Isolation and identification of mortalities in golden grey mullet (

Betanodavirus causing mass Liza aurat) is the Caspian Sea

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Abstract

Outbreaks causing mass mortalities have been observed in the golden grey mullet Liza auratus) population in southern coastal area of the Caspian Sea since 2002. The clinical signs include neurological abnormalities, such as erratic swimming behavior, spiral swimming, bellyup at rest and over inflation of the swim bladder. Three hundred and twelve samples of moribund fish with sizes that ranged from 15-26 cm showing abdominal enlargement were collected from 2006-2008. No pathogenic bacteria and parasite metfound. The observed histopathological changes in 29 fish were necrosis and vacuolation of the brain, optic nerve and retina. Cytopathological effects of nodavirus characterized by vacuolation were observed in the SSN-1 cell line at 25°C, 5 d after inoculation with the filtered supernatant of the brain and eye of four affected fish. The recovered virus from cell line exhibited 10 TCID per mL when titrated. Indirect immunoflorescent antibody test showed nodavirus antigens in the retina and postive-CPE SSN-1 cells. Also, a RT-PCR product of approximately 289 bp was amplified from the brain and retina of the 23 samples, including the SSN-1 positive samples. This is the first report of the isolation and characterization of nodavirus from golden grey mullet from the Caspian Sea and, therefore, possible infections in other valuable species of the Caspian Sea warrant further studies.

Introduction

The causative agent of VNN disease is a non-enveloped bisegmented single-stranded positive sense RNA

Viral nervous necrosis (VNN) causes viral nodavirus with size from 25-30 nm (Moetial ., 1992). encephalopathy and retinopathy (VER) in a variety of Golden grey mullet L(iza auratus) is one of the cultured and wild marine fish, including sea basses valuable commercial species in the Caspian Sea having calcalifer), flounder (Paralichthys olivaceus) sea about 50% of annual catching rate of bony fish in the bream (Sparus aurata), turbot (Scophthalmus Caspian Sea. This species has been faced with severe maximus), European sea bas (Dicentrarcus labrax), disease outbreaks causing high morbidity and mortality Atlantic halibut (Hippoglossus hippoglossus), in the Caspian Sea since several years ago but no groupers (Epinephelus adara, E. fuscogutatus, E.etiological agent has been reported so falt aband malabaricus, E. moara, E. tauvina, E. coioides), jackRehannandeh, 2002) This paper describes the isolation (Pseudocaranx dentex), parrotfish (Oplegnathusand characterization of a nodavirus from wild-caught fasciatus) flatfish (Verasper mosera)nd sturgeon sp. moribund golden grey mullet studied during 2006in different parts of world (Mundayet al ., 2002; 2008. Therefore, this is the first report of the isolation Athanassopouloret al ., 2004: OIE, 2006). Affected of nodavirus from golden grey mullet in the Caspian fish show a variety of neurological abnormalities, suchSea. as erratic swimming behavior, spiral swimming, whirling and belly-up at rest, resulting in severeMaterials and Methods morbidity and mortality (OIE, 2003). The most common histopathological changes observed arEish samples

necrosis and vacuolation of the brain and retina (Morsample collection

et al., 1992; Compost al ., 1994; Cleit al ., 1997, 2003). Samples were collected from 312 wild-caught

moribund golden grey mullets from coastal waters in L isopropanol. Samples were then centrifuged at the southern part of the Caspian Sea from 2006-20082,000 g for 10 min at 4°C, and the isopropanol was then Samples of gills, skin, fins, intestine, swimbladder.decanted. The pellets were washed with 0.5 mL 75% kidney, liver, brain and eye were processed forethanol and spun down at 7,500 g for 5 min at 4°C to parasitological and bacteriological analyses, as well ascover the RNA pellet. The ethanol was decanted and virus isolation, reverse transcriptase polymerase chaime dried pellet was dissolved in 200 µL DEPC-treated reaction (RT-PCR) analysis, histological andwater. RT- PCR and nested PCR were performed using serological studies. IQ2000[™] VNN detection kit (Farming Intelligene Tech.

Corp., Taiwan). We added 8 µL of RT-PCR reaction mixture containing 7.0 µL RT-RC pre-mix reagents,

Parasitological and bacteriological examinations

The external and internal organs, including the0.5 µL IQzyme[™] and 0.5 µL reverse transcriptase gills, skin, fins, intestine and swim bladder, underwenenzyme mixed with 2 µL of the extracted RNA sample or parasitological examinations under a light compoundstandard. The RT-PCR reaction was performed at 42°C microscope (Olympus). Also, samples of kidney and/or 30 min, 94°C for 2 min then continued for 15 cycles at liver were aseptically cultured on tryptic soy agar that 94°C for 20 s, 62°C for 20 s and 72°C for 30 s using was supplemented with 1.5% NaCl and incubated athermocycler (Biored). The final extension was at 72°C 26°C for 72 h. for 30 s and the samples were kept at 20°C for a further

30 s. For nested PCR reactions, a 15µL of nested PCR reaction mixture containing 14 µL nested PCR pire-m

PBS at 37°C for 30 min in a humid chamber after

Histopathological analysis

Samples of brain, spinal cord, eyes and swimand 1 µL IQzyme DNA polymerase was added to each bladder from moribund fish were fixed in 10% bufferedtube containing RT-PCR product. formalin and were processed using routine histological

procedures before being sectioned into m-thick Indirect florescent antibody test (IFAT) Serial tissue sections were deparaffinized and slices prior to staining with hematoxylin and eosin rehydrated to phosphate buffer saline (PBS) at pH 7.2. (H&E). The sections were then treated with 0.1% trypsin in

Virus isolation

The samples of brain and eve tissues of moribundvashing with cold PBS and treated with a rabbit antifish were homogenized in Eagle's minimumeestical nodavirus monoclonal antibody (Mab, Aquatic medium (EMEM) by grinding the tissues. The Diagnostics Ltd., Scotland) at 37°C for 30 min in a homogenates (10% w/v) were centrifuged at 4000 x burnid chamber. Samples were then rinsed four times at 4°C for 15 min and the supernatants were filtered with PBS-Tweer80 (PBST). The sections were then through a 0.45 µm filter. Virus isolation trials were treated with a commercially available fluorescein performed by inoculation of the filtrates onto a isothiocyanate€conjugated anti-rabbit Ig antibody at confluent SSN-1 cell line monolayer (Frericensal. ,37°C for 30 min, rinsed with PBST, and were then 1991) in minimum essential medium with 10 % fetalexamined under a fluorescent microscope. bovine serum (FBS) and were incubated at 25°C for 1Additionally, the supernatant of brain and eye tissues days. If no CPE was observed, three blind passagemere inoculated into the SSN-1 cell line in 96-well were then undertaken after freezing telescat -70°C plates, which were fixed after 7 d with cold acetone, and before thawing them. incubated with 2% Triton-X 100 for several minutes, washed with PBST (PBS containing 0.05% Tween 80),

Virus detection by RT-PCR

and thertreated with the antibody as mentioned above. The total RNA from samples of brain, spinal cord,

eve, swim bladder and the supernatant of SSN-1 celResults

inoculated with tissue filtrates were extracted using

IQ2000[™] RNA extraction solution (Farming Intelligene Macroscopic observations

Tech. Corp., Taiwan). A reagent based upon a During the fishing period, the affected fish with modification of the guanidium salt-phenol€chloroformbody weights that ranged from 80250 g exhibited method (Chomczynski and Sacchi, 1987) was useptrominent clinical signs, including lethargy, dark according to the instructions of the manufacture coloration of the skin, abnormal swimming behavior, instructions. Firstly, a volume of 500 µL RNA extraction belly-up positioning, and whirling movements. No solution containing phenol was added to 20 mg brain ogross lesions were observed in necropsy except for the eye sample. Samples were then ground and left at roogmlargement of the swim bladder (Figure 1). temperature for 5 min. A 100 µl of chloroform was

added, vortexed for 20 s and centrifuged for 15 minParasitological and bacteriological studies Approximately 200 µL of the upper clear aqueous phase No pathogenic bacteria and parasites were were transferred into new 0.5 mL tube containing 20@bserved in the affected fish.

Histological examination

fish species in the Caspian Sea. The most remarkable

Histological analysis revealed the presence of linical signs observed in the affected fish were erratic vacuolations in the brain, spinal cord, optic nerve and wimming behavior, belly-up at rest, hyperinflation of granular layer of the retina (Figures 2, 3, 4, and 5). Mosthe swim bladder and abdominal distention. Isolation of the examined fish had degenerative and vacuolated a nodavirus from the moribund fish was achieved lesions in the retina. The extent of vacuolation of theusing the SSN-1 cell line. The isolated virus was retinal and brain tissue varied between fish. In the braindentified as a fish nodavirus based on the results of and spinal cord, vacuolation was mostly observed inhistopathological, IFAT, RT-PCR and electron the grey matter. In some cases, the optic nerve showed croscopy examinations performed on both naturally extensive vacuolation. infected brain and eye tises, and the CPE of cell

Virus isolation by cell culture

cultures. Histopathological investigations in the naturally infected fish revealed severe vacuolation,

Six days after inoculation of tissuenspales on the necrosis and degeneration of nerve cells in the retina SSN-1 cell line at 25°C, a cytopathic effect (CPE)and brain.

developed slowly in some areas of cell line. Rounded, The clinical signs and histopathological changes, granular and vacuolar cells developed, and this effectarticularly lesions of vacuolation and necrosis in the spread to the other areas of the cell sheet, but not to tbeain, spinal cord and retina, were consistent with those entire sheet (Figures 6 and 7). More CPE werethat have been described in other VNN-affected fish observed in the first and second passages (Figuresspecies (Fukudaet al ., 1996; Nakaet al ., 1995; Munday et al., 1997; Munday, 2002). Remarkable and 9).

histopathological signs were seen in retina and brain tissues of these affected fish species. Nopædoarl

Electron microscopy examination

Electron microscopy analysis revealed (2009) recently described a range of histopathological vacuolation and aggregation of some virus particles gins caused by nodavirus infection, including with a size that ranged from 25-30 nm in the cytoplasnhyperplasia and necrotic changes in the gas glandular of the affected erve cells of the retinal layer (Figure epithelium of the swim bladder of affected spotted coral groupers P(lectropomus maculates). However, in 10).

RT-PCR analysis

this current study, we could not see any histopathological changes in the swimbladder of Gel electrophoresis of the nested RT-PCR productaffected_iza auratus. Extensive CPE characterized by vacuolation was

of the deproteinized nucleic acids extracted from the naturally infected fish and the CPE positive cellsobserved in SSN-1 cedultures 5 d after inoculation revealed a band that was 289 bp in size in samplesith filtered homogenates from the brain and eye of collected from the eye and brain tissues (Figure 11) moribund fish. CPE was also seen in the further Also, two bands with sizes of 289 and 479 bp were seepassages. CPE developed initially as some areas of in the PCR products of some eye and brain samplers unded granular cells and slowly spread to other areas. Electron microscopy examination of the CPE-positive (Figure 12).

IFAT

cell cultures revealed vacuolation and aggregation of some virus particles with a size of between 25-30 nm in

Using IFAT, a clear fluorescence was detected in the cytoplasm of nerve cells of the retinal layer. The the retina of some affecteds is samples that were size of viral particles detected in infedteells in this positive for nested RT-PCR amplicons and study were in agreement with those described for sea histopathological examinations (Figure 12). Also, viralbass nodavirus cultured in SSN-1 cells (Frerethal proteins were initially detected in infected SSN-1 cells1996) and Atlantic halibut (Danneviet al ., 2000). 7 d after inoculation (Figure 13). Recovered viral particles exhibited 4 log TCID mL

Discussion

when titrated in the SSN-1 cell line. In this study, the cultivation of L. auratus nodavirus at 20°C was not performed. However, Dannevettal . (2000) were able

The annual catching rate of grey mullet from the cultivate Atlantic halibut nodavirus in SSN-1 cells at Caspian Sea was above 6000 tm in 2002 and, it has o'C. It is difficult to determine whether temperatures decreased to below 3000 tm in 2008 (Department delow 25°C are optimator viral growth because the Fish Stock Assessment Management, Iranian FishericsSN-1 cells grow poorly at temperatures less than 20°C Organization). This indicates that the level of mullet(Danneviget al., 2000). However, natural infection of production has reduced by 50% (Frig 14). The L. aurausoccurred within the range of 11-18°C, and present study describes the causative agent associated er researchers have also reported the isolation of with the mass mortality of wild golden grey mullet, nodavirus from other species at water temperature of which is one of the most economically important bony12.5-26°C (Frerichset al ., 1996; Thiereyt al ., 1999;

Figure 1: Swim bladder enlargement in grey mullet obtained from the Caspian Sea.

Figure 4: Section of eye tissue of the affected L. auratus showing vacuolation in retina (arrow; H&E, x40).

Figure 2: Section of spinal cord of the affected L. auratus showing vacuolation (arrow; H&E, x100).

Figure 5: Lower magnification of brain tissue of the L. auratus showing vacuolation (arrow; H&E, x40).

Figure 3: Lower magnification of optic nerve tissue of the L. auratus showing extensive vacuolation (arrow; H&E, x40)

Figure 6: Vacuolation in the SSN1 cell line 6 d after inoculation of filtrated supernatant from the eye tissue of affected L. auratus (x200).

Figure 7: Vacuolation in the SSN1 cell line 7 d after inoculation of filtrated supernatant of eye tissue of affected L. auratus (x200).

Figure 10: Transmission electron micrograph of the section of eye tissue of the affected L. auratus showing nodavirus particles (25-30 nm in size) in the retina (arrow; x 7,650).

665bp →

289bp

Figure 8: Vacuolation in the SSN1 cell line 5 d after the first subcultivation of filtrated supernatant of eye tissue from affected L. auratus (x200).

Figure 9: Vacuolation in the SSN1 cell line 5 d after the second

subcultivation of filtrated supernatant of eye tissue from affected L.

Figure 11: Agarose gel electrophoresis of a nested RT-PCR product showing 289 bp for the amplicon of the test sample. Lane M = molecular weight marker; N = negative control (no template); P = positive control; Lanes 1, 3 and 5 = brain samples; Lanes 2, 4 and 6 = eye samples. The template cDNA was prepared from naturally infected fish.

848bp	667bp
630bp	479bp
333bp	289bp

Figure 12: Agarose gel electrophoresis of the nested RT-PCR product. Lane M = molecular weight marker; N = negative control (no template); P = positive control; Lanes 1 and 2 = non-infected SSN-1 cells showing 665 bp products; Lane 3 = infected SSN-1 cells showing 289 bp and 665 bp products; Lane 4 = infected SSN-1 cells showing 289 bp and 479 bp products.

Int.J.Vet.Res. (2010), 4; 3: 201-208

auratus (x200).



13a

Figure 14. IFAT showing nodaviral antigens in the SSN1 cell line 6 d post-inoculation (x 400).

Figure 15: Annual catching rate of grey mullet from the Caspian Sea, (Department of Fish Stock Assessment Management, Iranian Fisheries Organization, 2009).



Figure 13: IFAT showing fluorescent nodavirus particles (arrow) in the retina of affected L. auratus (IFAT, x 40).

Danneviget al., 2000). The water temperature of the demonstrate the virulence level of this isolated Caspian Sea at the time of sample collection was in the tanodavirus in some species of the Caspian Sea, range of 11-18°C, which indicates that the viral including mullet and sturgeon.

particles are able to infect the host at a wide range of In conclusion, our histopathological, virological water temperatures. In vitro studies also supported seromolecular findings confirm that VNNV is the evidence for viral resistance to temperaturecause of mass mortality in grey mullet in the Caspian fluctuations; for example, in a study by Ceta al . Sea. However, more investigations are required to (1999), the GNNV isolates of the virus were resistant transsess the virulence level of the isolated virus in this heating at 56°C, which caused CPE at 24-32°C.

In a study by Aspehauget al . (1999), VNN was susceptible hosts, wide geographical distribution a reported from some cold water species of fish.high losses reported in other species, further Therefore, the available data on the effect of investigations are required to study the epizootiology temperature on nodavirus infections under in vitro and and genotyping of this isolated Betanodavirus in the in vivo conditions may display some variations among Caspian Sea. Preliminary data from these studies show different viral isolates in marine and freshwaterthat both sturgeons and guppy are susceptible to the environments (Furusavet al ., 2006; Jithendetal .yirus under vitro conditions (data not shown). 2006). The virulence level of nodavirus in a study is a susceptible to the virulence level of nodavirus in the virulence level of nodavirus is

not clear and requires further investigations. Also, it is Acknowledgements

not clear whether this nodavirus is able to affect other

Caspian Sea species, particularly the hyighaluable This work was supported by a grant from the species of sturgeons. Further studies are in progress Research Council of the University of Tehran and

13b

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