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Early Detection of Dengue Virus by Use of Reverse Transcription-Recombinase Polymerase Amplification

Boon-Teong Teoh, Sing-Sin Sam, Kim-Kee Tan, Mohammed Bashar Danlami, Meng-Hooi Shu, Jefree Johari, Poh-Sim Hooi, David Brooks, Olaf Piepenburg, Oliver Nentwich, Annelies Wilder-Smith, Leticia Franco, Antonio Tenorio, Saizal Abubakar

A method for the rapid diagnosis of early dengue virus (DENV) infection is highly needed. Here, a prototype reverse transcription-recombinase polymerase amplification (RT-RPA) assay was developed. The assay detected DENV RNA in <20 min without the need for thermocycling amplification. The assay enabled the detection of as few as 10 copies of DENV RNA. The designed RT-RPA primers and exo probe detected the DENV genome of at least 12 genotypes of DENV circulating globally without cross-reacting with other arboviruses. We assessed the diagnostic performance of the RT-RPA assay for the detection of DENV RNA in 203 serum samples of patients with clinically suspected dengue. The sera were simultaneously tested for DENV using a reverse transcription-loop-mediated isothermal amplification (RT-LAMP) assay, quantitative RT-PCR (qRT-PCR), and IgM- and IgG-capture enzyme-linked immunosorbent assays (ELISA). Acute DENV infection was confirmed in 130 samples and 61 of the samples (46.9%) were classified as viremic with qRT-PCR. The RT-RPA assay showed good concordance (κ of 0.723) with the RT-LAMP and qRT-PCR assays in detecting the dengue viremic samples. When used in combination with ELISA, both the RT-RPA and RT-LAMP assays increased the detection of acute DENV infection to 95.7% (%45/47) in samples obtained within 5 days of illness. The results from the study suggest that the RT-RPA assay is the most rapid molecular diagnostic tool available for the detection of DENV. Hence, it is possible to use the RT-RPA assay in a laboratory to complement routine serology testing for dengue.

Dengue is one of the most prevalent mosquito-borne viral diseases in the tropics and subtropics. It is estimated that at least 3.6 billion people are at risk of contracting the infection (1). The spectrum of the illness of dengue ranges from mild dengue fever (DF) to severe and fatal dengue hemorrhagic fever (DHF) and dengue shock syndrome (DSS) (2). Dengue viruses (DENV) are the causative agents of dengue (3). There are at least four known antigenically distinct DENV serotypes, DENV-1, DENV-2, DENV-3, and DENV-4 (4), and each serotype contains several phylogenetically distinct genotypes (5). The virus is transmitted following mosquito bites of viremic febrile patients; the viremia phase usually corresponds to the first 3 days of illness (6). Therefore, the early detection of viremic individuals is important not only for patient management but also for early and immediate implementation of appropriate vector-control measures (7).

Molecular techniques to detect the presence of DENV genomic RNA sequences are gradually being accepted as routine procedures for the early detection of DENV infection. The reverse transcription-PCR (RT-PCR) and real-time quantitative RT-PCR (qRT-PCR) are the two most commonly used methods (8–12). However, the requirement for costly nucleic acid amplification instruments and the high level of skill needed for these methods have limited their use to well-equipped laboratories. The introduction of the isothermal nucleic acid amplification method, which obviates the need for any major instrumentation, could change this situation, hence allowing greater access to the nucleic acid amplification method and its wider application in regions where dengue is endemic and resources may be limited (13).

We have previously demonstrated the use of reverse transcription-loop-mediated isothermal amplification (RT-LAMP) for the detection of DENV RNA in a clinical setting (14). More recently, the recombinase polymerase amplification (RPA) assay has emerged as a novel alternative isothermal amplification method for the detection of nucleic acid (15). The RPA assay was reported to take <20 min to perform and requires a constant temperature of only 37°C to 42°C. Both the RPA and RT-RPA assays have been used for the detection of various other infectious agents (15–36), but they have not been implemented for the detection of DENV. In the present study, we describe the development and application of the RT-RPA assay for the detection of DENV infection in a diagnostic laboratory setting using freshly obtained samples from dengue-suspected patients.
**MATERIALS AND METHODS**

**Dengue viruses.** Twenty-two reference DENV strains were used in this study; 11 isolates from Malaysia and 11 isolates from Spain. The Malaysian DENV strains included four genotypes of DENV-1 (genotypes I, II, and III and sylvatic) (37, 38), two genotypes of DENV-2 (Asian I and Cosmopolitan) (39), three genotypes of DENV-3 (genotypes I, II, and III) (40), and two subgenotypes of DENV-4 (subgenotypes Ia and Iib) (10). All of the Malaysian dengue viruses were archived in the Department of Medical Microbiology repository; the 11 dengue viruses obtained from Spain were from the Arbovirus and Imported Viral Diseases, National Centre for Microbiology, Institute of Health Carlos III. The DENV strains from Spain included three isolates of DENV-1 (all genotype III or American/African) (41), three isolates of DENV-2 (Cosmopolitan, American/Asian, and African sylvatic) (42), three isolates of DENV-3 (one genotype I and two genotype III), and two isolates of DENV-4 (all genotype II). These viruses were from collections from Brazil, Paraguay, Venezuela, Thailand, Pakistan, India, South America, Senegal, and the Philippines (see Table S1 in the supplemental material).

**Clinical samples.** The study was approved by the University of Malaya Medical Center (UMMC) Medical Ethics Committee (Ethics Committee/Institutional Review Board [IRB] reference number 908.11). A total of 203 serum samples from 203 patients clinically suspected to be infected with DENV at the UMMC during the period from June to August 2013 were obtained for this study. The majority of the patients from this cohort were adults, and the median age of the patients was 29 years (range, 3 to 79 years). The serum samples were divided in the laboratory for simultaneous testing by the serology and nucleic acid amplification assays. A retrospective study was performed to review the patients’ medical records for the date of illness onset.

**RNA extraction.** The total RNA was extracted from 140 µl of infected culture supernatant or patient serum samples using the QiAamp viral RNA minikit (Qiagen, Germany). All of the RNA extractions were performed according to the manufacturer’s protocol. The RNA was eluted in 60 µl of nuclease-free water and stored at −80°C until needed.

**Design of DENV-specific RT-RPA assay primers and exo probe.** The DENV-specific primers and exo probe used for the RT-RPA assay were designed from the highly conserved 3′-untranslated region (UTR) consensuses of the genomes of all four DENV serotypes according to the criteria described previously (43). Each 3′-UTR consensus was derived from all of the genotypes of each DENV serotype (see Fig. S1 in the supplemental material). The coverage of the RT-RPA primers and exo probe was validated by evaluating the assay using viral RNA extracted from the different reference DENV strains, as described above.

**RT-RPA assay.** A DENV-specific TwistAmp RT exo lyophilized kit was supplied as strips of eight reactions in vacuum-sealed pouches (TwistDx, Ltd., Cambridge, United Kingdom). The RT-RPA was performed in a final reaction volume of 50 µl containing a lyophilized pellet dissolved with 45 µl of TwistAmp RT exo rehydration buffer and 5 µl of the extracted RNA template. The lyophilized RPA pellet contained an optimized blend of enzymes and primers and a fluorescein probe. The enzyme concentrations were based on standard commercial RPA pellets for exonuclease detection (TwistAmp Exo) (15) containing Moloney murine leukemia virus reverse transcriptase for combined RT-RPA (18). All of the enzyme levels were optimized for performance with dengue-specific oligonucleotides. Each reconstituted reaction mixture contained 0.63 mM DenMPF19, 0.84 mM DenMPR11, 0.63 mM DenMPR32, 0.12 mM DenMPEexoPFAM, and 5 U RNase inhibitor (Fermentas, USA). The customized rehydration buffer contained magnesium acetate [Mg(OAc)2], potassium acetate [KOA], Tris-acetate (pH 8.3), and polyethylene glycol (PEG) (35,000) and was engineered for a 5-µl input volume per 50-µl reaction.

A positive control using 1,000 copies (determined by qRT-PCR) of DENV RNA extracted from the culture supernatant and a negative control (nuclease-free water) were included in each run. The RT-RPA reaction mixtures were incubated at 40°C for 20 min with a brief mixing of the reaction mixtures at 230 s after the start of the incubation. The 6-carboxyfluorescein (FAM) fluorescence of RT-RPA was generated via the endonuclease-mediated cleavage of the TwistAmp exo probe at the tetrahydrofuran abasic site mimic. The fluorescence signal was measured using a Twista real-time fluorometer (TwistDx, Ltd.). In this study, the Twista real-time fluorometer was obtained at a cost almost 4 and 6 times lower than the potential market costs of devices used for RT-LAMP and qRT-PCR, respectively. Twista Studio software, version 2.06.06 (TwistDx, Ltd.) was used to interpret the fluorescence data with a combined threshold and signal slope analysis. The threshold limit was the mean fluorescence intensity (mV) plus 3 standard deviations of that mean. The slope limit was set at 70 mV/min. The RT-RPA reaction was deemed positive when the measured data were higher than the threshold and slope limit within the defined measurement time (adaptable).

**Detection limit and cross-reactivity of RT-RPA assay.** The detection limit of the RT-RPA assay for all of the four DENV serotypes was assessed using a panel of serially diluted viral RNAs (1,000, 100, 50, and 10 copy numbers) extracted from the culture supernatant. The RT-RPA detection limit test was repeated eight times. The viral RNA was quantified using the geosig real-time qRT-PCR DENV detection kit ( PrimerDesign, Ltd., United Kingdom) as previously described (14, 44). The cross-reactivity of the DENV RT-RPA primers and exo probe was evaluated using the Japanese encephalitis virus (JEV), Chikungunya virus (CHIKV), and Sindbis virus (SINV), all of which were obtained from the Department of Medical Microbiology repository (45, 46). The CHIKV and SINV were isolated from human and mosquito, respectively. No record was available for the origin of the JEV.

**Evaluation of RT-RPA assay.** The single-tube RT-RPA assay for the detection of DENV RNA was clinically assessed in 203 serum samples freshly obtained from dengue-suspected patients. The samples were also screened for the presence of DENV RNA using the RT-LAMP and qRT-PCR assays as previously described (14). The qRT-PCR assay was used as a reference for the detection of DENV RNA in the samples. In addition, the samples were simultaneously screened for the presence of anti-DENV IgM and IgG using dengue IgM- and IgG-capture enzyme-linked immunosorbent assay (ELISA) kits (Standard Diagnostics Inc., Republic of Korea), respectively. The test results for RT-RPA, RT-LAMP, qRT-PCR, and ELISA were analyzed and compared. Acute DENV infection was confirmed by the positive detection of DENV RNA with qRT-PCR or the presence of anti-DENV IgM detected with an ELISA. The presence of anti-DENV IgG concurrent with the positive detection of the DENV RNA genome indicated secondary acute DENV infection. The serum sample that tested positive for anti-DENV IgG only was identified as a reflection of a previous DENV infection. The samples that tested positive by the RT-RPA or RT-LAMP assays but negative by qRT-PCR were considered false-negative tests.

**Statistical analysis.** All of the statistical analyses were performed using IBM SPSS Statistics, version 21 (IBM Corporation, NY, USA). A probit analysis was performed to calculate the detection limit of the RT-RPA assay at a 95% probability level. The degrees of agreement among the RT-RPA, RT-LAMP, and qRT-PCR test results were measured with the kappa value (κ). A chi-square test (McNemar’s exact test, two-tailed) was performed to compare the sensitivities of all of the molecular and serological methods used in this study. In a P value of <0.001 was used to suggest significant results. The diagnostic performances of the RT-RPA and RT-LAMP assays compared with that the qRT-PCR assay were calculated using the web-based Centre for Evidence-Based Medicine Statistics Calculator (http://ktclearinghouse.ca/cebms/toolbox/statscalc).

**RESULTS**

Design of DENV-specific RT-RPA assay primers and exo probe. Three RT-RPA primers and one exo probe were designed to allow the simultaneous detection of all four DENV serotypes in a single-tube reaction (Table 1; see also Fig. S1 in the supplemental material). The developed RT-RPA primers and exo probe detected all of
the 22 reference DENV strains obtained in Malaysia (see Fig. S2A to D in the supplemental material) and those from the collection available at the Arbovirus and Imported Viral Diseases, National Centre for Microbiology, Institute of Health Carlos III, Spain (see Fig. S2E to G in the supplemental material).

Detection limit and cross-reactivity of the RT-RPA assay. The detection limits of the RT-RPA assay for all of the four DENV serotypes were determined using a panel of serially diluted viral RNAs extracted from infected cell culture supernatants with known copy numbers of 1,000, 100, 50, and 10 (quantified by qRT-PCR). The detection limit of the RT-RPA assay at the 95% probability level was 11 DENV genome copies (probit analysis, qRT-PCR). The detection limit of the RT-RPA assay at the 95% confidence interval was 10 DENV genome copies (Fig. 3). Using the present parameters, 3 DENV genome copies/ml in viremic patients were beyond the detection limit of qRT-PCR (10 RNA copies, equivalent to 2.93 log_{10} RNA copies/ml of serum).

Evaluation of the RT-RPA assay. The RT-RPA assay for the detection of DENV RNA was evaluated by performance of the assay on 203 freshly obtained serum samples from patients with clinically suspected dengue. The performance of the RT-RPA assay was compared to that of the RT-LAMP and qRT-PCR assays. Out of the 203 samples, acute DENV infection was confirmed in 130 samples (dengue IgM- or qRT-PCR-positive samples). Eight dengue IgG- and qRT-PCR-positive samples tested negative by dengue IgM ELISA.

TABLE 2  Summary of dengue detection in serum samples (n = 203) from patients with clinically suspected dengue at UMMC using RT-RPA, RT-LAMP, and qRT-PCR assays and IgM and IgG ELISA

<table>
<thead>
<tr>
<th>Assay and results</th>
<th>Positive (n = 91)</th>
<th>Negative (n = 112)</th>
<th>Positive (n = 96)$^a$</th>
<th>Negative (n = 107)</th>
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<tbody>
<tr>
<td>RT-RPA</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Positive (50$^b$)</td>
<td>17 (34.0)</td>
<td>33 (66.0)</td>
<td>16 (32.0)</td>
<td>34 (68.0)</td>
</tr>
<tr>
<td>Negative (153)</td>
<td>74 (48.4)</td>
<td>79 (51.6)</td>
<td>80 (52.3)</td>
<td>73 (47.7)</td>
</tr>
<tr>
<td>RT-LAMP</td>
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<td></td>
<td></td>
</tr>
<tr>
<td>Positive (51$^c$)</td>
<td>13 (25.5)</td>
<td>38 (74.5)</td>
<td>14 (27.5)</td>
<td>37 (72.5)</td>
</tr>
<tr>
<td>Negative (152)</td>
<td>78 (51.3)</td>
<td>74 (48.7)</td>
<td>82 (53.9)</td>
<td>70 (46.1)</td>
</tr>
<tr>
<td>qRT-PCR</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Positive (61$^d$)</td>
<td>22 (36.1)$^e$</td>
<td>39 (63.9)$^d$</td>
<td>22 (36.1)$^e$</td>
<td>39 (63.9)</td>
</tr>
<tr>
<td>Negative (142)</td>
<td>69 (48.6)$^f$</td>
<td>73 (51.4)</td>
<td>74 (52.1)</td>
<td>68 (47.9)</td>
</tr>
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$^a$ Asterisks indicate a phosphothioate link in the backbone to inhibit exonuclease degradation; dT-FAM, thymidine nucleotide carrying fluorescein; dSpacer, tetrahydrofuran basic site mimic; dT-BHQ1, thymidine nucleotide carrying black hole quencher; 3′C3-Spacer, C3 spacer at the 3′ end to block elongation.

$^b$ Sixteen IgG-positive dengue samples tested negative by qRT-PCR and dengue IgM ELISA.

$^c$ Three RT-RPA-positive samples tested negative by qRT-PCR.

$^d$ Five RT-LAMP-positive samples tested negative by qRT-PCR.

$^e$ Acute DENV infection was confirmed in 130 samples (dengue IgM- or qRT-PCR-positive samples).

$^f$ Eight dengue IgG- and qRT-PCR-positive samples tested negative by dengue IgM ELISA.

The RT-RPA assay detected the DENV genome in 63.8% (30/47) of the acute dengue samples obtained within 5 days of illness, compared with 72.3% (34/47) by the qRT-PCR assay and 61.7%...
(29/47) by the RT-LAMP assay (Fig. 4). When used in combination with the dengue IgM and IgG ELISA, both the RT-RPA and RT-LAMP assays showed significant increases ($P < 0.001$) in positive detection to 95.7% (45/47) and 97.9% (46/47), respectively, in comparison with use of the dengue IgM and IgG ELISA alone, which had a positive detection of only 55.3% (26/47).

**DISCUSSION**

In the present study, a rapid single-tube RT-RPA assay was developed for real-time isothermal detection of DENV genomic RNA in clinical specimens. We demonstrated that the RT-RPA assay detected a panel of 22 globally distributed DENV strains. There was no cross-reactivity of the RT-RPA assay with a number of other common arboviruses. The RT-RPA assay was as sensitive as the RT-LAMP and qRT-PCR assays for DENV detection in viremic patient serum samples.

This is the first study describing the application of the RT-RPA assay for the detection of DENV infection in clinically diagnosed patient samples. The detection limit of the RT-RPA assay (50 copies of DENV RNA) was similar to that of RT-RPA assays previously reported for other RNA viruses; the detection limits ranged from 10 to 100 copies of viral RNA (18, 21, 23, 24, 26, 34). The performance of the RT-RPA assay was superior to that of the RT-LAMP assay we previously developed for the detection of DENV (14), which had a detection limit of 100 DENV RNA copies. It is worth mentioning that the sensitivity of the RT-LAMP assay for DENV detection varied when it was evaluated on different patient cohorts. In this study, the sensitivity of the RT-LAMP assay was only 75.4%, which was lower than that of the previously assessed RT-LAMP assay (92.5%). The variation could be due to random
effects in the target amplification that occurred when the viral titer of the samples was below the detection limit of the assay. As previously mentioned, 100% reproducibility was only achieved in samples with at least 100 copies of viral RNA (14).

In the current study, the RT-RPA assay was superior to the RT-LAMP assay for the detection of DENV in patient serum samples. The qRT-PCR assay, however, was still the most sensitive method, especially for detecting a low viral RNA titer (<50 copies). The possibility of false-positive detections, especially in those samples with low copy numbers, was unlikely, as the qRT-PCR used in this study used the TaqMan probe to increase the specificity of the assay (47). Most of the viremic samples that were misdetected by the RT-RPA and/or RT-LAMP assays had viral loads of <50 RNA copies. We showed here that the low viral load was accompanied with a rise in the antibody titers. More than 80% of the misdetected viremic samples were positive when detected by IgM/IgG ELISA, and these samples were mainly among those obtained beyond day 4 of illness (data not shown). When used in combination with an ELISA, the RT-RPA and RT-LAMP assays still performed as well as qRT-PCR for increasing the diagnostic coverage of febrile dengue patients. However, the false-positive detection of the RT-RPA and RT-LAMP assays could reflect the false-negative detection of qRT-PCR, possibly due to mismatches of qRT-PCR primers and probes for certain DENV strains. This possibility needs to be verified using virus isolation.

The application of the molecular detection of DENV genomes...
primarily depends on the time of sample collection following the onset of disease. In this study, we demonstrated that the qRT-PCR, RT-RPA, and RT-LAMP assays were useful for the diagnosis of dengue in febrile patients during the first 5 days of illness. The use of nucleic acid amplification assays, however, is not promoted for samples obtained after day 5 of illness, which comprised 58% of the total samples used in our study. At this stage of the infection, there is already a significant rise in the antibody levels. Therefore, knowing the exact day of illness is desirable for choosing the appropriate method for dengue diagnosis to be cost-effective, especially in low-resource countries where dengue is endemic.

Although the nucleic acid-based diagnostics for DENV infection are more widely available in referral hospitals than in clinics, the majority of the febrile patients in countries where dengue is endemic prefer to seek early medical care from community clinics at their own convenience (48). Patients are usually only referred to hospitals for the confirmation of DENV infection in severe cases. It is shown in our study that >80% of the dengue patients presenting to the hospital were already beyond day 4 of illness. Most of the dengue-suspected patients from the community near the UMMC therefore came to the hospital long after the onset of fever (49). Many of these patients were already in the convalescent phase of dengue. Alternatively, the low percentage of early-phase samples obtained could be a reflection of the unavailability of molecular diagnostics for dengue at the hospital, and, therefore, blood samples were not taken from patients exhibiting fever for <4 days. Hence, the implementation of nucleic acid amplification assays such as RT-RPA for routine dengue diagnosis in hospitals and clinics could improve the early detection of viremic dengue patients. The early detection of DENV infection would in turn improve the recognition of severe dengue warning signs, such as impending intravascular leakages (49).

During the viremic phase of DENV infection, patients harbor tremendous amounts of viruses in their blood. The virus can be transmitted following mosquito bites. It was reported that the 50% mosquito infectious dose (MID$_{50}$) for DENV ranged from 6.27 to 7.52 log$_{10}$ RNA copies/ml of human plasma (50). In our study, 50 and 100 DENV RNA copies were equivalent to 3.63 and 3.93 log$_{10}$ RNA copies/ml of serum, respectively. The RT-RPA (detection limit of 50 RNA copies) and RT-LAMP (detection limit of 100 RNA copies) assays (14) can therefore still be useful for detecting the potential viremic patients who are contagious. This information is crucial for the immediate implementation of preventive measures and precautionary behavior to prevent mosquito bites. In our sample cohort, high viral loads (up to 5.53 log$_{10}$ RNA copies/ml) were found in some patients up to day 9 of illness, which was comparable to earlier findings (51, 52). This also implies that it is possible for DENV to be transmitted by patients even after the defervescence phase. However, the DENV RNA detected in these convalescent patients did not necessarily reflect the presence of infectious viruses; it may have been just the residual genome nucleic acid.

Although it is still in an early stage of development, the RT-RPA assay reported here has several advantages compared to the RT-LAMP assay, including (i) faster assay run time (<20 min for RT-RPA instead of ≥60 min for RT-LAMP), (ii) relative ease of performance (the freeze-dried RT-RPA ready-to-use reaction mixture was designed to simplify the operator workload and minimize pipetting errors), and (iii) lower energy consumption (40°C for RT-RPA versus 63°C for RT-LAMP). These advantages make the RT-RPA assay an attractive diagnostic test for the routine detection of DENV RNA, especially in resource-limited health care centers, as it can be easily performed even by less-skilled staff members. In addition, the RT-RPA assay is more amenable to scaling down for point-of-care applications than the RT-LAMP assay because it requires less energy input (53, 54).

It is possible that the performance of the DENV RPA assay could be improved from that observed in this study. The presence of divergent cations such as Mg$^{2+}$ and how they are added to the reaction could be optimized to limit the opportunity for spurious amplification, which would improve the assay performance and perhaps eliminate the false-negative results, particularly in samples with low titers. However, this is beyond the scope of this investigation, as in its present form the assay was proven to be as good as or comparable in sensitivity to other methodologies. The encouraging findings from this study, however, justify further improvement of the RT-RPA assay for eventual clinical evaluations and applications.

In summary, the RT-RPA assay developed in our study is specific and simple to perform. The RT-RPA assay has a rapid run time (<20 min) and can detect diverse DENV strains at viral titers up to as few as 10 RNA copies. The RT-RPA assay can greatly enhance the diagnostic coverage of suspected dengue cases, complementing serological assays and improving the diagnosis of dengue in rural clinics and health care facilities in regions where dengue is endemic and resources are limited.

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David Brooks, Olaf Piepenburg, and Oliver Nentwich are employees of TwistDx, Ltd., the owner and producer of the RPA technology.

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