Prokaryotic expression and purification of grass carp reovirus capsid protein VP7 and its vaccine potential

Yongxing He¹, Qian Yang¹, Hongxu Xu², Hao Wu¹, Fangyuan Wu¹ and Liqun Lu¹*

¹Key Laboratory of Aquatic Genetic Resources and Utilization /Ministry of Agriculture, Shanghai Ocean University, 201306 Shanghai, China.
²Department of Laboratory Medicine, The First Affiliated hospital of Sun Yat-sen University, Guangzhou 510080, China.

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The 11 dsRNA fragmental genome of grass carp reovirus (GCRV) is enclosed in five inner core proteins and two outer capsid proteins. The Glutathione S-transferase (GST) fusion protein expression vector pGEX-4T-3 was employed to clone and express of GCRV outer capsid gene vp7, which was amplified by reverse transcription-polymerase chain reaction (RT-PCR) from infected Grass carp. The recombinant GST-fusion protein rVP7 was induced by 1 mM IPTG in Dh5α and confirmed by SDS-PAGE and Western blot assays using both anti-GST-tag and anti-VP7 monoclonal antisera. An expected 52-kDa rVP7 was highly expressed, and was mainly exhibited in the formation of the inclusion body. After purification, rVP7 was intraperitoneally injected to the experimental mice to produce anti-rVP7 polyclonal serum. In vitro microneutralization assay indicated that polyclonal antibody against rVP7 could neutralize GCRV, and suggested that rVP7 had the potential to be used as subunit vaccine against GCRV infection. The present study paved the way for further characterization of the immunogenicity of viral outer capsid protein VP7 in grass carp Ctenopharyngodon idellus and could be based to develop antibody or antigen detection assays for GCRV pathogen.

Key words: Grass carp reovirus, VP7 protein, prokaryotic expression, western blot, microneutralization.

INTRODUCTION

Grass carp reovirus (GCRV), as a prototype member of aquareovirus, is a pathogen causing pandemic hemorrhage disease of the grass carp Ctenopharyngodon idellus and identified as the most causative pathogenic aquareovirus with high mortality (Qiu et al., 2001; Cheng et al., 2008). For the reason that grass carp C. idellus is the first cultured fish species, efficient and economic preventative strategy of GCRV infection is urgently needed in China.

Due to the lack of study on the immunogenicity and function of GCRV proteins, information on the structural proteins of GCRV was mainly deduced from its mammalian counterpart mammalian orthoreoviruses (MRV). Among the structural proteins of GCRV, VP7 and VP5 are the outer capsid proteins of the virus. VP7 is encoded by the S10 gene fragment and composed of a total of 276 amino acid residues (Benavente et al., 2007). The 3D structure and genome sequences demonstrated that there is a higher level of sequence homology in structural proteins between GCRV and MRV (Cheng et al., 2010), but vp7 gene of GCRV and MRV shares the most divergent sequences and lowest amino acids identity (Jaafar et al., 2008; Cheng et al., 2008; Attoui et al., 2002).

In the past several decades, progress has been made in the research on the grass carp hemorrhagic disease in China. The reverse transcription-polymerase chain reaction (RT-PCR) techniques have been developed for pathogen diagnosis (Li et al., 1999; Seng et al., 2004; Zhang et al., 2008, 2010). The most important advances about GCRV includes the biochemical and ultra-structural characteristics of GCRV and molecular strategies on diagnosis of GCRV (Ma et al., 2008; Guo et al., 2010). Vaccination has proven to be a very useful strategy in controlling pathogens in aquaculture, particularly bacterial pathogens (Heng et al., 2011). Furthermore, polyclonal antibodies against μ1 and 63 were reported able to...
neutralize Mammalian reovirus in vitro (Chandran et al., 1999). Current vaccines include inactivated vaccine and attenuated vaccine, whose features were the high costs and difficulty in administering it in aquatic environment. Subunit and oral vaccine remain urgent and urgently desired against GCRV for the fish cultivation industry (Rangel et al., 1999; Heng et al., 2011). Additionally, Zhang et al. (2008) had expressed His-tagged VP7 protein in *Escherichia coli* and develop the protein-based detection assay for GCRV, but the vaccine potential of VP7 remians unknown. In this paper, we investigated the vaccine potential of GST-fused rVP7 through in vitro microneutralization assay.

**MATERIALS AND METHODS**

**Virus isolation and propagation**

GCRV infected grass carp fingerlings (3 to 4 months of age) were obtained from a fish farm in Jiangsu, China. GCRV isolates from infected fingerling grass carp with typical hemorrhagic symptoms (dead and moribund fish fingerlings) were inoculated in the *C. idellus* kidney cell (CIK). CIK cells were maintained as stock cultures in DMEM (Dulbecco's Minimal Essential Media) and replated 2 days before infection assays.

**Viral RNA extraction and RT-PCR**

GCRV purification and RNA extraction followed the reported methods (Seng et al., 2002). RT-PCR assay was carried out by using total RNA from tissue infected with GCRV as template. To obtain virus RNA from infected tissue, genomic dsRNA was extracted from purified GCRV using a Trizol method (Invitrogen). For preparing RNA from virus-infected cell culture, 0.5 to 1.0 ml of virus-infected cell supernatant was pelleted by centrifugation at 8000 rpm for 15 min to pellet the virus-cell suspension for further total RNA extraction. Total RNA was obtained according to the manufacturer’s instructions and was resuspended in DEPC-treated water and stored at −80°C until use.

For RT-PCR, 500 ng of total RNA were used as template for a two-step reverse transcription reaction. The reaction mixture (20 µl) contained 5 µl 10× reaction buffer, 1 µl of the oligo(dT)12-18 primer, 2.5 µl of dNTPs (2.5 mM each), 2 µl of dithiothreitol (10 mM), 1 µl of RNAase inhibitor (40 U/µl), 1 µl of M-MLV reverse transcriptase (200 U/µl) and 1 µl of first-strand cDNA synthesis mixture. The reactions were performed as follows: 5 min at 94°C for pre-denaturation, followed by 34 cycles each consisting of denaturation at 94°C for 30 s, annealing at 50°C for 30 s and extension at 72°C for 30 s, with the final extension at 72°C for 10 min by using a DNA thermal cycler. Form each reaction amplicons were analyzed in 1.0% (w/w) agarose gels in TAE buffer for 30 min and followed by UV-light transillumination and image capture.

**Constructions of expression vector**

Cloning sites BamH1 and Sal1 were selected according to the multiple cloning sites of the Glutathione S-transferase (GST) fusion protein expression vector pGEX-4T-3 (GE Healthcare) and VP7 gene sequence. Recombinant plasmid was confirmed by restriction enzyme digestion and sequencing (Co., Ltd. Shanghai Sangon). The correct recombinant plasmid was named as pGEX-4T-vp7 containing the full length VP7 cDNA was identified according to the size of inserted segment by PCR and was purified using a plasmid DNA purification miniprep kit (Promega).

**Cultivation and induction condition of the recombinant protein**

*E. coli* strain DH5α was cultured in Luria-Bertani medium (LB) containing 100 µg/ml ampicillin and grown overnight at 37°C and 180 rpm. The pre-inocula were then transferred to 500 ml LB medium containing ampicillin of the same concentration at a ratio of 1:10. Expression of GST fusion protein was induced with 0.1 mM isopropyl-β-d-thiogalactopyranoside (IPTG) for 4 h at 37°C.

**Expression and purification the recombinant protein**

The GST-rVP7 contained in inclusion bodies was extracted as described previously (Zhang et al., 2008; Pathak, 2008) with some modifications. Briefly, after ultrasonic treatment, the sediment was collected at 4°C by centrifugation at 12000 rpm for 15 min. The samples (post-induction samples, host bacteria transformed with empty expression vector) were suspended with equal volume of 2 x SDS lysis buffer (Takara). The suspensions were incubated for 5 min at 100°C with stirring. The resulting cell lysate was centrifuged at 12000 rpm for 10 min. Then the extracted *E. coli* proteins were resolved by SDS-PAGE on vertical slab gels (5% stacking and 12% resolving gel). Protein bands were stained with 2.5% Coomassie Blue R-250 for visualisation with control wide molecular range markers (Takara) as molecular estimates. Protein sample of GCRV virions were also run here for comparison purposes. The recombinant protein was purified with the PAGE gel extraction kit (Co., Ltd. Shanghai Sangon). The purified recombinant protein was subjected for further SDS-PAGE analysis.

**Western blotting analysis**

For western blotting, the gel was soaked in transfer buffer (20 mM tris-HCl, 196 mM glycine, 40%(v/v) methanol, pH 8.0). The resolved proteins were electro-transferred on to the nitrocellulose membrane according methods described previously (Yin et al., 2004; Kayali et al., 2008). The membrane was then blocked in phosphate-buffered-saline (PBS) supplemented with 5% milk powder and diluted rat monoclonal IgG (1:2000) or diluted polyclonal antibody (1:300). Color development was performed by incubating with alkaline phosphatase-conjugated anti-mouse IgG (Sigma), followed by adding substrates: bromocholoroindolyl phosphate (BCIP) and nitroblue tetrazolium (NBT), or with alkaline phosphatase-conjugated anti-mouse IgG (Sigma), followed by adding substrates: bromocholoroindolyl phosphate (BCIP) and nitroblue tetrazolium (NBT), or with alkaline horseradish peroxidase- conjugated anti-Rabbit IgG (Ding National Bio Co., Ltd), and followed by adding diaminobenzidine (DAB) (Co., Ltd. Shanghai Sangon) substrate.

**Antisera preparation**

Four female and four male 4-week-old Balb/c mice were purchased from the Animal Holding Unit (AHU) of the Second Military Medical University (Shanghai, China). The mice were injected and followed the way as (Yu et al., 2008; Ma et al., 2009) intra-peritoneally (i.p.) with an emulsion of purified protein and Freund’s complete adjuvant (FCA) (1:1 in volume). Two weeks later, a mixture of (1:1 in volume) of purified protein and Freund’s incomplete adjuvant was administered. Finally, two subsequent i.p. and intravenous (i.v.) injections of purified protein were administered at 2 weeks interval. The mice were then bled 3 days after the last injection. Serum was collected at 4°C for overnight and stored until further use.

**Microneutralization assay**

Neutralization effect was measured using GCRV viruses to infect CIK cells as described previously (Yang et al., 2007; Cohen et al., 2007). Briefly, after series of two-fold diluted heat-inactivated serum
samples (1:20 to 1:2560) were loaded in 96-well tissue culture plates, about 50 TCID50 GCRV were added, and the plates with virus-serum dilution mixtures were incubated at 27°C for 48 h to allow neutralization to take place. Mock-infected cells served as positive control here, while infected cells with pre-immunized control serum served as negative control. The assay was performed in triplicate and the viral cytopathic effect (CPE) was observed every 6 h. To visualize the results, the plate was stained with crystal violet-formaldehyde stain (0.013% crystal violet, 2.5% ethanol, and 10% formaldehyde in 0.01 M PBS). The crystal violet solution was then removed to a designated waste container. Finally, the cell monolayer in each well was rinsed with approximately 200 µl, 1×PBS for image capture.

**RESULTS**

**Extracted RNA used for GCRV segment amplification**

To obtain the coding sequence of VP7 protein from total RNA extracted from virus-infected cells, RT-PCR was performed by using specific primer pairs for VP7 gene. Figure 1 showed that fragment up to 0.9 kb in length could be detected by RT-PCR from the total RNA sample.

**Expression and purification of recombinant protein**

When the Vp7 gene was cloned into the prokaryotic expression vector pGEX-4T-3, it was confirmed by restriction enzyme digestion and sequence analysis (data not show). To obtain the over-expressed recombinant protein, bacteria containing pGEX-4T-vp7 was induced by IPTG at 4 h. A protein of 52 kDa in molecular weight was induced by IPTG, which was consistent with the expected size of rVP7 (Figure 2, Lane 3). We found that the rVP7 protein mainly exists in the form of inclusion bodies rather than in the supernatant. Then we used a protein purification kit (Co., Ltd. Shanghai Sangon) to purify rVP7 from PAGE gel. Lane 4 of Figure 2 demonstrated the purified rVP7 for polyclonal antibody production.

**Western blot analysis with rVP7 and polyclonal anti-rVP7**

Polyclonal antibody, anti-rVP7, was produce in mice by using the purified rVP7 as immunogen. To test the specificity of the antiserum, extracts of the rVP7- expressing bacteria was immunoblotted with anti-rVP7 serum (Figure 3a), as well as the monoclonal anti-GST antiserum as control (Figure 3b). It appeared both anti-GST and anti-rVP7 could specifically recognize rVP7 (Lane 3 of Figures
Figure 3. Western blotting analysis of the rVP7 and viral VP7. Proteins from 12% PAGE gels were transferred to nitrocellulose. (a) membrane was probed with anti-GST monoclonal antiserum and detected by NBT-BCIP substrate, Lane 1: Molecular weight maker; Lane 2: total cell extract of uninduced bacterial; Lane 3: total cell extract of bacterial after 4 h induction with 1mM IPTG at 37°C; (b) membrane was probed with anti-rVP7 polyclonal antiserum and detected by DAB substrate. Lane 1: Molecular weight maker; Lane 2: total cell extract of uninduced bacterial; Lane 3: total cell extract of bacterial after 4 h induction with 1 mM IPTG at 37°C; (c) membrane was probed with anti-rVP7 polyclonal antiserum and detected by DAB substrate. Lane 1: Molecular weight maker II; Lane 2: cellular extract of GCRV infected cells.

3a and b), in contrast to the uninduced bacteria (Lane 2 of Figures 3a and b). The anti-rVP7 could detect VP7 from the GCRV-infected cells (Figure 3c), which confirmed the viral origin of rVP7.

**Microneutralization assay and virus titration**

As described previously, the anti-VP7 polyclonal antibody was able to recognize the viral VP7 protein. An *in vitro* microneutralization assay was performed to measure the neutralizing ability of this polyclonal antibody. Figure 4a indicated that the anti-rVP7 antibody had strong neutralizing capability (1:640) for GCRV. For more accuracy, the supernatant of the infected cells in 96 well plates was collected for viral titration assay. The viral production reached 10^7 TCID50/ml for the cells infected with GCRV when mixed with anti-VP7 sera of 1:1280 and 2560 dilution, while low viral progeny of 10^2-10^3 TCID50/ml was observed for the wells with anti-VP7 sera ranging from a dilution of 1:20 to 1:640 (Figure 4b).

**DISCUSSION**

Up to date, aquaculture in the world suffers from the frequent outbreak of epidemic disease. The grass carp reovirus is identified and regarded as the most pathogenic aquareovirus (Zhang et al., 2010). GCRV particle is composed of 7 proteins, VP1-VP7. Among the 7 structural proteins, the VP1, VP2, VP3, VP4 and VP6 are involved in forming the viral inner core, while the remaining VP5 and VP7 proteins comprise the outer capsid shell of the virus (Cheng et al., 2008). Among all the above proteins, only the outer capsid protein VP5 is seen to be reported as a subunit vaccine (He et al., 2011).

Current inactivated and attenuated vaccines have been limited because their high costs and inconvenience (Heng et al., 2011). Based on its neutralization ability, the passive administration of neutralizing outer capsid VP7 polyclonal antibodies could provide an immediate treatment strategy for emergency prophylaxis and treatment.
Serum dilution

Virus titration

Log TCID₅₀/ml

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<th>Dilution</th>
<th>Value</th>
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<tr>
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<tr>
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OFFENDING COMMAND: .buildcmap

STACK:

-dictionary-
/WinCharSetFFFF-V2TT9BF4ACCAt
/CMap
-dictionary-
/WinCharSetFFFF-V2TT9BF4ACCAt