A diagnostic one-step real-time reverse transcription polymerase chain reaction method for accurate detection of influenza virus type A

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Abstract

Introduction: Influenza A is known as a public health concern worldwide. In this study, a novel one-step real-time reverse transcription polymerase chain reaction (rtRT-PCR) assay was designed and optimized for the detection of influenza A viruses.

Material and methods: The primers and probe were designed based on the analysis of 90 matrix nucleotide sequence data of influenza type A subtypes from the GenBank database of the National Center for Biotechnology Information (NCBI). The influenza virus A/Tehran/5652/2010 (H1N1 pdm09) was used as a reference. The rtRT-PCR assay was optimized, compared with that of the World Health Organization (WHO), and its analytical sensitivity, specificity and reproducibility were evaluated. In total, 64 nasopharyngeal swabs from patients with influenza-like illness (ILI) and 41 samples without ILI symptoms were tested for the virus, using conventional cell culture, direct immunofluorescence antibody (DFA) methods, and one-step rtRT-PCR with the designed primer set and probe and the WHO’s.

Results: The optimized assay results were similar to the WHO’s. The optimized assay results were similar to WHO’s, with non-significant differences for 10–10^3 copies of viral RNA/reaction (p > 0.05). It detected 10 copies of viral RNA/reaction with high reproducibility and no cross reactivity with other respiratory viruses. A specific cytopathic effect was observed in 6/64 (9.37%) of the ILI group using conventional culture and DFA staining methods; however, it was not seen in non-ILI. Also, the results of our assay and the WHO’s were similar to those of viral isolation and DFA staining.

Conclusions: Given the high specificity, sensitivity and reproducibility of this novel assay, it can serve as a reliable diagnostic tool for the detection of influenza A viruses in clinical specimens and lab experiments.

Key words: influenza A virus, one-step real-time polymerase chain reaction, detection.

Introduction

Influenza A viruses are one of the major human and animal pathogens, with a global distribution and known as an important public health problem [1, 2]. Previous studies reported that most influenza viruses which have been circulating extensively among the human population since 2003 are of type B and two subtypes H1N1 and H3N2 of type A.
Material and methods

Virus strain and culture

Influenza virus strain A/Tehran/5652/2010 (H1N1 pdm09), as a reference strain, was kindly provided by Dr. T. Mokhtari Azad (School of Public Health, Tehran University of Medical Sciences). The virus was propagated in Madin-Darby canine kidney (MDCK) cells [23]. The cells were examined on a daily basis for the presence of a specific cytopathic effect (CPE). After the appearance of a CPE, the direct immunofluorescence antibody (DFA) test with specific antibody against influenza A virus nucleoprotein was performed to confirm the presence of the virus, using a commercially available kit (Dakocytomation Ltd; Ely, UK) and according to the manufacturer’s instructions. Moreover, nucleic acid was extracted from a 200 µl volume of cell lysate using a viral RNA isolation kit (Invitek, Berlin, Germany) for further experimentations.

Primer design

The oligonucleotide primers and probe were designed according to the 90 available M nucleotide sequence data of different subtypes of influenza type A viruses, from the GenBank database of the National Center for Biotechnology Information (NCBI). The subtypes included 20 H1N1, 20 H2N2, 20 H3N2, 10 H9N2, 10 H5N1, and 10 H7N3 from all around the world. The sequences were analyzed by the Lasergene sequence analysis software package (DNASTAR, Madison, WI, USA), and the primer set and probe were designed based on the conserved sequences. The probe was labeled with 6-carboxyfluorescein (FAM), as a fluorescent reporter dye, and black hole quencher-1 (BHQ-1), at the 5′ and 3′ ends, respectively. Besides cell culture inoculation and DFA, a provided World Health Organization (WHO) rtRT-PCR with the InFA primer-probe set (Table I) was also applied to compare its sensitivity with that of the designed primer and probe set for detecting serially diluted concentrations (10–106) of influenza virus type A RNA in three replicates, based on the recommended protocol [24]. All the primer sets and probes were synthesized by AnaSpec (Fremont, CA, USA).

Optimization of one-step real-time polymerase chain reaction

The reaction was performed using the specific primer set and probe designed to detect influenza virus type A. The 25-µl PCR reaction consists of 12.5 µl 2X Reaction Mix (Invitrogen, Carlsbad, CA, USA), 0.2 µM of each primer (FluAF and FluAR), and 0.1 µM probe, 0.5 µl of SuperScript III Platinum One-Step rtRT-PCR System (Invitrogen, Carlsbad, CA, USA), 0.5 µl of 25 µM ROX reference dye, 5 µl of RNA sample and distilled water. The amplification process was performed in a StepOnePlus Real-Time PCR System instrument (Applied Biosystems, USA), according to the cycling protocol: 50°C for 25 min (reverse transcription) and 95°C for 10 min (DNA polymerase activation), followed
by 50 cycles of 95°C for 15 s (denaturation) and 55–60°C (55, 56, 57, 58, 59, 60) for 60 s (annealing and extension).

**Construction of the plasmid containing the target M sequence of the influenza virus type A**

The designed primer set was used to construct the standard control plasmid containing the target M sequence of the influenza A virus. The extracted RNA was used for the synthesis of the first strand cDNA using M-MuLV reverse transcriptase (Fermentase, Lithuania), and PCR was performed using pfu DNA polymerase (Fermentas, Lithuania) in a 25 µl reaction. The amplified products were cloned into the pTZ57R/T vector and transformed into the *Escherichia coli* DH5α competent cells (TaKaRa Biotechnology Co., Dalian, China) using a commercially available cloning kit (Fermentase, Lithuania). To confirm the recombinant vectors, positive clones were selected, their plasmids were sequenced with the M13 primers (Shanghai Sangon, China) in the ABI 3130 Genetic Analyzer using the BigDye Terminator v3.1 Cycle Sequencing Kit (Applied Biosystems, USA), and the confirmed plasmid concentration and its copy number per µl of sample were calculated [25]. The resulting plasmid was used as the template for the subsequent analytical studies.

**Analytical sensitivity, reproducibility, and specificity**

To detect the sensitivity of the influenza A virus one-step rRT-PCR method, 10-fold serial dilutions in the range 10^1–10^6 copies/µl of the respective recombinant plasmid were prepared and used. To determine the reproducibility of the assay, all the samples were tested in four separate experiments in two replicates of each run for inter- and intra-assay variability. All the mean values, standard deviations (SD) and coefficient of variations (CV) were calculated using SPSS for Windows software (version 16, SPSS Inc., Chicago, IL, USA). The specificity of the method was tested against influenza virus type B, mumps, measles, rubella, Herpes simplex virus 1 and 2, Cytomegalovirus, and Epstein-Barr virus. The primer set and probe were also analyzed using the nucleotide BLAST in the NCBI gene bank to find any potential cross-reactivity with other microorganisms.

**Handling and preparation of clinical samples**

A total of 64 nasopharyngeal swabs in viral transport medium from individuals with influenza-like illness (ILI) and 41 samples without ILI symptoms were collected in Namazi hospital, southern Iran, during 2011–2012 and sent to the laboratory immediately. Nucleic acid was extracted from a 200 µl volume of each sample using a viral RNA isolation kit (Invitek, Berlin, Germany) and used for one-step rRT-PCR with the designed primer set and probe and those of WHO. In addition, 0.2 ml of each sample was inoculated on the monolayer MDCK cell culture. The cells were daily observed for the CPE and, after a week, lysate of the cell cultures were used for DFA, using a commercially available kit (Dakocytomation Ltd; Ely, UK).

**Results**

**Optimization of the one-step real-time reverse transcription polymerase chain reaction assay condition**

FluAF and FluAR were designed to amplify a 111 bp fragment of the M gene of influenza virus type A. It was successfully cloned into the pTZ57R/T vector. The qRT-PCR was performed optimally in different annealing temperatures (55–60°C), using our designed primer set and probe, which yielded similar results. As revealed, the results of the optimized one-step TaqMan
rtRT-PCR were consistent with those from the WHO, when using the same reference RNA sample in serially diluted concentrations (Table II).

Study of the sensitivity, reproducibility, and specificity of the influenza virus type A one-step real-time reverse transcription polymerase chain reaction assay

The sensitivity and reproducibility of the present assay were assessed by serial 10-fold dilutions (10^1–10^6 copies/μl) of the constructed plasmids which were amplified using this one-step rtRT-PCR method in independent and repeated experiments. A standard curve was plotted by amplification of 10-fold diluted (10^1–10^6 copies/μl) constructed plasmids versus the threshold cycle (CT) numbers and analysis, using StepOne Software Version 2.1 (Applied Biosystems, Foster City, CA, USA). The detection limit of the target M gene of influenza virus type A was approximately 10 copies per microliter of reaction. The linear regression analysis of the results exhibited a high value of the correlation coefficient ($R^2 = 0.997$), indicating the precise and reliable consistency of the replicates (Figure 1). The reproducibility of the assay was confirmed by high intra-CV and inter-CV as 0.39–1.20% and 0.38–1.23%, respectively (Table III). The experiments for the evaluation of specificity of the present one-step rtRT-PCR assay with the designed primer set and probe indicated no positive fluorescent signal, when testing the other respiratory viruses. Moreover, based on the blasting search in the NCBI database, no cross reactivity was observed between the respective primer set and probe and other microorganisms’ genome sequences.

<table>
<thead>
<tr>
<th>Copy number</th>
<th>Ct-value (mean ± SD for indicated assay) P-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>FluA</td>
<td>InfA</td>
</tr>
<tr>
<td>10^6</td>
<td>15.10 ±0.21</td>
</tr>
<tr>
<td>10^5</td>
<td>19.36 ±0.18</td>
</tr>
<tr>
<td>10^4</td>
<td>23.29 ±0.16</td>
</tr>
<tr>
<td>10^3</td>
<td>27.85 ±0.26</td>
</tr>
<tr>
<td>10^2</td>
<td>30.44 ±0.29</td>
</tr>
<tr>
<td>10^1</td>
<td>34.97 ±0.79</td>
</tr>
</tbody>
</table>

Figure 1. Amplification plot and standard curve of the one-step rtRT-PCR assay for the detection of 10-fold dilutions of the recombinant plasmid containing the target M gene
Evaluating the one-step real-time reverse transcription polymerase chain reaction assay using clinical specimens

Of the 64 nasopharyngeal swabs from patients with ILI, the specific influenza A CPE was observed in 6/64 (9.37%) samples; however, it was not detected in non-ILI ones (0/41). The results of DFA staining revealed the same results. Comparison between our one-step rtRT-PCR assay and conventional methods showed that its results were similar to those of viral isolation and DFA staining. In addition, the results of the assay using our primer set and probe and the WHO’s were the same (Table IV).

Discussion

Viral isolation by conventional culture methods currently serves as the gold standard for the detection and typing of influenza viruses, but important limitations including the laborious and time-consuming process, as well as low sensitivity for the detection of some subtypes of the virus, still exist [26, 27]. Besides, the risk of emergence and human pandemic potential of new influenza A viruses has prompted the attempts to develop more new rapid, sensitive and specific detection methods based on genetic recombination technology and molecular techniques for the most effective management and prevention of the respective infections [28].

In the present study, a novel one-step rtRT-PCR assay was designed to accurately differentiate the influenza virus type A from other respiratory ones. It was demonstrated that the assay was highly sensitive and could achieve a detection limit as low as 10^1 copies per microliter, which was more sensitive than earlier multiplex rtRT-CR assays for simultaneous detection of influenza virus type A and other subtypes or types. A previous study on the simultaneous detection of influenza type B virus and influenza A virus subtypes H5N1, H3N2, and H1N1 using a multiplex rtRT-PCR assay showed a sensitivity of approximately 10^1–10^2 copies per microliter of reaction [29]. Jiang et al. reported that the sensitivity of the designed rtRT-PCR method for the detection of a novel influenza A (H1N1) virus was 200 copies of in vitro-transcribed target RNA [30]. The sensitivity of a recently developed multiplex rtRT-PCR system for the detection of influenza type A and B viruses and subtype H1N1 by Huber et al. was found to be 3.5 × 10^2 RNA copies per PCR reaction [31]. However, the detection limit of the present study was almost the same as that of the single rtRT-PCR assay for the detection of influenza A virus, studied earlier [32]. The CDC rtRT-PCR Swine Flu Panel has been shown to detect 5 copies of RNA per reaction [33]. Moreover, the results of the present study revealed the same amplification curve in all of the used annealing temperatures (55–60°C). This may indicate the high efficiency of the designed primers and probe. Also, the current primers and probe were designed based on sequence data at least 10% of which belong to isolates from Middle East countries including Iraq, Iran, Turkey, Afghanistan, Pakistan, Kuwait, and Qatar. This could increase the specificity of the assay for the detection of domestic viruses circulating in the region.

The efficiency and performance of rtRT-PCR systems are strongly dependent on the quality of the primer sets and probes applied for the amplification of nucleic acid target regions. Development of a rtRT-PCR assay with higher sensitivity and specificity for the detection of influenza A viruses presents a big challenge because of the high mutation frequencies among the sequences of the virus segments. In the present study, oligonucleotide primer set and probe were designed based on the more conserved region of the M segment of the virus genome; however, some degenerative nucleotides were also included. As the results showed, the positivity cycles of the test were similar in all
used annealing temperatures, indicating the high efficiency of the assay. Moreover, considering the results of inter- and intra-CV, good reproducibility of the test was confirmed. The lack of cross reactivity with other organisms is indicative of high specificity of the test.

Previous studies revealed varying rates of sensitivity for the detection of influenza viruses in clinical specimens using conventional methods. It was reported that the sensitivity of the rapid influenza diagnostic test (RIDT) for the detection of influenza A (H1N1) pdm09 varied between 18% and 69%. The sensitivity results for DFA were also different and ranged from 47% to 93% [34–36]. In the present study, a great similarity was observed between the results of the conventional methods (culture and DFA), our designed assay and those of the WHO. This may be due to the limitation (culture and DFA), our designed assay and those of the WHO. This may be due to the limitation of the number of clinical samples evaluated. The positive results for the assay may increase if more samples are examined, compared with the conventional methods.

In conclusion, given the high specificity, sensitivity and reproducibility of this novel one-step real-time reverse transcription polymerase chain reaction method, this can serve as a potential reliable diagnostic tool for the detection of influenza type A both in clinical specimens and lab experiments. However, clinical sensitivity and specificity using more clinical specimens should be evaluated.

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Conflict of interest

The authors declare no conflict of interest.

References