Identification and characterization of two cleavage fragments from the *Aquareovirus* nonstructural protein NS80

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*Aquareovirus* species vary with respect to pathogenicity, and the nonstructural protein NS80 of aquareoviruses has been implicated in the regulation of viral replication and assembly, which can form viral inclusion bodies (VIBs) and recruit viral proteins to its VIBs in infected cells. NS80 consists of 742 amino acids with a molecular weight of approximately 80 kDa. Interestingly, a short specific fragment of NS80 has also been detected in infected cells. In this study, an approximately 58-kDa product of NS80 was confirmed in various infected and transfected cells by immunoblotting analyses using α-NS80C. Mutational analysis and time course expression assays indicated that the accumulation of the 58-kDa fragment was related to time and infection dose, suggesting that the fragment is not a transient intermediate of protein degradation. Moreover, another smaller fragment with a molecular mass of approximately 22 kDa was observed in transfected and infected cells by immunoblotting with a specific anti-FLAG monoclonal antibody or α-NS80N, indicating that the 58-kDa polypeptide is derived from a specific cleavage site near the amino terminus of NS80. Additionally, different subcellular localization patterns were observed for the 22-kDa and 58-kDa fragments in an immunofluorescence analysis, implying that the two cleavage fragments of NS80 function differently in the viral life cycle. These results provide a basis for additional studies of the role of NS80 played in replication and particle assembly of the *Aquareovirus*.

**KEYWORDS**  
*Aquareovirus*; nonstructural protein NS80; cleavage fragments; subcellular localization; functional analysis

**INTRODUCTION**

Viruses belonging to the family *Reoviridae* are currently classified into 15 genera and can infect a wide variety of organisms, including vertebrates, invertebrates, and plants. These viruses are non-enveloped and contain a genome with 9–12 segments enclosed within a multiple protein-shelled core. Aquareoviruses isolated from aquatic animals, including finfish and crustaceans, are classified into seven species (*Aquareovirus A* to *Aquareovirus G*) based on their genome identities (Mertens et al., 2011). Generally, aquareoviruses are minimally pathogenic in aquatic cultures. However, Grass carp reovirus (GCRV, a member of *Aquareovirus C*) can cause fatal epidemic hemorrhagic disease in fingerling and yearling grass carp, and has been regarded as the most highly pathogenic aquareovirus (Rangel et al., 1999). Therefore, GCRV serves as an ideal model for studying the replication and pathogenesis of aquareoviruses.  

Similar to the structure of other reoviruses in the family *Reoviridae*, GCRV virion is a multilayered icosahedral symmetry with 12 pentameric turrets that reside on the surface of the core or at the five-fold axis of the intact particle (Cheng et al., 2008; Zhang et al., 2010). The eleven-segmented double-stranded RNA genome encodes 7 structural proteins (VP1–VP7) and 5 nonstruc-
tural proteins (NS16, NS26, NS31, NS38, and NS80) (Attoui et al., 2002). The viral core that serves as a transcriptase particle in vivo is composed of 5 proteins (VP1–VP4 and VP6), and also possesses transcriptional activity in vitro (Smith et al., 1980; Yang et al., 2012; Yan et al., 2014). VP5 and VP7 make up the outer capsid required for viral entry into host cells during infection (Zhang et al., 2010). In addition, the nonstructural proteins are involved in the viral replication cycle (Fan et al., 2009a; Fan et al., 2010; Cai et al., 2011; Shao et al., 2013; Yan et al., 2015). Phylogenetic and three-dimensional structural analyses have revealed that there is a close evolutionary relationship between aquareoviruses and ortheoviruses, including mammalian ortheoviruses (MRV) and avian reoviruses (ARV) (Shaw et al., 2003; Miller et al., 2003; Broering et al., 2004; Sharpe et al., 1982; Parker et al., 2002; Becker et al., 1970; Fields et al., 1971; Brookes et al., 1993; Broering et al., 2004). VIBs are thought to provide a physical scaffold to concentrate viral components and related cellular factors so that viral replication and nascent particle assembly proceed at specific sites in the cytoplasm. A single μNS of MRV can form VIBs in transfected cells and can recruit five core structural proteins (λ1, λ2, λ3, μ2, and σ2), the nonstructural protein σNS, and cellular clathrin (Sharpe et al., 1982; Parker et al., 2002; Becker et al., 2003; Miller et al., 2003; Broering et al., 2004; Tousis-Otero et al., 2004; Miller et al., 2010; Ivanovic et al., 2011). The nonstructural protein NS80 of GCRV, the analogue of the μNS protein of MRV and ARV, consists of 742 amino acids with a molecular weight of approximately 80 kDa (Fan et al., 2009b; Fan et al., 2010). Previous studies in our lab have demonstrated that NS80 can form VIBs in singly expressed or infected cells. In addition, NS80 can retain viral proteins (VP1–VP4 and VP6) in VIBs, and associate with the nonstructural protein NS38 and newly synthesized viral RNAs in infected cells (Shao et al., 2010; Shao et al., 2013; Yan et al., 2015). Additionally, the C-terminal regions, including His569 and Cys571, in the coiled-coil region of NS80 are crucial for VIB formation (Shao et al., 2013).

The μNS protein, which is encoded by the M3 gene of MRV and ARV, expresses two isoforms in infected cells (Wiener et al., 1989; Tousis-Otero et al., 2004). Similarly, a shorter fragment with NS80 expression has also been found in GCRV-infected cells (Fan et al., 2009b; Fan et al., 2010; Shao et al., 2013; Yan et al., 2015). These results suggest that NS80 presents different forms or structural conformations in infected cells to regulate viral replication and assembly at different life cycle stages. As a step toward understanding the role of NS80 in the viral life cycle, we characterized NS80 and its related fragment, including their subcellular localization.

MATERIALS AND METHODS

Cells, virus, and antibodies
Ctenopharyngodon idellus kidney (CIK) cells and fathead minnow (FHM) cells were grown at 28 °C in Eagle’s Minimum Essential Medium (Invitrogen, Carlsbad, USA) and Medium 199 (M199; Gibco BRL, Rockville, USA), respectively, and both media were supplemented with 10% fetal bovine serum (FBS). BHK-21 (baby hamster kidney) cells and Vero cells were cultivated in Dulbecco’s modified Eagle medium (Gibco BRL, Rockville, MD, USA) supplemented with 10% FBS. The GCRV–873 strain used in this study was maintained in our laboratory, and virus propagation was described previously (Fang et al., 1989).

The polyclonal antibody (pAb) against GCRV NS80 C-terminal amino acid sequences (aa 335–742), named α-NS80C, was generated in our laboratory and is described elsewhere (Fan et al., 2009b), and the pAb against NS80 (1–334), termed α-NS80N, was prepared following previous methods (Fan et al., 2009b). The mouse anti-FLAG monoclonal antibody (mAb) was purchased from Shanghai Genomics (Shanghai, China) and mouse mAb against β-actin was purchased from Santa Cruz Biotechnology (Santa Cruz, California, USA). Rabbit anti-poly-ubiquitination pAb was purchased from Proteintech Co. (Wuhan, China). Alkaline phosphatase-coupled goat anti-rabbit IgG or goat anti-mouse IgG were purchased from Sigma-Aldrich (St. Louis, USA) and Alexa Fluor 488 or 568 donkey anti-rabbit IgG (H+L) antibodies were purchased from Invitrogen Co.

Plasmids and site-directed mutagenesis to obtain mutant NS80
The recombinant plasmids pCI-neo-NS80, pCI-neo-NS80 (130–742), and pNS38-GFP-NSP5 were constructed previously (Shao et al., 2013). The plasmids with FLAG-tagged NS80 at the N-terminus (FLAG-NS80) and C-terminus (NS80-FLAG), pCI-neo-NS80 (1–190) and pCI-neo-NS80 (386–742), were built using pCI-neo-NS80 as a template and then cloned into the pCI-neo vector (Promega, Madison, USA). In addition, based on a bioinformatics analysis using PeptideMass (http://web.expasy.org/peptide_mass), three plasmids with mutated NS80 carrying Arg to Ala substitutions at residues 137, 194, and 219 were also generated. The three mutants were generated using the plasmid pCI-neo-NS80 as a template with the Site-directed Gene Mutagenesis Kit (Beyotime Company, Haimen, China). The mutantogenic oligonucleotide primer pairs are shown in Table 1. The
correctness of the recombinant plasmids was confirmed by sequencing (Invitrogen Biotechnology Inc., Shanghai, China).

**Infection**

GCRV infection with sensitive cells was performed as previously described (Fang et al., 1989). Briefly, CIK or FHM cells were plated one or two days in advance and infected with GCRV-873 virus stock at a multiplicity of infection (MOI) of 1–5. After 45 min of absorption, unabsorbed viruses were removed and cells were washed with phosphate-buffered saline (PBS, 137 mmol/L NaCl, 2.7 mmol/L KCl, 8.1 mmol/L Na$_2$HPO$_4$, 1.5 mmol/L KH$_2$PO$_4$; pH 7.3). They were further incubated with fresh medium supplemented with 2% FBS. The virus-infected cells were collected at various time points to detect NS80 protein expression by immunoblotting (IB) as described previously (Shao et al., 2013). The relative intensity of IB band was quantified using Image J software.

**Transfection and immunofluorescence assay**

The transfection with recombinant plasmids and immunofluorescence (IF) assay were conducted following established methods, as previously described (Guo et al., 2013). In brief, cells were transfected with the indicated plasmid DNA and Lipofectamine 2000 transfection reagent according to the manufacturer’s instruction (Invitrogen). At 24-h post-transfection (p.t.), cells were fixed and subjected to fluorescence microscopy. Transfected cells were fixed for 20 min at room temperature in 4% paraformaldehyde, washed three times with PBS, and permeabilized for 10 min with 0.2% Triton X-100. They were then washed and blocked three times for 2 h at 37 °C in PBS containing 5% bovine serum albumin. Then, cells were incubated with primary antibodies at 37 °C for 2 h. Three washes were conducted and secondary antibodies were added. After incubation at 37 °C for another 1 h, cells were washed three times with PBS and stained with Hoechst for 5 min. Finally, samples were examined with an Olympus-IX51 inverted fluorescence microscope and images were obtained and processed using Adobe Photoshop (Adobe Systems, San Jose, USA).

**IB analysis**

For the IB analysis, both infected and transfected cell lysates were collected, pelleted, and resuspended in PBS before they were resolved by sodium dodecyl sulfate-polyacrylamide gel (SDS-PAGE). They were then transferred to polyvinylidene fluoride membranes using a semi-dry transfer cell (Bio-Rad, Hercules, CA, USA) for 45 min at 100 mA. Membranes were blocked with 5% bovine serum albumin and then incubated with NS80 pAbs or mouse anti-FLAG mAb as primary antibodies. Alkaline phosphatase-coupled goat anti-rabbit or goat anti-mouse IgG were used as the secondary antibodies. Finally, reactive bands were detected with nitroblue tetrazolium (NBT) /5-bromo, 4-chloro, 3-indolylphospho- (BCIP) alkaline phosphatase substrate solution.

**RESULTS**

An approximately 58-kDa fragment was detected in infected and transfected cells

In a previous immunological assay using α-NS80C in infected cells, we found that a short polypeptide exhibits NS80 expression (Fan et al., 2009b). However, whether the small fragment is related to NS80 expression in different infected cells and its precise molecular mass are unclear. To resolve these issues, additional infection and transfection experiments in various cell types were performed. Two specific protein bands were detected in virus-infected and transfected cells in an IB analysis with α-NS80C. As shown in Figure 1A, a specific band of approximately 80 kDa was detected in virus-infected cells, which corresponded to the expected size of NS80. A slightly shorter polypeptide with a molecular mass of approximately 58 kDa was also detected using α-NS80C. The two specific bands were also detected in transfected cells (Figure 1B). These results suggested that the 58-
The 58-kDa fragment is cleaved from the amino terminus of NS80

To determine whether the short polypeptide was generated by cleavage, rather than via an alternative start codon, recombinant constructs expressing FLAG-tagged NS80 at the N- or C-terminus were constructed. Each of the two plasmids was transfected into BHK cells, and the cell lysates were subjected to an IB analysis with an anti-NS80C antibody. As shown in Figure 2A, a band corresponding to the full-length NS80 and a slightly shorter band were detected for the three missense mutants, indicating that the three mutated sites were not related to the 58-kDa fragment. To determine whether the three mutant proteins were substantially misfolded and targeted for degradation by the ubiquitin-proteasome system in transfected cells, colocalization assays for the three NS80 mutants using polyubiquitination antibody were performed. As shown in Figure 2B, none of the three mutant proteins appeared to be aggregated or strongly colocalized with conjugated ubiquitin in transfected cells. These results suggested that the three mutated sites in the N-terminal region of NS80 were unrelated to the 58-kDa fragment.

Infection time and dose influence the accumulation of the short NS80 fragment

After excluding the possibility that the short NS80 fragment was a transient product of protein degradation, the expression of NS80 at various time points after infection was examined in order to understand the timing of cleavage events. Cells were infected with GCRV at an MOI of 2.5, and collected at various time points. As shown in Figure 3A, the band of the short polypeptide could be recognized at 9 h p.i., which was earlier than the observed detection time for 1 MOI infection in previous report (Yan et al., 2015). However, the expression of the full-length NS80 protein could be detected at 6 h p.i., which was consistent with previous results (Yan et al., 2015). This observation suggested that the shorter fragment is generated from the cleavage of NS80. To determine the relationship between the amount of the 58-kDa fragment and infection dose, the expression of NS80 in GCRV-infected cells at an MOI of 1.5 was examined (IB data not shown). The relative intensities of the 58-kDa fragment for 3 different MOIs (1, 1.5, and 2.5) were evaluated. As shown in Figure 3B, the relative amount of the 58-kDa fragment was related to infection time and dose. As the infection progressed, the intensity of the 58-kDa polypeptide bands increased slightly 12h p.i., and remained at a stable level 15h p.i.. These results suggested that infection time and dose influence the amount of the 58-kDa fragment of NS80.
Identification of cleavage fragments of NS80

Figure 2. Three mutated protease-sensitive sites are not related to the 58-kDa fragment. (A) Immunoblotting analysis of the expressed mutant NS80 protein. BHK cells were transfected with 4 µg/well pCI-neo empty vector, pCI-neo-NS80, pCI-neo-NS80<sup>R137A</sup>, pCI-neo-NS80<sup>R194A</sup>, pCI-neo-NS80<sup>R219A</sup>, and pCI-neo-NS80 (130–742). Cell lysates were resolved by 10% SDS-PAGE and subjected to an immunoblotting analysis with α-NS80C or β-actin mAb. Arrow indicates the full-length NS80, and the asterisk indicates the 58-kDa NS80 fragment. (B) Ubiquitination analysis of the expressed mutant NS80 protein in transfected cells. Vero cells transfected with plasmids pCI-neo-NS80, pCI-neo-NS80<sup>R137A</sup>, pCI-neo-NS80<sup>R194A</sup>, and pCI-neo-NS80<sup>R219A</sup> were immunostained with α-NS80C and then counterstained with rabbit anti-polyubiquitination pAb, followed by staining with Alexa Fluor® 488 donkey anti-rabbit IgG (Green) and Alexa Fluor® 568 donkey anti-rabbit IgG (red). Nuclei were stained with Hoechst (blue).
FLAG mAb. As shown in Figure 4A, using the anti-FLAG mAb, the full-length NS80 and a ~22-kDa protein band were detected when FLAG was tagged at the N-terminus of NS80 (FLAG-NS80), but a ~22 kDa was not found for C-terminal-tagged NS80 (NS80-FLAG), except for the full length NS80 and an approximately 58-kDa protein band. As expected, there were no detectable bands for the expression of non-tagged-NS80 and the empty vector. These results suggested that the 58-kDa fragment is C-terminal NS80, and the 22-kDa fragment might be another product from the N-terminus of NS80.

To further prove that the 58-kDa fragment is C-terminal NS80, and the 22-kDa fragment might be another product from the N-terminus of NS80.

Figure 3. Infection time and dose may influence the amount of the short NS80 fragment. (A) CIK cells infected with GCRV at an MOI of 2.5 were collected at different time points to detect the expression of NS80 by immunoblotting with α-NS80C or β-actin mAb. Arrow indicates the full-length NS80, and the asterisk indicates the 58-kDa NS80 fragment. (B) The relative intensity of the 58-kDa NS80 fragment was quantified using Image J. CIK cells were infected with GCRV at three different MOI (1, 1.5, and 2.5), and the relative intensity of the shorter fragment was determined with respect to β-actin and the maximum value was evaluated.

Subcellular localization of the 58-kDa polypeptide is different from 22-kDa fragment of NS80

The aforementioned results showed that NS80 could be cleaved into two fragments (22 kDa and 58 kDa) in infected and transfected cells. To determine the subcellular localization of the two NS80-related fragments, the localization of the 22-kDa or/and 58-kDa fragments in transfected cells was examined by IF assays with the pCI-neo-FLAG-NS80 (1–190) plasmid expressing the N-terminal NS80 protein and the pCI-neo-NS80 (386–742) plasmid expressing the C-terminal NS80 protein. As shown in Figure 5A, pCI-neo-NS80 (1–190) exhibited a diffuse subcellular distribution using α-NS80N, while pCI-neo-NS80 (386–742) appeared in typical globular inclusion structures in the cytoplasm using α-NS80C. To confirm the transfection results, the expression of NS80 in infected cells was further examined by IF using α-NS80N and α-NS80C. As shown in Figure 5B, many typical globular inclusion structures were detected in infected cells using both α-NS80N and α-NS80C, but a diffuse subcellular distribution was only detected using α-NS80N. These results suggested that the two cleavage fragments had distinct subcellular localizations, indicating that they might have different functions in viral replication.

In our previous study, we found that NS80 interacts...
Figure 5. Subcellular localization of the 58-kDa polypeptide is different from that of the 22-kDa fragment of NS80. (A) Vero cells transfected with pCI-neo-NS80 (1–190) or/and pCI-neo-NS80 (386–742) were fixed at 24 h p.t. for immunofluorescence assays with α-NS80N and α-NS80C, respectively, followed by Alexa Fluor® 488 donkey anti-rabbit IgG (Green). Nuclei were stained with Hoechst (blue). (B) FHM cells infected with GCRV were fixed at 12 h p.i. and stained with α-NS80N and α-NS80C, followed by Alexa Fluor® 488 donkey anti-rabbit IgG (Green). Nuclei were stained with Hoechst (blue). (C) Vero cells were co-transfected with NS38-GFP-NSP5 and either pCI-neo-NS80(1–190) or pCI-neo-NS80(386–742). At 24 h p.t., cells were fixed and then stained with α-NS80N and α-NS80C, followed by Alexa Fluor® 568 donkey anti-rabbit IgG (red). Nuclei were stained with Hoechst (blue).
with NS38 in infected and transfected cells (Shao et al., 2013). Additionally, the single rotavirus NSP5 is able to form viral factories in transfected cells (Miller et al., 2010). Accordingly, the NSP5-based protein–protein interaction platform including NS38-GFP-NSP5 with the plasmids NS80 (1–190) or/and NS80 (386–742), respectively. As shown in Figure 5C, N-terminal NS80 fully colocalized with NS38-GFP-NSP5, while no association was detected between the C-terminal NS80 and NS38-GFP-NSP5. Therefore, there is an interaction between NS38 and the 22-kDa fragment, but not the 58-kDa fragment. These results demonstrated that the 22-kDa and 58-kDa fragments of NS80 play different roles in viral replication.

**DISCUSSION**

Previous studies have indicated that reoviruses generate inclusion bodies in the cytoplasm during infection. The nonstructural protein μNS of MRV and its analogue proteins also interact with other viral or cellular proteins involved in viral replication and particle assembly. Moreover, these interactions between μNS and multiple viral proteins promote the maintenance within VIBs for efficient viral replication. Recent studies in our lab have indicated that Aquareovirus NS80 can induce inclusion structures in infected or transfected cells and interact with other viral proteins (Becker et al., 2003; Tousis-Otero et al., 2004; Broering et al., 2005; Brandariz-Nunez et al., 2010). These interactions between μNS and multiple viral proteins promote their maintenance within VIBs for efficient viral replication. Recent studies in our lab have indicated that Aquareovirus NS80 can induce inclusion structures in infected or transfected cells and interact with other viral proteins (Becker et al., 2003; Tousis-Otero et al., 2004; Broering et al., 2005; Brandariz-Nunez et al., 2010). These observations suggest that viruses use a variety of strategies to modify primary translation products for efficient viral replication and particle assembly.

The μNSC of MRV forms inclusion bodies. However, based on RNA interference-based trans-complementation approaches, μNSC is incapable of restoring MRV growth in cultured cells when μNS is absent (Kobayashi et al., 2006; Arnold et al., 2008; Kobayashi et al., 2009). We recently found that the C-terminal region of NS80 is indispensable for VIB formation, and the N-terminal region of Aquareovirus NS80 is required for interactions with viral proteins and viral replication (Shao et al., 2013; Zhang et al., 2016); accordingly, the N- and C-terminal regions of NS80 may play different roles in viral replication and particle assembly. In this study, we detected 22-kDa and 58-kDa NS80 fragments, as well as the full-length NS80 with NS80 expression in infected and transfected cells. The 22-kDa N-terminal NS80 showed a diffuse subcellular distribution that differed from the 58-kDa C-terminal NS80 and full-length NS80 phenotypes, suggesting that NS80 presents different forms at different replication stages for efficient replication and assembly in the viral life cycle. The precise cleavage site of the NS80 fragment and the molecular mechanism by which it mediates viral replication are currently under investigation.

In summary, the origin of two NS80 fragments and their related subcellular distributions were investigated. The 58-kDa product was derived from the cleavage of NS80. This cleavage yielded an approximately 58-kDa carboxy-terminal protein and a 22-kDa amino-terminal polypeptide, which exhibited distinct distributions in transfected and infected cells. To our knowledge, this is the first study to indicate that the 58-kDa fragment is derived from the cleavage of the NS80 protein. These results contribute to our understanding of the molecular events involving NS80 and the roles of different forms in interactions with other viral or cellular proteins involved in viral replication.
in viral replication and particle assembly.

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COMPLIANCE WITH ETHICS GUIDELINES

The authors declare that they have no conflict of interest. This article does not contain any studies with human or animal subjects performed by any of the authors.

AUTHOR CONTRIBUTIONS

QF designed the experiments. QXC, ZJ carried out the experiments. QXC, ZJ, GH, FXZ and QF analyzed the data. QXC, QF wrote the paper. All authors read and approved the final manuscript.

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