Methods and Applications

Establishment and Application of Heminested RT-PCR Assay for Detection of Mosquito-Borne Flavivirus — Guizhou Province, China, 2018

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ABSTRACT

Introduction: Mosquito-borne flavivirus can lead to serious infectious diseases worldwide and cause high mortality and disability. In order to strengthen epidemiological investigation of flavivirus and meet the needs of early warning of diseases, a simple, rapid, and sensitive detection method needs to be established to prevent and control mosquito-borne diseases.

Methods: Using the NS5 gene of flavivirus in GenBank, 3 universal primers targeting the conserved regions were designed. The complementary DNAs (cDNAs) of Japanese encephalitis virus (JEV), dengue virus (DENV), West Nile virus (WNV), and yellow fever virus (YFV) were used as the template to optimize the reaction conditions. A heminested reverse-transcriptase polymerase chain reaction (hnRT-PCR) was established to verify the sensitivity and specificity of this method and to detect field-caught mosquitoes.

Results: Our results showed that this method exhibited better specificity, higher sensitivity, and the ability to detect multiple viruses simultaneously. The lowest detection limit of JEV, DENV-2, YFV, and WNV was 3×10^4 , 3×10^6 , 3×10^5 , and 3×10^4 copies/ μ L, respectively. Mosquito-borne flavivirus was successfully detected in the field-caught mosquito samples using the method established in this study.

Discussion: The hnRT-PCR method established in this study can be employed for the rapid detection of flavivirus and provide technical support for early and rapid diagnosis of mosquito-borne flavivirus.

INTRODUCTION

Mosquito-borne viruses can be transmitted through mosquito bites on sensitive vertebrates and cause natural-focus infection diseases worldwide (1). Flavivirus is one of the most important mosquito-borne viruses, including Japanese encephalitis virus (JEV), dengue virus (DENV), West Nile virus

(WNV), yellow fever virus (YFV), etc., which can lead to fever, headache, hemorrhagic fever, and encephalitis (2). Flaviviruses therefore are responsible for considerable morbidity and mortality and have become a serious public health problem worldwide (3).

It was reported that dengue fever was endemic in more than 100 countries worldwide, threatening the health of more than 2.5 billion people (4). JE had occurred everywhere in China except for Xinjiang, Tibet, and Qinghai, and it was the most harmful mosquito-borne infectious disease with high morbidity and mortality (3). WNV was imported into New York city in 1999 and spread widely in the neighboring states for 5 years (5). In recent years due to global warming, the rapid development of tourism, and continuous changes in the ecological environment, the incidence of mosquito-borne diseases has been on the rise, which brings new challenges to the prevention and control of mosquito-borne diseases (6). Therefore, efficient detection and diagnosis of mosquito-borne viruses is urgently needed to prevent and control mosquito-borne diseases. Currently, conventional flavivirus diagnostic methodologies include virus isolation and culturing methods, serological methods, and nucleic acid detection methods. However, virus culture is relatively laborious and time-consuming, and serological methods lack of specificity and are prone to cross-reactions between viruses (7-8). Compared with traditional virus isolation and serological methods, nucleic acid detection technology has unique advantages in early detection of mosquito-borne viruses.

METHODS

Primer Design

The complete genomic sequences of mosquito-borne flavivirus were downloaded from the GenBank

database and aligned using the Clustal W program (Table 1). A stretch of nucleotides conserved in the strains was identified and the primers were designed using the Primers Express software. The primer sequences and characteristics are shown in Table 2.

Viruses, Viral RNA Extraction, and Reverse Transcription

Viral strains, including JEV, DENV-2, YFV, WNV, Sindbis virus (SINV), and chikungunya virus (CHIKV), were provided by the National Institute for Viral Disease Control and Prevention of China CDC. Viral RNA was extracted using the Rneasy MiNi Kit (QIAGEN, Germany) and used directly complementary DNA (cDNA) synthesis using QuantiNovaTM Reverse Transcription Kit (QIAGEN, Germany) according to the manufacturer's recommendations. Flaviviruses NS5 gene fragments amplified from cDNA employing oligonucleotide primers shown in Table 2. amplicons were purified (TransGen, Beijing, China) and then each cDNA was cloned into the pEASYR-T1 Simple Cloning Vector (TransGen, Beijing, China) and transformed into E. coli Trans1-T1 cells. Plasmid DNAs were purified using the EasyPureR Plasmid MiniPrep Kit (TransGen, Beijing, China) according to the manufacturer's instructions. The DNAs were quantified by a NanoDrop-1000 spectrophotometry (Thermo Fisher Scientific, USA). The copy numbers of the DNA were calculated based on the concentration, and 10-fold serial dilutions of this DNA from 109 to 100 copies per reaction were used as a standard in all heminested reverse transcriptase polymerase chain reactions (hnRT-PCR).

Sensitivity and Specificity of the hnRT-PCR Assay

The analytical sensitivity of the hnRT-PCR assay was determined by quantification with external standards using serially diluted plasmids (10^9 – 10^0 copies/ μ L) containing JEV, DENV-2, WNV, and YFV. For specificity testing, we used cDNA of JEV, DENV-2, YFV, WNV, SINV, and CHIKV. We also used mosquito samples spiked with cDNA of JEV, DENV-2, YFV, and WNV, and the cDNA of mosquito samples was used as control.

Field-caught Mosquitoes

Mosquito samples were collected at different sites in Guizhou Province between July to August 2018. The

collected mosquitoes were frozen and identified quickly and then pooled by species into groups of up 50 individuals. Viral RNAs were extracted and reverse transcribed into cDNA and the hnRT-PCR was performed using designed primers shown in Table 2.

RESULTS

In order to determine the sensitivity of the hnRT-PCR, we examined the lowest detection limits with external standards employing serial dilutions of quantified plasmid DNAs. The lowest detection limits in the first round PCR for JEV, YFV, and WNV were 3×10^{7} , 3×10^{9} , and 3×10^{7} copies/µL. DENV-2 had never been detected. In the second round PCR, there was a substantial increase in sensitivity and the lowest detection limits of JEV, DENV-2, YFV, and WNV were 3×10^4 , 3×10^6 , 3×10^5 , and 3×10^4 copies/µL, respectively (Figure 1). Therefore, through two rounds of amplification, the sensitivity of hnRT-PCR was increased 10,000 times, which greatly improved the sensitivity and made up for the possibility that conventional PCR could not be detected. For specificity testing, we used cDNA of JEV, DENV-2, YFV, WNV, SINV, and CHIKV. We also used mosquito samples spiked with cDNA of JEV, DENV-2, YFV, and WNV (JEV+mosquito cDNA, DENV-2+mosquito cDNA, YFV+mosquito cDNA, and WNV+mosquito cDNA), and cDNA of mosquito samples as control. The results showed that only the target bands appeared in the flavivirus and that no bands appeared in the control, suggesting that the test was 100% specific for the detection of flavivirus (Figure 2).

Overall, a total of 96 pools of mosquitoes belonging to 4 species of 3 genera were used to examine the sensitivity of the hnRT-PCR assay. Among all mosquitoes collected, 34% were Culex tritaeniorhynchus, 2.3% were Culex pipiens pallens, 24.7% were Anopheles sinensis, and 39% were Armigeres subalbatus. JEV was detected in 2 mosquito pools out of 96 from Culex tritaeniorhynchus and Armigeres subalbatus. One was identified as type JEV- I isolated from Culex tritaeniorhynchus, and the other was identified as type JEV-III isolated from Armigeres subalbatus.

DISCUSSION

Flaviviruses are single-stranded RNA viruses with

TABLE 1. Sequence alignment of oligonucleotide XF-F1, XF-F2, and XF-R with 36 flavivirus NS5 gene conserved regions.

Sequence of oligonucleotide ^{*,†}								
Region XF-F1 (8916-8938)		XF-F2 (8964-8985)	XF-R (9153-9178)					
Primer	AACATGATGGGVAARMGWGARAA	AARGGMAGYMGNGCHATHTGGT	GTRTCCCANCCDGCDGTRTCATCNGC					
Nakayama	AACATGATGGGAAAAAGAGAGAA	AAGGGAAGCAGGGCCATTTGGT	GCTGATGATACCGCCGGGTGGGACAC					
SA(A)	G	A A CA- G C T	T T C C G C					
Beijing-1	A AA- A A	A A CA- G C T	T T C T G C					
GP78	A AA- A G	A A CA- G C C	T C C C G C					
HVI	A AA- A G	A A CA- G C T	T T C C G C					
JaGAr01	A AA- A G	A A CA- G C T	T T C C G C					
JaOArS982	A AA- A G	A A CA- G C T	T C C C A C					
K94P05	A AA- A G	A A CA- G C T	C C C C G C					
SA-14	A AA- A G	A A CA- G C T	T T C C G C					
HawO3663	A GA- A G	A A TC- C A A	A C A C A C					
71/02GZ	A GA- A G	A A TC- C A A	A C A C A C					
Nauru Island	A GA- A G	A A TC- C A A	A C A C A C					
16681	A AA- A G	A C CA- A C A	C C C A A C					
New Guinea C	A AA- A G	A C CA- A C A	C C C A A C					
PUO-218	A AA- A G	A C CA- A C A	C C C A A C					
H87	C GA- A G	A C TA- G T A	T C A C T C					
80-2	C GA- A G	A C TA- G T A	T C A C T C					
P4	A AC- T G	G A CC- A A C	T C A A C C					
P75-215	A AC- T G	A A CC- G A T	T C A A T C					
11070	A AC- T G	G A CC- A A C	T C A A C C					
serum	A GA- A G	G A CA- A C T	T C A T C C					
FtC-3699	A GA- A G	G A CA- A C T	T C A T C C					
385-99	A GA- A G	G A CA- A C T	T C A T C C					
ARC13-12	A GA- A G	G A CA- A C T	T C A T C C					
Chin-01	G GA- A A	G A CA- A C A	T C A T C C					
33-G8	A GA- A G	A C CA- A C C	T T C A C C					
HNY1999	A GA- A G	G A CA- A C T	T C A T C C					
17D vaccine	G AA- A G	G A CC- T C A	G C C T A C					
French viscerotropic	G AA- A G	G A CC- T C A	G C C T A C					
17D vaccine	G AA- A G	G A CC- T C A	G C C T A C					
17DD	G AA- A G	G A CC- T C A	G C C T A C					
17D-213	G AA- A G	G A CC- T C A	G C C T A C					
85-82H	G AA- A G	A A CC- T C C	G T C T G C					
French viscerotropic	G AA- A G	G A CC- T C A	G C C T A C					
Trinidad 79A	G AA- A G	G A CC- T C A	G C C T A C					
French viscerotropic	G AA- A G		G C C T A C					

^{*}N=A+G+C+T, V=G+A+C, R=A+G, M=A+C, W=A+T, Y=C+T, H=A+T+C, D=G+A+T.

[†] DNA sequence were obtained from the GenBank databases. Accession numbers are as follows: Nakayama, EF571853.1; SA(A), D90195.1; Beijing-1, L48961.1; GP78, AF075723.1; HVI, AF098735.1; JaGAr01, AF069076.1; JaOArS982, M18370.1; K94P05, AF045551.2; SA-14, M55506.1; HawO3663, DQ672564.1; 71/02GZ, EF025110.1; Nauru Island, U88535.1; 16681, U87411.1; New Guinea C, AF038403.1; PUO-218, AF038402.1; H87, M93130.1; 80-2, AF317645.1; P4, AY648301.1; P75-215, EF457906.1; 11070, M14931.2; serum, AY646354.1; FtC-3699, KR868734.1; 385-99, AY842931.3; ARC13-12, KM012188.1; Chin-01, AY490240.2; 33-G8, M12294.2; HNY1999, AF202541.1; 17D vaccine, X03700.1; French viscerotropic, U21056.1; 17D vaccine, NC_002031.1; 17DD, U17066.1; 17D-213, U17067.1; 85-82H, U54798.1; French viscerotropic, U21055.1; Trinidad 79A, AF094612.1; French viscerotropic virus, U21056.1.

TABLE 2. Nucleotide sequences and positions of primers used in the heminested RT-PCR (hnRT-PCR) assay for detection of mosquito-borne flavivirus.

	Primer	Sequence (5'-3')*	Nucleotide position	Annealing temperature (℃)	Length (bp)
1 st round PCR	XF-F1	AACATGATGGGVAARMGWGARAA	8916-8938	52	263
	XF-R	GTRTCCCANCCDGCDGTRTCATCNGC	9153-9178		
2 nd round PCR	XF-F2	AARGGMAGYMGNGCHATHTGGT	8964-8985	54	215
	XF-R	GTRTCCCANCCDGCDGTRTCATCNGC	9153-9178		

*N=A+G+C+T, V=G+A+C, R=A+G, M=A+C, W=A+T, Y=C+T, H=A+T+C, D=G+A+T.

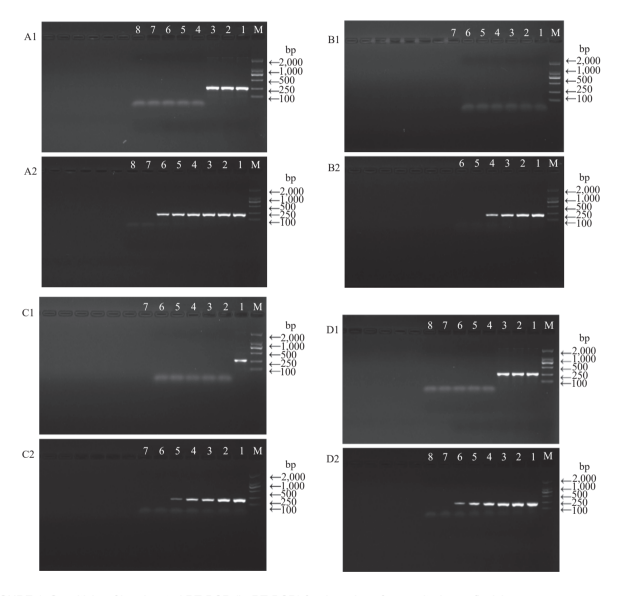


FIGURE 1. Sensitivity of heminested RT-PCR (hnRT-PCR) for detection of mosquito-borne flavivirus. A, B, C, and D show the lowest limit for JEV, DENV-2, YFV, and WNV, respectively; 1 and 2 denote the lowest limit for the first and second round PCR, respectively. The numbering 1–8 refers to 10⁹–10³ copies/μL and negative control, respectively. M: DL2000 DNA markerr.

various species and are widely distributed worldwide. Most flaviviruses are pathogens causing natural infectious diseases, which can lead to fever, hemorrhage, and encephalitis with a high mortality rate (9). In this study, four flaviviruses were screened based on the epidemic situation, pathogenicity, and risk of disease, including JEV, DENV-2, WNV, and YFV. Among them, JEV and DENV-2 are the

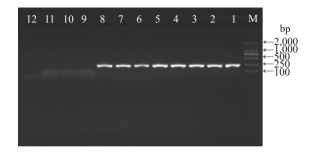


FIGURE 2. Specificity of heminested RT-PCR (hnRT-PCR) for detection of mosquito-borne flavivirus. The numbering 1–12 refers to JEV, DENV-2, YFV, WNV, JEV + mosquito cDNA, DENV-2 + mosquito cDNA, YFV + mosquito cDNA, WNV + mosquito cDNA, mosquito cDNA, SINV, CHIKV, and negative control, respectively. M: DL2000 DNA marker.

common mosquito-borne viruses, and associated outbreaks occur almost every year in China. Therefore, the rapid and accurate high-throughput detection method for flavivirus is of great significance for the prevention and control of mosquito-borne diseases.

The traditional methods for detecting mosquito-borne viruses are isolation and culturing method, but virus culturing is relatively time-consuming and cannot be used for early diagnosis of disease. Serological detection is a classical virus detection method, which is mainly used to detect specific antibodies or antigens in specimens, and it is indispensable in the diagnosis and identification of virus infection. But serological methods lack of specificity and are prone to cross-reactions between viruses. Traditional species-specific RT-PCR and real-time RT-PCR based on Taq-Man Probe are rapid and sensitive, but only detect single pathogens.

Wu et al. (10) established a method for screening samples by multiplex real-time RT-PCR and the lower limit of detection of this method was 11 copies/µL. However, compared with singleplex real-time PCR, multiplex real-time PCR requires multiple pairs of primers and probes in a single reaction tube and, therefore, will have to account for competition between reactants resulting in significantly increased difficulty of primer design. Gao et al. (11) used the established RT-PCR method to detect JEV, and the lower limit of detection was 10 pg/µl but could only detect a single flavivirus.

In this study, a new hnRT-PCR method for the simultaneous detection of various flaviviruses was established. The performance of the new hnRT-PCR method was evaluated including specificity and sensitivity. In the specificity testing, only the target

bands appeared in the flavivirus, and no bands appeared in control, which indicated high specificity in detecting mosquito-borne flavivirus. In the sensitivity testing, the lowest detection limits of JEV, DENV-2, YFV, and WNV were 3×10^4 , 3×10^6 , 3×10^5 , and 3×10⁴ copies/μL, respectively. When testing fieldcaught mosquito samples, 2 strains of JEV were detected by the method established in this study. Other flaviviruses had not been detected, which may be related to the species and quantity of mosquitoes collected, and the epidemic situation of the virus. However, considering that the variety and high genetic diversity of flavivirus, whether the hnRT-PCR method established in this study had a good detection effect for all flavivirus viruses needs to be further validated by other flavivirus viruses. In summary, the hnRT-PCR established in this study possessed both high specificity and sensitivity and could simultaneously be used for detection of multiple flaviviruses. This can be effectively used to detect the pathogen of flavivirus and provide technical support for early and rapid diagnosis of mosquito-borne flavivirus.

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REFERENCES

- Weaver SC, Reisen WK. Present and future arboviral threats. Antiviral Res 2010;85(2):328 – 45. http://dx.doi.org/10.1016/j.antiviral.2009. 10.008.
- Erlanger TE, Weiss S, Keiser J, Utzinger J, Wiedenmayer K. Past, present, and future of Japanese encephalitis. Emerg Infect Dis 2009; 15(1):1 – 7. http://dx.doi.org/10.3201/eid1501.080311.
- Gould E, Pettersson J, Higgs S, Charrel R, de Lamballerie X. Emerging arboviruses: why today? One Health 2017;4:1 – 13. http://dx.doi.org/ 10.1016/j.onehlt.2017.06.001.
- Tang KF, Ooi EE. Diagnosis of dengue: an update. Expert Rev Anti-Infect Ther 2012;10(8):895 – 907. http://dx.doi.org/10.1586/eri.12.76.
- 5. Asnis DS, Conetta R, Teixeira AA, Waldman G, Sampson BA. The west Nile virus outbreak of 1999 in New York: the flushing hospital

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- experience. Clin Infect Dis 2000;30(3):413 8. http://dx.doi.org/10.1086/313737.
- Franca RFO, Neves MHL, Ayres CFJ, Melo-Neto OP, Filho SPB. First international workshop on Zika virus held by Oswaldo Cruz foundation FIOCRUZ in Northeast Brazil March 2016-a meeting report. PLoS Negl Trop Dis 2016;10(6):e0004760. http://dx.doi.org/10.1371/ journal.pntd.0004760.
- 7. Jarman RG, Nisalak A, Anderson KB, Klungthong C, Thaisomboonsuk B, Kaneechit W, et al. Factors influencing dengue virus isolation by C6/36 cell culture and mosquito inoculation of nested PCR-positive clinical samples. Am J Trop Med Hyg 2011;84(2):218 23. http://dx.doi.org/10.4269/ajtmh.2011.09-0798.
- 8. Lu Z, Lu XJ, Fu SH, Zhang S, Li ZX, Yao XH, et al. Tahyna virus and human infection, China. Emerg Infect Dis 2009;15(2):306 9. http://dx.doi.org/10.3201/eid1502.080722.
- Yang CF, Chen CF, Su CL, Teng HJ, Lu LC, Lin C, et al. Screening of mosquitoes using SYBR Green I-based real-time RT-PCR with groupspecific primers for detection of Flaviviruses and Alphaviruses in Taiwan. J Virol Methods 2010;168(1 – 2):147 – 51. http://dx.doi.org/ 10.1016/j.jviromet.2010.05.006.
- 10. Wu WH, Wang J, Yu N, Yan JY, Zhuo ZH, Chen MY, et al. Development of multiplex real-time reverse-transcriptase polymerase chain reaction assay for simultaneous detection of Zika, dengue, Yellow fever, and chikungunya viruses in a single tube. J Med Virol 2018;90(11):1681 – 86. http://dx.doi.org/10.1002/jmv.25253.
- Gao ZQ, Yue BF, He ZM. Development and application of RT-PCR assay of *Japanese encephalitis* virus. Chin J Pharm Anal 2010;30(10): 1928-31. http://d.wanfangdata.com.cn/periodical/ChlQZXJpb2RpY2 FsQ0hJTmV3UzIwMjAxMjI4Eg95d2Z4enoyMDEwMTAwMjgaCDF 6Y2tsem9t. (In Chinese).