This is a repository copy of Serine phosphorylation of the hepatitis C virus NS5A protein controls the establishment of replication complexes.

White Rose Research Online URL for this paper:
http://eprints.whiterose.ac.uk/83736/

Version: Published Version

Article:

https://doi.org/10.1128/JVI.02995-14

Reuse
Unless indicated otherwise, fulltext items are protected by copyright with all rights reserved. The copyright exception in section 29 of the Copyright, Designs and Patents Act 1988 allows the making of a single copy solely for the purpose of non-commercial research or private study within the limits of fair dealing. The publisher or other rights-holder may allow further reproduction and re-use of this version - refer to the White Rose Research Online record for this item. Where records identify the publisher as the copyright holder, users can verify any specific terms of use on the publisher’s website.

Takedown
If you consider content in White Rose Research Online to be in breach of UK law, please notify us by emailing eprints@whiterose.ac.uk including the URL of the record and the reason for the withdrawal request.
Serine Phosphorylation of the Hepatitis C Virus NS5A Protein Controls the Establishment of Replication Complexes

Douglas Ross-Thriepland,* Jamel Mankouri, Mark Harris
School of Molecular and Cellular Biology, Faculty of Biological Sciences, and Astbury Centre for Structural Molecular Biology, University of Leeds, Leeds, United Kingdom

ABSTRACT
The hepatitis C virus (HCV) nonstructural 5A (NS5A) protein is highly phosphorylated and involved in both virus genome replication and virion assembly. We and others have identified serine 225 in NS5A to be a phosphorylation site, but the function of this posttranslational modification in the virus life cycle remains obscure. Here we describe the phenotype of mutants with mutations at serine 225; this residue was mutated to either alanine (S225A; phosphoablatant) or aspartic acid (S225D; phosphomimetic) in the context of both the JFH-1 cell culture infectious virus and a corresponding subgenomic replicon. The S225A mutant exhibited a 10-fold reduction in genome replication, whereas the S225D mutant replicated like the wild type. By confocal microscopy, we show that, in the case of the S225A mutant, the replication phenotype correlated with an altered subcellular distribution of NS5A. This phenotype was shared by viruses with other mutations in the low-complexity sequence I (LCS I), namely, S229D, S232A, and S235D, but not by viruses with mutations that caused a comparable replication defect that mapped to domain II of NS5A (P315A, L321A). Together with other components of the genome replication complex (NS3, double-stranded RNA, and cellular lipids, including phosphatidylinositol 4-phosphate), the mutation in NS5A was restricted to a perinuclear region. This phenotype was not due to cell confluence or another environmental factor and could be partially transcomplemented by wild-type NS5A. We propose that serine phosphorylation within LCS I may regulate the assembly of an active genome replication complex.

IMPORTANCE
The mechanisms by which hepatitis C virus replicates its RNA genome remain poorly characterized. We show here that phosphorylation of the viral nonstructural protein NS5A at serine residues is important for the efficient assembly of a complex that is able to replicate the viral genome. This research implicates cellular protein kinases in the control of virus replication and highlights the need to further understand the interplay between the virus and the host cell in order to develop potential avenues for future antiviral therapy.

Hepatitis C virus (HCV) currently infects an estimated 170 million individuals worldwide and in 85% of cases establishes a chronic infection, which typically progresses to liver cirrhosis and hepatocellular carcinoma (1). The virus has a single-stranded RNA genome that codes for an ~3,000-amino-acid polyprotein that is cleaved co- and posttranslationally into 10 mature viral proteins: the core and the E1 and E2 envelope proteins, which make up the viral particle, followed by the viroporin p7 and the nonstructural 2 (NS2) protein, which possesses autoprotease activity. The remaining nonstructural (NS) proteins, NS3, NS4A, NS4B, NS5A, and NS5B, are necessary and sufficient for genome replication (2). With the advent of the full-length clone of a geno-type 2a isolate (JFH-1) able to undergo the complete virus life cycle in cell culture, significant progress has been made in understanding how the different viral proteins contribute to the processes of genome replication and virus assembly. In this regard, NS5A has been shown to play a critical role in both of these processes and is known to perturb numerous host pathways in favor of virus persistence.

The 5’ untranslated region (UTR) of the viral genome contains an internal ribosome entry sequence (IRES) that allows viral proteins to be translated in a cap-independent fashion immediately after the viral genome is delivered into the cytoplasm. HCV proteins then rapidly recruit and remodel endoplasmic reticulum-derived membranes to form a membranous web (MW), a subcellular structure comprised of single-membrane, double-membrane, and multimembrane vesicles (SMVs, DMVs, and MMVs, respectively) that are enriched in viral (e.g., NS3, NS5A, and NS5B) and host cell (e.g., VAP-A) proteins (3, 4). By analogy to other positive-sense RNA viruses, this enrichment of viral NS proteins, the presence of double-stranded RNA (dsRNA), and the observation that isolated membrane fractions retain replication activity have led to the proposal that the MW is the site of viral genome replication (3). In Huh7 cells, the MW is extensively distributed throughout the cytoplasm, correlating with the observed subcellular distribution of the NS5A and NS3 proteins as discrete...
puncta throughout the cytoplasm. The formation and mainte-
nance of the MW have been shown to be dependent on the ac-
tivation of the phosphatidylinositol 4-kinase (PI4K) type III alpha
(PI4KIIIα) isoform by NS5A and the subsequent elevated produc-
tion of phosphatidylinositol 4-phosphate (PI4P) by this lipid ki-
rase (5–7). Additionally, lipid droplets (LDs) are a host organelle
responsible for the normal cellular storage of neutral lipids and
have been shown to be central to the HCV life cycle. While the site
of virus assembly has not been elucidated, it is known that both the
core and NS5A proteins coat the surface of LDs, displacing the
host factor adipocyte differentiation-related protein (ADRP). In
order for virus genomes that have been newly synthesized in the
MW to be packaged into virions, the core protein-coated LDs are
thought to be recruited to the MW, possibly in an NS5A-directed
manner, in order to allow the encapsidation of viral RNA (8, 9).

NS5A is a highly phosphorylated protein comprised of three
domains and tethered to membranes by an N-terminal amphi-
pathic helix. Domain I is highly structured and sufficient for NS5A
to dimerize, while domains II and III are intrinsically disordered
but have recently been shown to have elements of transient sec-
ondary structure throughout (10–12). These three domains are
linked by low-complexity sequences (LCSs) (13); a serine-rich
sequence (LCS I) links the first two domains, while a proline-rich
sequence (LCS II) links the last two domains. NS5A exists in two
different phosphorylated forms that can be readily resolved by
SDS-PAGE. These forms, termed the basal phosphorylated and
hyperphosphorylated species, are thought to represent different
functional pools of NS5A involved in different stages of the virus
life cycle, but as yet there is little direct evidence to support this
hypothesis. Until recently, the sites of NS5A phosphorylation had
not been unambiguously mapped, their locations having been in-
fected from mutagenesis data. Recently, however, we and others
have used mass spectrometry to demonstrate that NS5A, purified
from cells actively replicating the HCV subgenomic replicon
(SGR), is extensively phosphorylated within LCS I, containing at a
minimum seven phosphorylation sites (14–16). We also showed
that NS5A contains phosphorylation sites in domain I, domain II,
and LCS II. Previous work to characterize the role of LCS I phos-
phorylation has focused almost entirely on a role in virus replica-
tion and/or assembly/release, and numerous studies have shown
several serines to be important for viral replication, but in a geno-
type-dependent manner. These studies have, however, not ex-
plored the mechanism through which the phosphorylation of LCS
I might be acting.

We show here that the phosphorylation of serine 225 within
LCS I is critical for the correct localization of viral proteins during
infection. Ablation of serine 225 phosphorylation results in a 10-
fold reduction in replicative fitness. We show using confocal mi-
roscope that this phenotype is concomitant with a dramatic re-
localization of both cellular and viral factors known to participate
in genome replication. This phenotype is shared with mutants
with other serine mutations in LCS I, further illustrating the com-
plexity of phosphorylation in the region of NS5A.

MATERIALS AND METHODS

Plasmids. DNA constructs of and the full-length pFH-1 virus (17) and
the luciferase subgenomic replicon (SGR-luc-JFH-1) (18) were used
throughout the study. Previously, the unique restriction sites flanking
NS5A, BamHI and AfeI, were introduced into both the full-length virus
and SGR-luc-JFH-1 constructs; these constructs are denoted mFH-1 and
mSGR-luc-JFH-1, respectively, and have shown been to have no effect on
virus genome replication or assembly and release (19). Site-directed mu-
tagenesis of NS5A was performed with a QuiKChange kit (Stratagene) and
a LITMUS28i (NEB) subclone containing an NsiI/HindIII fragment of the
mFH-1 cDNA. Then, mutant NS5A was cloned into either mSGR-luc-
JFH-1 or mFH-1 via flanking BamHI/AfeI sites, and correct insertion was
confirmed by sequencing. For the insertion of the SNAP and CLIP tags
into NS5A, the gene was amplified from the pSNAPI and pCLIPf vectors
(NEB) using primers that insert a flexible linker sequence (GSGSS) on
either side of the insertion (primer sequences are available upon request).
This amplicon was inserted into the aforementioned LITMUS28i sub-
clone by BclI digestion, and correct insertion was verified by sequencing.
The NS5A-SNAP or NS5A-CLIP region was then inserted into mSGR-
luc-JFH-1 via flanking RsrII/AfeI sites. For plasmid expression, mutated
NS5A-coding sequences were subcloned into the pCMV (NS3-5B) vector
as described previously (20). All mutations were verified by sequencing.

Cell culture. Huh7 cells were cultured in Dulbecco’s modified Eagle’s
medium (DMEM; Sigma) supplemented with 10% fetal bovine serum
(FBS), 100 IU penicillin/ml, 100 µg streptomycin/ml, and 1% nonessen-
tial amino acids in a humidified incubator at 37°C with 5% CO2.

Electroporation of replicon and virus RNA constructs. The prepara-
tion of in vitro transcripts and electroporations for both mSGR-luc-JFH-1
and mFH-1 were conducted as described previously (20). In brief, 2 × 106
Huh7 cells in diethyl pyrocarbonate (DEPC)–phosphate-buffered sa-
line (PBS) were electroporated with 5 µg of in vitro RNA transcripts using
a square-wave protocol at 260 V for 25 ms. Subsequently, cells were re-
suspended in complete DMEM and seeded at 1 × 104 cells/cm2 culture
area in either 12- or 6-well dishes.

Virus titration. Supernatants were titrated on Huh7 cells in a 96-well
format as follows. Clarified virus supernatant was serially diluted 2-fold in
complete DMEM before addition of 100 µl to cells that had been seeded 8
h earlier at 8 × 105 cells/well (final volume, 200 µl). The counting of
infected cells was automated by acquisition and analysis using an
Incucyte ZOOM platform. The virus titer was determined by averaging
the positive cell count from 3 or more adjacent counted wells that gave
the corresponding titers.

SDS-PAGE/Western blotting. Cells were washed twice in PBS, lysed
in 1× Glasgow lysis buffer (GLB; 1% [vol/vol] Triton X-100, 120 mM
KCl, 30 mM NaCl, 3 mM MgCl2, 10% [vol/vol] glycerol, 10 mM Pipes
[piperazine-N,N’-bis(2-ethanesulfonic acid)]-NaOH, pH 7.2, with pro-
tease and phosphatase inhibitors), and clarified by centrifugation at
2,800 × g for 5 min at 4°C, before determining and normalizing the
protein concentration by a bicinchoninic acid assay (Pierce). Following
separation by SDS-PAGE, proteins were transferred to a polyvinyliden-
difluoride (PVDF) membrane and blocked in 50% (vol/vol) Odyssey
blocking (OB) buffer (LI-COR) in PBS. The membrane was incubated
with primary antibodies overnight at 4°C, followed by incubation with
secondary antibodies for 2 h at room temperature (RT); both primary
and secondary antibodies were prepared in 50% OB buffer. The primary
antibodies used were anti-NS5A (sheep, 1:5,000) (21), anti-NS3 (sheep,
1:2,000; prepared in-house), and anti-glyceraldehyde-3-phosphate dehy-
drogenase (anti-GAPDH; mouse, 1:2,000; Sigma). Secondary antibodies
were anti-goat (800 nm) or anti-mouse (700 nm), which were used at
1:5,000, prior to imaging fluorescence using a LI-COR Odyssey 6 infra-
red imaging system. Quantification of fluorescently labeled Western blots
was carried out using Image Studio (v3.1) software (LI-COR) and a back-
ground subtraction method.

Immunofluorescence and confocal microscopy. Cells were washed
once in PBS before fixation for 20 min in 4% (wt/vol) paraformaldehyde
(PFA). The cells were subsequently permeabilized in 0.2% (vol/vol) Tri-
ton X-100–PBS before immunostaining with the antibody denoted be-
low. The primary antibodies used were anti-NS5A (sheep, 1:1,000), anti-
NS3 (sheep, 1:300; prepared in-house), anti-dsRNA (mouse, 1:200;
Scicons), and anti-PI4P (mouse, 1:100; Echelon). Various fluorescently
conjugated secondary antibodies were used at 1:1,000 (Life Technology).
Phenotype of NS5A Serine Phosphorylation

Note that when conducting immunostaining for dsRNA, all reagents were prepared to be RNase free by DEPC treatment. Lipid droplets were stained using the BODIPY(558/568)-C12 dye at 1:1,000 (Life Technology), which was added at the same time as the fluorescent secondary antibody. Confocal microscopy images were acquired on a Carl Zeiss LSM 700 inverted microscope, and postacquisition analysis was conducted using either Zeiss Zen 2012 or Fiji (v1.49) software (22).

In-cell labeling of the SNAP/CLIP tag. HuH7 cells harboring SNAP/CLIP replicons were fixed in 4% PFA and permeabilized in 0.2% Triton X-100–PBS to which 0.9 mM CaCl$_2$ and 0.5 mM MgCl$_2$ were added (PBS$^{++}$). The SNAP and CLIP substrates were dissolved to 1 mM in dimethyl sulfoxide, before a 1 μM SNAP/CLIP labeling solution was prepared in PBS$^{++}$ containing 10% (vol/vol) FBS and added to cells for 2 h at RT. For immunostaining of NS5A, antibody was added to SNAP/CLIP labeling solution, followed by secondary antibody staining, as detailed above.

Quantification of lipid droplets. For quantification of the LD spatial arrangement, images were acquired by use of the same acquisition parameters described above, but a variable gain was used to ensure the correct exposure. The spatial coordinates of the LDs were determined using the FindFoci function of the GDSC plug-in for Fiji software, with the nuclear envelope being manually outlined (utilizing DAPI [4',6-diamidino-2-phenylindole] staining as a reference), and the coordinates were generated by Fiji. The distance from each lipid droplet to the nuclear envelope was then determined using trigonometry. LD spatial distribution data were generated for 12 randomly selected cells for each JFH-1 virus variant, and data were combined into a box-and-whisker plot (see Fig. 3E).

Statistics. Data sets were analyzed using Student’s t test, assuming a two-tailed, unequal variance to determine statistically significant differences from the results for the wild type (WT) ($n = 3$ or greater throughout).

RESULTS
Phosphorylation of serine 225 is important for genome replication but not assembly of infectious virus. Previously, we identified serine 225 to be one of a number of serines within the LCS I of NS5A that is phosphorylated during the normal course of infection. To investigate the role of serine 225 phosphorylation in the virus life cycle, we generated either a phosphoablatant alanine substitution or a phosphomimetic aspartic acid substitution at this residue in the context of both the mSGR-luc-JFH-1 and mJFH-1 constructs by site-directed mutagenesis, as described previously (16). HuH7 cells were then electroporated with in vitro transcribed of mSGR-luc-JFH-1, and genome replication was followed over 72 h by determination of luciferase activity. As seen previously (16), the phosphoablatant mutation S225A resulted in a 10-fold reduction in genome replication efficiency, while the phosphomimetic mutation S225D had no discernible effect (Fig. 1A). To investigate whether the phosphorylation of serine 225 had additional effects on the release of infectious virus, in vitro transcripts of the mJFH-1 virus with the S225A/D mutations were electroporated into HuH7 cells, and the titer of the released virus was followed over 144 h by a focus-forming assay. Figure 1B shows that the S225A mutant virus had about a 10-fold reduction in infectious virus compared to the WT, while the S225D mutant virus had titers indistinguishable from those of the wild type. This reduction in the titer of released S225A mutant virus correlated with the impairment of replication, in line with the findings of previous studies showing that genome replication is the rate-limiting step in the release of infectious virus (23). These data illustrate that, as previously concluded (16), phosphorylation of serine 225 is important for viral genome replication but not required for the release of infectious virus.

Cell lysates from infected cells were collected at 96 and 144 h postelectroporation (hpe), normalized for protein concentration, and analyzed by SDS-PAGE/Western blotting. It was found that there was a reduction in the level of cellular NS5A to a degree comparable to the reduction in the level of virus replication (Fig. 1C); quantification of the Western blot confirmed this (Fig. 1D). However, when the ratio of hyperphosphorylation to basal phosphorylation was determined, it was found that the S225A mutant exhibited a reduction in the extent of hyperphosphorylation at both early and late time points (Fig. 1D). This correlated with the involvement of the LCS I in hyperphosphorylation previously reported by us and others (14–16).

It was noted that when the protein concentration was normalized for SDS-PAGE, cells infected with the S225A mutant virus were growing at a higher rate than wild-type virus- and S225D mutant virus-infected cells (data not shown). To account for this, the titer of the released virus at 96 and 144 hpe was normalized to the cellular protein concentration, used here as a surrogate for cell number, and it was observed that the S225A mutant still exhibited an approximately 10-fold reduction in the amount of virus released compared to that for the wild type (Fig. 1E). To ensure that cell confluence (known to negatively affect HCV replication) was not affecting the titers of the released virus, electroporated cells were also passaged at 96 hpe, and at 144 hpe, the titer of the released virus was determined and normalized to the cellular protein concentration (Fig. 1E). As noted above, the S225A mutant exhibited a 10-fold reduction in the amount of virus released compared with that for the wild type.

Ablation of serine 225 phosphorylation alters the subcellular distribution of NS5A. When virus was titrated by a focus-forming assay, the subcellular distribution of NS5A with the S225A mutation [NS5A(S225A)] was observed to be markedly different from that of the wild-type protein. Typically in wild-type JFH-1-infected cells, NS5A was present throughout the cytoplasm in discrete puncta; however, in cells infected with the S225A mutant virus, NS5A localized into defined, condensed structures. To investigate this further, HuH7 cells were electroporated with in vitro transcripts of mJFH-1 virus (the wild type and the S225A and S225D mutants), fixed at 96 hpe, and immunostained for NS5A as previously (16) (Fig. 1F). NS5A localized into distinct puncta; however, in cells infected with the S225A mutant virus, NS5A localized into defined, condensed structures. To investigate this further, HuH7 cells were electroporated with in vitro transcripts of mJFH-1 virus, and genome replication was followed over 72 h by determination of luciferase activity. As seen previously (16), the phosphoablatant mutation S225A resulted in a 10-fold reduction in genome replication efficiency, while the phosphomimetic mutation S225D had no discernible effect (Fig. 1A). To investigate whether the phosphorylation of serine 225 had additional effects on the release of infectious virus, in vitro transcripts of the mJFH-1 virus with the S225A/D mutations were electroporated into HuH7 cells, and the titer of the released virus was followed over 144 h by a focus-forming assay. Figure 1B shows that the S225A mutant virus had about a 10-fold reduction in infectious virus compared to the WT, while the S225D mutant virus had titers indistinguishable from those of the wild type. This reduction in the titer of released S225A mutant virus correlated with the impairment of replication, in line with the findings of previous studies showing that genome replication is the rate-limiting step in the release of infectious virus (23). These data illustrate that, as previously concluded (16), phosphorylation of serine 225 is important for viral genome replication but not required for the release of infectious virus.

Cell lysates from infected cells were collected at 96 and 144 h postelectroporation (hpe), normalized for protein concentration, and analyzed by SDS-PAGE/Western blotting. It was found that there was a reduction in the level of cellular NS5A to a degree comparable to the reduction in the level of virus replication (Fig. 1C); quantification of the Western blot confirmed this (Fig. 1D). However, when the ratio of hyperphosphorylation to basal phosphorylation was determined, it was found that the S225A mutant exhibited a reduction in the extent of hyperphosphorylation at both early and late time points (Fig. 1D). This correlated with the involvement of the LCS I in hyperphosphorylation previously reported by us and others (14–16).

It was noted that when the protein concentration was normalized for SDS-PAGE, cells infected with the S225A mutant virus were growing at a higher rate than wild-type virus- and S225D mutant virus-infected cells (data not shown). To account for this, the titer of the released virus at 96 and 144 hpe was normalized to the cellular protein concentration, used here as a surrogate for cell number, and it was observed that the S225A mutant still exhibited an approximately 10-fold reduction in the amount of virus released compared to that for the wild type (Fig. 1E). To ensure that cell confluence (known to negatively affect HCV replication) was not affecting the titers of the released virus, electroporated cells were also passaged at 96 hpe, and at 144 hpe, the titer of the released virus was determined and normalized to the cellular protein concentration (Fig. 1E). As noted above, the S225A mutant exhibited a 10-fold reduction in the amount of virus released compared with that for the wild type.

Ablation of serine 225 phosphorylation alters the subcellular distribution of NS5A. When virus was titrated by a focus-forming assay, the subcellular distribution of NS5A with the S225A mutation [NS5A(S225A)] was observed to be markedly different from that of the wild-type protein. Typically in wild-type JFH-1-infected cells, NS5A was present throughout the cytoplasm in discrete puncta; however, in cells infected with the S225A mutant virus, NS5A localized into defined, condensed structures. To investigate this further, HuH7 cells were electroporated with in vitro transcripts of mJFH-1 virus (the wild type and the S225A and S225D mutants), fixed at 96 hpe, and immunostained for NS5A as previously (16) (Fig. 1F). NS5A localized into distinct puncta; however, in cells infected with the S225A mutant virus, NS5A localized into defined, condensed structures. To investigate this further, HuH7 cells were electroporated with in vitro transcripts of mJFH-1 virus, and genome replication was followed over 72 h by determination of luciferase activity. As seen previously (16), the phosphoablatant mutation S225A resulted in a 10-fold reduction in genome replication efficiency, while the phosphomimetic mutation S225D had no discernible effect (Fig. 1A). To investigate whether the phosphorylation of serine 225 had additional effects on the release of infectious virus, in vitro transcripts of the mJFH-1 virus with the S225A/D mutations were electroporated into HuH7 cells, and the titer of the released virus was followed over 144 h by a focus-forming assay. Figure 1B shows that the S225A mutant virus had about a 10-fold reduction in infectious virus compared to the WT, while the S225D mutant virus had titers indistinguishable from those of the wild type. This reduction in the titer of released S225A mutant virus correlated with the impairment of replication, in line with the findings of previous studies showing that genome replication is the rate-limiting step in the release of infectious virus (23). These data illustrate that, as previously concluded (16), phosphorylation of serine 225 is important for viral genome replication but not required for the release of infectious virus.
virus, LDs clustered close to the nucleus, whereas in the context of either wild-type or S225D mutant infection, the LDs were more evenly distributed throughout the cytoplasm. In uninfected cells, the distribution of LDs more closely resembled that in wild-type- or S225D mutant-infected cells (Fig. 3D).

To quantify this LD redistribution, we acquired images of 12 randomly selected cells for the wild type and each virus mutant and quantified both the number of LDs and their distance from the nuclear membrane. With this quantitative approach, we saw that in comparison to uninfected cells, wild-type virus-infected cells exhibited a broader distribution throughout the cytoplasm, whereas in the context of the S225A mutation, there was a significant shift of LDs toward the nucleus (Fig. 3E). Interestingly, however, cells infected with virus with the S225D mutation had an intermediate phenotype that resembled that of uninfected cells, where there was a slight shift of LDs toward the nucleus, indicating that this mutation was not quite fully able to recapitulate the effect of phosphorylation at serine 225, at least with respect to the LD distribution (Fig. 3E). With these data, we were also able to interrogate the total number of LDs per cell. There was about a 2-fold increase in the number of LDs in cells infected with either wild-type virus or the S225D mutant viruses compared to the number in naive cells (Fig. 3E). This was not the case for the S225A mutant, where the number of LDs per cell for S225A mutant-infected cells was broadly comparable to that for uninfected cells (Fig. 3E), suggesting that NS5A(S225A) is unable to increase the formation of LDs upon infection.

The viral protease/helicase NS3 is another key component of genome replication complexes; therefore, to further investigate the potential redistribution of these complexes caused by the S225A mutation, we costained infected cells with antibodies to both NS5A and NS3. As expected, NS3 extensively colocalized with NS5A in all cases, such that in the context of NS5A(S225A) it exhibited a restricted perinuclear distribution, whereas for both wild-type NS5A and NS5A with the S225D mutation [NS5A (S225D)], NS3 was extensively distributed throughout the cytoplasm (Fig. 4A). A similar phenotype was observed for the distribution of double-stranded RNA (dsRNA), a hallmark of viral genome replication that can be detected using a specific antibody, J2 (24). As shown in Fig. 4B, in cells infected with mJFH-1, the dsRNA antibody detected small, bright puncta, typically of a uniform size and with a distribution throughout the cytoplasm. In
Lastly, we investigated the distribution of a host cell factor, the phospholipid phosphatidylinositol 4-phosphate (PI4P). It has been well established that infection by HCV and many other RNA viruses results in the increased production of PI4P lipids, and it is widely accepted that this is one of the mechanisms through which HCV is able to induce the formation of the membranous web (5–7). Using an antibody specific to PI4P, we showed that in cells infected with wild-type or S225D mutant viruses, PI4P levels were elevated in comparison to those in uninfected cells (data not shown) and that there was a strong colocalization between NS5A and PI4P, with PI4P typically forming small irregular puncta (Fig. 4C). In contrast, cells infected with the S225A mutant virus showed a dramatic reduction in the abundance of PI4P puncta and their relocation to a perinuclear distribution along with NS5A (Fig. 4C).

From the findings of this confocal microscopy and immunofluorescence analysis, we conclude that the phosphoablative mutation S225A in NS5A is sufficient to induce the relocation of numerous factors associated with HCV genome replication. It is important to note that we did not observe a loss of colocalization of NS5A with these factors, reflecting the fact that NS5A(S225A) retained the ability to replicate and release infectious virus.

The relocalization of NS5A(S225A) and viral/host factors is not a result of reduced HCV replication. It was important to establish that the phenotype of reduced HCV replication was specifically the result of the loss of phosphorylation at serine 225 and not the result of the lower levels of replication (and, so, viral proteins) that are observed with virus containing NS5A with the S225A mutation. To achieve this, we utilized two previously characterized mutations within domain II of NS5A, P315A and L321A, that had previously been shown to impair (approximately 10-fold) virus replication similarly to S225A (20).

As expected, an approximately 10-fold reduction in the amount of virus that was released was exhibited for both the mutant virus containing NS5A with the P315A mutation [NS5A (P315A)] (Fig. 5A) and the mutant virus containing NS5A with the L321A mutation [NS5A(L321A)] (data not shown); this amount was comparable to that achieved with virus with the S225A mutation, confirming that the S225A, P315A, and L321A mutations impair virus replication to similar levels. Both mutant viruses exhibited a reduction in the levels of NS5A expression similar to that seen for virus with the S225A mutation (Fig. 5B and C and data not shown) and exhibited a ratio of NS5A hyperphosphorylation to basal phosphorylation similar to that for the wild type (Fig. 5C and data not shown) but different from that for both S225 mutants, consistent with the previously proposed notion that S225 is a site of phosphorylation in the hyperphosphorylated species of NS5A (16). Therefore, if the phenotype of NS5A relocalization was the result of reduced replication, it would be expected that both the P315A and L321A mutants would show the same phenotype.

Huh7 cells infected with mutant viruses with the NS5A (P315A) (Fig. 5D) or NS5A(L321A) (data not shown) mutation were analyzed by confocal microscopy for the abundance and distribution of NS5A and LDs. Reassuringly, the distribution of NS5A in cells infected with these mutants was indistinguishable from that in cells infected with the wild type, with no evidence of the restriction to a perinuclear distribution. Quantitative analysis revealed no significant difference between the wild type and the two mutants with regard to the distance of the LDs from the nu-
clear membrane (Fig. 5E). Similar results were seen for the distribution of NS3 and PI4P (Fig. 5D), as well as for that of dsRNA (data not shown). It is noteworthy that although the distribution of PI4P lipids in NS5A(P315A) virus-infected cells was similar to that in wild type-infected cells, there was an obvious reduction in their quantity, suggesting that this mutation might have an independent effect on NS5A-mediated PI4K activation. This observation notwithstanding, we conclude that the effect of the S225A mutation on the distribution of replication complex components cannot be explained by a reduction in replicative fitness; rather, it is specifically the loss of phosphorylation at serine 225 that is responsible for this phenotype.

The S225A mutant phenotype does not correlate with a loss of hyperphosphorylation. Although our original observation was made during routine virus titration experiments, we had not screened an entire panel of mutants with serine mutations within LCS I. Given the complexity of the phosphorylation patterns within this region and the potential role of S146 in regulating hyperphosphorylation (16), we therefore generated a complete panel of mutants with either phosphomimetic or phosphoablative mutations in all 8 serines in LCS I and S146 and examined their phenotypes by both virus titration and immunofluorescence. Consistent with the findings of our previous analysis (16) and those of Masaki et al. (15), most mutations had no effect on virus titer; however, the S225A, S232A, and S235D mutant viruses all exhibited a significant reduction in titer (approximately 10-fold) (Fig. 6A). Cells infected with the S232A and S235D mutants also shared the restricted perinuclear distribution of NS5A and LD

FIG 3 Analysis of lipid droplet redistribution in infected cells. Huh7 cells were electroporated with in vitro transcripts of either wild-type mJFH-1 (A) or the NS5A(S225A) (B) or NS5A(S225D) (C) mutant or were not infected (D), and cells were seeded onto coverslips and incubated for 48, 72, and 96 hpe prior to fixation, immunostaining for NS5A and LDs, and imaging by confocal microscopy. Uninfected control cells were also processed at 96 h postseeding. Spatial data for LDs were determined from 12 cells for each virus using the GDSC plug-in for Fiji. (E) These data were used to determine the number of LDs per cell as well as the distance of each LD from the nuclear envelope. Box-and-whisker plots show the 10 to 90% distribution. ***, significant difference (P < 0.05) from the results for the wild type.
FIG 4 Colocalization of NS5A phosphorylation mutants with host and viral factors. Huh7 cells were electroporated with in vitro transcripts of either wild-type mJFH-1 or the NS5A(S225A) or NS5A(S225D) mutant and seeded onto coverslips. At 96 hpe, cells were fixed and immunostained for NS5A and either NS3 (A), dsRNA (B), or the phospholipid PI4P (C). An NS5B GND (nonreplicating) control was included for the dsRNA staining. The colocalization scatter plots on the right are of an area of 200 by 200 μm containing >10 cells and show the correlation of absolute values (x and y axes, pixel numbers). The correlation value (r) is shown for each image.
As demonstrated previously (15), the S229D mutation did not result in the production of any infectious virus; however, the S229D mutant exhibited limited RNA replication and protein expression (15, 16), and the distribution of both NS5A and LDs in S229D mutant-infected cells was again restricted to a perinuclear region. We conclude that the phosphorylation of different serines within LCS I can have opposing effects on the subcellular distribution of NS5A.

Lastly, we examined the phenotype of a double mutant, the S146D and S225D mutant, that we had previously shown replicated as well as the wild type but in which hyperphosphorylation was almost completely abolished (16). The distribution of NS5A in cells infected with this mutant was comparable to that in cells infected with the wild type (see Fig. S1 in the supplemental material), confirming that the phenotype of a restricted perinuclear distribution for the S225A mutant could not be explained by a defect in hyperphosphorylation. As an aside, none of the mutants (including the S225A mutant) exhibited any significant differences in sensitivity to daclatasvir (DCV) from that of the wild type. The 50% effective concentrations (EC50s) ranged from 10 to 57 pM (in our hands, a wild-type JFH-1 replicon had an EC50 of 57 pM), and although treatment with DCV reduced the abundance of NS5A, it did not alter the widespread cytoplasmic distribution (data not shown).

The phenotype of the S225A mutant is not the result of cell culture conditions. To enable us to interrogate further the phenotype of the S225A mutant, we utilized the SNAP and CLIP tag (NEB) technology. This complementary tag system allows the specific conjugation of either benzylguanine or benzylcytosine derivatives, typically fluorescently labeled, to the SNAP or CLIP tag, respectively. This approach has recently been described for NS5A (25), and we reasoned that it would enable us to differentially label wild-type NS5A and the S225A and S225D mutants within the same population of cells. The advantage of the SNAP and CLIP tags over the traditional green fluorescent protein (GFP) and red fluorescent protein tags is a combination of the flexibility offered with the SNAP/CLIP system and the high sequence similarity between the SNAP and CLIP tags.

The SNAP/CLIP tags were initially introduced into the wild-type mSGR-luc-JFH-1 replicon (19) at position 431 in NS5A (domain III), as this site has previously been shown to tolerate large insertions (Fig. 7A). We confirmed that the insertion of either tag

![Fig 5](https://example.com/fig5.png)

**Fig 5** An NS5A domain II mutation that impairs replication does not have the same redistribution phenotype as NS5A(S225A). Huh7 cells were electroporated with in vitro transcripts of either wild-type mJFH-1 or the NS5A(S225A), NS5A(S225D), or NS5A(P315A) mutant. (A) The titer of the released virus was determined at 96 hpe by a focus-forming assay. The virus titer was normalized to the cellular protein concentration in order to account for differences in cell growth rates. (B) At 96 hpe, cells were lysed and analyzed by SDS-PAGE/Western blotting for NS5A and GAPDH. (C) Total NS5A levels and the ratio of hyperphosphorylation to basal phosphorylation were quantified from fluorescent Western blots. (D) At 96 hpe, electroporated cells were fixed and immuno-stained for NS5A and either LDs, PI4P, or NS3, as described in the legend to Fig. 4. (E) Spatial data for LDs were determined from 12 cells for each virus using the GDSC plug-in for Fiji. Box-and-whisker plots show the 10 to 90% distribution. ***, significant difference (P < 0.05) from the results for the wild type.
had no discernible effect on the replication kinetics of the mSGR-luc-JFH-1 replicon compared to those of the untagged wild-type replicon (Fig. 7B). As the SNAP and CLIP tags share very high sequence similarity yet retain different substrate specificities, it was important to establish that the insertion of the tag at an internal position did not introduce conformation changes that affected the substrate specificity. As such, cells harboring the mSGR-luc-JFH-1-NS5A-SNAP (NS5A-SNAP) replicon were fixed at 72 hpe, immunostained for NS5A, and labeled with a CLIP-tetramethyl-rhodamine (TMR) fluorophore. The converse, where the mSGR-luc-JFH-1-NS5A-CLIP (NS5A-CLIP) replicon was labeled with the SNAP-TMR fluorophore, was also carried out. Figure 7D

FIG 6 Mutants with other serine mutations in LCS I share the condensed redistribution phenotype with the S225A mutant. (A) Huh7 cells were electroporated with in vitro transcripts of either wild-type mJFH-1 or the indicated mutants. The titer of released virus was determined at 96 hpe by a focus-forming assay. (B) At 96 hpe, electroporated cells were fixed and immunostained for NS5A and LDs, as described in the legend to Fig. 3. Only the results for the three mutants that shared the phenotype are shown here for comparison with the wild type. See Fig. S1 in the supplemental material for the complete data set. (C) Summary table of mutant phenotypes. n.a., not available (did not replicate). (D) Amino acid sequence of LCS I. *, serine residues with a redistribution phenotype; DI, DII, and DIII, domains I, II, and III, respectively.
shows that both the SNAP and CLIP tags retained both catalytic activity and specificity toward the appropriate SNAP/CLIP substrate, with SNAP-TMR labeling only NS5A-SNAP and CLIP-TMR labeling only NS5A-CLIP. The corresponding immunostaining for NS5A also showed a very high degree of correlation, confirming that there was no subpopulation of either NS5A-SNAP or NS5A-CLIP that was not labeled with the SNAP/CLIP substrates.

We proceeded to use the SNAP/CLIP system to establish whether there was a hitherto unknown environmental factor that was inducing the phenotype of the S225A mutant. To do this, we generated mSGR-luc-JFH-1-NS5A(S225A)-SNAP [NS5A(S225A)-SNAP] and mSGR-luc-JFH-1-NS5A(S225D)-CLIP [NS5A(S225D)-CLIP] replicons. As shown in Fig. 7B, these two mutants retained their replication phenotypes (a 10-fold impairment of replication for the S225A mutant and wild-type replication for the S225D mutant) when expressed as SNAP/CLIP fusions. Analysis of cell lysates by Western blotting showed an abundance and hyperphosphorylation/basal phosphorylation ratios of NS5A comparable to those observed for the untagged mutants (Fig. 7C). As seen in the context of virus-infected cells (Fig. 3 and 4), cells harboring the NS5A(S225A)-SNAP replicon exhibited both a reduced abundance and a restricted perinuclear distribution of NS5A (Fig. 7E) in comparison to those for cells harboring either the wild-type (SNAP tagged) or the NS5A(S225D)-CLIP replicon.

To confirm that the S225A phenotype was not due to an environmental factor, such as differences in cell confluence, we electroporated either mSGR-luc-JFH-1-NS5A-CLIP or mSGR-luc-JFH-1-NS5A(S225A)-SNAP replicon RNAs into Huh7 cells and coseeded the electroporated cells into single cultures. By confocal microscopy, we observed that within the same population of cells, in the same field of view, those harboring the wild-type NS5A-CLIP replicon had a normal distribution of NS5A, while those harboring the NS5A(S225A)-SNAP replicon continued to show the restricted NS5A localization (Fig. 8A).

To further investigate the NS5A(S225A) phenotype, we utilized the SNAP/CLIP tag technology to ask if NS5A(S225A) could be transcomplemented by wild-type NS5A. In our experience, the simultaneous introduction of two replicons into a single cell is a rare event, for reasons that are not clear, particularly given the ratio of RNA molecules to cells in a typical electroporation experiment. However, after coelectroporation of mSGR-luc-JFH-1-NS5A-CLIP and mSGR-luc-JFH-1-NS5A(S225A)-SNAP replicon RNAs into Huh7 cells, we did observe some cells that were stained with both SNAP-505 and CLIP-TMR fluorophores, consistent with the presence of both actively replicating replicons in a single cell. In these cells, we observed two distinct phenotypes: either the presence of wild-type NS5A-CLIP was sufficient to restore the localization of NS5A(S225A)-SNAP to that of the wild type, resulting in a broad distribution of both and high levels of colocalization (Fig. 8B), or the distribution of NS5A(S225A)-SNAP replicon continued to show the restricted NS5A localization (Fig. 8C).

**DISCUSSION**

We present here the results of our investigation of the phenotype of mutants with mutations at residue serine 225 in NS5A of JFH-1. We propose that the phosphorylation of serine 225 is important for genome replication; mutation of this residue to alanine...
phenotype of NS5A serine phosphorylation

How might the altered distribution of the mutant NS5A be explained? A clue may come from the experiments examining the distribution of PI4P lipids (Fig. 4C). PI4P lipids are produced by PI4K, and previous work from other laboratories (5–7) has demonstrated that the PI4KIIα isoform is required for HCV genome replication. Indeed, PI4KIIα has been shown to bind to NS5A, with the binding site mapping to a short region at the C terminus of domain 1 (residues 202 to 210) (26), and disruption of this interaction had a profound effect on replication (100-fold reduction), blocked the stimulation of PI4P formation, and led to a collapse of the membranous web. In this regard, the restricted distribution and clustering of NS5A seen in the context of the S225A mutation are reminiscent of the NS5A localization seen in cells silenced for expression of PI4KIIα (7). We therefore propose that the replication defect of S225A may be explained in part by the inability of this NS5A mutant protein to effectively recruit and activate PI4KIIα. This simplistic explanation is complicated by the results of experiments in which we expressed NS5A in the context of an NS3-5B polyprotein from a cytomegalovirus promoter-driven plasmid vector (data not shown). In this situation, the nonstructural proteins can form a replication complex, but in the absence of a cognate RNA template, no genome replication can occur. Intriguingly, under these conditions the S225A phenotype is not apparent: the distribution and abundance of both NS5A and PI4P in transfected cells are very similar for both the wild type and the S225A mutant. Thus, we would further propose that the function of S225 phosphorylation in regulating interactions between NS5A and PI4KIIα (and/or other cellular components) is manifest only in the context of a replication complex that is actively replicating the viral genome. What is difficult to reconcile is the observation that inhibiting the interaction between NS5A and PI4KIIα increased the hyperphosphorylation of NS5A (26), concomitant with a reduction in virus genome replication. In contrast, our data show a loss of hyperphosphorylation of S225A (Fig. 1D), and indeed, our previous data (16) failed to indicate a correlation between hyperphosphorylation and replication. This discrepancy serves to highlight the complexity of NS5A phosphorylation, particularly regarding the multiple phosphorylation events within LCS I. There is a need to fully characterize both the kinases involved and the consequences of phosphorylation to clarify the function of these posttranslational modifications in the virus life cycle.

An alternative explanation for the restricted distribution of the S225A mutant NS5A is that it was impaired in its ability to traffic

(LCS I either positively (phosphorylation at S225 and S232) or negatively (phosphorylation at S229 and S235) regulating the formation and distribution of replication complexes. This conclusion also fits well with the concept of a phosphorylation-mediated switch of NS5A functions between replication and virus assembly, although the mechanics of such a switch remain elusive. Intriguingly, S232 corresponds to the residue which was the target for a dramatic culture adaptation (20,000-fold) in the context of genotype 1b replicons (S2204I in polyprotein numbering). Despite the conservation of the sequence of LCS I between HCV genotypes, these observations suggest that this does not manifest in a concomitant conservation of function, as neither we (Fig. 6A) nor others (15) have seen any major effect of mutations at this residue on JFH-1 virus replication. In the future, it will be important to compare the sites of phosphorylation and phenotypes in HCV isolates with a range of genotypes to address these questions.

FIG. 8 The redistribution of NS5A(S225A) is not the result of an environmental factor and can be partially transcomplemented by wild-type NS5A. (A) Huh7 cells electroporated with either mSGR-luc-JFH-1-NS5A-CLIP or mSGR-luc-JFH-1-NS5A(S225A)-SNAP RNAs were coseeded onto the same coverslip, and at 96 hpe, cells were fixed and labeled with either SNAP-505 (green) or CLIP-TMR (red), before imaging by confocal microscopy. (B, C) In vitro transcripts of mSGR-luc-JFH-1-NS5A-CLIP and mSGR-luc-JFH-1-NS5A(S225A)-SNAP were pooled prior to coelectroporation into Huh7 cells and seeding onto coverslips. At 96 hpe, cells were fixed and labeled with SNAP-505 and CLIP-TMR, prior to imaging by confocal microscopy. (B, C) Example of an Huh7 cell showing colocalization between wild-type NS5A(CLIP) and NS5A(S225A)-SNAP. (C) Example of an Huh7 cell showing a difference in cellular localization between NS5A(CLIP) and NS5A(S225A)-SNAP. The colocalization scatter plots at the bottom right of panels B and C are of individual cells and show the correlation of absolute values.

(S225A) resulted in a relocalization of NS5A and other components of the HCV genome replication complex (NS3, dsRNA, and PI4P lipids) to a perinuclear distribution, in comparison to either wild-type or a phosphomimetic (aspartic acid) substitution (S225D). This altered subcellular localization was accompanied by a 10-fold reduction in genome replication compared with the wild type. A 26-kb fragment of the JFH-1 genome, including the NS3-5B polyprotein, is actively replicating the viral genome. What is difficult to reconcile is the observation that inhibiting the interaction between NS5A and PI4KIIIα increased the hyperphosphorylation of NS5A (26), concomitant with a reduction in virus genome replication. In contrast, our data show a loss of hyperphosphorylation of S225A (Fig. 1D), and indeed, our previous data (16) failed to indicate a correlation between hyperphosphorylation and replication. This discrepancy serves to highlight the complexity of NS5A phosphorylation, particularly regarding the multiple phosphorylation events within LCS I. There is a need to fully characterize both the kinases involved and the consequences of phosphorylation to clarify the function of these posttranslational modifications in the virus life cycle.

An alternative explanation for the restricted distribution of the S225A mutant NS5A is that it was impaired in its ability to traffic
around the cytoplasm. In an elegant set of experiments using both tetracysteine and GFP-tagged NS5A-containing viruses, it was recently shown that NS5A was present in two classes of motile structures. The majority were relatively static, but some exhibited rapid, long-range, and sporadic movement throughout the cytoplasm (25). The rapid movement was shown to involve the microtubule network and was dynein dependent. It is thus formally possible that NS5A(S225A) is impaired in interactions with microtubules or dynein, preventing its movement around the cytoplasm and leading to its accumulation in the perinuclear region. However, we do not believe that this is the case because live cell imaging of either wild-type or S225A/D mutant viruses in which NS5A was Emerald-GFP (emGFP) tagged did not reveal any gross differences in the motility of NS5A-positive structures (see Videos S1 to S3 in the supplemental material). In addition, a defect in dynein-dependent trafficking might intuitively be expected to result in a distribution around the periphery of the cell, as dynein moves toward the minus end of microtubules, which is usually at the cell center.

The use of the SNAP/CLIP tagging system allowed us to differentially label NS5A(S225A)-SNAP and wild-type NS5A-CLIP within one population of cells, thereby visualizing in the same field of view cells that were harboring either S225A or wild-type replicons. Importantly, this analysis demonstrated that the distribution of NS5A(S225A) was still markedly different from that of the wild-type protein (Fig. 7E), illustrating that the altered localization of NS5A was not due to an environmental factor. However, the SNAP/CLIP system also revealed that the presence of wild-type NS5A was able to complement the S225A defect in only a proportion of cells. This suggests that, additional, yet undefined, stochastic factors, such as the stage of the cell cycle or the availability of cellular factors, may contribute to the observed phenotype.

In conclusion, this study highlights an important role of phosphorylation in regulating the function of NS5A. Clearly, there is much to do to define the mechanisms underpinning this effect; in particular, we need to understand the molecular interactions that are modulated by the phosphorylation of NS5A at serine 225 and other serines in LCI I. While a number of candidates, such as PI4KIII, are already apparent, there are many other NS5A-interacting proteins that should be considered, and a proteomics approach to compare the interactome of both the wild-type and mutant forms of NS5A is an appropriate way forward. Such experiments are under way in our laboratory.

ACKNOWLEDGMENTS

We thank John McLauchlan (Centre for Virus Research, University of Glasgow) for the SGR-luc-JFH-I construct, Takaji Wakita (National Institute for Infectious Diseases, Tokyo, Japan) for pFH-I, Thomas Pietzschmann (Hannover, Germany) for the NS3 antibody, and Joseph Shaw (Leeds, United Kingdom) for help with the daclatasvir EC50 analysis. Finally, we thank Volker Lohmann and Dave Rowlands for insightful comments on this work.

This work was funded by a Wellcome Trust Senior Investigator Award to M.H. (096670/MA). J.M. is a Royal Society University Research Fellow (grant number UF100419). The confocal microscope used for live cell imaging of HCV-infected cells was funded by a Royal Society equipment grant (grant number RG110306).

REFERENCES


