Optimization Of Real-Time Reverse Transcriptase Polymerase Chain Reaction For Detection Of Dengue Virus

Kaunara Ally Azizi (kaunara_azizi@yahoo.com)
TFNC https://orcid.org/0000-0003-2346-9445
Arnold J Ndaro
Kilimanjaro Christian Medical Centre
Athanasia Maro
Kilimanjaro Clinical Research Institute
Adonira T Saro
Kilimanjaro Christian Medical Centre
Reginald Kavishe
Kilimanjaro Christian Medical University College

Research note

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Abstract

**Objective** Rapid and accurate laboratory confirmatory is very essential for control measures of dengue virus infections. However, many cases of dengue virus infections in most of the hospitals remain undiagnosed due to presence of other febrile illnesses with overlapping symptoms and lack of specificity in most of laboratory diagnostic methods. This study was set to optimize conditions for real time reverse transcriptase polymerase chain reaction (RT-PCR) for detection of dengue virus by using rapid and simple nucleic acid extraction method.

**Results** The real time RT-PCR technique was successfully optimized using simple and rapid method for purification of nucleic acid, 'boom method'. The technique works better when performed in a two-step procedure and can works well with all range of real time PCR machines. The optimized real time RT-PCR used in the present study is a valuable and reliable technique for routine diagnosis of dengue. Further investigation on the cost effectiveness in adopting this technique for routine screening and monitoring of the dengue infection should be done.

**Introduction**

Dengue virus (DENV) is arthropod-borne virus that is prevalent in tropical and sub-tropical regions. Main vectors for the virus are mosquitoes of the *Aedes* genus, including *Aedes egypti* and *Aedes albopictus* [1, 2]. DENV belong to the family *Flaviviridae*, genus *Flavivirus*. The virus has a positive-sense, enveloped, single-stranded RNA genome of approximately 11 kb in length which have been categorised into four serotypes, DENV 1–4. DENV 2 reported to cause the most epidemics followed by DENV 1[3, 4, 5].

In recent year's DENV have caught worldwide attention because of increase in the frequency of major epidemics [1]. Globally, 50–100 million DENV infections occur annually, accounting for 20,000–70,000 deaths per year. An estimated 2.5 billion people are at risk of infection. Dengue hemorrhagic fever (DHF) and dengue shock syndrome (DSS) account for about 250,000–500,000 cases each year. Children under 5 years of age are mainly affected. In hyperendemic Asian countries, 22–292 per 1,000 children are infected each year [6, 7, 8]. Dengue epidemics have been reported in African countries since 19th century and all four DENV serotypes have been isolated [3, 9]. The recent outbreak of DENV in Sub-Saharan Africa ascertain that febrile illnesses are now becoming the next threat to the population living in malaria endemic areas [10]. Early 2013, outbreak of dengue was reported in Somalia and Kenya where majority of cases were found in the Indian Ocean Coastal town of Mombasa. Most recently, outbreak was reported in Dar es Salaam, Tanzania. In 2014 dengue outbreak, a total of 2,129 suspected case and 1,018 confirmed cases were reported, where as in 2019 outbreak about 3,000 suspected cases with 71.4% confirmed cases and 2 deaths were reported [11, 12].

Many cases of DENV infections in most of the hospitals in Sub-Saharan Africa remain undiagnosed due to lack of accurate diagnostic technique. Every so often febrile illness are treated as presumptive malaria without proper laboratory diagnosis [3, 13, 14]. The clinical manifestation of DENV infections may be
confused with those caused by chikungunya fever and other febrile illness, but unlike chikungunya and other febrile illness, DENV infections is associated with DHF and DSS [15].

Virus culture is often regarded as gold standard method for laboratory diagnosis of DENV but is time consuming and must be performed under biosafety level 3 conditions. Viral culture is rarely done in routine clinical diagnosis as these facilities are not widely available [1, 16]. Serological diagnosis based on capture IgM and IgG ELISA are reliable can only detect IgM around 5 days from the onset of illness and in addition strong antibody cross-reactivity occurs among members of the family which may confuse interpretation of the results [1, 17, 18].

To date, polymerase chain reaction (PCR) methods have been suggested as appropriate laboratory diagnostic technique of DENV infections [1, 14]. Real-time reverse transcriptase PCR (RT-PCR) has been developed as an accurate diagnostic technique at an early stage of infection of several arboviruses. Advantages of Real time RT-PCR over other diagnostic methods including, higher sensitivity, higher specificity and rapidity [2, 16, 19, 20]. The aim of this study was to optimize real-time RT-PCR for detection of DENV infections by using rapid and simple nucleic acid extraction method.

**Materials And Methods**

**Clinical Samples**

A total of 208 blood samples were collected from febrile children (aged 2 months to 12 years) admitted at KCMC hospital between October 2013 to April 2014

**Control Samples**

Four positive samples and one internal extraction control obtained from National Health Laboratory Quality Assurance Training Centre (NHL-QATC) were included to validate the assay.

**Extraction of viral RNA**

RNA was extracted from blood sample by using boom extraction method as described by Boom *et al.*, [21]. The extracted RNA was stored at -80°C for further use.

**Reverse Transcription**

RNA was transcribed into cDNA by using QuantiTect-Reverse-Transcription kit (Qaigen, German) following manufacturer's protocol.

**Primer and probe sequence**

Primers and Probes were adopted from Pongsiri *et al.*, [2] and purchased from Applied Biosystems, UK (Table 1).
Table 1
Primer and probe sequence used in the assay

| Name  | Primer/Probe sequence | Length (kb) | Mol. Weight | Dye          |
|-------|-----------------------|-------------|-------------|--------------|
| DenvF | GAC TAG YGG TTA GAG GAG ACC | 21          | 6528        |              |
| DenvR | GHR GAG ACA GCA GGA TCT CTG | 21          | 6501        |              |
| DenProb | AAG GAC TAG MGG TTA GWG GAG ACC C | 25          | 9430        | 6-FAM MGBNFQ |

Real Time RT-PCR optimization

Assay conditions were optimized using various primer and probe concentrations, thermal conditions and different real time PCR systems. The concentrations of primers and probes and thermal profile were optimized to increase the sensitivity and specificity. The assay was optimized in a one-step and two-step real time RT-PCR.

In a one-step real time RT-PCR, the 10 µl reaction volume contained 0.2 µl Moloney Murine Leukemia Virus Reverse Transcriptase (M-MLV RT) (Invitrogen, Carlsbad CA), 5 µl of TaqMan Gene Expression master mix (Applied Biosystems, USA), 0.25 µM of dengue forward and reverse primers, 0.125 µM of dengue probe and nuclease free water. Large volume assay were performed with the AgPath-ID one step RT-PCR kit (Applied Biosystems, USA). The 25 µl reaction volumes contained 12.5 µl of 2x RT-PCR Buffer, 0.1 µM of dengue forward and reverse primers, 0.05 µM of dengue probe, 1 µl of 25 RT-PCR enzyme mix, 1 µl RNA sample and nuclease free water.

In a two-step real time RT-PCR, the 10 µl reaction volume contained 5 µl of TaqMan Gene Expression master mix (Applied Biosystems, USA), 1 µl of cDNA, 0.25 µM of dengue forward and reverse primers, 0.125 µM of dengue probe and nuclease free water. The large volume, 20 µl reaction volume contained 10 µl of TaqMan Gene Expression master mix, 2 µl of cDNA, 0.25 µM of dengue forward and reverse primers, 0.125 µM of dengue probe and nuclease free water.

Thermal conditions were optimised in both one-step and two-steps real time RT-PCR. In a one-step RT-PCR the thermal conditions were; reverse transcription at 50 °C for 30 min, initial denaturation 95 °C for 10 min followed by 40 cycles of 95 °C for 15 sec and 60 °C for 60 sec. In two-step RT-PCR the thermal conditions were; UDG incubation at 50 °C for 2 min, enzyme activation 95 °C for 10 sec, followed by 45 cycles of 95 °C for 15 sec and 60 °C for 1 min.

Assay conditions were also optimised in different real time PCR systems namely Stratagene® Mx3000P™ and Applied Biosystems® ViiA™7 real time PCR system. Positive controls and no template control were
included in our assay for each run, and a run was validated if the no template control did not exhibit fluorescent signal that cross the threshold line. A positive result was considered when the fluorescent signal crossed the threshold (i.e. exceeds background level).

**Results**

**Optimization of real time RT-PCR**

Various primer and probe concentrations, thermal conditions and real time RT-PCR systems were evaluated. Primer concentrations ranging from 5 µM to 10 µM and probe concentrations ranging from 3 µM to 10 µM were evaluated; final concentrations of 0.25 µM of each primer and 0.125 µM of probe were found to be the most sensitive in the assay.

Thermal profile efficiencies were compared, with annealing temperatures ranging from 55 °C to 60 °C. Annealing temperature was maintained at 60 °C which was found to be the best and thus had the best specificity for the assay.

TaqMan Gene Expression master mix kit (Applied Biosystems, USA) and the AgPath-ID one step RT-PCR kit (Applied Biosystems, USA) were also compared in the assay. TaqMan Gene Expression master mix kit (Applied Biosystems, USA) was found to be most efficient in both one-step and two step-methods. The 10 µl reaction volume containing 5 µl of TaqMan Gene Expression master mix, 1 µl of cDNA, 0.25 µM of each dengue forward and reverse primers, 0.125 µM of dengue probe and nuclease free water was found to be efficiency in our assay.

Reaction efficiencies were similar in both one-step and two-step RT-PCR methods but the sensitivity of the two-step RT-PCR method was slightly better than one-step RT-PCR method. There was a roughly 3-cycle difference between one-step and two step method which indicates 8 fold change in detection. The calculation is done by the formula $2^{\Delta CT}$ where 2 is used as an assumption of doubling in every cycle.

The technique was successfully optimized using two different PCR thermocyclers namely, Stratagene® Mx3000P™ and Applied Biosystems® ViiA™7. Figures 1 and 2 shows all positive controls can be amplified with both real time PCR systems but with different fluorescence detections efficient.

**Extraction of viral RNA**

Nucleic acid were purified from blood samples in less than 1 hour. Nucleic acid concentration and purity were assessed and quantified by NanoDrop UV Visible Spectrophotometer.

**Validation of dengue diagnostic real time PCR**

Validation and determination of performance of our assay were performed on control samples from NHL-QATC, Dar es Salaam. All four positive control samples were tested positive for DENV by the real time RT-
PCR assay. The 100% correlation of the results demonstrated 100% sensitivity of the real time RT-PCR assay.

**Clinical samples**

PCR screening of the febrile children revealed that none of them were positive for DENV. In the present study, the prevalence of DENV infections among febrile children admitted at KCMC hospital Paediatric Department was found to be 0.0%.

**Discussion**

We report optimization of real time RT-PCR assay for detection of DENV nucleic acid. The real time RT-PCR assay optimised has been shown to be sensitive enough to detect all of the four serotypes of DENV. The assay was rapid and accurate that can be performed in four hours including nucleic acid extraction. The rapidity and accurateness of the real time RT-PCR is essential in facilitating management and initiation of suitable therapy.

While the reaction efficiencies were similar there was a difference in sensitivity of detection between one-step and two-step real time RT-PCR methods. The lower CT value expressed in two-step method was roughly 3-cycle differences (8-fold difference in detection levels) indicating two-step method was more sensitive than one-step method. Since the same PCR conditions was used, the difference was probably due to reverse transcriptase reaction conditions and different enzymes used which were likely correlated to reduced sensitivity of detection in one-step real time RT-PCR method.

The assay was sensitive enough to be used with a simple RNA extraction method such as the boom method. Boom extraction method proved to work effectively by produced purified viral RNA reliable for amplification. Boom extraction method seem compatible for accurate real time RT-PCR and can be used at any setting. To our knowledge this is the first study to investigate on boom extraction method in extraction of DENV genome in East Africa. Investigation on extraction methods is important information in selecting the proper method of genomic extraction.

Our study demonstrated that none of the study participants had tested positive for DENV infection by real time RT-PCR. The 0% prevalence of DENV infection revealed by the present study could be explained by absence or very minimal circulation of DENV in Northern Tanzania as a results of climate factors which influences dengue ecology. Community behaviour factors such as environmental management could also count for the reported prevalence. To date there is no any epidemiological data that indicates outbreak of DENV infections in Northern Tanzania. In spite of the fact that we did not had any children tested positive by the real time RT-PCR, our findings support that other arboviruses are circulating in Northern Tanzania as most of the study participants presented with signs and symptoms that are closely related to arboviral infections. As arboviral infections are common cause of febrile illness and routinely misdiagnosed as malaria, diagnosis of arboviral could be tested by rapid and accurate diagnostic methods such as real time PCR so as to understand the contribution of these infections to febrile illness.
Conclusion

The optimized technique utilize a simple RNA extraction method such as the boom method and can works well with all range of real time PCR machines. Following the recent outbreak of dengue in Sub-Saharan Africa, further investigation on the cost effectiveness in adopting this technique for routine screening and monitoring of the infection should be done.

Limitation

We have optimised this technique in Northern Tanzania where there is very minimal/ no circulation of DENV. Using this technique in high DENV infection rate could be a promising avenue for routine screening and monitoring DENV infections.

Abbreviations

DENV: Dengue virus
DHS: Dengue Hemorrhagic Fever
DSS: Dengue Shock Syndrome
KCRI: Kilimanjaro Clinical Research Institute
KCMC: Kilimanjaro Christian Medical Centre
NHL-QATC: National Health Laboratory Quality Assurance Training Centre
NTC: No template control

Declarations

Ethics approval and consent to participate: Ethical approval for this study was granted and approved by the Kilimanjaro Christian Medical University College Ethics and Research Committee (CRERC) with a certificate No. 612. Written informed consent was obtained from parents or legal guardian of the children before inclusion in the study.

Availability of data and material: Not Applicable

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Competing interest: We declare that we have no conflict of interest.

Contribution: KA and RK involved in conception and design. AN, AM and AS provided input into the study design and laboratory analysis. KA and RK completed the final draft of the manuscript. All authors read and approved the final manuscript.

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Figures
Figure 1

Figure 2

Amplification of dengue controls using Stratagene Mx3000P DENV=dengue virus positive control; NTC=no template control