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Replication of murine coronavirus requires multiple cysteines in the endodomain of spike protein

Jinhua Yang, Jun Lv, Yuyan Wang, Shuang Gao, Qianqian Yao, Di Qu, Rong Ye *

Shanghai Medical College, Fudan University, Shanghai 200032, China

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ABSTRACT

A conserved cysteine-rich motif located between the transmembrane domain and the endodomain is essential for membrane fusion and assembly of coronavirus spike (S) protein. Here, we proved that three cysteines within the motif, but not dependent on position, are minimally required for the survival of the recombinant mouse hepatitis virus. When the carboxy termini with these mutated motifs of S proteins were respectively introduced into a heterogeneous protein, both incorporation into lipid rafts and S-palmitoylation of these recombinant proteins showed a similar quantity requirement to cysteine residues. Meanwhile, the redistribution of these proteins on cellular surface indicated that the absence of the positively charged rather than cysteine residues in the motif might lead the dramatic reduction in syncytial formation of some mutants with the deleted motifs. These results suggest that multiple cysteine as well as charged residues concurrently improves the membrane-associated functions of S protein in viral replication and cytopathogenesis. © 2012 Elsevier Inc. All rights reserved.

Introduction

Coronaviruses are a family of enveloped, positive-sense RNA viruses that principally cause respiratory or enteric infections in humans and a broad range of animals (Holmes, 2003; Masters, 2006; Siddell, 1995). Mouse hepatitis virus (MHV), a member of the group II of coronaviruses (now called genus Betacoronavirus) (Carstens and Ball, 2009), causes neurovirulence and hepatitis in rodents, such as mice and rats (Weiss and Navas-Martin, 2005). MHV has been used extensively for in vivo studies of neurovirology and mechanisms of coronavirus entry and assembly (Masters, 2006). Three membrane proteins, including the spike (S), envelope and membrane proteins, are incorporated into the envelope of MHV virions (Masters, 2006; Siddell, 1995). The S protein is a 180 kDa, N-glycosylated, type I membrane protein which is generally cleaved into separate N-terminal S1 and C-terminal S2 subunits. The main component of the S protein is the ectodomain (Ecto) in which a few functional domains have been clearly described. These include the receptor-binding domain from S1 as well as the fusion peptide and two heptad repeats (HR1 and HR2) from S2. The Ecto is responsible for the cellular tropism and membrane fusion in coronavirus infection (Masters, 2006; Siddell, 1995). The minor part of the S protein is composed of a transmembrane domain (Tm) and an endodomain (Endo).

There are up to ten highly conserved cysteine residues located between the Tm and the Endo of all coronavirus S proteins (Bos et al., 1995). Previous studies have revealed that the cysteine-rich (CR) motif of S proteins is essential for virion assembly and membrane fusion (Bos et al., 1995; Bosch et al., 2005; Chang et al., 2000; Lopez et al., 2008; Takeda et al., 2003; Zhang et al., 2000), paramyxovirus Haemagglutinin (Chen et al., 2005; Jin et al., 1996; Kordyukova et al., 2006). The modification also occurs on the cysteine residues in the cytoplasmic tails of other viral membrane proteins, such as influenza virus Hemagglutinin (Chen et al., 2005; Jin et al., 1996; Kordyukova et al., 2008; Takeda et al., 2003; Zhang et al., 2000), paramyxovirus F (Branigan et al., 2006), vesicular stomatitis virus G (Whitt and Rose, 1991), hepatitis C virus core (Majeau et al., 2009), HIV-1 envelope (Rouso et al., 2000), Sindbis virus E1 and E2 (Smit et al., 2001) and baculovirus gp64 (Zhang et al., 2003). This modification enables...
the association of viral proteins with lipid rafts, which is essential for viral infectivity (McBride and Machamer, 2010; Zhang et al., 2000). However, the mechanism of S-palmitoylation on viral membrane proteins is not clearly understood.

In this study, the functional importance of the cysteine residues in the MHV S Endo for viral replication was investigated by reverse genetics. There was no single “key” cysteine residue required in the motif for virus replication. However, at least three cysteine residues were found to be required for the survival of recombinant MHVs. Immunofluorescence analysis of the heterogeneous hook proteins (HK) expressed by recombinant MHVs (Yang et al., 2011; Ye et al., 2004) showed that the dramatic reduction of the cytopathic effect (CPE) of recombinant MHVs with a partial deletion of motif did not involve these cysteines. This effect might result from the deletion of some of the charged residues in the motif, which decreased the amount of S protein on the surfaces of infected cells. Furthermore, both incorporation into detergent-resistant membrane fractions (DRM) and the S-palmitoylation level of these recombinant protein were examined. Results showed that an association into lipid rafts and S-palmitoylation of the recombinant protein with a deletion motif containing six cysteines not only was detected but also exhibited a similar level to that of wild-type motif. However, the protein with a deletion motif containing only two cysteines could not be detected.

Results

No single specific cysteine in the S protein endodomain was essential for virus replication

There are nine cysteines in the CR motif of the MHV S protein (Lopez et al., 2008). To examine the effects of these residues on virus replication, a series of recombinant MHVs was constructed with mutated S Endo by RNA targeted recombination (Fig. 1A). The deletion of all 18 residues of the CR motif (Mut-d1), the deletion of 10 residues of the CR motif near the Tm (Mut-d2), and the substitution of all nine cysteine residues of the CR motif with alanine residues (Mut-p1) were lethal for the respective recombinant MHVs (Fig. 1B).

However, the deletion of 8 residues in the CR motif near the Endo (Mut-d3) and all point mutations of 1–2 cysteine residues (Mut-p2 to -p7) did not impair the survival of recombinant MHVs (Fig. 1B). The size of plaques produced by all survived mutants was significantly smaller than that of the wild-type MHV (p < 0.001) (Fig. 1C). The statistical analysis showed that the partial deletion mutant (Mut-d3) produced smaller plaques compared with other point mutants (Mut-p2 to -p7) that could be further divided into three groups from small to large: Mut-p4, -p5, and -p7; Mut-p2 and -p3; Mut-6. The tendency might be associated with the importance of these mutated cysteines in the motif. We observed that there was no significant difference between one and two cysteine substitution. Also, the substitution with alanine, serine, or glycine showed the similar effect. Notably, the boundary of the plaques formed by Mut-d3 seemed to be much fuzzy and the plaque size of Mut-p7 had a bigger deviation (Fig. 1B). These results indicated that no unique cysteine is crucial for virus survival but some cysteines may play more important roles in the function of the S protein.

Multiple cysteines in the S endodomain were concurrently responsible for virus replication

Previous studies have shown that when the CR motif of the S protein was mostly deleted (leaving only four or five cysteines), the production of recombinant MHVs was not affected, but the cell–cell fusion was decreased dramatically (Yang et al., 2011; Ye et al., 2004). To determine the exact number of cysteines required for the survival of MHV, mutant viruses with only one to six cysteines left in the CR motif of S gene were constructed (Fig. 2A). Meanwhile, in order to facilitate the detection of viral survival, gene 4 was replaced by an EGFP reporter in these mutants (Fig. 2A). L2 cells were infected with supernatants of the various mutant MHVs produced by targeted RNA recombination and visualised by microscopy at 24 h post infection (hpi). The recombinant MHVs containing one to four cysteine residues in the motif (MH54GFP-1C to -4C) failed to induce any visible syncytia, and the recombinant MHVs with five or six cysteine residues in the motif (MH54GFP-5C and -6C) induced only light syncytia formation (Fig. 2B). Fluorescent microscopy showed that the MH54GFP-1C and -2C mutant viruses failed to express EGFP, that the MH54EGFP-3C virus expressed a low level of EGFP, and that the MH54GFP-4C, -5C and -6C mutants expressed a high level of EGFP (Fig. 2C). These results indicated that the replication of the virus depends on the number of cysteine residues in the CR motif and that at least three cysteines in the motif are required for the viability of the recombinant MHVs.

Deletion of the cysteine-rich motif impaired the localisation of the S protein to the cell surface

The substitution of the cytoplasmic tail of the MHV S protein, which contains the CR motif, with the cytoplasmic tail of the VSV G protein resulted in abolition of cell–cell fusion but did not interfere with expression of the S protein on the cell surface (Bos et al., 1995). The replication of coronaviruses is critically affected by the cellular localisation of the S protein. The less S protein that is expressed on the cell surface, the weaker is the extent of cell–cell fusion (Bosch et al., 2003). We previously found that membrane-anchored HK protein and cytoplasmic EGFP were differentially located within the cell (Yang et al., 2011).

In this study, the distribution of the HK proteins expressed by four recombinant MHVs (Fig. 3A) was further explored by sequential staining to the living and permeabilized cells with two anti-HA antibodies. Two HK proteins, the HKM protein with wild-type MHV S Tm/Endo and the HKP protein with a Tm and a truncated Endo of platelet-derived growth factor receptor (PDGFR) without cysteine residues, were largely localised on the cellular surface (upper two panels in Fig. 3B). However, HK-2C and HK-6C with the MHV S Tm but partially deleted CR in the Endo were found to be principally distributed in the cytoplasm (bottom two panels in Fig. 3B). These results indicated that the sequence containing charged residues rather than the number of cysteines in the motif is essential for the accumulation of S protein on the plasma membrane.

Multiple cysteines in the endodomain enhance the incorporation of S protein to DRMs

DRMs are membrane subdomains that are resistant to solubilisation with cold non-ionic detergents, such as Triton X-100 (Lingwood and Simons, 2007). The extractability of infected Neuro-2a cells was examined by treatment with 1% Triton X-100 in cold TNE buffer. DRMs were separated by flotation ultracentrifugation. All of four HK proteins were distributed in the higher density fractions (fractions 7 to 10) (Fig. 4). Both HKM and HK-6C were also found to diffuse into the lower density fractions (fractions 3 to 4), where the DRM maker protein caveolin-1 was distributed. However, the distribution of HKP and HK-2C to the lower density fractions was significantly reduced (Fig. 4). These results showed that the HK proteins with the wild type CR motif and the CR motif mutant containing only six cysteines were associated with lipid rafts to a higher degree than those recombinant proteins with CR motifs with zero or two cysteines.
Multiple cysteines in the endodomain enhance the S-palmitoylation of the S protein

S-palmitoylation of the MHV S protein has been shown to focus on the carboxyl terminal cysteine residues and is necessary for virus assembly (Bosch et al., 2005; Thorp et al., 2006). The S-palmitoylation of some membrane-proximal cysteine clusters in the SARS CoV S protein may be important for S-mediated cell fusion, and it may also be necessary for the partitioning of the S protein into DRMs (McBride and Machamer, 2010; Petit et al., 2007). Hydroxylamine treatment (Drisdel et al., 2006; Gonzalo and Linder, 1998; Valdez-Taubas and Pelham, 2005) has been used to detect the S-palmitoylation level of HK proteins expressed in recombinant MHVs in Neuro-2a cells (Fig. 5A). The electrophoretic mobility of HK proteins was examined after treatment with or without hydroxylamine. The mobility of the HKM and HK-6C proteins was significantly altered, whereas that of the HK-2C and HKP mutants was not (Fig. 5B). These results indicated that the level of S-palmitoylation is correlated with the number of cysteine residues in the CR motif of the S protein endodomain. Moreover, the level of S-palmitoylation is consistent not only with the survival of the recombinant MHV (Fig. 2) but also with the association of S protein in lipid rafts of the infected cells (Fig. 4).

Discussion

Both the Tm and the Endo of the S protein are essential for virion assembly and membrane fusion of coronavirus (Bos et al., 1995; Bosch et al., 2005; Chang et al., 2000; Petit et al., 2007; Shulla and Gallagher, 2009; Thorp et al., 2006; Ye et al., 2004). The Tm is an efficient membrane anchor because of its integration of hydrophobic residues with lipids of membranes. Some charged residues of the Endo have been shown to direct the S protein to specifically assemble into virions via their interactions with membranes or other structural proteins (Bosch et al., 2005; Ye et al., 2004). The cysteines in the Endo have been shown to enhance the trafficking and membrane binding of S proteins to cells through the S-palmitoylation of the S proteins (Bos et al., 1995; McBride and Machamer, 2010; Petit et al., 2007; Thorp et al., 2006).

To analyse the effects of the cysteine residues in the unique CR motif on the replication of murine coronavirus, two sets of recombinant MHVs were generated by targeted RNA recombination (Masters and Rottier, 2005; Masters et al., 2006). One set, in which substitution or deletion mutations of the CR were directly located in the S protein endodomain, was in the plasma membrane and on the cell surface. However, the mutants with truncated CR motifs (HK-2C and -6C) were distributed in the cytoplasm (Fig. 3). This redistribution of recombinant proteins might be attributed to the deletion of some of the positively charged residues other than the cysteines in the motif. The charged residues, located on the cytoplasmic tail of transmembrane proteins, generally serve as the targeting and sorting signals to different membrane compartments, including the plasma membrane (Itin et al., 1995). Further investigation of the charged residues in the CR motif of the MHV S will be needed to address this issue.

The number of cysteines in the CR motif of the MHV S protein was shown to directly correlate with its trafficking to lipid rafts of the cell membrane (Fig. 4), and the correlation is also consistent with the levels of S-palmitoylation of the recombinant HK proteins (Fig. 5). The recombinant proteins with either a wild-type CR motif (HKM) or a CR motif containing six cysteine residues (HK-6C) could be S-palmitoylated and incorporated into DRMs. The mutants with either two cysteines (HK-2C) or no cysteines (HKP) in the CR motif could neither be S-palmitoylated nor be incorporated into DRMs.

Three 3-cysteine clusters in the motif might guarantee the S-palmitoylation, which is beneficial to the association of the S protein to the lipid rafts during viral membrane fusion and virion assembly (Fig. 6). Both trafficking between membranes signalized by the

Previous studies have shown that the CR motif of the S protein Endo is associated with the ability of the S protein to induce syncytia (Bos et al., 1995; Bosch et al., 2005; Chang et al., 2000; Thorp et al., 2006; Ye et al., 2004). In this study, the cysteines in the CR motif were mutated either alone or in combination with alanine, glycine, or serine, and no specific cysteine was found to be crucial for the survival of the recombinant MHVs (Fig. 1). The difference in the plaque morphology among these survived mutants implied that some cysteines may play more important roles in viral replication. We found that the plaques of Mut-p7 and Mut-d3 were uneven and slightly smaller than those of Mut-p2 and -p3, but the plaques of Mut-p6 were slightly larger than those of the other mutants (Figs. 1B and C). The two cysteines in this region were more important for the incorporation of S proteins into virions, which consolidated previous results (Shulla and Gallagher, 2009; Ye et al., 2004). On the other hand, the deletion of the CR motif (Mut-d1), the partial deletion of the motif near the Tm (Mut-d2), and the mutation of all nine cysteines to alanines (Mut-p1) were lethal for the recombinant MHVs, which indicated that six cysteines close to the Tm might be critical for survival of recombinant viruses (Fig. 1B).

Three cysteines in centre of CR motif (Mut-3 and Mut-4) might play more important role in viral replication (Figs. 1B and C). Based on the above results, we hypothesise that the multiple cysteines in the motif may account for the function of S proteins. To verify this hypothesis, some deletion mutations of the motif retained one to six cysteines were introduced into recombinant MHVs in which the EGFP reporter gene was substituted for gene 4 (Fig. 2A). These results indicated that only the recombinant MHVs with three or more cysteines were viable (Fig. 2C). However, the ability to induce syncytia by the recombinant MHVs with only three to six cysteines in S protein CR motif was dramatically decreased in L2 cells (Fig. 2B). The decreased CPE was consistent with results of previous studies (Yang et al., 2011; Ye et al., 2004).

The subcellular distribution of recombinant HK proteins in infected Neuro-2a cells provided an explanation for this phenomenon. The distribution of the wild type S Endo (HKM) or PDGFR Endo (HKP) mutants was in the plasma membrane and on the cell surface. However, the mutants with truncated CR motifs (HK-2C and -6C) were distributed in the cytoplasm (Fig. 3). This redistribution of recombinant proteins might be attributed to the deletion of some of the positively charged residues other than the cysteines in the motif. The charged residues, located on the cytoplasmic tail of transmembrane proteins, generally serve as the trafficking and sorting signals to different membrane compartments, including the plasma membrane (Itin et al., 1995). Further investigation of the charged residues in the CR motif of the MHV S will be needed to address this issue.

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A

| Mutations | Transmembrane Domain | Endodomain |
|-----------|-----------------------|------------|
| MEV WT    | EMVNEWYVWLLGAVCVCVLLLFFIACCTCQ0SQCFCGCQHCDYEGH0QSDIVVSH hed |             |
| Mut-d1    | EMVNEWYVWLLGAVCVCVLLLFFI----------------------DEYEGH0QSDIVVSH hed |             |
| Mut-d2    | EMVNEWYVWLLGAVCVCVLLLFFI----------------------DEYEGH0QSDIVVSH hed |             |
| Mut-d3    | EMVNEWYVWLLGAVCVCVLLLFFIOCCCTCQ0SQCFCGCQHCDYEGH0QSDIVVSH hed |             |
| Mut-p1    | EMVNEWYVWLLGAVCVCVLLLFFI-------------------------------DEYEGH0QSDIVVSH hed |             |
| Mut-p2    | EMVNEWYVWLLGAVCVCVLLLFFI-------------------------------DEYEGH0QSDIVVSH hed |             |
| Mut-p3    | EMVNEWYVWLLGAVCVCVLLLFFI-------------------------------DEYEGH0QSDIVVSH hed |             |
| Mut-p4    | EMVNEWYVWLLGAVCVCVLLLFFI-------------------------------DEYEGH0QSDIVVSH hed |             |
| Mut-p5    | EMVNEWYVWLLGAVCVCVLLLFFI-------------------------------DEYEGH0QSDIVVSH hed |             |
| Mut-p6    | EMVNEWYVWLLGAVCVCVLLLFFI-------------------------------DEYEGH0QSDIVVSH hed |             |
| Mut-p7    | EMVNEWYVWLLGAVCVCVLLLFFI-------------------------------DEYEGH0QSDIVVSH hed |             |

B

WT: Lethal, no plaques
-d1: Lethal, no plaques
-d2: Lethal, no plaques
-d3: Lethal, no plaques
Mock: Lethal, no plaques
-p1: Lethal, no plaques
-p2: Lethal, no plaques
-p3: Lethal, no plaques
-p4: Lethal, no plaques
-p5: Lethal, no plaques
-p6: Lethal, no plaques
-p7: Lethal, no plaques

C

Plaque diameter (mm)

| Mutants | Plaque diameter (mm) |
|---------|----------------------|
| WT      | 1.938 ***            |
| -d3     | 2.140 ***            |
| -p2     | 2.380 ***            |
| -p3     | 2.400 ***            |
| -p4     | 2.630 ***            |
| -p5     | 2.860 ***            |
| -p6     | 3.090 ***            |
| -p7     | 3.220 ***            |

\( t = 11.67 \) ***
\( t = 14.00 \) ***
\( t = 15.23 \) ***
\( t = 7.06 \) ***
\( t = 15.00 \) ***
Fig. 2. Effects of the number of cysteines in the S protein endodomain on the production and cytopathogenesis of recombinant MHVs. (A) The diagram represents part of the genomic organisation of the EGFP-replaced mutant MH54GFP derived from MHV A59 in which the mutated region in the S gene is marked with a full triangle (▲). The sequence alignment shows the deletion mutations with the numbers of cysteine residues (MH54GFP-1C to -6C) replacing the cysteine-rich motif. (B) L2 cells infected with recombinant MHVs (24 hpi) were visualised under contrast microscope (with magnification 400×) and showed no syncytia (-1C to -4C), small syncytia (-5C and -6C), and large syncytia of wild-type MHV (12 hpi). (C) L2 cells infected with recombinant MHVs (24 hpi) newly produced by RNA-targeted recombination were visualised under fluorescent microscope (with magnification 100×), which showed no cells infected (-1C to -2C), a few cells infected (-3C), more cells infected (-4C to -6C); small syncytia (-5C and -6C) and large syncytia of wild-type MHV (12 hpi).
positively charged residues and S-palmitoylation as well as lipid raft incorporation induced by the cysteine residues were primarily uncovered in this study.

Materials and methods

Plasmid construction

The donor RNA transcription vectors used in this study were derived from pMH54, pMH54GFP, pHKM and pHKP (Das Sarma et al., 2002; Kuo et al., 2000; Ye et al., 2004). The construction strategy we used has been previously described (Yang et al., 2011). Briefly, pMH54 and pMH54GFP transcription vectors were digested with MluI and SbfI, and a set of homogenous fragments with point or deletion mutations in the CR motif were inserted. These fragments were generated by splicing overlap extension-PCR (Aiyar et al., 1996) using pMH54 DNA as a template that contained the endodomain sequence of MHV S gene.

Both pHKM and pHKP were previously constructed (Ye et al., 2004). The recombinant hook protein (HK) gene contained a DNA sequence encoding a signal peptide from the immunoglobulin light chain, an HA tagged-scFv ectodomain and a Tm/Endo from either the MHV S gene or a truncated PDGFR gene (Masters et al., 2006; Yang et al., 2011; Ye et al., 2004). For the generation of pHK-2C and pHK-6C, pHKM as the receptor vector was digested with SalI and AsclI and two homogenous fragments with mutations in the cysteine-rich motif were inserted. All DNA manipulations were performed according to standard molecular cloning procedures (Sambrook and Russell, 2001).
or recombinant MHVs, monolayers of cells at 60–80% confluence were inoculated at a multiplicity of infection (MOI) of 0.1–1.0.

Production of recombinant MHVs

Recombinant MHVs were produced according to the procedure of targeted RNA recombination developed by Kuo et al. (2000) and Masters and Rottier (2005). fMHV is a chimeric MHV A59 strain virus in which the ectodomain of the S gene is replaced with that of feline infectious peritonitis virus. This replacement results in the fact that recombinant virus is grown in feline FCWF cells but not in murine L2 cells. Donor RNA transcription vectors were linearised by digestion with Pael and transcribed into mRNAs using the mMESSAGE mMACHINE T7 Ultra kit (Ambion). Approximately 10 μl of each recombinant transcript was prepared for transfection of ~10⁷ of FCWF cells. The fMHV-infected cells were briefly digested with a low concentration of trypsin at 2 h post-infection (hpi). RNA transfection was performed using a Gene Pulser Xcell electroporation system (Bio-Rad) with one pulse at settings of 975 μF and 0.3 kV. The infected and transfected FCWF cells were then added to 6-well cell-culture clusters seeded with L2 monolayers. Culture supernatants containing recombinant MHVs were collected at 24, 48 and 72 hpi. Recombinant MHVs were plaque-purified twice on L2 cells, and all mutations were verified by RT-PCR and sequencing.

Plaque assay

L2 cells were grown in 60-mm dishes and infected with 1 ml of viruses at dilutions of 10⁻², 10⁻⁴, 10⁻⁵ and 10⁻⁶ for 2 h at 37 °C. Seven milliliters of 0.95% agar containing 50 μg/ml neutral red (Sigma-Aldrich) was overlaid onto the cells. Plaques were picked at 48 hpi. For visualisation of the plaques, 3 ml of 0.9% agar containing 50 μg/ml neutral red (Sigma-Aldrich) was overlaid onto the plate. Plaques were photographed on a white light transilluminator (Upland) with a digital camera after incubation for 6 to 8 h at 37 °C.

Confocal immunofluorescence

Neuro-2a cells were grown on glass coverslips coated with poly-L-lysine (Sigma-Aldrich) to ~60% confluence and then infected with recombinant MHV that expressed HKM, HKP, HK-2C, or HK-6C. At 16–18 hpi, the cells were washed with PBS and incubated with a rat mAb to the HA tag (Roche) for 1 h at 37 °C for staining the HK proteins on the cell surface. Then cells were fixed with 4% paraformaldehyde (Sigma-Aldrich) in PBS for 20 min and permeabilized with 0.1% Triton X-100 in PBS for 5 min at room temperature. Fixed cells were blocked with 3% BSA and then incubated with a mouse mAb to the HA tag (Roche) overnight at 4 °C. The cells were then incubated with a FITC-conjugated antibody to rat IgG and a TRITC-conjugated antibody to mouse IgG (Jackson ImmunoResearch) for 1 h at room temperature. At last the cells were washed three times in PBS and mounted with Mowiol antifade reagent (Sigma-Aldrich). Fluorescent images were obtained using a Leica TCS-SLP laser-scanning confocal microscope and processed with LAS AF software (Leica).

Analysis of DRM association

DRM fractions were isolated using discontinuous density ultracentrifugation as previously described (Brown and London, 1998; Hooper, 1999; McBride and Machamer, 2010; Zhang et al., 2000). At 12 hpi, Neuro-2a cells were rinsed twice with ice-cold PBS and lysed on ice for 30 min in 1 ml of TNE buffer (10 mM Tris–HCl, 150 mM NaCl and 5 mM EDTA, pH 7.4) containing a protease inhibitor cocktail (Roche) and 1% Triton X-100. After passing through a 25-G needle 10 times, lysates were mixed with 1 ml of 80% (w/v) sucrose in TNE. The
mixture was overlaid with 4 ml of 35% sucrose in TNE and then with 1 ml of 5% sucrose in TNE. Finally, 3 ml of TNE was overlaid on the 5% sucrose solution. After centrifugation for 18 h at 285,000 × g at 4 °C (SW40 rotator, Beckman-Coulter), 1.0 ml fractions were collected from the top to bottom of the gradient and analysed by 10% SDS-PAGE and Western blotting with mAbs to the HA tag (Roche) and to caveolin-1 (Abcam).

**Hydroxylamine treatment**

The levels of S-palmitoylation of recombinant proteins were analysed by a shift in the electrophoretic mobility after treatment with hydroxylamine (Drisdel et al., 2006; Gonzalo and Linder, 1998; Valdez-Taubas and Pelham, 2005). In brief, infected monolayers of Neuro-2a cells were washed twice with ice-cold PBS and dissolved in 300 μl of IPP buffer (50 mM Tris–HCl, 150 mM NaCl, 1.0% Nonidet P-40, pH 8.0) containing a protease inhibitor cocktail. Cell lysates were held on ice for 5 min and then clarified by centrifugation at 12,000 × g for 5 min at 4 °C. Clarified lysates were mixed with 15 μl of Anti-HA Affinity Matrix (Roche) by inversion overnight at 4 °C. The matrix was precipitated by centrifuging at 12,000 × g for 30 s, and the supernatant was carefully removed. The matrix was washed with 1 ml of cold IPP buffer three times. The final rinsed matrix was suspended in 60 μl of 2× SDS sample buffer and heated at 95 °C for 5 min. The immunoprecipitated proteins were divided into two parts. One was mixed with an equal volume of 2 M hydroxylamine (Sigma-Aldrich) in water at room temperature for 1 h, and the other was mixed with an equal volume of water as a control. The samples were separated by SDS-PAGE with 4–20% gradient gel (Bio-Rad) and characterised by Western blotting using an anti-HA tag mAb (Roche).

**Gel electrophoresis and Western blotting**

Protein samples were separated by 10% SDS-PAGE and transferred to PVDF membrane using a Mini Trans-blot Electrophoretic Transfer Cell (Bio-Rad) according to the laboratory manual (Sambrook and Russell, 2001). The membranes were blocked with 5% nonfat milk in PBST (PBS containing 0.1% (v/v) Tween-20) for 1 h at room temperature followed by an incubation with a mAb to the HA tag (Roche) overnight at 4 °C. After washing three times in PBST, membranes were incubated with an HRP-conjugated secondary antibody (GE, Amersham) in 5% nonfat milk-PBST for 1 h at room temperature and washed again with PBST. Signals were visualised using the ECL detection system (GE, Amersham) according to the manual instructions.

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