INTRODUCTION

Taura syndrome virus (TSV) was first reported from Penaeus vannamei in Ecuador in 1992 (Lightner 1995). TSV disease has spread widely via infected broodstock throughout America and Southeast Asia, causing severe economic losses in the shrimp aquaculture industry in the last decade (Brock 1997, Tu et al. 1999, Robles-Sikisaka et al. 2002, Chang et al. 2004). TSV is known to cause high mortality in P. vannamei, P. schmitti, and P. seiferus, while sparing P. stylirostris, P. monodon, P. japonicus, P. duorarum, P. chinensis, and P. aztecus (Brock 1997, Overstreet et al. 1997).

TSV infection leads to 60–90% mortality in P. vannamei; the survivors become chronically infected but present no symptoms (Lightner 1996).

TSV, a member of the Dicistroviridae, contains a single-stranded, positive-sense RNA genome that is about 10,205 nt long and encodes 2 non-overlapping open reading frames, ORF1 and ORF2 aura syndrome.

Phylogenetic analysis based on the nucleotide sequences of CP2 suggested that there are 3 lineages, namely the Americas, Belize, and SE Asia, among the various geographic TSV isolates (Robles-Sikisaka et al. 2002).
In the CP1 and CP2 regions, low genetic variation (0 to 0.24% for CP1 and 0 to 0.35 or 0 to 5.6% for CP2 nucleotide sequences) was observed among the isolates analyzed. Due to the low-fidelity nature of the RNA-dependent RNA polymerase, new isolates of TSV that are able to replicate freely in a new species, such as the one described by Chang et al. (2004) in *Penaeus monodon*, are expected to keep emerging.

Sensitive and specific detection of the causative agents is a prerequisite for effective disease prevention and management. Methods developed for TSV detection and characterization include histological analyses, immunostaining, *in situ* hybridization, reverse transcription–polymerase chain reaction (RT-PCR), and real-time RT-PCR (Lightner 1995, Mari et al. 1998, Nunan et al. 1998, Poulos et al. 1999, Castagna et al. 2004, Tang et al. 2004). Among the methods described above, diagnosis of potential TSV carriers or TSV outbreaks at early stages relies mainly on sensitive and specific platforms, such as the nested and real-time RT-PCR (Romano et al. 1995, Nunan et al. 1998, Tang et al. 2004). However, because of the extensive technical training and expensive equipment involved, these systems have not been widely adopted by farmers for sensitive and timely on-site diagnosis of TSV.

With the advantage of requiring only simple heating devices, various isothermal nucleic acid amplification methods providing detection sensitivity and specificity comparable to those of PCR have been under intensified development lately (Demidov 2002, Cook 2003, Savan et al. 2005, Zhang et al. 2006). These methods include rolling-circle amplification, loop-mediated isothermal amplification, and nucleic acid-based amplification (NASBA), to name a few. NASBA is a single-step, enzyme-based RNA amplification reaction (Kievits et al. 1991, Deiman et al. 2002), employing 3 enzymes: avian myeloblastosis virus reverse transcriptase (AMV-RT), ribonuclease-H, and T7 RNA polymerase. Two target-specific primers, including one with a 5′-extension containing the promoter sequence for T7 RNA polymerase, have been designed to amplify the RNA target in exponential kinetics through double-stranded cDNA intermediates. The amplicons are mainly single-stranded RNA, with a polarity opposite that of the target. Since the RNA synthesis step does not require a primer, the maximal yield of the NASBA amplification process could exceed the initial primer levels by at least 1 order of magnitude (Deiman et al. 2002). Being extremely rapid, highly sensitive, and specific (Lanciotti & Kerst 2001), NASBA is most suitable for amplification of single-stranded RNA targets and has been applied to the detection of RNA viruses, such as human immunodeficiency virus, avian influenza virus, and astrovirus, to name just a few (Hibbitts et al. 2003, Tai et al. 2003, Lambert et al. 2005, Yao et al. 2005, Brown 2006).

For aquaculture animal pathogens, NASBA has been reported for the detection of nodaviruses, in which a real-time NASBA was developed (Starkey et al. 2004). Since real-time detection requires expensive apparatus and reagents, such assays are unsuitable for the purpose of simple on-site detection. The ultimate aim of this study was to develop a quick and easy point-of-operation diagnostic system for pathogens, by coupling isothermal NASBA with solid-phase platforms that are user friendly and affordable for amplicon detection. Therefore, we report here on the development of NASA assays for the detection of TSV, as well as the development of postamplification solid-phase-based detection platforms through the use of target-specific capture and detection probes for NASBA products. The sensitivity and specificity of this system were compared to those of RT-nested PCR.

**MATERIALS AND METHODS**

**Sample collection.** *Penaeus vannamei* samples were collected from farms in Taiwan. TSV-positive samples were identified by using the IQ2000 TSV Detection and Prevention System (IQ2000 TSV DPS; Farming IntelliGene). Gill-associated virus (GAV)- and infectious myonecrosis virus (IMNV)-positive shrimp samples were kindly provided by Dr. Peter Walker (CSIRO, Australia) and Dr. Donald V. Lightner (University of Arizona, USA), respectively. Yellow head virus (YHV)-positive samples were collected from the SE Asian countries.

**Nucleic acids.** Although TSV appears to be present in the lymphoid organ at the highest levels, it has been shown that pleopod sampling provides sufficient degrees of sensitivity for TSV detection, allowing non-invasive screening of costly broodstock (Tang et al. 2004). RNA was extracted from the pleopods or gills of shrimp samples using the RNA Extraction Kit (Farming IntelliGene), following the protocols provided with the kit. The plasmid constructs, pTSV containing a 350 bp fragment from the TSV VP2 gene region and pGFP containing the green fluorescent protein gene, were purified using the Midi-V100 Ultrapure Plasmid Extraction System (Viogene-Biotek).

**Primers and oligonucleotide probes.** The sequences of the oligonucleotide primers for the NASBA and the probes for detection are listed in Table 1. The cDNA sequences encoding the TSV capsid proteins of different lineages available in GenBank (AF277378, AF277675, AF406789, AY355309, AY355310, AY355311, AY590471, AY826052, and AY826056) were aligned
transcription was carried out at 60°C for 30 min, following the reaction with 50 µl of light mineral oil, reverse acetate, and 8% (w/v) glycerol, pH 8.2. After overlaying manganese acetate, 5 mM bicine, 11.5 mM potassium of the dNTPs, 2.5 U of rTth DNA polymerase, 2.5 mM of the genome. The 50 µl reaction also contains 300 µM each (Table 1), which amplify a 231 bp sequence of the TSV tissue and 0.46 µM each of Primers 9195 and 9992 tions that include 10 µl of the total RNA extracted from Kit (Applied Bioscience) was used for the RT-PCR reac-tions that include 10 µl of the total RNA extracted from Kit (Applied Bioscience) was used for the RT-PCR reac-

**RT-nested PCR.** For RT-nested PCR analysis, the IQ2000 TSV, YHV, GAV, and IMNV DPS were adopted for the detection of the corresponding viruses. RNA extracts and standards were diluted in yeast tRNA. Briefly, 2 µl of RNA samples was mixed with 7 µl of First PCR Premix, 0.5 µl of RT enzyme, and 0.5 µl of Taq DNA polymerase. The cDNA was synthesized at 42°C for 30 min, followed by 15 of the first PCR cycles. Subsequently, 14 µl of Nested PCR Premix and 1 µl of Taq DNA polymerase were added to the first PCR reaction, followed by 30 cycles of PCR. The products were separated by electrophoresis in a 1.5% agarose gel, stained in ethidium bromide, visualized under UV light, and documented by using the AlphaImager Imaging System (Alpha Innotech).

**OIE TSV RT-PCR.** The sampling and TSV RT-PCR detection procedures were carried out as described in an OIE (Office International des Epizooties) manual (OIE 2003). Briefly, the GeneAmp EZ rTth RNA PCR Kit (Applied Bioscience) was used for the RT-PCR reactions that include 10 µl of the total RNA extracted from tissue and 0.46 µM each of Primers 9195 and 9992 (Table 1), which amplify a 231 bp sequence of the TSV genome. The 50 µl reaction also contains 300 µM each of the dNTPs, 2.5 U of rTth DNA polymerase, 2.5 mM manganese acetate, 5 mM bicine, 11.5 mM potassium acetate, and 8% (w/v) glycerol, pH 8.2. After overlaying the reaction with 50 µl of light mineral oil, reverse transcription was carried out at 60°C for 30 min, fol-

and compared. Design of the primers was based on conserved sequences within the VP2 gene. A standard BLASTN nucleotide-nucleotide search was performed to confirm the specificity of the designed primers. Primers were purchased from BioBasic (Canada) or synthesized at Farming IntelliGene, and purified through high-affinity purification or gel purification before use. The TSVF3-biotin primer was labeled with biotin at its 3’-end.

The sampling and TSV RT-PCR detection procedures were carried out as described in an OIE (Office International des Epizooties) manual (OIE 2003). Briefly, the GeneAmp EZ rTth RNA PCR Kit (Applied Bioscience) was used for the RT-PCR reactions that include 10 µl of the total RNA extracted from tissue and 0.46 µM each of Primers 9195 and 9992 (Table 1), which amplify a 231 bp sequence of the TSV genome. The 50 µl reaction also contains 300 µM each of the dNTPs, 2.5 U of rTth DNA polymerase, 2.5 mM manganese acetate, 5 mM bicine, 11.5 mM potassium acetate, and 8% (w/v) glycerol, pH 8.2. After overlaying the reaction with 50 µl of light mineral oil, reverse transcription was carried out at 60°C for 30 min, followed by inactivation at 94°C for 2 min. The subsequent PCR reaction included 35 cycles of denaturation at 94°C for 45 s and annealing/extension at 60°C for 45 s and a final extension step at 60°C for 7 min; 10 µl of the amplified products were analyzed in a 2.0% agarose gel in 0.5 × TBE (Tris, boric acid, ethylene diamine tetra-acetic acid [EDTA]) and visualized as described above.

**NASBA.** The NASBA amplicons derived from Primers T7 TSVR2 and TSVF2 were expected to be 209 nt long. Reaction conditions for NASBA were modified slightly from those described by Jean et al. (2002a). Briefly, the NASBA reactions were performed in a final 20 µl reaction volume containing 2 µl of the extracted RNA, 40 mM Tris-HCl (pH 8.5), 15 mM MgCl$_2$, 50 mM KCl, 15% (v/v) dimethyl sulfoxide, 5 mM dithiothreitol, 0.5 mM of each deoxyribonucleoside triphosphate, 2 mM of each nucleoside triphosphate, 50 µg µl$^{-1}$ bovine serum albumin, and 0.2 µM of each primer. The mixture without enzymes was incubated at 65°C for 5 min and 42°C for 5 min before the addition of 2.5 U of RNase H (USB), 12 U of the RNase inhibitor (Takara), 40 U of the T7 RNA polymerase (USB), and 8 U of AMV-RT (Promega), followed by a designated incubation period at 41°C. Yeast tRNA (40 ng µl$^{-1}$) was used to dilute RNA samples, to avoid losses of targets at extremely low concentrations.

**Denaturing agarose gel electrophoresis and Northern blotting.** NASBA amplicons were denatured in 1× RNA sample buffer (95% formamide, 0.5 mM EDTA) at 65°C for 2 min, and subsequently resolved in a 1.5% agarose gel containing 3.7% formaldehyde. After staining with ethidium bromide, the bands were visualized under UV light and subsequently transferred to Hybond-N membrane (Amersham Biosciences) in 20× SSPE overnight. After 1 wash with 5× SSPE at room temperature, the membrane was UV cross-linked for 2 min in HL-2000 HybriLinker (UVP) and dried at 63°C. Hybridization was carried out in Dr. Hyb buffer (DR. Chip) containing 0.025 µM biotin-labeled TSP-3 at 63°C for 30 min. After 2 washes in 1× SSPE at 42°C, the membrane was rinsed with 0.5% (v/v) blocking reagent (Roche Diagnostics) in Buffer I (100 mM Tris-HCl, pH 8.0, 150 mM NaCl, 0.3% [v/v] Tween 20) at room temperature, followed by the addition of strepavidin-alanine phosphatase conjugate (Promega) diluted 1:2000 in Buffer I containing 0.5% (w/v) blocking reagent. The incubation was carried out at room temperature for 10 min. After 1 wash with 1× SSPE and 1 with Buffer II (100 mM Tris-HCl, pH 9.5, 100 mM NaCl, 50 mM

<table>
<thead>
<tr>
<th>Primer or probe</th>
<th>Sequence</th>
<th>Function</th>
</tr>
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<tbody>
<tr>
<td>TSVF2</td>
<td>5’CATTGTGGCTGTGGAATAA 3’</td>
<td>NASBA primer</td>
</tr>
<tr>
<td>T7 TSVR2</td>
<td>5’AAATCTTACAGCTACTATTAGGAGGAAGAAGCAGATAATTTAGGT 3’</td>
<td>NASBA primer with T7 promoter</td>
</tr>
<tr>
<td>TSVF3-biotin</td>
<td>5’CCTGGCAATGGGATGACTGATGCGATGCTCTGAAGCTTAGCTA 3’</td>
<td>Detection primer</td>
</tr>
<tr>
<td>TSVR5</td>
<td>5’CTCATAGCGTTGACTTTCG 3’</td>
<td>Capture primer</td>
</tr>
<tr>
<td>TSVR4</td>
<td>5’ACGCTTCAAGCTGCAAGTTCTCATCTCAAT</td>
<td>Hybridization control capture primer</td>
</tr>
<tr>
<td>9195</td>
<td>5’TCAATGAGAGCTTTGG 3’</td>
<td>OIE RT-PCR primer</td>
</tr>
<tr>
<td>9992</td>
<td>5’AAGTACAGACCGCGCGCTT 3’</td>
<td>OIE RT-PCR primer</td>
</tr>
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Table 1. Nucleotide sequences of primers and probes used in this study. The T7 promoter sequence is underlined (NASBA: isothermal nucleic acid-based amplification; OIE: Office International des Epizooties)
MgCl₂), color was developed by the addition of 0.5 mg ml⁻¹ NBT and BCIP in Buffer II. Color was allowed to develop in the dark for no more than 10 min. The reaction was stopped by rinsing the membrane with water.

**Dot blotting.** The NASBA products in 1× RNA sample buffer and the DNA controls in TE buffer were denatured at 95°C for 2 min, diluted in 6× SSPE buffer, and spotted on the Hybond-N filter. The subsequent UV cross-linking, hybridization, and detection procedures were carried out as described above for Northern blotting. Different concentrations of the pTSV plasmid DNA were included as positive controls for the detection steps.

**Microtiter plate detection of NASVA products.** NASBA amplicons were captured by an amplicon-specific primer coated on the wells of the FluoroNunc microtiter plate Module MaxiSorp Surface and detected by a TSV-specific, biotin-labeled primer. Briefly, 0.5 nmol of the capture probes, namely TSVR4 and TSVR5, were immobilized on the well in coating buffer (0.045 M Na₂CO₃/NaHCO₃), dried at 63°C for 5 min, and subsequently UV cross-linked (1200 × 100 µJ cm⁻²). The wells were sequentially washed in Buffer I and water. Then, 5 µl of the NASBA products mixed with 45 µl of 0.025 µM biotin-labeled TSP-3 in the hybridization buffer was added to the wells of the microtiter plate which were coated with capture probe in advance. After 30 min of hybridization at 63°C, the wells were washed 2 times with 1× SSPE at room temperature. Subsequently, the wells were blocked with Buffer I containing 0.5% (w/v) blocking reagent at room temperature, followed by the addition of strepavidin-alkaline phosphatase conjugate diluted 1:2000 in Buffer I containing 0.5% (w/v) blocking reagent at room temperature for 10 min. After 1 wash with 1× SSPE and 1 with Buffer II, 50 µl of 0.5 mg ml⁻¹ NBT and BCIP in Buffer II was added. Color was allowed to develop in the dark for no more than 10 min before the reaction was stopped by rinsing the wells with water.

**RESULTS**

**Detection of TSV by NASBA**

The *Penaeus vannamei* samples collected in Taiwan were first screened for TSV infection on the basis of the high sensitivity of the IQ2000 TSV DPS. RNA prepared from these samples was used as the template for the following NASBA tests. Using analysis by denaturing agarose gel electrophoresis, a single band corresponding to the expected size of 209 nucleotides (Fig. 1A) was detected only from the TSV(+) shrimp RNA extract, and not from a TSV(−) shrimp RNA extract, yeast tRNA, or ddH₂O. The band was confirmed by Northern blotting using the TSVF3-biotin probe (Fig. 1B) to represent the TSV NASBA amplicons. The specificity of the Northern-blotting analysis was validated by the presence of signals only with the pTSV DNA control, but not with pGFP, a plasmid with unrelated sequences (Fig. 1B).

**Dot-blotting analysis of TSV NASBA products**

Based on the Northern-blotting analysis of the TSV NASBA amplicons, no cross-reactive signals could be picked up by the biotin-labeled, TSV-specific primer (Fig. 1B). In order to simplify the detection protocols for the TSV NASBA amplicons, a dot-blotting analysis was developed. The results (Fig. 1C) showed that using the TSV-specific TSVF3-biotin probe, positive signals were obtained only from the NASBA reaction of the TSV(+) RNA and pTSV, the positive control DNA for the detection steps, among the samples tested. In

![](image)
the following experiments, different concentrations of the positive hybridization control, pTSV, were included to ensure that satisfactory degrees of detection sensitivity were reached.

### Optimal incubation time for TSV NASBA

Several conditions for TSV NASBA were varied to optimize its performance. For example, the reaction was stopped after different periods of time and analyzed for the production of TSV NASBA amplicons (Fig. 2). Based on the results, no signals were derived from yeast tRNA and ddH2O alone after 120 min. On the other hand, after 30 min at 41°C, TSV NASBA produced enough copies of amplicons from the TSV-positive RNA to be vaguely detected by dot blotting. However, the NASBA amplicons could be detected without ambiguity after 50 min of incubation. The intensity of the signal increased with time, suggesting that the production of NASBA amplicons increases with incubation time between 50 and 120 min of incubation. Therefore, the NASBA reactions were carried out at 41°C for 90 to 120 min in the following tests.

### Comparison of sensitivity between NASBA/dot blotting and RT-nested, PCR-based TSV assays

To evaluate the feasibility of developing NASBA as an on-site TSV detection platform, the performance of the TSV NASBA/dot-blotting system was compared with a commercially available TSV RT-nested PCR system, the IQ2000 TSV DPS. This system has been applied to routine SPF (specific pathogen-free) broodstock screening at numerous large-scale shrimp farms. First, comparative analysis between the IQ2000 TSV DPS and the OIE TSV RT-PCR detection system, described as one of the standard screening methods for TSV in the ‘Manual of diagnostic tests for aquatic animals (OIE 2003), was performed to verify the sensitivity of the IQ2000 TSV DPS. A TSV(+) shrimp RNA was 10-fold serially diluted and subjected to both assays carried out in parallel. The results showed that the end point of detection was at around a 10^4-fold dilution with the OIE RT-PCR assay (Fig. 3A) and at about a 10^5-fold dilution with the IQ2000 TSV DPS (Fig. 3B), demonstrating that the latter is approximately 10 times more sensitive than the former.

With the sensitivity of the IQ2000 system confirmed, we compared the sensitivity of the TSV NASBA established by us side by side with that of the RT-nested, PCR-based IQ2000 TSV DPS. A series of 5-fold dilutions of TSV(+) RNA extract was prepared in a yeast tRNA solution, and the same tRNA was included as the negative control. Replicate analyses were performed. The NASBA/dot-blotting analysis gave significant positive amplicon signals at 6250-fold dilution (Fig. 4A), while the controls worked properly. With the IQ2000 TSV DPS carried out in parallel, amplification of the...
positive controls \(10^3, 10^2, \text{ and } 10^1\) copies) led to the production of the typical ladders of 1 to 3 TSV-specific PCR products (arrowheads; Fig. 4B). Within a 3-log range of target copies, higher amounts of targets would result in the production of more bands. Meanwhile, no products were detected in the negative control reaction (tRNA; Fig. 4B), and the band of around 650 bp (arrow; Fig. 4B) is the RT-PCR internal control signal derived from shrimp RNA. The end point of sample dilution to obtain positive RT-nested PCR signals fell on the 31 250-fold dilution (Fig. 4B), demonstrating that the TSV NASBA assay was about 5-fold less sensitive than the TSV RT-nested PCR assays.

For the subsequent TSV NASBA reactions in this study, serial dilutions of aliquots of the same TSV(+) RNA sample were included to monitor the sensitivity of the assays.

### Specificity of TSV NASBA

To further investigate the specificity of the TSV NASBA system, samples positive for several non-target RNA viruses were confirmed to be TSV-negative in advance. Fig. 5B,C shows an example of the screening. In this case, an IMNV(+) sample undiluted or 10-fold diluted \((1 \times \text{ or } 1/10 \times, \text{ respectively})\) produced IMNV-specific PCR product ladders using the IQ2000 IMNV DPS. The positive controls \((2 \times 10^3, 2 \times 10^2, \text{ and } 2 \times 10^1)\) diluted in yeast tRNA gave typical ladders of the IMNV-specific PCR products (Fig. 5B). When the same IMNV(+) sample was subjected to TSV detection by the IQ2000 TSV DPS, only the internal control signals were detected (arrow; Fig. 5C), indicating that the IMNV(+) RNA is TSV negative. Similarly, the YHV(+) and GAV(+) samples were confirmed to be TSV negative (Fig. 5C). When these YHV(+), GAV(+), and IMNV(+) RNA extracts were subjected to the TSV NASBA/dot-blotting analysis, no TSV NASBA products were detected (Fig. 5A), demonstrating that the TSV NASBA reaction established is indeed TSV specific.

### NASBA detection of TSV in shrimp specimens

A total of 23 samples from 4 bathes (FITC, CTC, TP, and YS) of *Penaeus van-
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**Sample Collection and Testing**

Samples of *namei* collected in Taiwan were tested by both TSV NASBA/dot-blotting analysis and IQ2000 TSV DPS (Table 2). In total, 8 of the 23 samples were determined to be TSV negative by both methods. Of the other 15 samples that were diagnosed as TSV positive by the IQ2000 TSV DPS, only 13 were designated as TSV positive by the TSV NASBA/dot-blotting analysis. The 2 samples designated TSV positive by the RT-nested PCR, but TSV negative by TSV NASBA/dot blotting were both categorized by the RT-nested PCR kit as 'weakly infected' (data not shown), i.e. these appeared to be samples with very low virus loads.

**Discussion**

NASBA is a robust amplification technology that is highly appropriate for RNA analyses and has been applied particularly to detecting various RNA viruses, as well as a number of other pathogens. This report describes the development of a NASBA/solid-phase detection assay for the rapid and sensitive detection of TSV in shrimp tissue specimens. The NASBA reactions were coupled with the simple colorimetric dot-blotting or microtiter plate method for the detection of TSV-specific amplicons (Figs. 1 & 6). No false-positive results were obtained with yeast tRNA and non-target shrimp RNA viruses, namely GAV, YHV, or IMNV (Fig. 5), demonstrating a high degree of specificity of the TSV NASBA.

Various formats, including solid-phase hybridization, electrochemiluminescence, molecular beacon, enzyme-linked gel assay, and fluorescence correlation spec-

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**End-point detection of TSV by microtiter plate hybridization**

As stated above, solid-phase detection is an attractive platform to be incorporated into a straightforward and cheap diagnostic system (Jean et al. 2002a). Toward this end, the dot-blotting detection protocols were modified and applied to a microtiter plate format, in which NASBA amplicons were detected by a combination of a TSV NASBA amplicon-specific capture probe and a biotin-labeled detection probe. Similarly, positive signals can be visualized by the formation of the purple-colored precipitates. Moreover, a capture probe specific for the biotin-labeled detection probe was fixed on each well (arrows; Fig. 6B) to serve as an internal control for the detection steps. Specific detection of the TSV NASBA RNA amplicons was confirmed by the presence of signals with the TSV(+) samples and the absence of signals with the GFP RNA produced *in vitro* (arrowheads; Fig. 6B). Both dot-blotting and microtiter plate detection systems led to positive detection of the TSV NASBA amplicons of the TSV(+) RNA diluted up to 6250-fold (Fig. 6A, B), indicating that these 2 simple systems offer similar degrees of detection sensitivity for the TSV NASBA amplicons.

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**Fig. 5. Specificity of the TSV NASBA reaction.** (A) TSV NASBA RNA of YHV(+), GAV(+), or IMNV(+) shrimp, together with 10x, 50x, 250x, 1250x, 6250x, or 31 250x dilutions of TSV(+) shrimp RNA, and the negative controls (yeast tRNA and DEPC-H2O) were amplified by TSV NASBA. The pTSV plasmid DNA was included as a positive control for the detection procedures. TSV-specific targets were detected by dot blotting as described in 'Materials and methods'. (B) IMNV RT-nested PCR amplification. The IMNV(+) RNA was identified by using the IQ2000 IMNV DPS. The IMNV(+) RNA, undiluted (1x) or diluted 10-fold (10x), as well as 2 × 10^3, 2 × 10^2 or 2 × 10^1 copies of the positive controls provided by the kit were analyzed. The products were analyzed by electrophoresis in a 1.5% agarose gel, stained with ethidium bromide, and visualized under UV light. (C) TSV RT-nested PCR. The YHV(+), GAV(+), IMNV(+) RNA samples, together with the negative (yeast tRNA and DEPC-H2O) and positive (TSV[+]) controls, were assessed by the IQ2000 TSV DPS (MW: 100 bp DNA molecular weight markers; tRNA: yeast tRNA; arrowheads: virus-specific PCR products; arrows: internal control signal)
...troscopy have been developed for the detection of NASBA products (Deiman et al. 2002, Jean et al. 2002b). The high costs of equipment, chemicals, and reagents needed to perform real-time assays are prohibitive to the application of most of these formats at shrimp farms. On the other hand, solid-phase hybridization systems, in general, do not require any elaborate tools or expensive apparatus for signal generation, detection, and interpretation. Furthermore, the single-stranded RNA amplicons of NASBA are especially compatible with hybridization-based detection systems using target-specific probes. To exclude the requirement of any instrument during the detection steps, the colorimetric alkaline phosphatase/BCIP/NBT system was adopted, to allow direct visualization of the signals. The dot-blotting system using a TSV-specific, biotin-labeled primer together with the strepavidin-alkaline phosphatase/BCIP/NBT signal amplification system was proven to be efficient for the detection of TSV NASBA products (Figs. 1, 2, 4, & 5). The sensitivity of the NASBA reaction reproducibly reached the same dilution end point (6250×; Figs. 5A & 6), providing evidence to support the reproducibility of the NASBA method. Moreover, colorimetric microtiter plate detection of NASVA products was later set up and shown to reach a sensitivity similar to that of the dot-blotting assay (Fig. 6).

The TSV NASBA assay showed levels of sensitivity comparable to those of the PCR assays (Fig. 4, Table 2). The sensitivity of the TSV NASBA/colorimetric dot-blotting method was determined using serial dilutions of TSV(+)-RNA extracts prepared from shrimp tissues, and the results were compared against the commercially available RT-nested PCR assay, the IQ2000 TSV DPS, which was demonstrated to be about 10-fold more sensitive than the standard RT-PCR assay, the OIE TSV RT-PCR, as expected (Fig. 3). The detection limits of the TSV NASBA/solid-phase detection system were reproducibly 5-fold less sensitive than the RT-nested PCR assay (Fig. 4). Previously, a NASBA/slot blot detection system using a biotinylated oligonucleotide probe reached detection sensitivity comparable to that of a RT-nested PCR/agarose gel electrophoresis detection system for enterovirus. (Heim & Schumann 2002, Guichon et al. 2004). Therefore, the sensitivity of the TSV NASBA can likely be improved through further adjustments of either the NASBA reaction or the detection protocols to reach higher levels of sensitivity. To enhance the sensitivity of the solid-phase detection of the NASBA amplicons, other signal amplification methods, such as branched DNA (Tsongalis 2006) or dendrimers (Kim et al. 2003) will be tested.

Results of TSV screening among different batches of shrimp samples by the TSV NASBA and the IQ2000 TSV DPS were in close agreement with each other (Table 2). Thirteen out of 15 TSV RT-nested, PCR-positive samples also tested positive in the TSV NASBA analysis (Table 2). It is likely that levels of TSV RNA in the 2 samples tested negative were below the detection limits of the TSV NASBA, since they were also categorized as ‘weakly infected’ by the RT-nested PCR assay (data not shown).

Molecular diagnosis of shrimp virus pathogens is particularly important given the difficulty encountered in establishing shrimp cell lines and isolating shrimp viruses from the specimens in tissue culture. Rapid and sensitive detection of TSV can facilitate proper and timely disease management measures to be implemented following the confirmation of TSV cases in the field. Coupling with the end-point solid-phase detection formats described in this study, the TSV NASBA assays can be completed straightforwardly in about 3 to 4 h, meeting the demands for speedy diagnosis of...
aquaculture pathogens at custom and import-control checkpoints and at the farms. The NASBA assay has also been shown to tolerate field contaminants, such as cloacal, pharyngeal, and anal swabs, fecal samples, cage sweepings, and blood, that could interfere with assays, such as ELISA and RT-PCR (Shan et al. 2003), also making it feasible for environmental sampling. Taken together, the isothermal TSV NASBA/solid-phase detection systems have satisfactory sensitivity and specificity, providing an easy and simple approach for TSV detection.

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