



# Evaluation of a novel TaqMan probe-based real-time polymerase chain reaction (PCR) assay for detection and quantitation of red sea bream iridovirus

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## Abstract

The red sea bream iridovirus (RSIV) belonging to genus *Megalocytivirus* is responsible for red sea bream iridoviral disease (RSIVD) in marine and freshwater fishes. Although several diagnostic assays for RSIV have been developed, diagnostic sensitivity (DSe) and specificity (DSp) of real-time polymerase chain reaction (PCR) assays are not yet evaluated. In this study, we developed a TaqMan probe-based real-time PCR method and evaluated its DSe and DSp. To detect RSIV, the probe and primers were designed based on consensus sequences of the major capsid protein (MCP) genes from megalocytiviruses including RSIV, infectious spleen and kidney necrosis virus (ISKNV), and turbot reddish body iridovirus (TRBIV). The probe and primers were shown to be specific for RSIV, ISKNV, and TRBIV-types megalocytiviruses. A 95% limit of detection ( $LOD_{95\%}$ ) was determined to be 5.3 viral genome copies/ $\mu$ L of plasmid DNA containing the MCP gene from RSIV. The DSe and DSp of the developed real-time PCR assay for field samples ( $n = 112$ ) were compared with those of conventional PCR assays and found to be 100% and 95.2%, respectively. The quantitative results for SYBR Green and TaqMan probe-based real-time PCR were not significantly different. The TaqMan probe-based real-time PCR assay for RSIV may be used as an appropriate diagnostic tool for qualitative and quantitative analysis.

**Keywords:** Red sea bream iridovirus, TaqMan probe, Real-time polymerase chain reaction (PCR), Diagnostic sensitivity, Diagnostic specificity

## Introduction

Red sea bream iridoviral disease (RSIVD) was originally identified in Japan in 1990 in red sea bream (*Pagrus major*) that showed severe anemia with spleen and kidney enlargement (Inouye et al., 1992). The causative agent of RSIVD was later found to be

*Megalocytivirus* of the Family Iridoviridae (Chinchar et al., 2017). According to the World Organisation for Animal Health (OIE), RSIVD is a major cause of mortality in freshwater and marine fish (OIE, 2019). Based on phylogeny of the major capsid protein (MCP) and adenosine triphosphatase (ATPase) genes, *Megalocytivirus* causing RSIVD is classified into three genotypes, red sea

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bream iridovirus (RSIV), infectious spleen and kidney necrosis virus (ISKNV), and turbot reddish body iridovirus (TRBIV) (Kurita & Nakajima, 2012). RSIV and TRBIV genotypes have been identified from farmed fishes for RSIVD as endemic in Korea since the 1990s (Do et al., 2005; Jeong et al., 2003; Kim et al., 2019; Oh et al., 2006).

Among several diagnostic assays for RSIVD, bioassays involving virus isolation using cell culture followed by indirect antibody testing or conventional polymerase chain reaction (PCR) amplification, direct PCR analysis, and sequencing have been classified as recommended methods for reasons of availability, utility, and diagnostic specificity (D<sub>Sp</sub>) and sensitivity in the Manual of Diagnostic Tests for Aquatic Animal (OIE, 2019). Real-time PCR assays allow for the detection of amplified signal for PCR amplicons in real-time using fluorescent dyes without the need for agarose-gel electrophoresis and also has advantages of high analyte sensitivity and specificity compared to that of conventional PCR (Arya et al., 2005; Klein, 2002). However, a real-time PCR assay for RSIV as a standard diagnostic assay is yet not provided in the Manual of Diagnostic Tests for Aquatic Animal (OIE, 2019). Several diagnostic assays for megalocytiviruses causing RSIVD based on real-time PCR have been developed and evaluated for their analyte sensitivity and specificity (Kim et al., 2014; Lee et al., 2020; Rimmer et al., 2012). However, their diagnostic sensitivity (D<sub>Se</sub>) as determined by the proportion of samples from known infected reference animals testing positive in an assay and their D<sub>Sp</sub> as determined by the proportion of samples from known non-infected reference animals testing negative in an assay have not been validated against the conventional PCR assays using field samples.

In the present study, a TaqMan probe-based real-time PCR assay that used a probe and primers designed based on consensus sequences of MCP genes from megalocytiviruses was developed and validated with its D<sub>Se</sub> and D<sub>Sp</sub> for RSIV being compared against conventional PCR assays. The utility of the new TaqMan probe-based real-time PCR assay for viral quantitative analysis was evaluated through comparison to that of a SYBR-Green real-time PCR assay using RSIV-positive samples.

## Materials and Methods

### Virus culture

For virus propagation, primary cells derived from rock bream (*Oplegnathus fasciatus*) were grown in L-15 medium (Gibco, Grand Island, NY, USA) containing 10% Premium Plus fetal

bovine serum (FBS; Gibco) and 1% antibiotic-antimycotic solution (Gibco) as described by Lee et al. (2020). The following megalocytiviruses isolated from primary cells were used in the study: RSIV subtype I (17SbTy strain) derived from sea bass (*Lateolabrax japonicus*), RSIV subtype II (17Rb24Gs strain) derived from rock bream (Kim et al., 2019), ISKNV-type (PGIV strain) derived from pearl gourami (*Trichogaster leeri*) (Jeong et al., 2008), and TRBIV-type (FLIV strain) derived from olive flounder (*Paralichthys olivaceus*) (Kim et al., 2014). Viral infections were conducted in 25-cm<sup>2</sup> flasks of 80%–90% cell confluency in L-15 medium supplemented with 5% FBS and 1% antibiotic-antimycotic solution at 25°C for 7 days. The supernatants were collected following the appearance of cytopathic effects of rounding cells and centrifuged at 500×g for 10 min and the clarified supernatants stored at –80°C until use.

### Primer and probe design

To design primers and a probe for detecting *Megalocytivirus* causing RSIVD (RSIV, ISKNV, and TRBIV-types), consensus sequences of 166 MCP genes retrieved from GenBank of the National Center for Biotechnology Information (NCBI) and 39 MCP genes of RSIVs identified from farmed fish in Korea (Kim et al., 2019). The sequences were aligned using BioEdit Ver 7.2.5 software (Fig. 1). Forward primer RSIV 1094F and reverse primer RSIV 1221R were designed according to consensus sequence base positions 1,094–1,112 and 1,199–1,221, respectively (Table 1). The TaqMan probe (RSIV 1177 probe) was designed according to base positions 1,199–1,221 of the consensus sequences and presented a 6°C higher melting temperatures ( $T_m$ ) than those of the forward and reverse primers. The probe was labeled with carboxyfluorescein (FAM) at the 5' end as a reporter. Black hole quencher-1 (BHQ-1) was used to label the 3' end of the probe as a quencher.

### Positive control plasmid construction

Viral nucleic acid of cultured RSIV 17Rb24Gs strain was extracted using a Patho Gene-spin<sup>TM</sup> DNA/RNA Extraction Kit (Intronbio, Seongnam, Korea) according to the manufacturer's protocol. PCR amplification of the MCP gene was performed using the primers described in a previous study (Kim et al., 2018) and shown in Table 1. To construct a recombinant plasmid for determination of analyte sensitivity and quantitative analysis as a positive control template, a 1,362 bp PCR amplicon was inserted into pGEM-T easy vector (Promega, Madison, WI, USA) and used to transform *Escherichia coli* DH5α cells. The

ATGTCTGCRA TCTCAGGTGC RAACGTAACC AGYGGGTTC A TCGACATCTC CGCKTTYGAT GCGATGGAGA CCCACTTGTA YGGCGGCGAC AATGCVGTGA 100  
 CMTACTTTGC CCGYGAGACC GTGCGBAGTT CCTGGTACAG CAARCTRCCC GTYACCCRTT CAAAACAGAC TGGCCAYGCV AAYTTYGGSC AGGAGTTTAG 200  
 TGTGACKGTG GCVAGGGGHG GCGACTACCT CATTAAAYGTG TGGCTGCGTG TYAAGATCCC CTCCATCACR TCCAGCAAGG ADAACAGCTA CATYCGCTGG 300  
 TGYGAYAATY TGATGCACAA YCTRGTKGAG GAGGTGTCGG TGTCAATYAA CGACCTGGTG GCACADACCC TSACCAGCGA GTTCCTYGC TTYTGGAAACG 400  
 CCTGCATGAT GCCHGGCAGC AAACARTCTG GCTAYAACAA GATGATTGGC ATGCGCAGCG ACCTGGTSKS CGGYATCACC AACGGYACAG CYATGCCYGC 500  
 YRYCTWCCTY AATTTGCCCA TYCCCTVTTT CTTYACYCGH GACACRDGCC TBGCRYTGCC YACYGTGTCT CTGCCRTACA AYGAGGTGCG CATCCACTTC 600  
 AAGCTGCGSC GCTGGGAGGA CCTGCTCATC AGCCAGAGCA VCCAGGCBGA CATGGCCATM TCGACBGTCR CCBTGGCHAA CATTRGCAAT GTARMACCCG 700  
 CRYTGACCAA YGKTKCYGK ATGGGCACYT ACGCYGTRCT GACAAGYAG GAGCGYGAG TBGTGGCCCA GTCYAGYCGT AGCATGCTCA TTGARCAGTG 800  
 YCAGGTGGCG CCYCGTGTG CYGTCACRCC BGYAGACAAT TCYYTGGTGC ATCTSGACCT SAGGTTAGT CAYCCYGTSA AGGCCTTGT CTTTGAGTM 900  
 AAGAAYTCA CYACCCAA CGTGCAAAGC AAYTACACYG CRGCCAGTCC CGTGTAAGTGC AACARCAAGG TSAATCTGCC WYTGMTGGCC ACCAATCCCC 1000  
 TGTCYAGGT GTCRCTCATT TACGAGAACA CCCTMGKCT CCACCAGATG GGAGTAGACT ACTTYACATC HGTCGACCCC TACTACTTTG CGC**CCAGCAT** 1100  
**GCCTGAGATG** GAYGGYGTYA TGACCTACTG YTATACSYTG GACATGGGCA ATRTCAACCC YATGGGYTCR ACCAAY**TAG GCCGCTGTC CAACG**TCACC 1200  
**CTGTCATGTA AGGTGTCGGA** CAAYGCMAG AMMACNGCRG CGGCGGYGG MRVCAACGGC WSCGGYTACA CRGTVGCCA AAAGTTTGAA CTGGTCGTTA 1300  
 TTGCWGTCAA CCACAACATM ATGAAGATTG CTGACGGYGC YGCAGGCTTC CCTATCCTGT AA 1362

**Fig. 1. Consensus sequences of the major capsid protein genes of megalocytiviruses including RSIV, ISKNV, and TRBIV-types.** The sequence data were used to design the probe and primers for the TaqMan probe-based real-time PCR assay. Nucleotide sequences of the forward primer (RSIV 1094F), reverse primer (RSIV 1221R), and probe (RSIV 1177 probe) were denoted as bold and colored boxes. RSIV, red sea bream virus; ISKNV, infectious spleen and kidney necrosis virus; TRBIV, turbot reddish body iridovirus; PCR, polymerase chain reaction.

**Table 1. Primers and probe used in this study**

Genomic regions	Primers	Sequence (5'–3')	Amplicon (bp)	Purpose	Reference
<i>Pst</i> I fragment	1-F	CTC AAA CAC TCT GGCTCA TC	570	Detection <sup>1)</sup>	Kurita et al. (1998)
	1-R	GCA CCA ACA CAT CTC CTA TC			
DNA polymerase gene	4-F	CGG GGG CAA TGA CGA CTA CA	568		
	4-R	CCG CCT GTG CCT TTT CTG GA			
Major capsid protein	RSIV-MCP1F	ATG TCT GCR ATC TCA GGT GC	1,362	Cloning <sup>2)</sup>	Kim et al. (2018)
	RSIV-MCP1362R	TYA CAG GAT AGG GAA GCC TGC			
	MqF	GGC GAC TAC CTC ATT AAT GT	141	Detection and quantification <sup>3)</sup>	Jin et al. (2014)
	MqR	CCA CCA GGT CGT TAA ATG A			
	RSIV 1094F	CCA GCA TGC CTG AGA TGG A	128	Detection and quantification <sup>4)</sup>	This study
	RSIV 1221R	GTC CGA CAC CTT ACA TGA CAG G			
	RSIV 1177 probe	FAM-TAC GGC CGC CTG TCC AAC G-BHQ1			

<sup>1)</sup> PCR assays for RSIVD in Manual of Diagnostic Tests for Aquatic Animal in OIE.

<sup>2)</sup> Construction of standard curve for quantitative analysis.

<sup>3)</sup> SYBR Green-based real-time PCR assay.

<sup>4)</sup> TaqMan probe-based real-time PCR assay.

RSIV, red sea bream virus; MCP, major capsid protein; RSIV, red sea bream virus; PCR, polymerase chain reaction.

plasmid containing the MCP gene was then purified from the transformed *E. coli* using GeneAll Plasmid SV Mini Kit (GeneAll Biotechnology, Seoul, Korea).

### Real-time PCR assay

Real-time PCR was performed with a 20 µL final volume reagent mixture containing 1 µL DNA, 200 nM of each primer

(RSIV 1094F and RSIV 1221R) and probe (RSIV 1177 probe), 10  $\mu$ L 2 $\times$  HS Prime qPCR Premix (Genet Bio, Nonsan, Korea), 0.4  $\mu$ L 50 $\times$  ROX dye, and 5.6  $\mu$ L nuclease-free water using a StepOne Real-time PCR system (Applied Biosystems, Waltham, MA, USA). After pre-denaturation at 95  $^{\circ}$ C for 10 min, amplifications were conducted with 40 cycles of 94  $^{\circ}$ C for 10 s (denaturation) and 60  $^{\circ}$ C for 35 s (annealing and extension).

### Analyte specificity and sensitivity analysis

To evaluate analyte specificity of the developed real-time PCR assay, DNA extracted from four different megalocytiviruses (RSIV subtype I, RSIV subtype II, ISKNV, and TRBIV) were used. For RNA viruses hirame rhabdovirus (HIRRV; ATCC VR-1391), viral nervous necrosis virus (VNNV; RGNNV type), and viral hemorrhagic septicemia virus (VHSV; IVa type) described by Kim et al. (2016), complementary DNAs (cDNAs) were synthesized using an UltraScript 2.0 cDNA Synthesis Kit (PCR Biosystems, London, UK) and then used as template for real-time PCR assays. DNA extracted from two bacteria species commonly isolated from marine fish in Korea, *Vibrio harveyi* isolated from olive flounder and *Streptococcus parauberis* (FP3287) from olive flounder, was obtained from the Korean Culture collection of Aquatic Microorganisms (KoCAM) of the National Institute of Fisheries Sciences (NIFS) and used as the template for real-time PCR assays. For evaluation of analyte sensitivity, the limit of detection (LOD) was determined using serial-diluted plasmid DNA containing the MCP gene (200, 100, 50, 20, 10, 1 viral genome copies/ $\mu$ L) in twelve replicates using the modified method of Uhlig et al. (2015).  $LOD_{95\%}$ , the lowest number of viral genome copies at which 95% of samples test positive, was calculated using Quodata software (<http://quodata.de>). The probability of detection (POD) graph was generated with a 95% confidence interval.

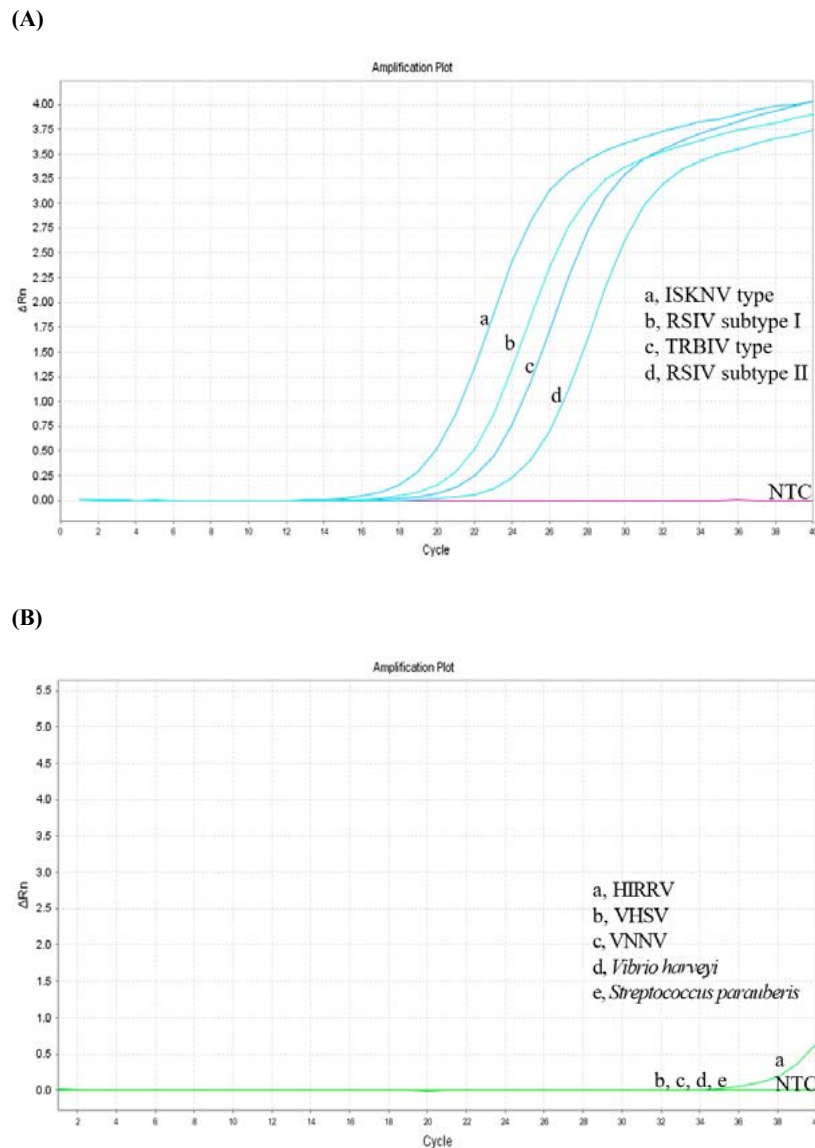
### Comparative analysis of real-time PCR and conventional PCR assays

DNAs obtained from 112 farmed fish spleens (10 mg/specimen) and/or kidneys (10 mg/specimen) were collected between 2012 and 2020 and used for analysis in the current experiment. The farmed fish included 58 olive flounder, 27 rock bream, 10 starry flounder (*Platichthys stellatus*), 8 red sea bream, 7 blackhead sea bream (*Acanthopagrus schlegelii*), 1 stone flounder (*Kareius bicoloratus*), and 1 Korean rockfish (*Sebastes schlegelii*) as detailed in Supplemental Table 1. DNA samples with 260/280 nm absorbance ratios between 1.8 and 2.0 were diluted to 50 ng/

$\mu$ L for use in the experiment. For the qualitative analysis, two OIE-recommended PCR assays (1F/1R and 4F/4R primer sets; Table 1) and the newly developed real-time PCR assay were carried out. The positive control sample was confirmed by positive results for both PCR assays and sequencing analysis. The DSE and DSp were determined as described by Jacobson (1998). For comparison of the quantitative results using the positive control samples, SYBR Green-based real-time PCR was performed using the primers targeting a fragment of the MCP gene (Jin et al., 2014). Briefly, the real-time PCR assay was performed using a reagent mixture containing 1  $\mu$ L DNA, 200 nM of each primer (MqF and MqR), 10  $\mu$ L 2 $\times$  Prime Q-Mastermix (Genet Bio), 0.4  $\mu$ L 50  $\times$  ROX dye, and 6.6  $\mu$ L nuclease-free water and amplified with a StepOne Real-time PCR System (Applied Biosystems). The cycling profile consisted of 94  $^{\circ}$ C for 10 min followed by 40 cycles of 94  $^{\circ}$ C for 10s, 62  $^{\circ}$ C for 15 s, and 72  $^{\circ}$ C for 20 s. A serial 10-fold dilution of positive control plasmid containing the MCP gene was used to generate a standard curve ranging from  $10^1$  to  $10^8$  viral genome copies/ $\mu$ L. Statistical significance between the SYBR Green and TaqMan probe-based real-time PCR assays in quantifying the number of viral genome copies was determined according to a paired *T*-test using GraphPad Prism (Ver. 8.4.3) software.  $p < 0.05$  were considered statistically significant.

## Results and Discussion

To develop a TaqMan probe-based real-time PCR assay for RSIV, specific primers and probe were designed based on 205 consensus sequences of MCP genes compiled from the NCBI database and Korean isolates. The MCP gene is one of the conserved regions that has been used for genotyping based on the phylogeny, as well as used for molecular diagnosis of the genus *Megalocytivirus* (Jin et al., 2014; Kim et al., 2019; Kurita & Nakajima, 2012; Lee et al., 2020; Song et al., 2008). The amplification plot demonstrated that specific amplicons were generated from RSIV subtype I, RSIV subtype II, ISKNV, and TRBIV-types megalocytiviruses (Fig. 2A). Other pathogens, including VHSV, VNNV, *S. parauberis*, and *V. harveyi*, were not amplified using the newly designed primers and probe (Fig. 2B). Fluorescent signals were detected with analysis of HIRRV cDNA after more than 35 cycles of PCR amplification; however, PCR amplicons were not observed with agarose-gel electrophoresis (data not shown). This result suggests the developed real-time PCR assay may be used in specific qualitative analysis for megalocytiviruses including RSIV, ISKNV, and TRBIV. The

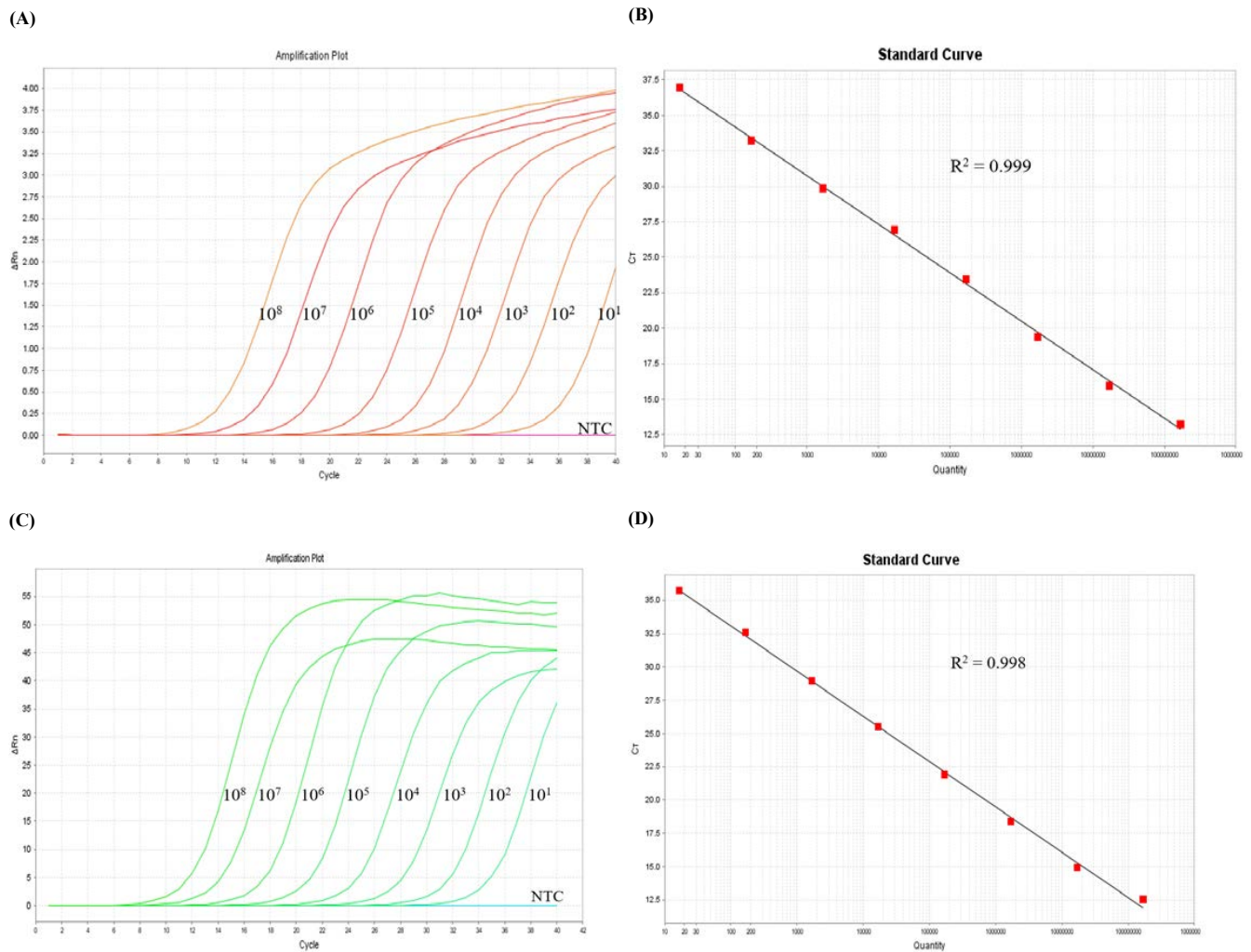


**Fig. 2. Analyte specificity of the TaqMan probe-based real-time PCR assay.** Amplification plots generated using (A) different megalocytiviruses (RSIV subtype I, RSIV subtype II, ISKNV, and TRBIV-types) and (B) other pathogens (VHSV, VNNV, HIRRV, *Streptococcus parauberis*, and *Vibrio harveyi*) as sources of PCR template. PCR, polymerase chain reaction; RSIV, red sea bream virus; ISKNV, infectious spleen and kidney necrosis virus; TRBIV, turbot reddish body iridovirus.

standard curve results showed high correlation coefficients for both the TaqMan probe PCR ( $r^2 = 0.999$ ; Fig. 3A and 3B) and the for SYBR Green PCR ( $r^2 = 0.998$ , Fig. 3C and 3D) within the range of  $1.68 \times 10^1$ – $10^8$  viral genome copies/ $\mu\text{L}$ . This suggested the positive control plasmid DNA could be used for both quantitative analysis and sensitivity analysis with a  $\text{LOD}_{95\%}$  for the lowest number of viral genome copies. In previous studies, detection limits were reported as  $1 \times 10^4$  viral genome copies/

rxn for an OIE recommended PCR assays (Rimmer et al., 2012) and  $10^2$ – $10^3$  viral genome copies/ $\mu\text{L}$  for a peptide nucleic acid (PNA)-based real-time PCR assay (Lee et al., 2020). Although comprehensive validations using artificially infected samples by infection stages are needed, the detection limit of the developed real-time PCR assay was 0.5  $\mu\text{g}/\mu\text{L}$  of DNA (corresponding to 2.5 viral genome copies/ $\mu\text{L}$ ) isolated from an RSIV-infected rock bream in the field (data not shown). Furthermore, the  $\text{LOD}_{95\%}$



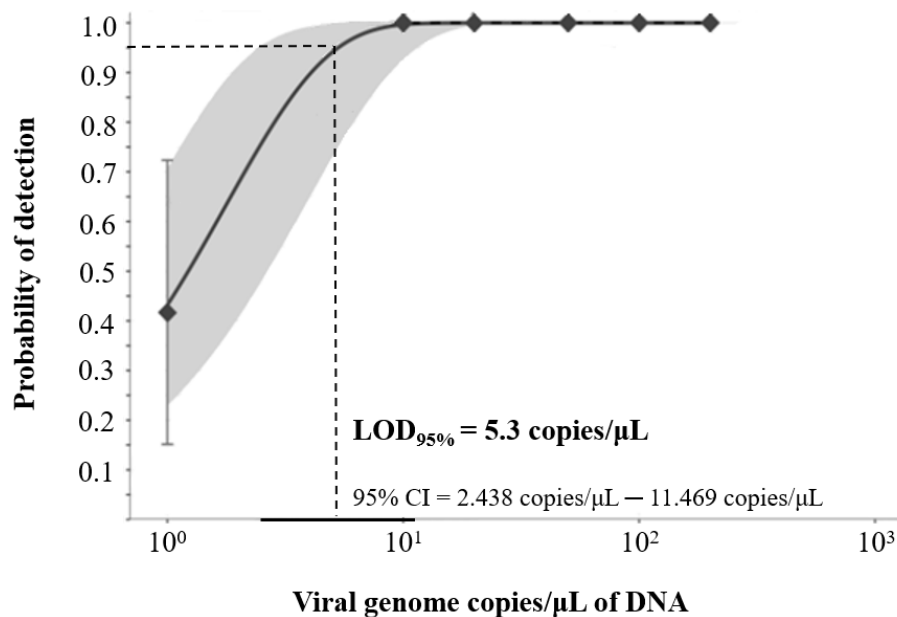


**Fig. 3. Analysis of real-time PCR assays.** Amplification curves of (A) TaqMan probe-based real-time PCR and (C) SYBR Green-based real-time PCR assays. Amplification of the positive-control plasmid DNA containing major capsid protein gene from RSIV at different quantities of viral genome copies ( $1.68 \times 10^1 \times 10^8$  viral genome copies/ $\mu\text{L}$ ). Standard curves of the TaqMan probe-based real-time PCR (B) and SYBR Green-based real-time PCR assay (D) showing linear relationships between viral genome copy number and the cycle threshold. PCR, polymerase chain reaction; RSIV, red sea bream virus.

was determined to be 5.3 viral genome copies/ $\mu\text{L}$  of positive plasmid DNA containing the MCP gene from RSIV (95% confidence interval, 2.438–11.469 viral genome copies/ $\mu\text{L}$ ; Fig. 4), suggesting this assay could be applied with samples possessing low viral levels (approximately 10 viral genome copies), such as environmental water samples, reservoir samples, and biopsy samples collected during the initial stages of infection.

The proportion of true positive and true negative samples evaluated by a diagnostic assay is identified according to DSe and DSp, respectively. Furthermore, DSe and DSp can be applied as indicators for verification of a diagnostic assay to estimate

the status of pathogen infection or exposure in an individual or group (Jacobson, 1998; Saah & Hoover, 1997). Detection results for a total of 112 samples using conventional PCR and the developed real-time PCR assays were in agreement in which 28 samples being true positives and 80 samples being true negatives (Table 2). Only 4 samples presented different results between the assays. Based on the formulas [DSe = true positive (TP) / TP + false negative (FN); DSp = true negative (TN) / TN + false positive (FP)] described by Jacobson (1998), DSe and DSp of the developed real-time PCR assay were 100% and 95.2%, respectively. This result indicated that the developed real-time PCR assay can



**Fig. 4. Probability of detection (POD) curve for the TaqMan probe-based real-time PCR assay.** The solid line denotes the POD means with the 95% confidence interval highlighted as the grey zone. PCR, polymerase chain reaction.

**Table 2. Detection results of samples using the PCR and TaqMan probe real-time PCR assays**

Comparison of the assays	Positive <sup>1)</sup>	Negative <sup>1)</sup>	TaqMan probe real-time PCR assay
Positive <sup>2)</sup>	28	4	32
Negative <sup>2)</sup>	0	80	80
PCR assays	28	84	112

<sup>1)</sup> Tested using the OIE-recommended PCR assays.

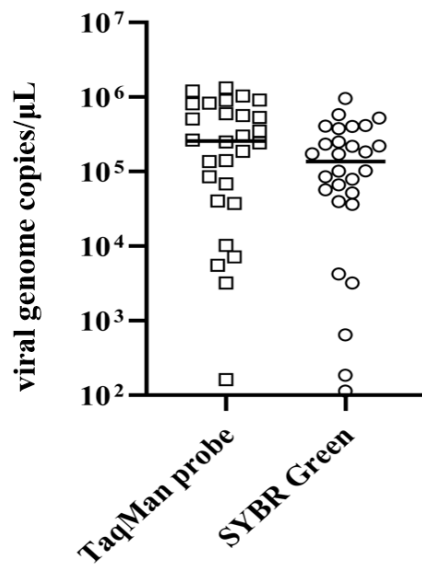
<sup>2)</sup> Tested using the TaqMan probe real-time PCR assay developed in this study.  
PCR, polymerase chain reaction.

be efficiently utilized for RSIV detection in parallel with conventional PCR assay.

Among fluorescent dyes generating amplification signals from PCR amplicons in real-time PCR assays, SYBR Green and the TaqMan probe are generally used for quantitative analysis (Arya et al., 2005). SYBR Green dye, which binds double-stranded DNA by intercalating between the DNA bases during the extension step of PCR amplification, has several advantages over TaqMan probe, including easier primer design and less expensive operation cost. However, SYBR Green-based assays may produce false-negative results caused by non-specific amplicons or primer-dimerization (Arikawa et al., 2008). TaqMan probes are oligonucleotide consisting of reporter and quencher dyes and demonstrate greater DNA binding affinity than that of SYBR Green (Wilhelm & Pingoud, 2003). Thus, the

TaqMan probe-based real-time PCR assay would be expected to be a more appropriate method of qualitative and quantitative virus analysis in terms of specificity and sensitivity. Results of the number of viral gene copies in the 28 RSIV-positive samples was not significantly different ( $p > 0.05$ ) between the SYBR Green and TaqMan probe-based real-time PCR assays (Fig. 5, Supplemental Table 2). This indicated the newly developed real-time PCR assay was sufficient for use in quantitative analyses.

In conclusion, we developed a specific and highly sensitive ( $LOD_{95\%}$ , 5.3 viral gene copies/ $\mu$ L) TaqMan probe-based real-time PCR assay for RSIV based on primers and probe designed using consensus sequences of MCP genes. Based on the comparative analysis to that of conventional PCR assays, the DSe and DS<sub>p</sub> of the developed real-time PCR assay for RSIV were 100% and 95.2%, respectively. Furthermore, the determi-



**Fig. 5. Quantitation of viral genome copies in RSIV-positive DNA samples using the TaqMan probe-based real-time PCR assay and SYBR Green-based real-time PCR assay.** Bars denote the mean of viral genome copies. RSIV, red sea bream virus; PCR, polymerase chain reaction.

nation of the number of viral genome copies in RSIV-positive samples did not significantly differ between the SYBR Green and TaqMan probe-based real-time PCR assays. These results suggest that our developed real-time PCR assay may be used as a robust diagnostic tool for qualitative and quantitative analysis of RSIV.

## Supplementary Materials

Supplementary materials are only available online from: <https://doi.org/10.47853/FAS.2021.e34>.

### Competing interests

No potential conflict of interest relevant to this article was reported.

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Not applicable.

### Availability of data and materials

Upon reasonable request, the datasets of this study can be available from the corresponding author.

### Ethics approval and consent to participate

This article does not require IRB/IACUC approval because there are no human and animal participants.

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