Detection of Tilapia Lake Virus (TiLV) in Healthy Fish from the Pre-Existing Disease Environment Using Different RT-PCR Methods

Patharapol Piamsomboon¹, Janenuj Wongtavatchai¹*,

¹Chulalongkorn University, Faculty of Veterinary Sciences, Department of Veterinary Medicine, Bangkok 10330, Thailand.

How to cite

Abstract
We investigated the presence of TiLV in different fish species from the water reservoir that TiLV was previously reported. The virus was detected in healthy wild tilapia (Oreochromis niloticus) and farmed barramundi (Lates calcarifer) that shared water supply. Detection of TiLV was performed using three RT-PCR assays. Semi-nested RT-PCR detected 9 positives, while the nested and one-step RT-PCR produced 7 and 4 positives, respectively. This study shows evidence that TiLV found in these two fish species is associated with the virus circulating in close proximity environment, nevertheless, it is deficient to indicate disease infection in these asymptomatic fish.

Introduction
Tilapia lake virus (TiLV), an emerging disease of tilapia, was first reported in Israel in 2011 (Eyngor et al., 2014). Subsequently, the disease was reported in tilapia culture regions worldwide, including Asia, Africa, South America (OIE, 2018) and most recently in the USA in 2019 (Ahasan et al., 2020). TiLV is designated to the family Amnoonviridae, a new genus Tilapinevirus and Tilapia tilapinevirus species (ICVT, 2018). The virus contains a negative-sense, single-stranded RNA (-ssRNA) and icosahedron envelop with the size of 55–100 nm diameter (Eyngor et al., 2014). TiLV was reported a causative agent of mass mortality within a month after transfer from the hatchery to the grow-out facility (Surachetpong et al., 2017). However, TiLV has also been detected in other fish species without clinical signs, such as wild river carp (Barbounmys schwanenfeldii) (Abdullah et al., 2018) and giant gourami (Osphronemus goramy) (Chiamkunakorn et al., 2019).

In the past decade, the production of barramundi or Asian seabass (Lates calcarifer) in Thailand has gradually increased from 12.5 metric tons in 2007 to 20.5 metric tons in 2017, while the Nile tilapia (Oreochromis niloticus) is also a major cultured fish species that shares a common cultured area (DOF, 2017). Recently, several viruses have been reported in both fish species, including betanodavirus causing viral nervous necrosis (VNN) disease (Glazebrook et al., 1990; Keawcharoen et al., 2015), and iridovirus causing infectious spleen and kidney necrosis virus (ISKNV) (Suebsing et al., 2016; Thanasaksiri et al., 2019).
As recommended by the World Organization for Animal Health (OIE), the confirmatory test methods for TiLV detection are one-step RT-PCR (Eyngor et al., 2014), nested RT-PCR (Tsofack et al., 2017) and semi-nested RT-PCR (Dong et al., 2017); however, validation of these three methods are suggested (OIE, 2018). We compared the TiLV detection using these three RT-PCR methods in fish species from the water source where TiLV has been detected. In addition, the TiLV detection was extended to farmed barramundi locating in close proximity to the TiLV-PCR positive wild fish.

Materials and Methods

Fish Sample

Observation for TiLV in freshwater fish residing in the natural reservoir was conducted from December 2019 to February 2020 in Chachoengsao province where TiLV was previously reported in farmed tilapia (Surachetpong et al., 2017). Three fish species, a total of 50 fish were collected from an irrigation canal (13°29′08″N 100°53′04″E). Samples were Nile tilapia, climbing perch (Anabas testudineus), and snakeskin gourami (Trichogaster pectoralis). In addition, two barramundi farms supplied with the same irrigation canal were sampled (10 barramundi/farm) for RT-PCR detection.

Sample Preparation and RNA Extraction

Fish were euthanized by immersion with 100 mgL⁻¹ clove oil (Aquanes®, Better Pharma, Thailand). The liver from each fish was collected. The tissue was kept in absolute ethanol at -20°C before to RNA extraction and PCR analysis. A total of 60–80 mg of tissue was prepared for total RNA extraction using the Trizol reagent (Invitrogen, USA) based on the manufacturer’s protocols, followed by RNA quantification with Qubit 3 Fluorometer® and Qubit® RNA assay kit (Invitrogen, USA). Samples of the extracted RNA with a RNA yield of 50–100 ng µL⁻¹ were used for the RT-PCR amplifications.

Detection of TiLV by RT-PCR

Detection of TiLV was performed by the one-step RT-PCR as described by Eyngor et al. (2014) using the 7450/150R/ME2 (5′-TATCACGTACTTTTCCGTGCGTACTC-3′) and ME1 (5′-GGTGGGCAAGGCATCTGCTCA-3′) primers, giving expected amplicon sizes of 250 bp. The RT-PCR mixture was composed of 12.5 µL of 2X reaction mix, 0.5 µL of forward and reverse primer (10 µM), 1 µL of SuperScript™ III RT/Platinum™ Taq Mix (Invitrogen, USA), 2 µL of RNA template (100–200 ng), and nuclease-free water to make a final volume of 25 µL. Thermocycling (Biometra, Göttingen, Germany) was performed at 50°C for 30 min for the first stage RT reaction to form cDNA followed by the second stage qPCR of 35 cycles at 94°C for 30 s, 60°C for 30 s, and 72°C for 30 s. The 415 bp gene fragment of the TiLV gene segment (Integrated DNA Technologies, Singapore) was used as a positive control and no template (nuclease-free water) was used as the negative control.

The nested and semi-nested RT-PCR detection of TiLV were also applied in all extracted RNA samples to compare the result. The nested RT-PCR was conducted using the Nested ext-1 (5′-TTATGCGACTTTCCTCGC-3′) and Nested ext-2 (5′-TTGCTCTGACACAGATGACC-3′) primers followed by the 7450/150R/ME2 and ME1 primers (Tsofack et al., 2017). For semi-nested RT-PCR, the primers Nested ext-1 and ME1 were used for the first step and the primers 7450/150R/ME2 and ME1 were used for the second step PCR (Dong et al., 2017). The first step RT-PCR mixture was as described above and the second step PCR was performed in a final volume of 25 µL containing 12.5 µL of 2X DreamTaq PCR Master Mix (Thermo Fisher Scientific, USA), 0.5 µL of 10 µM forward and reverse primers, 1 µL of the first round RT-PCR product, and 10.5 µL nuclease-free water. The same thermocycling protocol was as applied, except the cycle was decreased to 25 cycles for both steps. The expected amplicon size for both methods was 250 bp. All amplified samples were purified with a NucleoSpin Gel Extraction kit (Macherey-Nagel, Düren Germany), cloned into the pGEM-T easy vector, and submitted for DNA sequencing (Macrogen, Republic of Korea). The nucleotide sequence was aligned and compared with TiLV sequences available in the NCBI database using BLASTn search (https://blast.ncbi.nlm.nih.gov/Blast.cgi).

Results and Discussion

Detection of TiLV in Healthy Fish

A total of 70 healthy juvenile fish were tested, including 29 samples of Nile tilapia, 12 samples of climbing perch, 9 samples of snakeskin gourami collected from the natural water reservoir, and 20 samples of farmed barramundi (Supplementary table). From these, TiLV was detected in 7 Nile tilapia samples and in 2 barramundi samples (Figure 1). The farmed barramundi samples were examined because they shared a common water source with the tilapia that found to be TiLV positive. Some viral pathogens are known to be able to infect both tilapia and barramundi, including VNN (Glazebrook et al., 1990; Bigarré et al., 2009) and ISKNV (Suebsing et al., 2016; Thanasakiri et al., 2019). Although the risk factor of TiLV occurrence is not well-documented, it may share similar characteristics with closely related aquatic orthomyxoviruses, such as infectious salmon anemia virus (ISAV). The virus transmits between adjacent sites through water, fish movement, parasites and wild fish (OIE, 2019). On the contrary, an experimental transmission with intraperitoneal injection (I.P.) of TiLV showed that barramundi was not susceptible, and the virus was undetectable in liver tissues at 5 d post challenge (Jaemwimol et al., 2018). Even so, the
transmission study in rainbow trout (Oncorhynchus mykiss) with ISAV Atlantic salmon (Salmo salar) isolate revealed unsuccessful transmission via I.P. challenge whilst the transmission was achieved by cohabitation with the ISAV-infected salmon, mimicking the natural route of exposure (Snow et al., 2001a; Snow et al., 2001b).

This is the first report of TiLV detection in farmed barramundi and all TiLV positive samples in the present study showed 100% nucleotide sequence identity, both from barramundi (Thailand_BA_2019) and Nile tilapia (Thailand_NT_2019). The 250 bp sequence alignment of the TiLV fragment is shown in Figure 2. A BLASTn search of the NCBI database revealed 98.8% sequence similarity with the genome segment 3 of the Israeli isolate (Til-4-2011; GenBank accession no. KU751816.1). One barramundi sample gave an amplified product from a single step PCR, potentially indicating a substantial viral load in the tested tissue.

Also, detection of TiLV genomes in asymptomatic fish have been reported globally, such as in wild tilapia in Lake Victoria, Africa (Mugimba et al., 2018), wild river carp in Timah Tasoh Lake, Malaysia (Abdullah et al., 2018), cultured tilapia at fingerling and juvenile stages in Peru (Pulido et al., 2019), and in the adult stage in Thailand (Senapin et al., 2018). Natural outbreaks of TiLV infection associated with high mortalities mostly occur in fingerling and juvenile stages of tilapia (Eyngor et al., 2014; Ferguson et al., 2014; Surachetpong et al., 2017). From this fact, there is a possibility of TiLV inhabiting an aquatic environment and only affecting the susceptible young population under particular circumstances, such as an abrupt environmental change. In order to describe the disease condition, additional diagnosis including viral isolation and pathology are required with the positive PCR result.

**Comparison Between RT-PCR Methods for Detection of TiLV**

Three OIE cited RT-PCR assays were used to detect TiLV in our study. Among 70 healthy fish of 4 species, the one step RT-PCR revealed 4 positive samples, while the nested and semi-nested RT-PCR yielded 7 and 9 positive samples, respectively. The samples showed negative results in one step RT-PCR but yielding positive results in other two methods may indicate a latent or carrier status of those fish (Senapin et al., 2018). TiLV detection in apparently healthy (asymptomatic) fish may suggest low viral load in the fish tissues, therefore, screening of TiLV in healthy or wild fish without clinical symptoms should not rely solely on one step RT-PCR. Nevertheless, amplification error of samples detected by nested RT-PCR in our study may be resulted from the mismatch of the Nested ext-2 primer with the mRNA sequence of Nile tilapia (Dong et al., 2017).

**Conclusion**

Polyculture systems and clustering of aquaculture farms in one area are commonly found in many parts of the world, and would facilitate the movement of viruses, including TiLV, between fish species or farms. The present study detected TiLV in tilapia and barramundi that were reared in the same vicinity and shared a water source. Only semi-nested, nested, or higher sensitivity PCR assays are recommended for testing asymptomatic individuals. Nevertheless, the pathological and transmission potential of TiLV in PCR-positive asymptomatic fish cannot be confirmed solely on the detection of the viral genome. Histopathology or viral isolation should be included to confirm a positive case.
Ethical Statement

Not applicable

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Author Contribution

Patharapol Piamsomboon: Funding acquisition, conceptualization, investigation and drafting the original manuscript. Janenuj Wongtavatchai: Conceptualization, provided critical comments and final approval of the manuscript.

Conflict of Interest

The authors declare no conflict of interest.

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