INTRODUCTION

Lymphocystis disease virus (LCDV) is the causative agent of lymphocystis disease. It has a worldwide distribution and has infected over 125 species of fish belonging to 42 families including feral, cultured and ornamental fish. Lawler et al. (1977, 1978) reported the occurrence of lymphocystis in over 8 species of fish, including the Koran angelfish Pomacanthus semicirculatus (Cuvier), the queen angelfish Holacanthus ciliaris (Linnaeus), the Moorish idol Zanclus canescens (Linnaeus) and the foureye butterflyfish Chaetodon capistratus (Linnaeus), all of which are commercially important aquarium fishes. Giavenni (1982) found 71 cases of lymphocystis disease in 29 fish species in tropical saltwater aquarium fishes during a 2 year period. Furthermore, many other aquarium fish species, including squirrelfish Sargocentron punctatissimum, snakeskin gourami Trichogaster pectoralis, quillback rockfish Sebastes maliger, and glassfish Chanda ranga, have been reported as lymphocystis hosts (McCosker et al. 1976, Curry 1977, Richards 1977, Leibovitz 1980, Paperna et al. 1987, Anderson et al. 1988, Williams et al. 1996). Also, lymphocystis disease was found in the farm-reared tropical ornamental fish scalare Pterophyllum scalare and gourami Trichogaster spp. (Paperna et al. 2001). In China lymphocystis disease has been reported in about 10 species of cultured fish (Zhang et al. 1992, Qu et al. 1998, Sheng et al. 2006), but not in aquarium fish species until now.

In April 2004 white nodules were found developing on the fins of whitespotted puffer Arothron hispidus reared in the Qingdao Aquarium of China. The lesions were not observed in other fish species in the aquarium. To confirm the type of the disease several diagnostic studies were performed; as a result, we proved that it was lymphocystis disease and A. hispidus was a new host for LCDV.
MATERIALS AND METHODS

Arothron hispidus were reared in the aquarium tank at 25 ± 1°C in the Qingdao Aquarium of China and fed small fish and sleeve-fish. The diseased fish averaged about 25 to 30 cm in body length. The white nodules first appeared in April 2004, with the largest size being the 1 × 1 cm. They appeared smooth and dense and only occurred on the fish’s pectoral fins. The nodules gradually decreased in size and by the end of June only one small white dot was visible.

Parafin sectioning and histochemical staining. White nodules on the fins were dissected along with the connected skin, fixed with Bouin’s fixative, processed for routine paraffin sectioning (5 µm in thickness) and stained with H&E for histopathological observation. Histochemical methods were used to indicate the inclusion bodies (Mann’s method), DNA (Feulgen reaction) and polysaccharide (periodic acid-Schiff [PAS] reaction).

Ultrathin sectioning. Nodule samples were collected and fixed in 2.5% glutaraldehyde in phosphate-buffered saline (PBS) (0.1 mol l⁻¹, pH 7.4) for 2 h at 4°C, postfixed in 1% osmium tetroxide in PBS for 1 h at 4°C, dehydrated in an ascending ethanol series, infiltrated and embedded in Epon epoxy resin according to standard procedures. Ultrathin sections were double-stained with uranyl acetate and lead citrate, and observed and photographed with a Hitachi H-7000 transmission electron microscope (TEM).

Nested PCR assay for the detection of LCDV in Arothron hispidus. The skin samples of healthy A. hispidus and the white nodules of diseased fish were removed and the genomic DNA of the samples was extracted according to the method of Sambrook & Russell (2001). The DNA extracts from white nodules of A. hispidus were used for nested PCR for the detection of LCDV with the DNA extracts from the skin of healthy A. hispidus and the white nodules of diseased fish were used for nested PCR. The DNA extracts from the skin of healthy A. hispidus and the white nodules of diseased fish were used for nested PCR. The DNA extracts from the skin of healthy A. hispidus and the white nodules of diseased fish were used for nested PCR. The DNA extracts from the skin of healthy A. hispidus and the white nodules of diseased fish were used for nested PCR.

The primer sets for the PCR were designed based on the sequence of the major capsid protein (MCP) of LCDV from Paralichthys olivaceus (GenBank: AF126405, Xu et al. 2000). The primers P1 (5’-CAT CAT GGC TTT GAC AGC-3’) and P2 (5’-GGA TCA GCA ATA CCC-3’) were designed for the first round of nested PCR expected to generate a 384 bp fragment product; primers P3 (5’-TCC ACC GTC AAA GAT TAC-3’) and P4 (5’-CAA TTC CAC CGT CAA AGA-3’) were designed to amplify a 173 bp product for the second round of nested PCR.

The primary and secondary PCR of the nested PCR was performed on a GeneAmp PCR System 9700 (Applied Biosystems) under the same conditions: an initial denaturation at 94°C for 5 min, followed by 30 amplification cycles (94°C for 2 min, 49°C for 1 min and 72°C for 1 min) and a final extension step at 72°C for 5 min. In the primary PCR, the reaction mixture in a final volume of 50 µl contained 1 x reaction buffer (10 mmol l⁻¹ Tris-HCl pH 9.0, 50 mmol l⁻¹ KCl, 1.5 mmol l⁻¹ MgCl₂, 0.1% Triton X-100), 0.5 µmol l⁻¹ of each primer (P1/P2), 2 units Taq-polymerase (Sangon), 1.5 mmol l⁻¹ MgCl₂, 200 µmol l⁻¹ dNTP and 1 µg of DNA template. For the secondary PCR, 1 µl of 1:100 dilution of the reaction mixture from the primary PCR was used as the template; the PCR was carried out under the same conditions as in the first cycle, but in the presence of the primers P3/P4. Amplification products were electrophoresed in a 1.5% agarose gel using pBR322DNA/AluI (BBI) as a marker, and visualized with a UV transilluminator after ethidium bromide staining (1 mg ml⁻¹) (Amersco).

RESULTS

Histopathologic features of white nodules from Arothron hispidus

The white nodules on the fins of Arothron hispidus were composed of enlarged cells in the connective tissue of the dermis with the largest size being 400 µm in diameter. Abundant connective tissues were evident among these cells and many fibres were distributed in the connective tissue, which might be the reason why the nodules appeared dense and smooth. Infiltration by many inflammatory cells into the connective tissue and around the giant cells was observed. The enlarged cells had irregular nuclei in which the marginalized and condensed chromatin usually connected with the nuclear membrane. The cytoplasm was basophilic and stained unevenly in color by hematoxylin. Abundant strong basophilic dot-shaped substances were located at the cell periphery. A distinctive thick hyaline capsule appeared outside of the cell membrane (Figs. 1 & 2). The senile cells, which usually were surrounded by many inflammatory cells and fibres that formed a collar, eventually ruptured and released the cytoplasm, leaving a cavity containing cellular debris (Fig. 3). The areas where lymphocystis cells had disappeared were observed to be light in color (Fig. 4). No lymphocystis cells could be seen in the June samples.

Histochemical staining with Mann’s method stained the basophilic substances located at the cell periphery (as seen in H&E stained sections) blue, indicating they were inclusion bodies. The inclusion bodies, the chromatin and the cytoplasm of the giant cells were Feulgen-positive, indicating DNA was present (Fig. 5). The hyaline capsule stained purple with the PAS reaction, showing it contained polysaccharides (Fig. 6).
Fig. 1 to 8. Lymphocystis disease in *Arothron hispidus*. Fig. 1. Lymphocystis cells and fibre (Fi) in the connective tissue. Scale bar = 200 µm. Fig. 2. Enlargement of lymphocystis cell. Scale bar = 100 µm. Fig. 3. Ruptured senile lymphocystis cells (LCC) and inflammatory cells (arrow) surrounding them. Scale bar = 200 µm. Fig. 4. Light-colored areas (arrows) show where lymphocystis cells had ruptured and disappeared. Scale bar = 200 µm. Fig. 5. Feulgen reaction showing the Feulgen-positive cytoplasm, chromatin in the nucleus and inclusion bodies in lymphocystis cells (arrows). Scale bar = 150 µm. Fig. 6. Periodic acid-Schiff (PAS) reaction showing the positive-stained capsule (arrow). Scale bar = 200 µm. Fig. 7. Ultrastructure of lymphocystis cells showing virus particles (VP), inclusion body (IB) in the cytoplasm and the capsule (C) outside the cells. Scale bar = 2 µm. Fig. 8. Cellular residuals distribute in the connective tissue in June sample. Scale bar = 500 nm.
Under TEM numerous mature or immature virus particles about 200 nm in diameter and with hexagonal profiles were observed in the cytoplasm of the enlarged cells, but not in the nucleus (Fig. 7). High electron-dense inclusion bodies were located at the cell periphery with virions budding from the surface of the cell. Various electron-lucent vacuoles were present in the cytoplasm, especially at the cell periphery. The capsule outside the enlarged cells was homogeneous with low electron density. In the June samples many cellular residuals were distributed in the connective tissue and no viruses were observed (Fig. 8).

Detection of LCDV in Arothron hispidus by nested PCR

A nested PCR assay was tested using 10-fold dilutions of LCDV DNA from Paralichthys olivaceus. Primers P1/P2 for the first round of PCR amplified a 348 bp fragment (Fig. 9A) and primers P3/P4 for the second round of PCR produced a 173 bp fragment (Fig. 9B); the sizes of amplifications were consistent with those predicted. A minimum of $1 \times 10^{-4}$ µg of LCDV DNA were required to get a positive signal in the primary PCR and the detection limit was $1 \times 10^{-9}$ µg of viral DNA in the secondary PCR.

After the second round amplification of nested PCR, a 173 bp fragment was obtained from the DNA extracts from the April nodule samples with the same molecular weight as the positive control (Paralichthys olivaceus LCDV DNA), suggesting the existence of LCDV DNA in Arothron hispidus; no corresponding band was observed in the negative control (the skin DNA of healthy A. hispidus) and the DNA extracts from the June nodules (Fig. 10). Therefore, lymphocystis virus particles existed in the nodules of A. hispidus in April; the nodules were caused by the LCDV infection and they regressed by the end of June.

DISCUSSION

The results of histopathology, histochemistry and TEM revealed that the nodules on the fins of Arothron hispidus have the common histological features of lymphocystis disease such as cellular hypertrophy, enlarged nucleus, Feulgen-positive cytoplasmic inclusion bodies and cell enclosure by a distinctive PAS-positive hyaline capsule. Also, virus particles about 200 nm in diameter with hexagonal profiles in the cytoplasm of the enlarged cells were present, indicating that the white nodules in A. hispidus were lymphocystis lesions.

The MCP of iridoviruses is a suitable target for the study of viral evolution since it contains highly conserved domains (Tidona et al. 1998). The MCP of LCDV from Paralichthys olivaceus in China shows a high identity to those of LCDV-1 and other iridoviruses (Xu et al. 2000, He et al. 2001, 2002, Zhang et al. 2004). Based on the gene sequence of the MCP of LCDV (GeneBank: AF126405, Xu et al. 2000), 2 primer pairs were designed and applied in this study, and a 348 bp fragment in the first round PCR and a 173 bp fragment in the second round PCR were obtained. For the detection of the LCDV in the nodules of Arothron hispidus, the nested PCR amplified a 173 bp fragment in the April samples similar to that of LCDV DNA from P olivaceus, revealing the existence of LCDV. Therefore, the nested PCR assay using the primers derived from the MCP of LCDV from P. olivaceus is an effective method for the diagnosis of lymphocystis disease.
Sano et al. (1994) reported that the progression and regression of lymphocystis was closely related to rearing water temperature. Lesions were visible when water temperature was less than about 22°C in October and not visible when temperatures were greater than about 20°C in June and July. In both seasons when the animals were kept experimentally in water over 25°C, the lesions regressed remarkably in 3 to 5 wk. In China lymphocystis disease breaks out in fish throughout the year. Qu et al. (1999) found that the regression of lymphocystis seemed to be related to the course of the disease, and regression occurred 2 to 3 mo after the onset of the disease. In the present study, Arothron hispidus were reared at 25 ± 1°C and the water temperature did not change with the season. Lymphocystis nodules were present in early April and regressed by the end of June, which suggested that the occurrence of the disease was not strictly related to the rearing temperature, but related to the course of the disease as suggested by Qu et al. (1999).

Colorni et al. (1995) showed lymphocystis nodules to be present in the skin and sometimes in the internal tissues of red drum Sciaenops ocellatus. We have previously observed hypertrophic lymphocystis cells in the gill, submucosa of the intestine, spleen, peripheral portion and the surface of the head kidney, and the adjacent mesenteries of the liver and intestine in Paralichthys olivaceus (Sheng et al. 2006), but in the present study lymphocystis nodules were observed only on the pectoral fins of Arothron hispidus. Internal tissues were not sampled and further study is needed to confirm or deny the existence of internal nodules in this species.

Lymphocystis disease is viral, but the infection rate is increased by stress factors such as malnutrition, dense host population or water pollution (Sindermann 1996, Dethlefsen et al. 2000, Essbauer et al. 2004). The cause of lymphocystis disease in Arothron hispidus has not been determined definitively, but it may originate from virus particles in the water or food. Since there are no effective medications for lymphocystis disease, the best preventive measures are to remove the diseased fish from the population or keep them under optimal, stress-free conditions with good nutrition to maintain a healthy immune system.

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