Rapid and Sensitive Detection of Novel Avian-Origin Influenza A (H7N9) Virus by Reverse Transcription Loop-Mediated Isothermal Amplification Combined with a Lateral-Flow Device

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Abstract

A severe disease in humans caused by a novel avian-origin influenza A (H7N9) virus emerged in China recently, which has caused at least 128 cases and 26 deaths. Rapid detection of the novel H7N9 virus is urgently needed to differentiate the disease from other infections, and to facilitate infection control as well as epidemiologic investigations. In this study, a reverse transcription loop-mediated isothermal amplification combined with a lateral flow device (RT-LAMP-LFD) assay to rapidly detect H7N9 virus was developed and evaluated. The RT-LAMP primers were designed to target the haemagglutinin (HA) and neuraminidase (NA) genes of H7N9 virus. Results of 10-fold dilution series assays showed that analysis of RT-LAMP products by the LFD method was as sensitive as real-time turbidity detection, and that the analytic sensitivities of the HA and NA RT-LAMP assays were both 10 copies of synthetic RNA. Furthermore, both the assays showed 100% clinical specificity for identification of H7N9 virus. The performance characteristics of the RT-LAMP-LFD assay were evaluated with 80 clinical specimens collected from suspected H7N9 patients. The NA RT-LAMP-LFD assay was more sensitive than real time RT-PCR assay. Compared with a combination of virus culture and real-time RT-PCR, the sensitivity, specificity, positive predictive value, and negative predictive value of the RT-LAMP-LFD assay were all 100%. Overall, The RT-LAMP-LFD assay established in this study can be used as a reliable method for early diagnosis of the avian-origin influenza A (H7N9) virus infection.

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Introduction

Since February, 2013, a previously unrecognized novel avian-origin influenza A (H7N9) virus associated with human deaths has emerged in China [1]. Although the transmission of avian influenza virus subtypes H5, H7, and H9 to human has been reported early [2], it is the first time that N9 subtype influenza virus has infected human beings. Phylogenetic analysis has shown that all the genes of the novel H7N9 viruses are of avian origin, and are recombined from at least three influenza virus lineages [1,3]. Compared with avian influenza A (H5N1) virus, it seems to be easier for the novel H7N9 virus to transmit from animals to human. As of 2 May 2013, there have been 128 laboratory-confirmed cases of human infection with H7N9 virus including 26 fatalities [4]. Although different strains of the novel H7N9 virus have been isolated from poultry, birds and the environmental specimens (Data from Global Initiative on Sharing All Influenza Data, GISAID), the source of infection in most of the human cases still remains to be determined. So far, there has been no direct evidence of human-to-human transmission of this virus, however, the presence of mutations in the polymerase basic protein 2 (PB2) gene associated with improved replication of avian influenza viruses in mammals might indicate a certain propensity of the H7N9 virus to further adapt to humans [1,3]. Therefore, the potential for the novel influenza A (H7N9) virus to spread among the human population is still exist.

A rapid and sensitive methodology for the diagnosis of H7N9 infection is urgently needed to facilitate clinical care, infection control, and epidemiologic investigations. Methods based on PCR are more rapid and sensitive than traditional techniques including virus isolation and serological assays. Real-time RT-PCR is at present the powerful molecular diagnostic method for the novel influenza A (H7N9) virus infection [5,6], however, it requires expensive real-time PCR equipment and highly skilled technicians, which make this method not suitable for use in primary clinical
settings or for field use. The Loop-mediated isothermal amplification (LAMP) method allows amplification of DNA with high specificity and sensitivity at a constant temperature of 60–65°C [7]. As the reaction is conducted under isothermal conditions, it can be carried out with a simple water bath so that a thermal cycler is not required. LAMP can also be used to detect RNA template by the use of reverse transcriptase together with DNA polymerase [7,8]. To date, RT-LAMP methods have been developed to detect various RNA viruses [9–12].

LAMP products can be detected by agarose gel electrophoresis, by the use of spectrophotometric equipment to measure turbidity [13], or by visual inspection of turbidity or color changes [14,15]. As a result, non-specific amplification products may cause false positive results. To help overcome this problem, LAMP products can be identified by restriction endonuclease digestion [7] or by hybridization with specific probes [16]. To further simplify and speed up the process of the LAMP assay, amplicon detection by lateral flow device (LFD) was successfully applied [17]. In this study, a RT-LAMP amplification combined with a LFD (RT-LAMP-LFD) assay was developed for the rapid detection of novel avian-origin influenza A (H7N9) virus. The characteristics of simplicity (without the need for complex or expensive equipment), rapidity (results can be obtained within 1 h), and excellent sensitivity and specificity make this RT-LAMP-LFD method more suitable for use in low-equipment setting laboratory.

**Materials and Methods**

**Ethics statement**

Written informed consent for the use of the clinical specimens was obtained from all patients involved in this study. This study was approved by the Ethics Committee of the Jiangsu Provincial Center for Disease Prevention and Control.

**Viral isolates and clinical specimens**

H7N9 virus isolates were cultured in Madin-Darby canine kidney (MDCK) cells using standard techniques. An isolated A/Nanjing/1/2013 (H7N9) virus was used as the reference virus. Other genetically and clinically related virus isolates including seasonal influenza viruses (A/H1N1, A/H3N2, and B), 2009 swine-origin influenza virus A/H1N1, avian influenza viruses (A/H5N1 and A/H9N1), parainfluenza viruses (types 1, 2, 3, and 4), human coronaviruses (229E, OC43, HKU1, and NL63) and respiratory syncytial viruses (types A and B) were used as control viruses to assess the specificity of the RT-LAMP assay. A total of 80 respiratory clinical specimens (65 pharyngeal swabs, 7 sputa, and 8tracheal aspirates) were collected from suspected H7N9 patients in the acute phase of illness. All these isolates and specimens were stored at −80°C until use.

**Extraction of viral RNA**

The genomic viral RNA was extracted from 200 μl of samples by using a High Pure viral RNA kit (Roche Diagnostics, Mannheim, Germany) according to the manufacturer’s instructions. The RNA was eluted in a final volume of 100 μl of elution buffer and kept at −80°C until further analysis.

**Preparation of virus RNA standards**

The haemagglutinin (HA) and neuraminidase (NA) genes were amplified from the H7N9 virus strain A/Nanjing/1/2013 (H7N9) with primers containing T7 promoter sequence in the reverse sides (Table 1) and were in vitro transcribed with T7 RNA polymerase (TaKaRa Biotechnology Co. Ltd., Dalian, China) according to the manufacturer’s instructions. The synthetic RNA transcripts were then purified, quantified, mixed in equal-molar amounts, and ten-fold diluted ranging from 10⁷ to 10¹ RNA copies/μl.

** Primer design for RT-LAMP**

Two separate RT-LAMP assays (six primers each) were designed according to HA and NA gene sequences of the reference virus using a PrimerExplorer V4 program (http://primerexplorer.jp/e/). The feasibility and specificity of the primers were subsequently checked by BLAST search against published genome sequences of the novel H7N9 viruses from the GISAID EpiFlu database and against NCBI database. Details of the final primers are shown in Figure 1 and Table 2. All the primers were synthesized by Sangon (Shanghai, China).

**RT-LAMP reaction and product detection**

RT-LAMP was carried out with a RNA Amplification Kit (RT-LAMP) (Eiken China Co., Ltd., Shanghai, China) in a final volume of 25 μl containing 0.2 μM each of the outer primers F3 and B3, 2.0 μM each of the primers FIP and BIP, and 1.0 μM each of the primers LF and LB (labeled or not). The reaction mixture containing distilled water was used as negative controls. RT-LAMP amplification reactions were carried out at 60, 63, and 65°C for 60 min using a real-time turbidimeter (LA320C, Teranecs, Tokyo, Japan). The RT-LAMP products were also detected by a LFD (Ustar Biotech Co., Ltd., Hangzhou, China) as described previously [17]. The principle is primers LF and LB are conjugated to fluorescein isothiocyanate (FITC) and biotin, respectively, during a positive RT-LAMP reaction both tags are incorporated in the amplicons resulting in a visible test line. The LFD detection also utilizes a control line to indicate whether the unit has functioned correctly.

**Sensitivity of RT-LAMP by real-time turbidity detection and LFD**

To compare test sensitivity, RNA was extracted from the culture supernatants of H7N9 virus strain A/Nanjing/1/2013 (H7N9)-

| Table 1. Primers used for in vitro transcription of virus RNA standards. |
|---|
| **Name** | **Sequence (5′–3′)** | **Position** |
| HA-Forward | CTCGTTAATTGCCGCTGGTTGC | 15–35 |
| HA-Reverse | ACTCGTTAATTGCCGCTGGTTGC* | 1668–1649 |
| NA-Forward | TCTATGCACTTCAGCCACTG | 21–40 |
| NA-Reverse | ACTCGTTAATTGCCGCTGGTTGC* | 1373–1354 |

* T7 RNA polymerase promoter sequence.

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Table 2. Details of primers used for RT-LAMP assay.

| Target | Name   | Type               | Position | Sequence (5'-3') |
|--------|--------|--------------------|----------|-----------------|
| HA     | F3     | Forward outer      | 309-328  | TGTCTGTTATCCCTGGGAAAAT |
|        | B3     | Reverse outer      | 504-484  | AGCATTATCGTGGTAGCAG |
|        | FIP (F1c+F2) | Forward inner  | F1c: 405–381 | GTATGTGATCCATTGCTTCCGT |
|        |        |                    | F2: 331–351 | GTGAAATGAGAAGCTGGAGG |
|        | BIP (B1c+B2) | Reverse inner | B1c: 409–433 | GGAATAGAATACAGGARCAACA |
|        |        |                    | B2: 482–462 | AGGAAATTCTTTTCGTAAG |
|        | LF a   | Forward loop       | 374–353  | CCCGCTGTACTCTGAGATAATT |
|        | LB b   | Reverse loop       | 436–459  | GCATGAGGAGACTCGATCC |
| NA     | F3     | Forward outer      | 1097–1114| CGAGGTCTGATACGAGA |
|        | B3     | Reverse outer      | 1295–1276| TTATCYTCCTGGGTCTTCC |
|        | FIP (F1c+F2) | Forward inner  | F1c: 1188–1164 | AGCGTTTARTACAATTGTCTGACCT |
|        |        |                    | F2: 1117–1138 | TTAAGCTGAAATATGAGA |
|        | BIP (B1c+B2) | Reverse inner | B1c: 1191–1212 | CTGGACCTGATCTACAGTAC |
|        |        |                    | B2: 1266–1247 | YTCCACATAAAAAACACGTC |
|        | LF a   | Forward loop       | 1163–1139| TGAATGGGCTTTGTACTCATCTG |
|        | LB b   | Reverse loop       | 1217–1239| TGGACTATTGGGCTGARGGGAC |

*5'-Labeled with FITC when used in RT-LAMP-LFD assay.
*5'-Labeled with biotin when used in RT-LAMP-LFD assay.

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Figure 1. Schematic showing location of RT-LAMP primer binding sites within HA and NA genes. (A) HA RT-LAMP primer binding sites. Assay spans region from nucleotide 309–504 with reference to the HA gene sequence of the H7N9 virus strain A/Nanjing/1/2013 (H7N9). (B) NA RT-LAMP primer binding sites. Assay spans region from nucleotide 1097–1295 with reference to the NA gene sequence of the H7N9 virus strain A/Nanjing/1/2013 (H7N9).
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infected MDCK cells. Ten-fold serial dilutions of RNA were used as templates for RT-LAMP reactions at 63°C for 1 h. Besides real-time turbidity detection, the RT-LAMP products were simultaneously detected by LFD as described above.

Analytical sensitivity and specificity of the RT-LAMP-LFD

Ten-fold serial dilutions of synthetic H7N9 viral RNA transcripts of HA and NA genes (ranging from 10^7 to 10^1 RNA copies) were used to assess the detection limits of the HA and NA RT-LAMP-LFD assays. The specificities of the RT-LAMP-LFD assays were determined by analyzing the RNA extracts from various control viruses mentioned above and the reference virus.

Real-time RT-PCR detection of H7N9 virus

Real-time RT-PCR for detection of H7N9 virus was carried out using the primers and probes recommended by the WHO Collaborating Centre in Beijing [5] and the SuperScript III Platinum One-Step qRT-PCR Kit (Invitrogen) according to the instructions. The cycling conditions were composed of 20 min at 50°C, 2 min at 95°C, followed by 40 cycles with 94°C for 15 s and 60°C for 1 min, and a final extension cycle of 72°C for 10 min.

Evaluation of the RT-LAMP-LFD assay with clinical specimens

To evaluate the performance characteristics of the RT-LAMP-LFD assay, a reference standard which represents the combined

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**Table 3. Performance of RT-LAMP-LFD assay compared with the reference standard for detecting avian-origin influenza A (H7N9) virus.**

|                  | Reference standard | Performance characteristics (%) * |
|------------------|--------------------|----------------------------------|
|                  | Positive | Negative | Total | Sensitivity | Specificity | PPV | NPV |
| HA Positive      | 22       | 0        | 22    | 100%        | 100%        | 100%| 100%|
| HA Negative      | 0        | 58       | 58    |             |             |     |     |
| HA Total         | 22       | 58       | 80    |             |             |     |     |
| NA Positive      | 22b      | 0        | 22    | 100%        | 100%        | 100%| 100%|
| NA Negative      | 0        | 58       | 58    |             |             |     |     |
| NA Total         | 22       | 58       | 80    |             |             |     |     |

*PPV, positive predictive value; NPV, negative predictive value.

bOne HA-positive while NA-negative sample detected by real-time RT-PCR was further confirmed to be H7N9 infection by viral culture and sequencing.

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The LFD method which can provide a result within 5–10 min after detection limits obtained by RT-LAMP-turbidity detection. Thus, Figure 3 viruses. As shown in were tested against viral RNA extracts from various control LAMP-LFD assays to cross-react with other related viruses which copies of synthetic RNA. To assess the potential for the RT-LAMP-LFD assays, indicating the high specificities of both the assays.

Evaluation of the RT-LAMP-LFD with clinical specimens
To evaluate the performance characteristics of the HA and NA RT-LAMP-LFD assays, a total of 80 clinical specimens collected from suspected H7N9 patients in the acute phase of illness were subjected to RT-LAMP-LFD assay with the parallel analysis by the reference methods. The results showed that, of 80 specimens, 22 were positive for both HA and NA genes as detected by RT-LAMP-LFD, of which 21 were positive for both HA and NA genes as detected by real-time RT-PCR. The HA-positive but NA-negative specimen detected by real-time RT-PCR was further analyzed by viral culture and sequencing, and was verified as H7N9 virus infection, indicating the superior sensitivity of the RT-LAMP-LFD assay. Compared to the reference standard, the sensitivity, specificity, positive predictive value, and negative predictive value of both HA and NA RT-LAMP-LFD assays were all 100% (Table 3).

Discussion
Major uncertainties still exist with regards to the genetic variability and the pandemic potential of the novel avian-origin influenza A (H7N9) virus which warrant the development of new methods to detect this virus. In this study, we describe a sensitive RT-LAMP-LFD method for the specific detection of H7N9 virus for the first time. The optimal LAMP reaction conditions are 63°C for 60 min, which is more suitable for low-equipment setting laboratory and for on-site testing.

The use of the four specific primers and two loop primers targeting the HA or NA gene of H7N9 virus ensured high specificity of nucleic acid amplification. Both the HA and NA RT-LAMP-LFD assays showed 100% specificities for identification of H7N9 virus. Furthermore, loop primers could accelerate the LAMP reaction because they hybridize to the stem-loops, except for those loops that are hybridized by the inner primers and prime strand displacement DNA synthesis [18]. Although 60 min was used for H7N9 RT-LAMP reactions, most of the amplification reactions could be finished within 40 min (Figure 2). Thus, the RT-LAMP-LFD assay is faster than real-time RT-PCR.

The concordance of high analytical sensitivity between RT-LAMP and the most sensitive molecular methods for detection of pathogens has been previously reported [19–21]. Several possible factors may contribute to this fact. For example, the RT-LAMP reaction was less affected by the presence of various salts, was less
sensitive to inhibitors, and was able to tolerate the inhibitory effect of large amounts of templates [22]. In this study, the analytic sensitivities of the HA and NA RT-LAMP assays were both 10 copies of synthetic RNA. Compared to the reference standard, the sensitivity, specificity, positive predictive value, and negative predictive value of the RT-LAMP assays were all 100%.

The RT-LAMP results can be determined by agarose gel electrophoresis, by the use of spectrophotometric equipment to measure turbidity, by naked eye for the presence of a white precipitate derived from magnesium pyrophosphate, or by visualization of the RT-LAMP products under natural light or UV irradiation after adding SYBR green I or calcine dyes. However, due to the use of several primers, RT-LAMP generates a complex mixture of DNA products, and thus these product detection methods cannot distinguish specific and non-specific amplicons. Furthermore, assessment of turbidity or color with the unaided eye is potentially subjective, and there is always the possibility that a sample may be somewhat ambiguous to the naked eye when the concentration of virus is low [23]. Additionally, some dyes such as SYBR green I have adverse effect on LAMP amplification reaction. Gel electrophoresis has been found to be approximately 10×more sensitive than the SYBR green/naked eye inspection [24], nevertheless, electrophoresis after amplification increases the opportunity for product contamination. In the present study, we used a simple LFD utilizing a lateral flow strip housed in an enclosed, sealed plastic device to prevent the leakage of amplicons to objectively detect RT-LAMP products in approximately 5 min [17,25,26]. The use of LFD to detect RT-LAMP products not only makes the assay more specific, but also negates the need for electrophoresis equipment and DNA detection equipment. Our results showed that the LFD method was as sensitive as real-time turbidity detection. In some studies, only one labeled primer was used in LAMP reaction, another labeled probe was added into the LAMP amplicons to form double-labeled detectable products [27–29]. This would increase the chance of product contamination. In our study, both loop primers were labeled with tags such as FITC and biotin. The results obtained from this study and others [24,30,31] demonstrated that the usage of two labeled primers has no adverse effect on LAMP reaction.

In conclusion, these data show that a reliable RT-LAMP-LFD assay has been developed for the detection of novel avian-origin influenza A (H7N9) virus causing the current outbreak, which would facilitate the clinical care, infection control, as well as epidemiologic investigations. The RT-LAMP-LFD assay is specific and sensitive, and does not require expensive equipment. The use of the LFD provides a rapid and objective readout of the assay’s results and avoids cross-contamination. This RT-LAMP-LFD assay is especially useful in resource-limited situations such as primary care facilities.

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Author Contributions

Conceived and designed the experiments: YG BW LC. Performed the experiments: YG BW XQ KZ XG. Analyzed the data: YQ YZ ZS. Contributed reagents/materials/analysis tools: XG YQ YZ. Wrote the paper: YG MZ HW LC.

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