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Citation for published version:
Del-Pozo, J, Mishra, N, Kabuusu, R, Cheetham, S, Eldar, A, Bacharach, E, Lipkin, WI & Ferguson, HW 2016, 'Syncytial Hepatitis of Tilapia (Oreochromis niloticus L.) is Associated With Orthomyxovirus-Like Virions in Hepatocytes' Veterinary Pathology. DOI: 10.1177/0300985816658100

Digital Object Identifier (DOI):
10.1177/0300985816658100

Link:
Link to publication record in Edinburgh Research Explorer

Document Version:
Peer reviewed version

Published In:
Veterinary Pathology

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The final publisher version is available at http://dx.doi.org/10.1177/0300985816658100

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Syncytial hepatitis of tilapia (Oreochromis niloticus L.) is associated with orthomyxovirus-like virions in hepatocytes.

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ABSTRACT

Using transmission electron microscopy (TEM), the presented work expands on the ultrastructural findings of an earlier report on “syncytial hepatitis”, a novel disease of tilapia (SHT). Briefly, TEM confirmed the presence of an orthomyxovirus-like virus within diseased hepatocytes, but not within endothelium. This was supported by observing extracellular and intracellular (mostly intraendosomal), 60-100nm round virions with a trilaminar capsid containing up to 7 electron dense aggregates. Other patterns noted included enveloped or filamentous virions and virion-containing cytoplasmic membrane folds, suggestive of endocytosis. Patterns atypical for orthomyxovirus included the formation of syncytia and the presence of virions within perinuclear cisternae (suspected to be the Golgi apparatus). The ultrastructural morphology of SHT-associated virions is similar to that previously reported for Tilapia Lake Virus (TiLV). A genetic homology was investigated using available RT-PCR probes for TiLV and comparing clinically sick with clinically normal fish and negative controls. By RT-PCR analysis, viral nucleic acid was detected only in diseased fish. Taken together, these findings strongly suggest that a virus is causally associated with SHT, that this virus shares ultrastructural features with orthomyxoviruses, and that it presents with partial genetic homology with TiLV (190 nucleotides).

KEYWORDS

Fish, hepatitis, orthomyxovirus, syncytia, tilapia, virus, ultrastructure

INTRODUCTION

Tilapia are considered to be relatively resistant to many of the diseases that beset other farmed fish. Although they are targeted by the usual parasites commonly found in almost any intensively reared fish species and by some bacteria, notably
Streptococcus spp., viral diseases have not been frequently reported. However, viruses have been recently implicated in several large disease outbreaks with high levels of mortality in both farmed and wild tilapia.2,8,9,23 These outbreaks featured infection by betanodavirus and herpes-like viruses causing central nervous system alterations (neuropil vacuolation and meningoencephalitis, respectively), and a novel enveloped RNA-virus causing ocular, cutaneous, and meningeal pathology. A novel cause of tilapia mortality suspected of having a viral aetiology is syncytial hepatitis of tilapia (SHT), which has been described in Ecuador.9 This disease presents grossly with ascites, and histologically with hepatocellular lipoprotein accumulation, necrosis and syncytia formation, plus necrosis of gastrointestinal mucosa.9 Previously described examples of piscine viral diseases that feature hepatic necrosis include infectious salmon anaemia,24 viral haemorrhagic septicaemia in several species,10,12,19 channel catfish virus,21 adenovirus-like disease of cultured white sturgeon,11 and halibut reovirus.3 Interestingly, the latter also presents with syncytia formation in the liver, which is a feature of SHT.

Our previous report of SHT provided a preliminary description of the clinical presentation, histopathology, and ultrastructural alterations. These included virus-like particles within the cytoplasm of hepatocytes.9 In this work, we aim to provide additional, detailed information on some of the ultrastructural features of SHT-associated virions using transmission electron microscopy (TEM). Based on ultrastructural similarities, and using RT-PCR, we also set out to test the hypothesis that the virions noted in SHT may have genetic homology with those of TiLV.

**MATERIAL AND METHODS**

**Fish.** Details of the clinical presentation of these fish have been previously reported.9 Briefly, intensively-reared tilapia fingerlings (Oreochromis niloticus L.) were collected
for diagnostic work-up following several months of above-normal mortality. In the involved farm, only the own farm-bred tilapia “Chitralada” were affected (with up to 90% mortality), whereas cohabiting tilapia from an outsourced, genetically all male strain were not affected (basal mortality levels). Typically, clinical disease started shortly after transferring the fish into larger on-growing ponds; at that time, the average fish weight was approximately 3g.

**Case definition.** Grossly, these fish presented with darkening, exophthalmia, and ascites (clear fluid). The livers were grossly unremarkable. Histopathological lesions included necrotizing hepatitis with distinctive hepatocellular syncytial giant cell formation, and necrosis of gastric glandular epithelium and intestinal enterocytes (Figure 1).

**Transmission electron microscopy.** Ten fish consistent with the gross case definition were anaesthetised with clove oil, and the liver was dissected. Half of the liver was fixed in 10% neutral buffered formalin (NBF) and the other half fixed in 2.5% glutaraldehyde in a 0.1M sodium cacodylate buffer. The NBF sample was processed for histology to confirm the presence of the histological case definition, which was confirmed in all 10 fish. Samples fulfilling the case definition were selected for subsequent processing for TEM. Specimens destined for TEM were post-fixed in 1% osmium tetroxide in 0.1M sodium cacodylate for 45 min, then washed in three 10 min changes of 0.1M sodium cacodylate buffer. These samples were then dehydrated in 50%, 70%, 90% and 100% normal grade acetones for 10 min each, then for a further two 10-min changes in analar acetone. Samples were then embedded in Araldite resin. Sections, 1μm thick, were cut on a Reichert OMU4 ultramicrotome, stained with toluidine blue, and viewed in a light microscope. Ultra-thin sections, 60nm thick, were cut from selected areas and contrasted with uranyl
acetate and lead citrate. They were examined under a Philips CM120 transmission electron microscope, equipped with a Gatan Orius CCD camera.

**RT-PCR.** Samples of liver were collected from 3 groups of fish selected as follows: a) 17 clinically sick fish, b) 10 asymptomatic fish from the same farm, and c) 6 unexposed healthy control tilapia (also *Oreochromis niloticus*) from Grenada. Tilapia were deeply anaesthetized with tricaine methane sulphonate (MS-222 - Syndel) and a total spinal cord severance was performed post-anaesthesia. After dissection of the liver, all samples were individually placed in RNA-later (Qiagen). Samples were kept at -20°C while not in transport. The RNA extractions were performed using RNeasy Minikit (cat # 74104 Qiagen) as directed by the manufacturer. A one-step RT-PCR (Qiagen) assay was conducted using degenerate primers *NM-CLU7-SF1*, AGTTGCTTCTCAYAAGCCTGCTA and *NM-CLU7-SR1*, TCGTGCTACARCCAGGTTTACTT to amplify a ~245-nt region of the tilapia lake virus (Accession no. KJ605629). The RT-PCR conditions were 50°C for 30 min, 94°C for 15 min, and 35 cycles of 94°C for 30 s, 58°C for 30 s, and 72°C for 1 min, with a final extension of 72°C for 5 min. Amplicons were visualized by electrophoresis in 1.5% agarose gels stained with ethidium bromide under UV light. Negative controls for RNA extraction and RT-PCR assays (SGU fish and water) were included in each assay. PCR products were visualized on 2% agarose gel, purified with Purelink Gel Extraction kit (Invitrogen) and confirmed for target specificity by sequencing of both strands by Sanger Sequencing (GeneWiz, NJ). Negative controls for RNA extraction and RT-PCR assays (SGU fish and water) were included in each assay.
RESULTS

**RT-PCR.** All liver samples from clinically sick fish were RT-PCR positive (17/17) while no amplicons were detected in liver samples from asymptomatic fish (0/10) or any of the Grenada controls (0/6). After trimming the primer sequences 190nt long PCR fragments from seventeen positive Ecuadorian fish, samples were aligned with published tilapia lake virus fragment (Accession no. KJ605629). Sequences showed 98-100% nucleotide identity with reference sequences and with positive samples.

**Transmission electron microscopy.** Virions were seen both within the cytoplasm of hepatocytes as well as in the space of Disse. Although occasionally icosahedral in appearance (Figures 2-5), most virions noted were round and 60-100nm in diameter (mean=75nm). They presented with variably retained structural detail (Figure 5). When the structure was clearly visible, it featured a moderately electron-dense core containing several (up to 7) irregular electron dense aggregates, bounded by a trilaminar capsid.

Virions were seen to have at least two additional morphologies: round to oval enveloped structures, and filamentous/tubular forms (Figures 3 & 4). Enveloped virions were 70-110nm wide (average=100nm), they had a moderately electron dense core, a trilaminar capsid-like structure, and a 10-15nm thick moderately electron dense external band containing regularly spaced electron dense structures (envelope). Filamentous/tubular forms were less common, but when present they were 90-110nm wide, of varying length, and had a moderately electron dense core, a trilaminar capsid, and an envelope similar to that described above.

Multifocal invaginations of the cytoplasmic membrane containing virions were frequently seen at the sinusoidal pole of hepatocytes. This was occasionally
associated with an electron dense band below the membrane (vesicle formation with coating, Figure 7).

The intracellular location of the virions is shown in Figures 6 to 9. In most instances, virions were present within membranous intracytoplasmic structures morphologically consistent with endosomes, located below the plasma membrane and at the central region of the cytoplasm. Conversely, in perinuclear areas, virions were contained within prominent membrane-bound anastomosing cisternae. Morphologically, these resemble greatly enlarged Golgi apparatus (Figure 9). More rarely, virions were noted free floating within the cytoplasm.

Affected hepatocytes featured several additional pathological changes (data not shown). These included scattered, intracytoplasmic lattices of electron dense material, some of which were membrane-delimited. Other features included nucleolar dispersion, prominent nuclear pores, and intranuclear clusters of granular material. There was also multifocal hepatocyte necrosis, hepatocellular syncytial formation, hydropic degeneration, mitochondrial swelling and the presence of intracytoplasmic lamellar bodies.

**DISCUSSION**

The ultrastructural changes described here provide further evidence to support the previous suggestion of a viral aetiology for syncytial hepatitis of tilapia (SHT). Further confirmatory studies to isolate the virions and transmit this disease are required to confirm this. Several ultrastructural features of the virions shown in this paper are similar to those of orthomyxoviruses. Moreover, preliminary RT-PCR data support the fact that the virus noted in SHT presents with at least partial genetic homology
(190nt) with Tilapia Lake Virus (TiLV), a novel RNA virus causing mortality in tilapia in Israel.\textsuperscript{8}

SHT targets the gastro-intestinal tract as well as liver, but the ultrastructural component of this study has been restricted to the liver. Ultrastructurally, virions were noted only within hepatocytes and in the space of Disse, and not within the endothelium. This suggests tropism of the SHT virus for the epithelial, but not endothelial cell population in the liver. This ties in with the histological absence of sinusoidal disruption and haemorrhage, in contrast with infectious salmon anaemia (ISA), where the causative isavirus is endotheliotropic.\textsuperscript{15} A range of cell types has been shown to be infected by orthomyxoviruses. Influenza virus can infect respiratory epithelial cells, erythrocytes, leukocytes and platelets,\textsuperscript{5} whereas isavirus will target endothelium,\textsuperscript{15} but also leukocytes\textsuperscript{20} and gill lamellar epithelium.\textsuperscript{25} By contrast with SHT, isavirus has not been reported in necrotic hepatocytes,\textsuperscript{20,24} although the isavirus (orthomyxovirus) associated with haemorrhagic kidney syndrome in Canada infected renal tubular epithelial cells,\textsuperscript{4,18} suggesting that isavirus tropism for epithelial cells is possible in certain circumstances.

The ultrastructural features of the orthomyxoviral virion and replication cycle have been described in detail for several species, including influenza viruses,\textsuperscript{5} and isavirus.\textsuperscript{4,15} The ultrastructural features of the virions noted in SHT are similar to those described for the above orthomyxoviruses. The electron-dense aggregates noted in the core of SHT-associated virus are similar to those previously described for influenza virus, which have been associated with the segmentation of the viral genome.\textsuperscript{5} Moreover, extracellular tubular forms, similar to those noted in SHT, have been reported for both influenza virus\textsuperscript{5} and isavirus.\textsuperscript{15} The presence of an envelope at the extracellular stage has been described for influenza virus,\textsuperscript{5} isavirus,\textsuperscript{14,15} and
TiLV virus. By contrast, in our study, an envelope was not always seen in extracellular virions. The significance of this is unknown at this stage. Defective virions have been described for isavirus, and we cannot preclude the possibility of a similar situation in SHT-associated virus.

In our study there were multiple instances of cytoplasmic membrane invagination containing virions, with occasional electron-dense coating, suggestive of virion endocytosis. We suspect that the process we have noted is similar to that previously observed in other orthomyxoviruses, where both clathrin-dependent and clathrin-independent endocytic mechanisms may be involved. However, more work is required to ascertain this (e.g. studies using double labelling for clathrin and SHT-virions to assess hypothetical co-location). In influenza virus, endocytosis is followed by viral trafficking from early endosomes below the plasma membrane to perinuclear late endosomes. The latter have an acidic environment, and both influenza and isavirus have been shown to require a low pH step for release of their RNA into the cytoplasm. In our SHT study we noted intraendosomal virions below the plasma membrane, but did not detect perinuclear, late endosomes.

Exceptionally for RNA viruses, orthomyxoviruses require nuclear involvement during their replication. This results in ultrastructural nuclear changes, including increase in nucleolar density, followed by dispersion and presence of granular material throughout the nucleus. Sections of tilapia liver infected with SHT virus presented with nuclear dispersion, granular intranuclear material (occasionally clustered below the nuclear envelope), and prominence of nuclear pores. These nuclear features are similar to those described for orthomyxoviruses, although it is not feasible to discard the possibility that they are artefactual without more information on the features of the replication cycle of SHT-associated virions.
The final step in the viral replication cycle is exit from the host cell. In both influenza and isavirus, viral assembly takes place at the host cell cytoplasmic membrane by budding.\textsuperscript{5,15} This is followed by release of the enveloped virus particle from the cell, associated with formation of long microvilli. Regrettably, instances of viral assembly and exit were not captured in the sections of this study on SHT, although there was disarray of hepatocyte villi at the sinusoidal pole. Further work is needed to characterize this.

So far, we have described ultrastructural features of SHT-associated virions that match those of described orthomyxoviruses. There are, however, several noteworthy areas of divergence. Perhaps the most notable of these is the presence of virions within a perinuclear complex of anastomosing membranous structures, that are strongly suggestive of Golgi apparatus. The role and significance of this feature in the replication cycle of SHT-associated virus are unknown, but it is possible that this could represent an intracytoplasmic transport step of virions either towards the nucleus, or towards the cytoplasmic membrane. There is a role for the Golgi apparatus in the replication cycle of influenza virus, where it participates in the transport of a subset of viral proteins to the plasma membrane,\textsuperscript{5} but this does not feature intraluminal virions. Also characteristic of SHT is the formation of syncytia, which is not a feature of orthomyxoviral infections. Syncytia formation, driven by a fusion factor, is reported in paramyxoviruses.\textsuperscript{6} Finally, another noteworthy divergence is that in our sections we recorded free intracytoplasmic virions in SHT, which are not reported for orthomyxoviruses.\textsuperscript{8} Overall, overlapping features with orthomyxovirus suggest it is possible to hypothesize that SHT-associated virions are novel orthomyxoviruses, but there are enough divergences to cast doubts over this
possibility. Genome sequencing of SHT-associated virus is necessary to elucidate its taxonomy.

Above, we have discussed the ultrastructural features of SHT-associated virions. The published report of TiLV describes sparse electron-dense, 55-75nm virions extra- and intra-cellularly in E-11 cell culture. Further examination of the pictures published (picture 2F) reveals an ultrastructural morphology similar to SHT-associated virions (i.e. intraendosomal location, and capsid containing electron dense aggregates). These similarities led us to test the probes published for TiLV on SHT clinical cases (and negative controls). Revealingly, the RT-PCR results do indeed confirm partial genetic homology between the virus noted in TSH and TiLV (190nt). A complementary study has revealed further genetic homology. These results highlight future research possibilities. For example, TiLV was isolated in E-11, a cell line originating from snakehead (Ophicephalus striatus), as well as in a primary tilapia brain cell culture. E-11 is a clone of SSN-1, a cell obtained from whole fry (i.e. composed of a mixed population). It is therefore not unreasonable to assume that isolation of SHT virus may be also possible using E-11. This is despite possible differences in tissue tropism, as gross and histological diversity between TiLV and SHT also suggests. In TiLV, lesions include encephalitis and ophthalmitis with limited hepatic degeneration (hepatocellular pigment accumulation); conversely, in SHT there is hepatic necrosis with hepatocyte dissociation/syncytia formation and mucosal necrosis of gastro-intestinal tract. This diversity may indicate these are distinct viruses, although it is noteworthy that differences in tissue tropism of one virus are possible. One example of this is isavirus: early descriptions of ISA in Norway featured hepatic lesions, while in Canada lesions were almost exclusively renal. In fact, the Canadian presentation was called haemorrhagic kidney syndrome
before it was shown that the aetiology was isavirus, albeit a strain different to that found in Norway. Factors influencing tissue tropism include strain and age of fish and this may explain the enormous inter-strain variation in susceptibility in SHT (from 90% for a farm-bred strain -“chiralada”-, to 0% in a genetically all-male strain GMT). In this outbreak, SHT was seen only in fry, which died very quickly. By contrast, TiLV was recorded in much larger fish, which experimentally survived for 7-10 days post-infection. Variation in tissue tropism due to age differences has been reported in a viral disease of salmon, infectious pancreatic necrosis (IPN, caused by a birnavirus). In fry, IPN targets the liver, while in older fish the main lesions are pancreatic/gastrointestinal. Concluding, the genetic homology noted between SHT-associated virus and TiLV represents only the initial stage of research into the link between these two viruses.

Overall, the results of the present study suggest that the virus associated with SHT is ultrastructurally similar to an orthomyxovirus, and that it presents with partial genetic homology with TiLV (190nt). This homology has been confirmed further in a complementary study. Further questions to be resolved include isolation of the virus, transmission studies, and further study of its replication cycle and pathogenesis.

ACKNOWLEDGEMENTS

The authors would like to acknowledge Stephen Mitchell at the electron microscopy unit of the University of Edinburgh for his help with TEM processing and viewing of the samples.
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FIGURE LEGENDS

Figure 1. Syncytial hepatitis of tilapia (SHT). There are multifocal to coalescing areas of necrosis, frequent single hepatocellular necrosis with dissociation, syncytial cell formation (white arrows), and perivenular, lymphocytic, inflammatory infiltration. Haematoxylin & eosin (HE).

Figures 2-5. Ultrastructure, three distinct virion morphologies, SHT-affected tilapia.

Fig. 2. Non-enveloped free virions within the space of Disse (black arrow). Fig. 3: Free extracellular virion (white arrow), and intracellular non-enveloped virions within intracytoplasmic membrane bound vacuoles (black arrow). Fig. 4: Free extracellular virion (white arrow), and filamentous/tubular extracellular virions (black arrow).

*=hepatocyte, + = endothelium. Fig. 5: High magnification image of the most
commonly seen intracellular virion form, which lacks an envelope and may contain up to 7 electron-dense irregular aggregates within a core delimited by a trilaminar capsid. TEM, scale bars = 200nm.

**Figures 6-9.** Ultrastructure, liver; SHT-diseased tilapia. Fig. 6: Hepatocytes have multifocal invaginations of the cytoplasmic membrane. Note the electron dense band below the membrane surrounding one of the virions (arrow). Fig. 7. Intracytoplasmic virions are observed within membrane-bound structures consistent with endosomes below the plasma membrane (black arrow). Fig 8. Virions observed at the centre of
the cytoplasm. Fig. 9. Intracytoplasmic virions observed within perinuclear cisternae; *=nucleus. TEM, scale bars: 500nm.