Monoclonal antibodies developed for detection of an epizootic virus associated with mass mortalities of cultured scallop *Chlamys farreri*

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**ABSTRACT:** Recently, an enveloped, spherical RNA virus was identified as the causative agent of mass mortalities among adult scallop *Chlamys farreri*, which is cultured on the northern coast of China. Hybridomas were prepared from mice immunized with highly purified virions. Four stable hybridomas secreting monoclonal antibodies (MAbs) of IgG isotype were obtained after screening by means of enzyme-linked immunosorbent assay (ELISA) and immunofluorescence assay (IFA). The specificity of the MAbs to this virus was confirmed by immunogold electron microscopy (IEM). All the selected MAbs recognized epitopes on the envelope spikes of the virions. Subsequently, the MAbs were used for *in situ* immunofluorescent detection of the virus in Davidson’s fixed tissue sections. The results showed that the fluorescent cells were mostly observed in epithelia of different organs, but not in the epithelium of the digestive diverticulae. Cytopathological changes and focal lesions corresponding to virus-positive cells were clearly recognized in the affected epithelia, revealing a potential role of this virus in pathogenesis.

**KEY WORDS:** Scallop · *Chlamys farreri* · Acute virus necrobiotic disease · Monoclonal antibodies

**INTRODUCTION**

The scallop *Chlamys farreri* is widely farmed along the northern coast of China. In the 1980s, an abnormal occurrence of mass mortality among cultured and wild adults was initially reported in some regions of Shandong Province (Liu et al. 1992). Since the mid-1990s, the scallop farming industry in China has experienced a period of severe crisis due to ongoing outbreaks of mass mortality. Mortality occurred annually during late summer and could approach 90% within 5 to 8 d after the event began (Yu et al. 1998, Guo et al. 1999).

In a search for the causative agent, a number of tentative hypotheses have been proposed (Wang & Xiang 1999, Zhang & Yang 1999, Li et al. 2002, Xiao et al. 2003). More recently, supported by the ‘Key Project of Chinese National Programs for Fundamental Research and Development (973 program)’, systematic investigations including epidemiological surveys (X. H. Wang et al. 2002, He et al. 2003), RNA studies on the causative agent (Huang et al. 2002) and experimental infections (Ai et al. 2003) have been carried out by our research group since 2000. All results suggested that the causative agent is an unknown spherical virus. Based on the results obtained, the disease was named ‘acute virus necrobiotic disease (AVND)’ (Song et al. 2001).

Currently, the tissue distribution of the AVND virus and its role in pathogenesis are still obscure. Hence, the main goal of the present study was to develop reliable monoclonal antibodies (MAbs) against the AVND virus to allow sensitive detection of virus antigens and a simultaneous evaluation of histopathological changes in diseased scallops.

**MATERIALS AND METHODS**

**Virus collection and purification.** Moribund scallops *Chlamys farreri* (14 to 15 mo old) were collected during
mass mortalities in mid-July 2001 and 2002 from several private farms in Qingdao and Rizhao (Shandong Province of China). The gills and mantles were removed from the scallops and stored at −85°C before use. Virus purification by centrifugation was performed according to C. M. Wang et al. (2002). Tissues were homogenized in 8 volumes of TEN buffer (50 mM Tris-HCl; 10 mM EDTA; 360 mM NaCl, pH 7.8) and clarified by 2 low speed centrifugations (3500 × g for 15 min followed by 7500 × g for 15 min). All centrifugations were carried out at 4°C. The supernatants were then concentrated by sedimentation through a 6 ml cushion of 35% sucrose (w/w) at 113 000 × g for 90 min and the resulting pellets were resuspended in TEN buffer. After centrifugation at 8000 × g for 20 min, supernatants were layered onto a 30 to 60% (w/w) continuous sucrose density gradient. After centrifugation at 113 000 × g for 2.5 h, the opalescent virus containing band was collected. The virus was washed with TN buffer (50 mM Tris-HCl; 360 mM NaCl, pH 7.8) and resuspended in a small amount of TN buffer. The purity was checked by a transmission electron microscopy (TEM, JEOL JEM-1200) using 2% phosphotungstic acid (PTA, pH 7.0) as the negative stain. The concentration of viral proteins was determined using the procedure described by Bradford (1976).

Hybridoma production. Eight week old BALB/c mice were immunized intraperitoneally (i.p.) with purified virus (100 µg mouse−1) mixed with an equal volume of Freund's complete adjuvant (Gibco). Subsequent boosters with antigen in incomplete adjuvant were given on Days 14, 28, and 42. Three days before cell fusion, mice received the last immunization via intrasplenic injection (20 µg mouse−1) with the purified virus diluted in phosphate buffered saline (PBS), but without adjuvant. Antibody titers to AVND virus in the mice were checked by means of enzyme-linked immunosorbent assay (ELISA) as described below.

Spleen cells from the immunized mice were fused with NS-1 myeloma cells at a ratio of 5:1 using polyethylene glycol (PEG 2000, 50%). The fused cells were resuspended with RPMI-HAT (Gibco) medium and plated into 96-well microplates in the presence of normal mouse spleen cells serving as feeder cells. After about 1 wk of cultivation, HAT medium was partially replaced by HT (Gibco) medium and hybridoma supernatants were screened for antibody production from Day 10 onwards by ELISA. Positive hybridomas were subsequently cloned 3 times by limiting dilution. The immunoglobulin (Ig) isotype of the MAbs was determined using a Mouse Monoclonal Antibody Isotyping Kit (Sigma) according to the manufacturer's instructions.

Hybridoma supernatants with high antibody titers were collected and applied directly. Antibodies were also produced by injecting hybridomas into the peritoneal cavity of 10 wk old BALB/c mice. After hybridoma cell proliferation, ascitic fluids with high antibody titers were obtained within 2 to 4 wk.

ELISA. An ELISA method was standardized to screen the hybridoma supernatants. Briefly, the purified virus was solubilized with 1% SDS and diluted in carbonate-buffer (pH 9.6) (Lucht et al. 2003). Then, the virus antigen was coated to the solid phase of 96-well high binding EIA/RIA plates (Costa) overnight at 4°C. After removing the antigen fluid, nonspecific binding sites were blocked with 3% bovine serum albumin (BSA)-PBS, followed by incubation for 30 min at 37°C. The wells were emptied and hybridoma supernatants were added and incubated for 1 h at 37°C. After washing the plates 3 times for 5 min by immersion in PBST (0.05% Tween-20 in PBS), HRP-conjugated goat anti-mouse IgG (H&L) (Sigma; 0.15% H2O2) was applied for 15 min at room temperature. The reaction was stopped by 2 M H2SO4 and the optical density (OD) of each well was read at 492 nm using a plate reader (Labsystems Mutiskan MK3).

Positive controls (P) corresponded to immunized mouse sera (diluted 1:16 000), obtained 3 d before cell fusion. Homogenate (diluted 1:1000) of healthy scallop tissues (mantle and gill) served as negative controls (N). These healthy scallops were sampled in spring 2003 and were examined by TEM to confirm viral-free status. Supernatants of OD_{492 nm} P/N ≤ 2.1 were selected for further cloning and subsequent testing by immunofluorescence assay (IFA).

IFA on fixed sections. Diseased (moribund) scallops were collected from a private farm during mass mortality in 2003. Healthy scallops serving as negative controls were the same as above. Mantles, gills, and parts of kidney, gonad, hepatopancreas, stomach and intestine were dissected, fixed with cold Davidson's alcohol-formalin-acetic solution for 24 to 48 h, and stored in 70% ethanol solution before use. The specimens were processed using a routine laboratory procedure (Lighter 1996). For immunofluorescence staining, sections were deparaffinized in xylene and rehydrated in graded ethanols, followed by washing 3 times with PBST. The sections were incubated with hybridoma supernatants in a moist chamber for 1 h at 37°C. After washing 3 times with PBST, the sections were incubated with FITC-conjugated goat anti-mouse IgG (H&L) (diluted 1:10) in a dark moist chamber for 1 h at 37°C. After extensive washing, the slides were mounted in 50% glycerol and examined with a fluorescence microscope (Olympus B ×50).
Immunogold electron microscopy (IEM). Formvar-coated grids were placed on to 25 to 30 µl purified virus aliquots for 1 min. All subsequent steps were performed by floating the grids with the specimen side down on aliquots of solutions, in a moist chamber with parafilm at its base at room temperature (Gowen et al. 2003). After washing 3 times with PBS, nonspecific binding was controlled by incubation with 3% BSA-PBS for 30 min. The grids were then exposed for 1 h to the hybridoma supernatants. The grids were washed 3 times with PBS and then transferred to drops of 10 nm colloidal gold-conjugated goat anti-mouse IgG (whole molecule) (Sigma, G7777; diluted 1:10 in 3% BSA-PBS) for 1 h. After all labeling steps, the grids were washed 3 times for 5 min each time with PBS and 3 times for 1 min with distilled water. Finally, the virus was stained with 2% PTA and viewed by TEM.

RESULTS

Virus purification

Virions purified by sucrose gradient centrifugation are shown in Fig. 1. The virions were spherical, and possessed a bilaminal envelope with dense spikes on its surface. They were about 130 to 170 nm in diameter, while the nucleocapsids were 90 to 140 nm.

Production and screening of hybridomas

The fused cells were cultured in 4 96-well plates. After cultivation in HAT medium for 2 wk, hybridomas survived in 249 wells and the fusion rate was about 65%. In the initial screening by ELISA, 12 wells containing hybridomas were capable of secreting antibodies that reacted with purified virus preparation. After subcloning 3 times to ensure monoclonality, indirect IFA was carried out to confirm their specificity to AVND virus. Ten of the 12 MAbs reacted with mantle and other tissues of diseased (moribund) scallops but not with any tissues of healthy scallops (negative controls). The other 2 MAbs reacted with hemocytes of diseased and healthy scallops (data not shown). Of the 10 MAbs that reacted with diseased tissues in IFA, 4 MAbs (2B3, 3C8, 3G7 and 4C7) were eventually chosen for further study because of their strong reactivity and high stability in culture (Table 1).

IEM

For confirmation purposes, the 4 selected MAbs were examined by immune electron microscopy and 10 nm colloidal gold was used as a marker to visually detect their combining positions on AVND virions. For all MAbs (2B3, 3C8, 3G7 and 4C7), specific labeling was observed on the envelope of AVND virions, indicating that the epitopes recognized by these 4 MAbs were on the envelope of the virus (Fig. 2).

Table 1. Characterization of selected monoclonal antibodies (MAbs). Immunoglobulin (Ig) class or subclass; reaction with acute virus necrobiotic disease (AVND) virus in ELISA and immunofluorescence assay (IFA). +: weak reaction; ++: strong reaction; +++: very strong reaction

<table>
<thead>
<tr>
<th>MAb</th>
<th>Subclass</th>
<th>Intensity of reaction in ELISA</th>
<th>IFA</th>
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<tbody>
<tr>
<td>2B3</td>
<td>IgG2b</td>
<td>+++</td>
<td>++</td>
</tr>
<tr>
<td>3C8</td>
<td>IgG1</td>
<td>+++</td>
<td>+++</td>
</tr>
<tr>
<td>3G7</td>
<td>IgG2b</td>
<td>++</td>
<td>+</td>
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<tr>
<td>4C7</td>
<td>IgG2b</td>
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Fig. 1. Negatively stained acute virus necrobiotic disease (AVND) virions purified from moribund Chlamys farreri by density gradient centrifugation. Intact virions with dense spikes on their envelope were clearly observed (arrow). Scale bar = 100 nm

Fig. 2. Monoclonal antibodies (MAB) specificity for viral surface epitopes demonstrated by immunogold labeling. The MAb used was 3C8. Scale bar = 100 µm
**AVND virus tissue distribution and histopathological changes**

Because MAb 3C8 demonstrated the strongest reactivity to AVND virus, it was used for immunofluorescence to detect the virus antigens in diseased scallops during mass mortality. In all cases (n = 30), virus-positive cells (fluorescent cells) were primarily observed in external (protective) epithelia and in the mucosal epithelia of the digestive tract (intestine, stomach). In certain cases, a few positive cells were also present in the connective tissue of some organs, e.g. mantle and gonad (data not shown). On transverse sections of whole mantle, the positive epithelial cells were distributed in a uniform pattern, i.e. dense aggregations of positive cells were only observed to be located in the inner epithelium of the mantle’s basal portion as well as in the basal portion of the gills (Fig. 3).

Cellular lesions associated with the positive cells were observed. The positive epithelial cells appeared swollen and disassociated from each other. A disorder or excessive sloughing phenomenon apparently related to the density of the positive cells was clearly recognized in the inner epithelium of the mantle’s basal portion. These histopathological changes were also observed in several organs, e.g. in the kidney and stomach (Fig. 4).

No fluorescence was seen in the sections from healthy scallops (serving as the negative control) and the blank control from which the primary antibody was omitted (data not shown).

**DISCUSSION**

Although viruses or virus-like particles have been observed ultrastructurally in many bivalves, direct
evidence linking viral infections with a specific disease is scarce (Elston 1997, Hine et al. 1997). Numerous studies have showed that bivalve molluscs can rapidly accumulate virus from contaminated water (Shieh et al. 1999, Muniain-Mujika et al. 2003). The most significant example may be infectious pancreatic necrosis virus (IPNV), a fish epizootic pathogen that can also be isolated from scallop Pecten maximus but which appeared to be non-pathogenic for adult scallops (Mortensen et al. 1992). In our study, the newly characterized MAbs improved our ability to diagnose infections caused by AVND virus and will make further investigations on the pathogenesis of this scallop etiologic virus possible.

Two screening methods, indirect ELISA and IFA, were used in our study. By ELISA, positive reaction occurred in 12 wells, and by IFA, in 10 wells. This discrepancy may be due to the antigen used in ELISA and the fact that the initial immunization was contaminated with host-derived proteins. This problem was also encountered in some previous studies attempting to develop MAbs to shrimp viral pathogens (Poulos et al. 1999, 2001, Zhan et al. 1999). Although the Western blot assay is a valuable testing format to characterize or confirm the specificity of the MAbs, it is also more time consuming and highly technical. The major drawback to this assay was the requirement that the virus preparation be highly purified. In our study, IEM was performed to further confirm the specificity of selected MAbs and visually detect their specific combining sites on AVND virions. Problems arising from impure virus preparations appear to have been circumvented. This protocol proved to be relatively simple and very rapid. The total time required to test a MAb was about 4 h.

In the present study, a rapid and sensitive IFA testing format was established based on the reliable MAbs and employed to detect the virus tissue distribution in scallop tissues. The results showed that most of the positive cells were located in epithelia of the diseased scallops, though only a few positive cells were simultaneously scattered in connective tissues of some heavily infected individuals. This suggested that AVND virus is epitheliotropic. Some sections were also stained with Harris hematoxylin and eosin (H&E) with the cytoplasm of the virus-positive epithelia cells appearing more basophilic than uninfected cells (data not shown).

No immunofluorescence was detected in the epithelium of the digestive diverticulae. It is worth pointing out, therefore, that all of the affected epithelial cells were those that in general communicated most directly with seawater. It is possible that when environmental conditions are favorable, the virus may replicate rapidly in infected epithelial cells and then upon rupture of these cells be released and transmitted to other scallops through the seawater. Epithelial disruption
would allow other opportunistic pathogens colonize the soft tissues, possibly contributing to acute mass mortality among the scallop population.

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