quantification by real-time RT-PCR**

20 μg of carrier RNA (Qiagen) was added to 150 ul of each fraction. RNA was extracted from each fraction using the Quick-RNA MiniPrep kit (Zymo) according to manufacturer’s recommendations. RNA was eluted using 35 μl H₂O.

cDNA was synthesized using an HCV replicon (Con1, genotypelb) antisense probe (nt 238-302; 5’-cttctggagccacataaact-3’) and AMV reverse transcriptase (Promega) followed by RNase A (Thermo Scientific) digestion. For the generation of RNA standards, cDNA was generated for a 2-fold serial dilution of in vitro transcribed HCV replicon RNAs using the above protocol.
For real-time quantitative PCR, we used HCV replicon specific primers (sense 13-30: 5'-aagcctgctagcagc-3', anti-sense 268-285: 5'-tactgctgctagcactc-3'). Real-time PCR was performed using a 2x HotSybr Real-time PCR kit (MCLAB) on a 7900HT Fast Real-time RT-PCR System (Applied Biosystems). Each sample was done in triplicates. The cDNA from the 2-fold serial dilution of HCV Replicon RNA was used to create a standard curve. Absolute amounts of viral RNA in fractions were calculated using this standard curve.

Lipid droplet isolation

Lipid droplets were isolated as described with minor changes [12]. Briefly, cells were scrapped in PBS, lysed in hypotonic buffer (50 mM HEPES, 1 mM EDTA and 2 mM MgCl2, pH 7.4) supplemented with protease inhibitors with 30 strokes in a tight-fitting Dounce homogenizer. After spinning 5 min at 1500 rpm, post-nuclear fractions were mixed with equal volumes of 1.5 M sucrose in isotonic buffer (50 mM HEPES, 100 mM KCl, 2 mM MgCl2) and placed at the bottom of SW55 Ti (Beckman) centrifuge tubes, overlaid with isotonic buffer containing 1 mM PMSF and centrifuged for 2 h at 100,000 g. Tubes were frozen at -80°C for 1 h. The top layer of floating lipid droplet fraction was cut off, and proteins of this fraction were precipitated with 15% trichloroacetic acid and 30% acetone, washed once with acetone and resuspended in urea loading dye (200 mM Tris/Cl, pH 6.8, 8 M urea, 5% SDS, 1 mM EDTA, 0.1% bromophenol blue, 15 mM DTT). 

CellTiter-Blue Cell Viability assay. The CellTiter-Blue Cell Viability Assay (Promega) was performed according to the manufacturer's recommendations. Briefly, naïve Huh7.5, and Huh7.5 containing shRNA Control or shRNA TIP47 were seeded in a 2-fold serial dilution in 96-well plates in 100 ul medium. Cells were incubated at 37°C for 1 hour, 20 ul CellTiter-Blue Reagent was added to each well, and the plates then further incubated at 37°C for 18 h. Absorbance was measured at 570 nm and 600 nm. Absorbance at 600 nm of medium only was deducted from absorbance at 570 nm of each time point.

Statistical analysis

Statistical analysis was performed using unpaired two-tailed student's t-test.

Accession numbers

The NCBI gene database (http://www.ncbi.nlm.nih.gov/gene) ID numbers for genes and transcripts in this paper are TIP47 (ID # 10226), Perilipin (ID # 5346), ADRP (ID # 123), MPR (ID # 3482), Rab9 (ID # 9367), Calectinulin (ID # 811), Rab5 (ID # 5868), Rab7 (ID # 7879).

Supporting Information

Figure S1 Localization of NS5A-Flag and W9A-NS5A-Flag in Huh7 Lunet cells. Indirect immunofluorescence of NS5A (green) in Huh7 Lunet cells transfected with DNA expression vectors for either NS5A-Flag, W9A-NS5A-Flag, or Δ31-NS5A-Flag (containing an N-terminal 31 amino acids deletion of NS5A). Lipid droplets were stained with LipidToxRed (red).

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