RESEARCH ARTICLE

Sensitive Visual Detection of AHPND Bacteria Using Loop-Mediated Isothermal Amplification Combined with DNA-Functionalized Gold Nanoparticles as Probes

Narong Arunrut1, Jantana Kampeera1, Sarawut Sirithammajak1, Piyachat Sanguanrut2, Porranee Proespraiwong3, Rungkarn Suebsing1, Wansika Kiatpathomchai1*

1 Bioengineering and Sensing Technology Laboratory, BIOTEC, National Science and Technology Development Agency (NSTDA), 113 Thailand Science Park, Phahonyothin Rd., Khlong Nueng, Khlong Luang, Pathum Thani 12120, Thailand, 2 Center of Excellence for Shrimp Molecular Biology and Biotechnology (CENTEX Shrimp), Faculty of Science, Mahidol University, Rama VI road, Bangkok 10400, Thailand, 3 Aquatic Animal Health Research Center, Charoen Pokphand Foods Public CO., LTD. Samutsakorn 74000, Thailand

* wansika@biotec.or.th

Abstract

Acute hepatopancreatic necrosis disease (AHPND) is a component cause of early mortality syndrome (EMS) of shrimp. In 2013, the causative agent was found to be unique isolates of Vibrio parahaemolyticus (VP) carrying binary Pir-like toxin genes PirvpA and PirvpB. In Thailand, AHPND was first recognized in 2012, prior to knowledge of the causative agent, and it subsequently led to a precipitous drop in shrimp production. After VP was characterized, a major focus of the AHPND control strategy was to monitor broodstock shrimp and post larvae for freedom from VP by nucleic acid amplification methods, most of which required use of expensive and sophisticated equipment not readily available in a shrimp farm setting. Here, we describe a simpler but equally sensitive approach for detection of VP based on loop-mediated isothermal amplification (LAMP) combined with unaided visual reading of positive amplification products using a DNA-functionalized, ssDNA-labeled nanogold probe (AuNP). The target for the special set of six LAMP primers used was the VP gene. The LAMP reaction was carried out at 65°C for 45 min followed by addition of the red AuNP solution and further incubation at 65°C for 5 min, allowing any gene amplicons present to hybridize with the probe. Hybridization protected the AuNP against aggregation, so that the solution color remained red upon subsequent salt addition (positive test result) while unprotected AuNP aggregated and underwent a color change from red to blue and eventually precipitated (negative result). The total assay time was approximately 50 min. The detection limit (100 CFU) was comparable to that of other commonly-used methods for nested PCR detection of VP and 100-times more sensitive than 1-step PCR detection methods (10^4 CFU) that used amplicon detection by electrophoresis or spectrophotometry. There was no cross reaction with DNA templates derived from non-AHPND bacteria commonly found in shrimp.
Introduction

Early mortality syndrome (EMS) refers to unusually high mortality in cultivated shrimp within approximately 35 days after stocking of rearing ponds. A component of EMS is acute hepatopancreatic necrosis disease (AHPND) that was first described as acute hepatopancreatic necrosis syndrome (AHPNS) in farmed pacific white shrimp (*Penaeus vannamei*) and giant or black tiger shrimp (*Penaeus monodon*) from China in 2009 [1,2]. The name was changed to AHPND when the causative agent was later discovered to be unique isolates of *Vibrio parahaemolyticus* (VP_AHPND) that carry a 69 kbp plasmid (pAP1) that contains binary Pir-like toxin genes *Pir*<sup>P</sup> A and *Pir*<sup>P</sup> B [3–6]. AHPND spread from China to Vietnam, Malaysia, Thailand, Mexico and the Philippines [7]. After its first appearance in Thailand on the eastern coast of the Gulf of Thailand in late 2012, shrimp production dropped from a high of approximately 600,000 metric tons in 2011 to less than 200,000 in 2014 (FishStat; Food and Agriculture Organization of the United Nations) from a combination AHPND mortality and farmer reluctance to stock ponds until a solution was found.

Recently, molecular tools such as polymerase chain reaction (PCR) based on targeting the toxin genes *Pir*<sup>P</sup> A and *Pir*<sup>P</sup> B have been reported for early detection and prevention of AHPND spread. Several one-step PCR methods target either the *Pir*<sup>P</sup> A gene or *Pir*<sup>P</sup> B gene [5, 6, 8, 9], while the AP4 nested PCR method targets both the *Pir*<sup>P</sup> A and *Pir*<sup>P</sup> B genes [10]. The total assay detection time for these methods may require more than 8–12 h including steps of enrichment (for the 1-step method), DNA extraction, PCR amplification and amplicon detection by electrophoresis. In contrast, loop-mediated isothermal amplification (LAMP) achieves synthesis of large amounts of DNA in a shorter time and in a simpler manner without sacrificing sensitivity or specificity, and it requires only a heating block or hot water bath rather than an expensive thermocycler.

LAMP amplicons are usually detected by agarose gel electrophoresis (LAMP-AGE), followed by staining with carcinogenic ethidium bromide as described in a recent VP_AHPND detection method [11]. To speed up the process by avoiding electrophoresis and to confirm the LAMP amplicons by a hybridization step, detection can be achieved using DNA-functionalized gold nanoparticles (AuNP) [12–15]. AuNPs chemically functionalized with alkyl thiol-terminated oligonucleotides are highly stable in saline solutions and can hybridize with complementary nucleic acids in a very selective and cooperative manner [16]. AuNPs exhibit characteristics of a surface plasmon resonance (SPR) absorption band in the visible light region with the spectrum dependent upon the inter-particle distance. Specifically, particle aggregation gives rise to a shift of the SPR absorption band and a concomitant red to purple-blue color change. This property has been utilized for solution-phase colorimetric detection of specific nucleic acid sequences [12–15]. In this manner, the detection method not only makes the amplicons visible by the unaided eye but also has the advantage of confirming their identity by DNA hybridization.

In this paper, LAMP was combined with use of an ss-DNA-labeled AuNP probe for detection of a *Pir*<sup>P</sup> A target gene sequence in DNA extracts from VP_AHPND bacteria or from shrimp infected with them. Visual detection of the LAMP amplicons by the unaided eye was based on their ability to hybridize with the complementary gold-bound ss-DNA and thus prevent the normal red to purple-blue color change that would otherwise occur by salt-induced aggregation of the gold particles, as shown in Fig 1. The LAMP method combined with amplicon
detection by AuNP has advantages over previously published methods for VPAHPND detection by PCR or LAMP followed by electrophoresis in terms of reduced assay time, amplicon confirmation by hybridization and use of simpler equipment (i.e., no need for a thermocycler, electrophoresis equipment or a UV trans-illuminator.

Materials and Methods

Bacterial strains and DNA preparation

A total of 89 bacterial isolates were used. These included 77 isolates of *V. parahaemolyticus* obtained from shrimp, shrimp pond water or shrimp pond sediments. After isolation, all were tested for ability to cause AHPND by the laboratory bioassay of Tran et. al. (2015) [3] and 50 isolates were found to cause AHPND (VPAHPND isolates) while 27 isolates did not (non-AHPND bacterial isolates), as previously reported [5,17]. An additional set of 7 isolates representing other *Vibrio* species commonly found in diseased shrimp in Thailand and 6 isolates representing non-*Vibrio* species were obtained from culture collections. A summary of these isolates is given in Table 1. *Bacillus subtilis* was included because it is often used as a shrimp probiotic, and the other 4 isolates were included because of our unpublished results indicating their possible presence in shrimp from EMS ponds. The isolates were stored at -80°C and re-streaked on suitable agar plates as previously described [18,19]. In brief, all *Vibrio* isolates were cultured on thiosulfate citrate bile salt sucrose agar (TCBS agar; Difco) containing additional 1.5% NaCl, while non-*Vibrio* isolates were cultured on tryptic soy agar (TSA; Difco) and incubated at 37°C overnight. Although it is customary to cultivate *Vibrio* pathogens of shrimp at 28–30°C for pathology studies with shrimp, the objective of our cultivation was to obtain DNA extracts only, and all the isolates we used grew sufficiently well at 37°C for this purpose. We selected 37°C for convenience related to limitations in incubator space. Bacterial DNA was extracted from a single loop of cells from these agar cultures using a Genomic DNA Purification Kit (Fermentas) according to the manufacturer’s protocol. The concentration and quality of the extracted DNA were analyzed by spectrophotometer at 260 and 280 nm and kept at -80°C until used.

Preparation of AuNPs

The colloidal solution containing AuNPs with an average diameter of 15 nm ± 3.5 nm was prepared as previously reported with minor modifications [15,20]. In brief, all glassware was thoroughly cleaned in *aqua regia* cleaning solution (three parts HCl and one part HNO₃), rinsed in double-distilled water and oven dried prior to use. In a 250 ml round-bottom flask, 100 ml of a 1 mM solution of HAuCl₄ (Sigma-Aldrich, USA) in double-distilled water was brought to a boil with vigorous stirring followed by the addition of 10 ml of 40 mM trisodium citrate...
Table 1. Bacterial isolates used in this study.

| Bacterial isolates | Bioassay | Amplification | Origin | Source |
|--------------------|----------|---------------|--------|--------|
|                    |          | NestedPCR     | LAMP-AGE | LAMP-AuNP |
| V. parahaemolyticus|          |               |         |         |
| 1D                 | ✓        | +             | +       | +       | P. vannamei, Centex |
| 3HP                | ✓        | +             | +       | +       | P. vannamei, Centex |
| 5HP                | ✓        | +             | +       | +       | P. vannamei, Centex |
| SA                 | x        | –             | –       | –       | Shrimp pond, DMST   |
| SB                 | x        | –             | –       | –       | Shrimp pond, DMST   |
| CHN                | ✓        | +             | +       | +       | P. vannamei, Centex |
| F1-CP              | ✓        | +             | +       | +       | P. vannamei, CP     |
| F2-CP              | ✓        | +             | +       | +       | P. vannamei, CP     |
| F3-CP              | ✓        | +             | +       | +       | P. vannamei, CP     |
| F4-CP              | ✓        | +             | +       | +       | P. vannamei, CP     |
| F5-CP              | ✓        | +             | +       | +       | P. vannamei, CP     |
| F6-CP              | ✓        | +             | +       | +       | P. vannamei, CP     |
| F7-CP              | ✓        | +             | +       | +       | P. vannamei, CP     |
| F8-CP              | ✓        | +             | +       | +       | P. vannamei, CP     |
| F9-CP              | ✓        | +             | +       | +       | P. vannamei, CP     |
| F10-CP             | ✓        | +             | +       | +       | P. vannamei, CP     |
| F11-CP             | x        | –             | –       | –       | P. vannamei, CP     |
| F12-CP             | x        | –             | –       | –       | P. vannamei, CP     |
| F13-CP             | x        | –             | –       | –       | P. vannamei, CP     |
| F14-CP             | x        | –             | –       | –       | P. vannamei, CP     |
| F15-CP             | x        | –             | –       | –       | P. vannamei, CP     |
| F16-CP             | x        | –             | –       | –       | P. vannamei, CP     |
| F17-CP             | x        | –             | –       | –       | P. vannamei, CP     |
| F18-CP             | x        | –             | –       | –       | P. vannamei, CP     |
| F19-CP             | x        | –             | –       | –       | P. vannamei, CP     |
| F20-CP             | ✓        | +             | +       | +       | P. vannamei, CP     |
| F21-CP             | x        | –             | –       | –       | P. vannamei, CP     |
| VP1-CP             | x        | –             | –       | –       | P. vannamei, CP     |
| VP2-CP             | x        | –             | –       | –       | P. vannamei, CP     |
| VP3-CP             | ✓        | +             | +       | +       | P. vannamei, CP     |
| VP4-CP             | ✓        | +             | +       | +       | P. vannamei, CP     |
| VP5-CP             | x        | –             | –       | –       | P. vannamei, CP     |
| VP6-CP             | ✓        | +             | +       | +       | P. vannamei, CP     |
| VP7-CP             | ✓        | +             | +       | +       | P. vannamei, CP     |
| VP8-CP             | ✓        | +             | +       | +       | P. vannamei, CP     |
| VP9-CP             | ✓        | +             | +       | +       | P. vannamei, CP     |
| VP10-CP            | x        | –             | –       | –       | P. vannamei, CP     |
| UN1-CP             | x        | –             | –       | –       | P. vannamei, CP     |
| UN2-CP             | x        | –             | –       | –       | P. vannamei, CP     |
| UN3-CP             | ✓        | +             | +       | +       | P. vannamei, CP     |
| UN4-CP             | ✓        | +             | +       | +       | P. vannamei, CP     |
| UN5-CP             | x        | –             | –       | –       | P. vannamei, CP     |
| UN6-CP             | ✓        | +             | +       | +       | P. vannamei, CP     |
| UN7-CP             | ✓        | +             | +       | +       | P. vannamei, CP     |
| UN8-CP             | x        | –             | –       | –       | P. vannamei, CP     |

(Continued)
| Bacterial isolates | Bioassaytest | Amplification | Origin | Source |
|-------------------|-------------|---------------|--------|--------|
|                   |             | NestedPCR | LAMP-AGE | LAMP-AuNP |      |
| UN9-CP            | ✓           | +         | +       | +       | P. vannamei | CP |
| UN10-CP           | ✓           | +         | +       | +       | P. vannamei | CP |
| CAAHRI1-CP        | ✓           | +         | +       | +       | P. vannamei | CP |
| CAAHRI2-CP        | ✓           | +         | +       | +       | P. vannamei | CP |
| CAAHRI3-CP        | ✓           | +         | +       | +       | P. vannamei | CP |
| CAAHRI4-CP        | ✓           | +         | +       | +       | P. vannamei | CP |
| CAAHRI5-CP        | ✓           | +         | +       | +       | P. vannamei | CP |
| CAAHRI6-CP        | ✓           | +         | +       | +       | P. vannamei | CP |
| CAAHRI7-CP        | ✓           | +         | +       | +       | P. vannamei | CP |
| CAAHRI8-CP        | ✓           | +         | +       | +       | P. vannamei | CP |
| CAAHRI9-CP        | x           | –         | –       | –       | P. vannamei | CP |
| CAAHRI10-CP       | ✓           | +         | +       | +       | P. vannamei | CP |
| CAAHRI11-CP       | ✓           | +         | +       | +       | P. vannamei | CP |
| CAAHRI12-CP       | x           | –         | –       | –       | P. vannamei | CP |
| CAAHRI13-CP       | x           | –         | –       | –       | P. vannamei | CP |
| CAAHRI14-CP       | x           | –         | –       | –       | P. vannamei | CP |
| CAAHRI15-CP       | ✓           | +         | +       | +       | P. vannamei | CP |
| CAAHRI16-CP       | ✓           | +         | +       | +       | P. vannamei | CP |
| CAAHRI17-CP       | ✓           | +         | +       | +       | P. vannamei | CP |
| CAAHRI18-CP       | ✓           | +         | +       | +       | P. vannamei | CP |
| CAAHRI19-CP       | x           | –         | –       | –       | P. vannamei | CP |
| CAAHRI20-CP       | ✓           | +         | +       | +       | P. vannamei | CP |
| CAAHRI21-CP       | ✓           | +         | +       | +       | P. vannamei | CP |
| CAAHRI22-CP       | ✓           | +         | +       | +       | P. vannamei | CP |
| CAAHRI23-CP       | ✓           | +         | +       | +       | P. vannamei | CP |
| CAAHRI24-CP       | ✓           | +         | +       | +       | P. vannamei | CP |
| CAAHRI25-CP       | ✓           | +         | +       | +       | P. vannamei | CP |
| CAAHRI26-CP       | x           | –         | –       | –       | P. vannamei | CP |
| CAAHRI27-CP       | ✓           | +         | +       | +       | P. vannamei | CP |
| CAAHRI28-CP       | x           | –         | –       | –       | P. vannamei | CP |
| CAAHRI29-CP       | ✓           | +         | +       | +       | P. vannamei | CP |
| CAAHRI30-CP       | ✓           | +         | +       | +       | P. vannamei | CP |
| V. vulnificus     |             |           |         |         |        |
| VVS4907001        | x           | –         | –       | –       | P. vannamei | DBSWU |
| VVS4907011        | x           | –         | –       | –       | P. vannamei | DBSWU |
| V. harveyi        |             |           |         |         |        |
| Centex639         | x           | –         | –       | –       | P. monodon | Centex |
| Centex1526        | x           | –         | –       | –       | P. monodon | Centex |
| V. alginolyticus  |             |           |         |         |        |
| DMST22082         | x           | –         | –       | –       | Stool (human) | DMST |
| DMST22084         | x           | –         | –       | –       | Food (human) | DMST |
| DMSC14800         | x           | –         | –       | –       | Seafood (human) | DMSC |
| B. subtilis       | x           | –         | –       | –       | Shrimp probiotic | Centex |
| Shewanella sp.    | x           | –         | –       | –       | P. vannamei | Centex |
| Rhodococcus fascians | x           | –         | –       | –       | Not specified | NCCB |

(Continued)
The solution turned deep blue immediately but later changed to a final wine-red. After this color change, boiling was continued for an additional 15 min before the heater was turned off and the colloidal AuNP solution was continuously stirred overnight. The resulting solution of AuNPs was characterized by an absorption maximum at 520 nm and it was stored in dark bottles at 4°C.

Preparation of DNA-labelled AuNP probes

An ssDNA probe sequence was designed to complement that which spanned the F1c-B1c region of the AHPND-LAMP amplicon. It was labelled with a thiol group at the 5'-end (Table 2). The DNA-labelled AuNPs were prepared as previously described [12–15] with slight modifications. In brief, 10 ml of the colloidal AuNP solution was incubated with 5 nmol (i.e., 50 μl of the 100 μM stock) of 5'-thiol-modified ssDNA probes at 50°C with shaking at 150 rpm for 22 h. Then, the solution was transferred to 1 ml of phosphate buffer (100 mM sodium phosphate buffer, PH 7.6) containing 1 M NaCl and 10% SDS and incubated under the same conditions for another 4 h. The AuNPs were pelleted by centrifugation at 20,000 g at 4°C for 30 min to remove the unbound ssDNA. The supernatant solution was removed and the pelleted DNA-labelled AuNPs were washed with 5 ml washing buffer (100 mM PBS, 100 mM NaCl, 0.01% SDS) and finally re-suspended in 1 ml of the same buffer and kept at 4°C until used.

LAMP primer design and optimization

A set of six primers was designed for LAMP to target eight distinct regions of the PirP'A gene of VP_AHPND isolates according to the sequence of GenBank accession no. KM067908.1 using Table 2. Primers and probe used for LAMP to detect VP_AHPND.

Table 2. (Continued)

| Bacterial isolates       | Bioassaytest | Amplification | Origin | Source |
|--------------------------|--------------|---------------|--------|--------|
|                          |              | NestedPCR     | LAMP-AGE | LAMP-AuNP |
| Delftia acidovorans      | x            | -             | -      | -      |
| Ralstonia solanacearum   | x            | -             | -      | -      |

(Sigma-Aldrich, USA). The solution turned deep blue immediately but later changed to a final wine-red. After this color change, boiling was continued for an additional 15 min before the heater was turned off and the colloidal AuNP solution was continuously stirred overnight. The resulting solution of AuNPs was characterized by an absorption maximum at 520 nm and it was stored in dark bottles at 4°C.
Primer Explorer ver. 4 (http://primerexplorer.jp/lamp4.0.0/index.html). The sequences of the primers and their locations are indicated in Table 2. All primers were synthesized by Bio Basic Inc., Canada. To determine the optimal temperature for the LAMP assay, reactions were performed on a thermal cycler (Gene Amp PCR System 2700, Applied Biosystems®) at 60, 63 and 65°C for 1 h, followed by 85°C for 7 min to terminate the reaction. The products were analyzed by 2% agarose gel electrophoresis (AGE). The LAMP reaction mixtures (25 μl) consisted of 0.2 μM outer primers (F3 and B3), 2 μM inner primers (FIP and BIP), 0.2 μM loop primers (LF and LB), 0.4 M betaine (USB Corporation, USA), 1.2 mM dNTPs (Promega, USA), 6 mM MgSO4 (Sigma-Aldrich, USA), 8 U Bst DNA polymerase large fragment with the 1x buffer supplied (New England Biolabs, USA) and the specified amount of template DNA. DNA extracted from uninfected shrimp samples and sterile water were included as negative controls, and DNA extracted from the VP_{AHPND} isolate 5HP [17] was used as a positive control. To test specificity of the LAMP primers, DNA templates from 80 bacterial cultures (Table 1) were used to test the LAMP assay followed by both AGE and AuNP probe analysis.

Optimization of the AuNP-probe hybridization step

The conditions for optimization of AuNP-probe hybridization were previously described [12–15] Briefly, hybridization for the detection of LAMP products was conducted in a total volume of 15 μl by mixing together the AuNP probe solution (5 nM) with the LAMP product solution at various ratios ranging from [1 μl AuNP solution: 9 μl product solution (1:9)] to [9 μl AuNP solution: 1μl product solution (9:1)] before incubation at 65°C for 5 min. After determining the optimum ratio, the conditions for salt-induced AuNP probe aggregation were determined using salt concentrations ranging from 3 to 666 mM MgSO4 in a fixed volume of 5 μl. Results were compared using LAMP amplicons obtained using DNA templates extracted from AHPND-bacterial culture isolate 5HP (positive control) and using distilled water and DNA extracted from shrimp infected with white spot syndrome virus (WSSV) (non-complementary target DNA) as negative controls. Color changes (red to blue color) were compared by the unaided eye and by UV-visible spectrum analysis (Thermo Fisher Scientific).

Detection of VP_{AHPND} by 1-step and nested PCR

The DNA extracted from bacterial samples was used as a template for PCR amplification by both 1-step and nested PCR detection methods targeting the PirvpA gene. Conditions for the AP3 1-step PCR method were similar to those described in a previous report [5] with some modifications. The 336-bp target fragment was amplified with primers F-AP3 and R-AP3 (Table 3) using the cycling protocol of 94°C for 5 min followed by 30 cycles of 94°C for 30 s, 55°C for 20 s and 72°C for 40 s and a final extension step at 72°C for 5 min.

The two-tube nested PCR method (AP4) was carried out as in a previous report [10] with some modifications. The first-step PCR target of 1269-bp was amplified using primers AP4-F1 and AP4-R1 (Table 3), and this amplicon was then used as the template for a second PCR reaction that yielded a 230-bp amplicon using primers AP4-F2 and AP4-R2 (Table 3). Each PCR reaction was conducted in a 25 μl reaction mixture containing 1 x PCR buffer, 0.2 mM dNTPs (Promega), 3 mM MgCl2 (Invitrogen), 1.5 units Taq DNA polymerase (Invitrogen), 0.2 μM each forward and reverse primer, and 2 μl of template DNA. The PCR amplification was conducted with an initial cycle at 94°C for 2 min followed by 30 cycles of 94°C for 20 s, 55°C for 30 s and 72°C for 90 s (for external primers) or 20 s (for internal primers) followed by an extension step at 72°C for 2 min.
Comparison of sensitivity between LAMP-AuNP and traditional PCR assays

The sensitivity comparison was carried out using tenfold serial dilutions of DNA extracted from pure cultures of VP_{AHPND} isolate 5HP at concentrations ranging from 10^7 to 1 CFU/ml and prepared as previously described [18,19] with some modifications. Briefly, a small number of cells from a single bacterial colony on TCBS agar was inoculated into 5 ml of tryptic soy broth (TSB; Difco) supplemented with 1.5% NaCl and incubated overnight at 37°C. Then, 50 μl of this TSB culture was transferred into a new tube containing 5 ml of TSB followed by incubation at 37°C with shaking at 250 rev min^{-1} to obtain mid-log phase cells (OD_{600} nm = 0.5). Tenfold serial dilutions of these cultures were prepared in phosphate buffered saline solution (PBS).

DNA template was prepared from these dilutions by transferring 100 μl from each dilution into a 1.5 ml microcentrifuge tube that was centrifuged at 15,000 g for 5 min. After removal of the supernatant, the pellets were subjected to DNA extraction using a Genomic DNA Purification Kit (Fermentas) according to manufacturer’s protocol. The resulting DNA extracts were used as templates for the LAMP-AuNP method and for the AP3 and AP4 PCR methods. The sensitivity tests were carried out in triplicate, and the last dilution that gave positive results with all three of the replicates was considered to be the detection limit for each method.

In parallel with the above, 100 μl aliquots of each dilution were spread on TSA supplemented with 1.5% NaCl (in duplicate) and the plates were incubated overnight at 37°C to determine bacterial counts. After colonies were visible, plates for counting were selected from dilutions that yielded 30–300 colony-forming units (CFUs) and these counts were used to calculate the CFU ml^{-1} of all the bacterial suspensions used.

Specificity of the LAMP-AuNP assay for VP_{AHPND} detection

To test for cross-hybridization using the AuNP probe with LAMP products from other pathogens, the amplicons from a VP_{AHPND} DNA template (this study) served as the positive control for analysis by the AuNP colorimetric assay (analyzed by naked eye and confirmed by UV-vis spectrophotometry). The results were compared with those obtained using LAMP or RT-LAMP amplicons obtained by comparable LAMP methods for the following non-shrimp pathogens *Mycobacterium tuberculosis* (TB) [21] and *Plasmodium* (Malaria) [22], and for the shrimp pathogens WSSV [23], YHV [12], IMNV [13], IHHNV [24], TSV [25], LSNV [26] and PemoNPV [27].

Evaluation of the LAMP-AuNP method for VP_{AHPND} detection in field samples

It has previously been recommended that shrimp and environmental samples to be tested for VP_{AHPND} using the AP3, 1-step PCR method should be subjected to a preliminary culture enrichment step to avoid false negative test results [5]. The subsequent AP4, nested PCR detection

### Table 3. Primers used for 1-step PCR (AP3 method) and nested PCR (AP4 method) for detection of VP_{AHPND}

| Primer name | Sequence (5’-3’) | Length (bp) |
|-------------|-----------------|-------------|
| F-AP3       | ATGAGTAACAATATAAAACATGAAC | 26          |
| R-AP3       | GTGGTAATAGATGTACAGAA | 21          |
| F1-AP4      | ATGAGTAACAATATAAAACATGAAC | 26          |
| R1-AP4      | ACGATTCTGACGTTCACCA | 20          |
| F2-AP4      | TTAGGAATACGGGACGTTGGG | 20          |
| R2-AP4      | GTTAGTCATGTGAGCACCCTTTC | 21          |

doi:10.1371/journal.pone.0151769.t003
method was developed for testing samples that could not be enriched before testing (e.g., samples preserved in alcohol or archived DNA samples) [10]. Thus, we wished to compare the test results of these two methods with the LAMP-AuNP method to determine whether it might be used for the same purpose as the AP4 method for DNA extracts derived directly from shrimp stomach tissue (i.e., no enrichment step). Samples including 20 black tiger shrimp and 10 whiteleg shrimp were arbitrarily selected from shrimp farms where some VP<sub>AHPND</sub> specimens had been previously detected. They were kindly provided by the Charoen Pokphand Foods (CPF) PCL Laboratory. The samples themselves were of unknown VP<sub>AHPND</sub> infection status. Stomach tissue from individual shrimp was used for DNA extraction as described above. The extracted DNA was then used as the template for VP<sub>AHPND</sub> detection by the LAMP-AuNP method and by the AP3 1-step and AP4 nested PCR methods. Then, the results for each method were compared.

**Results and Discussion**

**Optimization of the reaction for VP<sub>AHPND</sub> detection by LAMP**

Tests for the optimal LAMP reaction temperature using 100 ng of DNA template from the VP<sub>AHPND</sub> isolated 5HP revealed that 65°C gave a slightly better result by AGE analysis than 60 or 63°C (Fig 2). Thus, 65°C was selected as the standard assay temperature. When the LAMP reactions were conducted at 65°C for 30, 45 and 60 min using various concentrations of DNA template, the 45 and 60 min reaction times gave clear LAMP amplicon patterns of the same intensity while 30 min yielded pattern of lower intensity (data not shown). Thus, the shortest reliable time of 45 min was chosen as the standard assay time.

**AuNP synthesis and hybridization for detection of VP<sub>AHPND</sub> LAMP amplicons**

Successfully synthesized AuNPs gave a UV-Vis spectrum with one peak at 525 nm (Fig 3), while the ssDNA-labelled AuNPs absorbed at 530 nm (Fig 3). This wavelength shift confirmed that the preparation contained monodispersed AuNPs.
Tests of the effect on hybridization when using a 5 nM AuNP solution at variable ratios with a LAMP product solution revealed that the best hybridization and aggregation result after addition of 50 mM MgSO₄ was obtained using a mixture of the AuNP probe solution and LAMP product solution at a ratio of 5:5. This gave the clearest color difference between the positive red result and negative purple-blue result (Fig 4A).

After optimizing the ratio of the AuNP probe and LAMP product solutions, tests needed to optimize the MgSO₄ concentration to obtain the best visible test result from AuNP aggregation revealed that a final concentration of 50 mM was the most suitable (Fig 4B). It gave a distinct red positive result for VP_{AHPND}-LAMP amplicons and a clearly contrasted purple-blue negative result for the distilled water and non-target DNA probe negative controls. Lower concentrations of MgSO₄ (3 to 17 mM MgSO₄) resulted in false-positive results (pink color) with the negative controls, while higher concentrations (100 and 667 mM MgSO₄) resulted in false negative results (purple-blue/clear) with LAMP amplicons from VP_{AHPND} (Fig 4B). Therefore, the optimal MgSO₄ concentration chosen in this study was 50 mM.

Comparative sensitivity of LAMP-AuNP and PCR-electrophoresis methods
Using DNA extracts from ten-fold serial dilutions of VP_{AHPND} isolate 5HP, the LAMP-AuNP method was able to detect 5HP in a solution containing 100 CFU/ml (Fig 5A). This result showed similar sensitivity to LAMP-AuNP followed by UV-Vis analysis (Fig 5B), LAMP followed by AGE (Fig 5C) and nested-PCR using the AP4 method that amplifies a 230-bp fragment (Fig 6A). By contrast, the AP3, 1-step PCR method required a solution containing 10⁴ CFU/ml to obtain a positive result (Fig 6B). Another advantage of the LAMP-AuNP method was the short, total assay time of 50 min (45 min for LAMP, 5 min for hybridization and less than 1 min salt-induced aggregation). This compared to 90–120 min for the LAMP-AGE method, 3–5 h for the AP3, 1-step PCR method (excluding enrichment that might be required for natural samples) and 4–6 h for the AP4 nested-PCR method.

Specificity of LAMP-AuNP for VP_{AHPND} detection
Tests for cross hybridization using the AuNP probe with amplicons produced using LAMP methods for other shrimp pathogens (Fig 7A) gave no positive results using the AHPND AuNP probe
Instead, the LAMP-AuNP assay gave a red positive result only with the LAMP product from the VPAHPND isolate. The color of the positive LAMP-AuNP test result was stable over 30 min after salt addition, while LAMP products from other pathogens gave an immediate color change to purple-blue followed by precipitation of the aggregated probe to yield a colorless supernatant solution within 30 min (not shown). The results by UV-Vis detection for the red nanogold probe showed identical specificity results (Fig 7C) to those using the unaided eye (Fig 7B).

**Detection of VP**<sub>AHPND</sub> **using LAMP-AuNP, LAMP-AGE and nested PCR with DNA from pure cultures of various bacterial isolates**

Testing the specificity of LAMP-AuNP for detection of VP<sub>AHPND</sub> with DNA templates from 89 bacterial isolates (Table 1) revealed that both the LAMP-AuNP assay and nested PCR assay gave positive test results for all 50 VP<sub>AHPND</sub> isolates but negative results for the 40 non-AHPND isolates. The LAMP-AGE assay also gave positive test results for all the VP<sub>AHPND</sub> isolates and negative results for all the non-AHPND isolates.

**Comparison of LAMP-AuNP and PCR methods for detection of VP**<sub>AHPND</sub> **in field samples**

These tests were carried out using 30 samples of shrimp of unknown VP<sub>AHPND</sub> infection status arbitrarily selected from shrimp farms were shrimp infected with VP<sub>AHPND</sub> had been detected.
Fig 5. Sensitivity of the LAMP-AuNP assay for the detection of VP$	ext{}_{\text{AHPND}}$ using 10-fold serial dilution of DNA extracted from a culture of VP$	ext{}_{\text{AHPND}}$ isolate 5HP ($10^7$–1 CFU/ml). (A) Colorimetric results of LAMP followed by AuNP probe assay. (B) UV-visible spectrum analysis corresponding to the individual tubes in Fig 5A (measured after salt addition). (C) AGE results of LAMP reactions. Lane M: 2 log DNA marker and N: 100 ng of DNA extracted from normal shrimp.

doi:10.1371/journal.pone.0151769.g005
The purpose was to test whether the LAMP-AuNP method could be used similarly to the AP4 PCR method to detect VPAHPND in shrimp or other specimens without the culture enrichment step usually employed prior to using the AP3, 1-step PCR detection method. Using direct DNA extracts from the stomachs of the shrimp specimens as templates, 7 of the 30 shrimp (2 black tiger shrimp and 5 whiteleg shrimp) (23.3%) gave positive test results for VPAHPND using the 1-step PCR AP3 method. In contrast, 12 of the 30 samples (7 more for a total 40%) gave positive test results for VPAHPND by both the LAMP-AuNP and the AP4

Fig 6. Comparison of a sensitivity test carried out using total DNA template as in Fig 5 with traditional PCR methods. (A) Nested PCR followed by AGE (AP4 method). (B) 1-step PCR followed by AGE (AP3 method). Lane M: 2 log DNA marker and N: 100 ng of DNA extracted from normal shrimp.

doi:10.1371/journal.pone.0151769.g006

previously.
Fig 7. Comparison of results obtained using the VP<sub>AHPND</sub> AuNP hybridization probe with LAMP amplicons from VP<sub>AHPND</sub> (Lane 1) and other common pathogens (Lanes 2–10). (A) The result of agarose gel electrophoresis (AGE) of LAMP products from various pathogens. Lane M: 2 log DNA marker; Lane N: normal shrimp DNA as negative control; Lanes 2–10: TB, Plasmodium (Malaria), WSSV, YHV, IMNV, IHHNV, TSV, LSNV and PemoNPV, respectively. (B) Colorimetric result for the same LAMP products as in Fig 7A measured after salt addition. (C) UV-visible spectra analysis corresponding to the individual tubes in Fig 7B measured after salt addition.

doi:10.1371/journal.pone.0151769.g007
nested-PCR methods (Table 4). Together with the results on comparative sensitivity described in the preceding section above, these results showed that the LAMP-AuNP method has similar utility to the AP4 nested PCR method in detecting $V_{PAHNP}$ in samples where the target DNA is too low in concentration to be detected directly by a representative 1-step PCR method. This test was not carried out to determine the prevalence of such lightly-infected samples that might occur in field testing for $V_{PAHNP}$, but to show that the LAMP-AuNP method can be used to test samples that are not enriched before testing. At the same time, it is clear that the LAMP-AuNP was faster and simpler to use in farm-site laboratories than the AP4 method.

### Overall conclusions for the LAMP-AuNP method to detect $V_{PAHNP}$

The initial detection methods for $V_{PAHNP}$ were based on standard 1-step PCR detection of the pAP1 plasmid, but these methods gave a small percentage (~3%) of false positive results, probably due to absence of the Pir$^{PA}$ and Pir$^{PB}$ toxin genes on pAP1 plasmids of the bacteria being tested [5]. Subsequent 1-step PCR detection methods targeted either the Pir$^{PA}$ or Pir$^{PB}$ toxin [5, 6, 8, 9] and none of those reports included any false negative or false positive results for $V_{PAHNP}$ in the form of specimens that have tested positive or negative, respectively, for the Pir$^{PA}$ or Pir$^{PB}$ genes. In other words, all AHPND specimens positive for one of the toxins have also been positive for the other. This suggests that prevalence of the theoretically possible mutant specimens carrying only one or the other of the two toxin genes is very low and that for practical purposes only one of the toxin genes is sufficient for detection of $V_{PAHNP}$. Indeed, the AP4 nested PCR method that targets both toxins has given identical results to those obtained using the AP3 1-step PCR method with large numbers of samples, and the two methods differ only in higher sensitivity of the nested PCR method [10].

More recently, a LAMP-AGE method for detection of $V_{PAHNP}$ has been published [11], but compared to the LAMP-AuNP method, it has the disadvantages of requiring the use of electrophoresis equipment and lacking a hybridization step to confirm the specific nature of the LAMP amplicons. There are two other LAMP amplicon detection protocols (not yet reported for use in $V_{PAHNP}$ detection) that do not use electrophoresis but give color reactions visible to the naked eye, like the LAMP-AuNP method. However, both lack a hybridization step to confirm the nature of the amplicons. These are the calcein method [28] and the hydroxynaphthol blue method [29], both of which measure accumulation of the phosphate byproduct of the LAMP reaction and could give false-positive color reactions with non-specific LAMP amplicons [30]. Similarly, the LAMP amplicon detection methods that involve use of the fluorescent, DNA-intercalating dye SYBR Green I, would also give positive fluorescence signals with non-specific amplicons, and they have the added requirement for a fluorescence spectrophotometer [31]. Thus, high sensitivity, high specificity including amplicon confirmation by hybridization, relatively short analysis time and use of simple equipment (i.e., no thermocycler, no electrophoresis equipment and no spectrophotometer) are the key advantages of the LAMP-AuNP detection method for $V_{PAHNP}$. It is a rapid and relatively simple assay with

---

**Table 4. Comparison of detection results for $V_{PAHNP}$ in field samples using LAMP combined with AuNP, 1-step PCR (AP3 method) and nested PCR (AP4 method).**

| Type of sample        | No. of samples | 1-PCR (No. %) | Nested PCR (No. %) | LAMP-AuNP (No. %) |
|-----------------------|----------------|---------------|--------------------|-------------------|
| Whiteleg shrimp       | 10             | 2 (20.0)      | 4 (40.0)           | 4 (40.0)          |
| Black tiger shrimp    | 20             | 5 (25.0)      | 8 (40.0)           | 8 (40.0)          |
| Total                 | 30             | 7 (23.3)      | 12 (40.0)          | 12 (40.0)         |

DOI: 10.1371/journal.pone.0151769.t004
sensitivity comparable to that of traditional nested-PCR detection but suitable for confirmation of AHPND outbreaks in small, farm-scale laboratories.

Acknowledgments
This work was supported by grants from National Research Council of Thailand. The authors would also like to thank Aquatic Animal Health Research Center, Charoen Pokphand Co. Ltd, Thailand for providing some of the AHPND and non-AHPND bacterial isolates used in this study. Thanks also to Prof. T.W. Flegel for assistance in editing the manuscript.

Author Contributions
Conceived and designed the experiments: NA WK. Performed the experiments: NA JK SS. Analyzed the data: NA WK. Contributed reagents/materials/analysis tools: NA PS PP WK. Wrote the paper: NA RS WK.

References
1. Flegel TW. Historic emergence, impact and current status of shrimp pathogens in Asia. J Invertebr Pathol. 2012; 110: 166–173. doi: 10.1016/j.jip.2012.03.004 PMID: 22426934
2. Leano EM, Mohan CV. Early mortality syndrome threatens Asia’s shrimp farms. Glob Aquacult Advocate, July/August 2012: 38–39.
3. Tran L, Nunan L, Redman RM, Mohney LL, Pantoja CR, Fitzsimmons K, et al. Determination of the infectious nature of the agent of acute hepatopancreatic necrosis syndrome affecting penaeid shrimp. Dis Aquat Org. 2013; 105: 45–55. doi: 10.3354/dao02621 PMID: 23836769
4. Yang YT, Chen IT, Lee CT, Chen CY, Lin SS, Hor LI, et al. Draft genome sequences of four strains of Vibrio paraaerolymyticus, three of which cause early mortality syndrome/acute hepatopancreatic necrosis disease in shrimp in China and Thailand. Genome Announc. 2014; 2(5): e00816–14. doi: 10.1128/genomeA.00816-14 PMID: 25189578
5. Sirikharin R, Taengchaiyaphum S, Sanguanrut P, Chi TD, Mavichak R, Proespraiwong P, et al. Characterization and PCR Detection of binary, pir-like toxins from Vibrio parahaemolyticus isolates that cause acute hepatopancreatic necrosis disease (AHPND) in shrimp. PLoS One. 2015; 10: e0126987. doi: 10.1371/journal.pone.0126987
6. Lee CT, Chen IT, Yang YT, Ko TP, Huang YT, Huang JY, et al. The opportunistic marine pathogen Vibrio paraaerolymyticus becomes virulent by acquiring a plasmid that expresses a deadly toxin. Proceedings of the National Academy of Sciences of the United States of America. 2015; 112 (34): 10798–10803. doi: 10.1073/pnas.1503129112 PMID: 26261348
7. Thitamadee S, Prachumwat A, Srisala J, Jaroentak P, Salachan PV, Sritunyalucksana K, et al. Review of current disease threats for cultivated penaeid shrimp in Asia. Aquaculture. 2016; 452: 69–87. doi: 10.1016/j.aquaculture.2015.10.028
8. Han JE, Tang KFJ, Tran LH, Lightner DV. Photobacterium insect-related (Pir) toxin-like genes in a plasmid of Vibrio paraaerolymyticus, the causative agent of acute hepatopancreatic necrosis disease (AHPND) of shrimp. Dis Aquat Org. 2015; 113: 33–40. doi: 10.3354/dao02830 PMID: 25667334
9. Tinwongger S, Proespraiwong P, Thawonsuwan J, Sriwanayos P, Kongkumnerd J, Chaweepack T, et al. Development of PCR diagnosis for shrimp acute hepatopancreatic necrosis disease (AHPND) strain of Vibrio paraaerolymyticus. Fish Pathol. 2014; 49: 159–164.
10. Dangtip S, Sanguanrut P, Srisala J, Jaroentak P, Kiatpathomchai W, Nozaki R, et al. AP4 method for two-tube nested PCR detection of AHPND isolates of Vibrio paraaerolymyticus. Aquaculture. 2015; 437–438: 105–112. doi: 10.1016/j.aquaculture.2015.04.005
11. Koizumi K, Tinwongger S, Nozaki R, Kondo H, Hirono I. Detection of acute hepatopancreatic necrosis disease strain of Vibrio paraaerolymyticus using loop-mediated isothermal amplification. J Fish Dis. 2015. doi: 10.1111/jfd.12387
12. Jaroentak W, Anunut N, Kiatpathomchai W. Rapid and sensitive detection of shrimp yellow head virus using loop-mediated isothermal amplification and a colorogenic nanogold hybridization probe. J Virol Meth. 2012; 186: 36–42. doi: 10.1016/j.jviromet.2012.08.013
13. Anunut N, Kampeera J, Suebsing R, Kiatpathomchai W. Rapid and sensitive detection of shrimp infectious myonecrosis virus using a reverse transcription loop-mediated isothermal amplification and visual
colorogenic nanogold hybridization probe assay. J Virol Meth. 2013; 193: 542–547. doi: 10.1016/j.jviromet.2013.07.017

14. Seetang-Nun Y, Jaroenram W, Srirairatana S, Suebsing R, Kiathpathomchai W. Visual detection of white spot syndrome virus using DNA-functionalized gold nanoparticles as probes combined with loop-mediated isothermal amplification. Mol Cell Probes. 2013; 27: 71–79. doi: 10.1016/j.mcp.2012.11.005 PMID: 23211683

15. Suebsing R, Prombun P, Srisala J, Kiathpathomchai W. Loop-mediated isothermal amplification combined with colorimetric nanogold for detection of the microsporidian Enterocytozoon hepatopenaei in penaeid shrimp. J Appl Microbiol. 2013; 114: 1254–1263. doi: 10.1111/jam.12160 PMID: 23387348

16. Mirkin CA, Letsinger RL, Mucic RC, Storhoff JJ. A DNA-based method for rationally assembling nanoparticles into macroscopic materials. Nature. 1996; 382: 607–609. doi: 10.1038/382607a0 PMID: 8757129

17. Joshi J, Srisala J, Truong VH, Chen I-T, Nuangsaeng B, Suthienkul O, et al. Variation in Vibrio paraheamolyticus isolates from a single Thai shrimp farm experiencing an outbreak of acute hepatopancreatic necrosis disease (AHPND). Aquaculture. 2014; 428–429: 297–302. doi: 10.1016/j.aquaculture.2014.03.030

18. Yamazaki W, Ishibashi M, Kawahara R and Inoue K. Development of a loop-mediated isothermal amplification assay for sensitive and rapid detection of Vibrio parahaemolyticus. BMC Microbiol. 2008; 8:163. doi: 10.1186/1471-2180-8-163 PMID: 18823567

19. Prompamorn P, Sithigomgul P, Rukpratanpon S, Longyant S, Sridulyakul P, Chaisivuthangkura P. The development of loop-mediated isothermal amplification combined with lateral flow dipstick for detection of Vibrio paraheamolyticus. Lett Appl Microbiol. 2010; 52: 344–351.

20. He Y, Zhang S, Zhang X, Baloda M, Xu H, Xu H, et al. Ultrasensitive nucleic acid biosensor based on enzyme-gold nanoparticle dual label and lateral flow strip biosensor. Biosens Bioelectron. 2011; 26: 2018–2024. doi: 10.1016/j.bios.2010.08.079 PMID: 20875950

21. Kaewphinit T, Arunrut N, Kiathpathomchai W, Santiwatanakul S, Jaratsing P, Chansiri K. Detection of Mycobacterium tuberculosis by using loop-mediated isothermal amplification combined with a lateral flow dipstick in clinical samples. Biomed Res Int. 2013: 926230. doi: 10.1155/2013/926230 PMID: 23555102

22. Yongkiettrakul S, Jaroenram W, Arunrut N, Chareanchim W, Pannengpetch S, Suebsing R, et al. Application of loop-mediated isothermal amplification assay combined with lateral flow dipstick for detection of Plasmodium falciparum and Plasmodium vivax. Parasitol Int. 2014; 63: 777–784. doi: 10.1016/j.parint.2014.06.004 PMID: 25038579

23. Jaroernram W, Kiathpathomchai W and Flegel TW. Rapid and sensitive detection of white spot syndrome virus by loop-mediated isothermal amplification combined with a lateral flow dipstick. Mol Cell Probes. 2009; 23: 65–70. doi: 10.1016/j.mcp.2008.12.003 PMID: 19124071

24. Arunrut N, Prombun P, Saksmperome V, Flegel TW, Kiathpathomchai W. Rapid and sensitive detection of infectious hypodermal and hematopoietic necrosis virus by loop-mediated isothermal amplification combined with a lateral flow dipstick. J Virol Methods. 2011; 171: 21–25. doi: 10.1016/j.jviromet.2010.09.022 PMID: 20887752

25. Kiathpathomchai W, Jaroernram W, Arunrut N, Jitrapakdee S, Flegel TW. Shrimp Taura syndrome virus detection by reverse transcription loop-mediated isothermal amplification combined with a lateral flow dipstick. J Virol Methods. 2008; 153: 214–217. doi: 10.1016/j.jviromet.2008.06.025 PMID: 18662723

26. Arunrut N, Suebsing R, Wityayachumnarnkul B, Kiathpathomchai W. Demonstration of a very inexpensive, turbidimetric, real-time, RT-LAMP detection platform using shrimp Laem-singh virus (LSNV) as a model. PLoS One. 2014; 9: e108047. doi: 10.1371/journal.pone.0108047 PMID: 25255231

27. Nimthaphak T, Meemetta W, Arunrut N, Senapin S, Kiathpathomchai W. Rapid and sensitive detection of Peneaus monodon nucleopolyhedrovirus (PemoNPV) by loop-mediated isothermal amplification combined with a lateral-flow dipstick. Mol Cell Probes. 2010; 24: 1–5. doi: 10.1016/j.mcp.2009.09.004 PMID: 19818396

28. Tomita N, Mori Y, Kanda H, Notomi T. Loop-mediated isothermal amplification (LAMP) of gene sequences and simple visual detection of products. Nat Protoc. 2008; 3: 877–82. doi: 10.1038/nprot.2008.57 PMID: 18451795

29. Goto M, Honda E, Ogura A, Nomoto A, Hanaki K. Colorimetric detection of loop-mediated isothermal amplification reaction by using hydroxy naphthol blue. Biotechniques. 2009; 46: 167–72. doi: 10.2144/000113072 PMID: 19317660

30. Jiang YS, Li B, Milligan JN, Bhadra S, Ellington AD. Real-time detection of isothermal amplification reactions with thermostable catalytic hairpin assembly. J Am Chem Soc. 2013; 135: 7430–7433. doi: 10.1021/ja4023978 PMID: 23647466
31. Iwamoto T, Sonobe T, Hayashi K. Loop-mediated isothermal amplification for direct detection of *Mycobacterium tuberculosis* complex, *M. avium*, and *M. intracellulare* in sputum samples. J Clin Microbiol. 2003; 41: 2616–2622. doi: 10.1128/JCM.41.6.2616–2622.2003 PMID: 12791888