Reverse transcription recombinase-aided amplification assay for H5 subtype avian influenza virus

Suchun Wang†, Yang Li†, Fuyou Zhang†, Nan Jiang1,2, Qingye Zhuang3, Guanyu Hou1, Lijian Jiang1,2, Jianmin Yu1, Xiaohui Yu1, Hualei Liu1, Chenglong Zhao1,2, Liping Yuan1, Baoxu Huang1 and Kaicheng Wang1,4*

Abstract

Background: The H5 subtype avian influenza virus (AIV) has caused huge economic losses to the poultry industry and is a threat to human health. A rapid and simple test is needed to confirm infection in suspected cases during disease outbreaks.

Methods: In this study, we developed a reverse transcription recombinase-aided amplification (RT-RAA) assay for the detection of H5 subtype AIV. Assays were performed at a single temperature (39 °C), and the results were obtained within 20 min.

Results: The assay showed no cross-detection with Newcastle disease virus or infectious bronchitis virus. The analytical sensitivity was 10^3 RNA copies/μL at a 95% confidence interval according to probit regression analysis, with 100% specificity. Compared with published reverse transcription quantitative real-time polymerase chain reaction assays, the \( \kappa \) value of the RT-RAA assay in 420 avian clinical samples was 0.983 (\( p < 0.001 \)). The sensitivity for avian clinical sample detection was 97.26% (95% CI, 89.56–99.52%), and the specificity was 100% (95% CI, 98.64–100%).

Conclusions: These results indicated that our RT-RAA assay may be a valuable tool for detecting H5 subtype AIV.

Keywords: Avian influenza, H5 subtype, Reverse transcription recombinase-aided amplification assay, Single temperature, Rapid, Sensitive

Introduction

Influenza A virus (IAV), a member of the genus Influenza, family Orthomyxoviridae, contains eight single-stranded negative-sense RNA segments encoding at least 10 proteins [1, 2]. The IAV member avian influenza virus (AIV) can cause avian disease and public health problems. AIV can be classified into highly pathogenic avian influenza virus (HPAIV) and low pathogenic avian influenza virus (LPAIV) based on pathogenicity in chickens. The current HPAI H5N1 virus outbreak (from 2003 onwards) is, however, unprecedented in scale and geographic distribution, which have been prevalent among poultry in Asia, Europe and Africa. The viruses constantly undergo genetic drift and shift that permanently threatens the poultry industry and human health [3].

All LPAIV and HPAIV infections with H5 subtypes in poultry are notifiable to the World Organization for Animal Health [4]. Determination of the subtypes of AIV is of importance for the diagnosis of these infections, which can be achieved by analysis of the amino acid sequence of the cleavage site in the AIV hemagglutinin (HA) gene.
Besides, there are many methods, such as restriction enzyme cleavage patterns, probe hybridization and real time reverse transcription quantitative polymerase chain reaction (RT-qPCR) [5–8]. Based on the widespread availability of RT-qPCR technology in diagnostic laboratories, it has been used to identify H5 subtype HPAIV of the goose/Guangdong (gs/GD) lineage. Recently, rapid isothermal amplification techniques have been developed, such as loop-mediated isothermal amplification (LAMP) [7], recombinase polymerase amplification (RPA) [8], recombinase-aided amplification (RT-RAA) and strand displacement amplification (SDA) [9], and widely used in clinical detection. Among these rapid nucleic acid detection methods, reverse transcription recombinase-aided amplification (RT-RAA) is a rapid thermostatic nucleic acid amplification technology that utilizes a recombinant enzyme obtained from bacteria or fungi. At room temperature, the recombinant enzyme can tightly bind to the primer DNA to form polymer. When the primer recognizes the template DNA for a complementary sequence that perfectly matches it, with the help of a single-stranded DNA binding protein, it opens the double-stranded structure of the template DNA under the action of DNA polymerase, a new cDNA strand is formed, and the amplification product grows exponentially [10]. This technology has the characteristics of high sensitivity, stronger specificity and reliability.

In order to quickly and simply detect H5 subtype AIV and provide more sufficient preparation time for prevention and control, an RT-RAA assay was designed and its analytical specificity and sensitivity were evaluated in this study. The result suggested that RT-RAA assay meet the need of field testing, and presented a rapid and sensitive detection method that could be used as an alternative to animal inoculation or nucleotide sequencing.

Materials and methods

Ethics statement

This study was conducted according to the animal welfare guidelines of the World Organization for Animal Health and was approved by the Animal Welfare Committee of the China Animal Health and Epidemiology Center (CAHEC). CAHEC has permission to study the activities of HPAIV. The swab samples were collected after being granted permission by several relevant parties, including the Ministry of Agriculture and Rural Affairs of China, CAHEC, the relevant veterinary sections of the provincial and national governments, and the relevant farm owners.

Samples and extraction of viral nucleic acids

All the nucleic acid of AIVs, Newcastle disease virus (NDV) and infectious bronchitis virus (IBV) used in this study were identified in our laboratory (the National Avian Influenza Professional Laboratory in China Animal Health and Epidemiology Center) and were stored at −80 °C. The samples were centrifuged at 12,000 × g for 10 min and the supernatant from each sample was used for RNA extraction on the QIAxtractor platform using a cador Pathogen 96 QIAcube HT kit (Qiagen). The extracted RNA was stored at −80 °C for subsequent tests.

Preparation of viral RNA standard

An H5 subtype AIV viral RNA standard was developed using the reference strain A/duck/Yunnan/5310/2006(H5N1) (GenBank accession number CY030889). A 1776-bp fragment of the whole HA gene of H5 subtype AIV was constructed and digested with enzyme. The digested plasmid was purified with DNA recovery kit (Thermo Fisher Scientific) and transcribed in vitro with RNA Production System-T7 kit. The products were extracted and purified by Rneasy MiNi Kit to remove the heteroproteins and various ions in the system. The concentration of cRNA standard transcribed in vitro was accurately measured by Nano Drop nucleic acid quantitative analyzer, and the copy number was calculated. The cRNA standard was continuously diluted with RNase free H2O in tenfold gradient. The sensitivity of the RT-RAA was evaluated by real-time fluorescence detection, using the diluted cRNA standard ranging from 10^6 to 10^3 copies/μL.

Design of H5 RT-RAA primers and exo-probes

To detect H5 subtype AIV, a total of 4636 available HA gene sequences of H5 subtype AIV obtained from GenBank database were aligned, which contained the HA gene sequence of all currently circulating branches of H5 subtype AIV, and highly conserved regions were subsequently identified with Molecular Evolutionary Genetics Analysis (MEGA) software 6.0 [11] for the design the gene-specific primers and probes. Primers were designed using OLIGO 7 software [12] and showed no major nonspecific sequence similarities by BLAST analysis. Three H5 forward primers and five reverse primers were designed to select the best primers and probes in combination. The appropriate primers and probe combination were selected by sequence analysis, which were shown in Table 1. The 30th base at the 5’ end of the probe was labeled with the FAM fluorophore. The 30th base was connected to the abasic site tetrahydrofuran (THF). The 31st base was labeled with the BHQ1 quencher, and the 3’ end was modified by 3’ block. All the primers and probes were synthesized by Sangon Biotech.
RT-RAA for detection of H5 subtype AIV

The appropriate primers and exo-probes were screened, and the RT-RAA reaction was performed with an RT exo kit in 50 μL reaction mixture including the necessary enzymes and reagents for RT and DNA amplification in lyophilized pellets (Jiangsu Qitian Bio-Tech Co. Ltd.). The reaction mixture contained the following: 2 μL RNA template, 25 μL rehydration buffer, 15.7 μL deionized distilled water, 2.5 μL magnesium acetate, 2.1 μL each primer (10 μM) and 0.6 μL target-specific RT-RAA exo-probe (the probe concentration is 40 ng/μL). For amplification, the tubes were then transferred to a tube holder in an RT-RAA fluorescence detection device (QT-RAA-F7200; Jiangsu Qitian Bio-Tech Co. Ltd.) set at 39 °C for 20 min. Each run included nuclease-free water as a negative control. The results of the RT-RAA interpretation criteria were determined by the slope of amplification curve of the RT-RAA fluorescence detection device (QT-RAA-F7200), the test result is positive when the slope k value was greater than or equal to 20, in addition, the test result is negative.

Analytical sensitivity and analytical specificity of RT-RAA

Recombination plasmid of HA gene of H5 subtype has been constructed using the pEASY™-T5 Zero Cloning Kit. The plasmid was cleaved using a suitable restriction enzyme. The digested plasmid was purified with a DNA recovery kit, the concentration was measured with a Nano Drop nucleic acid quantifier, and then an appropriate amount was taken as a template according to its concentration, and the RNA Production System-T7 kit was used for in vitro transcription. The in vitro transcription products were extracted and purified with the Rneasy Mini Kit to remove impurities and various ions in the system. Serial tenfold dilutions of known concentrations of in vitro transcribed cRNA standards with RNase Free H2O.

A dilution range of H5 subtype HA plasmid standard was used to select the appropriate RAA primers and exo-probe combination. The selected appropriate primers and exo-probes were verified by H5 RT-RAA analysis. To determine the analytical sensitivity of the H5 RT-RAA assay, we detected the H5 subtype AIV molecular RNA standard over a dilution range of 10^7–10^1 copies/μL under the optimal RT-RAA conditions, with eight replicates for each dilution.

The analytical specificity assay of the RT-RAA for H5 subtype AIV was evaluated using four H5-positive AIVs (one strain in clade 2.3.2.1, one in clade 7, and 55 in clade 2.3.4.4), 10 other subtype AIVs (H1N2, H3N2, H4N2, H6N2, H7N3, H7N9, H9N2, H10N7, H11N9), two NDVs and two IBVs. These viruses are the main respiratory viruses affecting birds and were previously identified by our laboratory. The details of all the viruses tested are listed in Table 2.

| Table 1 | Primer and probe sequences used for RT-RAA and RT-qPCR assays |
|---------|---------------------------------------------------------------|
| Primer  | Sequence (5’–3’) | Size (bp) | Gene | Source |
| H5-F    | CAGTTTGACGGCTGTTGGAAGGAATTAAAYA | 32 | HA | This study |
| H5-R    | CTGTCRATACGGTTCGATGACATG | 32 | HA | This study |
| H5-P    | CTATGCTGTGCTCTTATATGCTGAACT/i6FAMdT//THF//iBHQ1dT//GGTCTCTGGAAATC[3-spacer] | 47 | HA | This study |
| H5 + 1456 | ACGTATGACTACCAACATAACTCAG | 25 | HA | [13] |
| H5-1685 | AGACGACGATCTACATG | 20 | HA | [13] |
| H5 + 1637 | FAM-TCAACAGTTGCGAGTGATCCATGCA-TAMRA | 24 | HA | [13] |

| Table 2 | Some information about the samples used for analytical sensitivity and analytical specificity assay in the study |
|---------|---------------------------------------------------------------|
| Sample  | Virus | HA subtype(clade) | H5 RT-RAA assay | H5 RT-qPCR assay |
| K144    | AIV   | H5N1(2.3.2.1)     | +               | +               |
| QD1     | AIV   | H5N2(7)           | +               | +               |
| G2324   | AIV   | H5N6(2.3.4.4)     | +               | +               |
| G2084   | AIV   | H5N6(2.3.4.4)     | +               | +               |
| Q221    | AIV   | H1N2             | –               | –               |
| X1330   | AIV   | H3N2             | –               | –               |
| P174    | AIV   | H4N2             | –               | –               |
| A1267   | AIV   | H6N2             | –               | –               |
| H7N3    | AIV   | H7N3             | –               | –               |
| 1605    | AIV   | H7N9             | –               | –               |
| X169    | AIV   | H9N2             | –               | –               |
| H9      | AIV   | H9N2             | –               | –               |
| T55     | AIV   | H10N2            | –               | –               |
| S82     | AIV   | H11N2            | –               | –               |
| ND      | NDV   | /                | –               | –               |
| JS1816  | NDV   | /                | –               | –               |
| M1      | IBV   | /                | –               | –               |
| HS2     | IBV   | /                | –               | –               |
Detection and evaluation of clinical samples by H5 RT-RAA

To test the efficacy of the method for clinical application, 420 throat and cloacal swabs and tissue samples whose hosts contained chickens, ducks, geese, pigeons and wild birds were collected from 29 live poultry markets and farms in six provinces. Throat and cloacal swabs were immediately placed in 1 mL antibiotic-containing PBS as described above and then stored at −80 °C until total nucleic acids were extracted with the above viral RNA extraction kit [14]. The tissue samples were sheared and ground after adding PBS, and their supernatants are extracted after centrifugation for nucleic acid extraction. We evaluated the performance of the RT-RAA assay in 420 avian clinical samples and compared it with the published RT-qPCR method for H5 subtype AIV. The primers and probe of the RT-qPCR assay are listed in Table 1 [13]. The RT-qPCR assay was conducted with the One Step PrimeScript RT-PCR Kit (Takara).

Statistical analysis

To determine the RT-RAA detection limit, a probit analysis was performed at a 95% confidence interval (CI), and the κ and p values of RT-qPCR and RT-RAA were calculated [15]. In addition, we calculated the sensitivity and specificity of RT-qPCR and RT-RAA for detection in clinical samples of poultry. All statistical analyses were performed in SPSS 21.0 (IBM).

Results

Analytical sensitivity of RT-RAA

The sequences of the appropriate H5 RT-RAA primers and exo-probe are listed in Table 1. The detection results of RT-RAA sensitivity assay are shown in Fig. 1. The primer and probe combinations designed by the present invention have a cRNA concentration of 10^6 copies/μL, 10^5 copies/μL, 10^4 copies/μL, 10^3 copies/μL, 10^2 copies/μL, 10^1 copies/μL, a fluorescence amplification curve appears. Therefore, the detection limit of H5 RT-RAA assay was 10^2 cRNA copies/μL (Table 3, Fig. 1).

Table 3 Assay data used for probit analysis to calculate the detection limits of H5 RT-RAA

| Copies/μL | No. of positive samples/No. of samples tested by the RT-RAA assays for detection of H5 subtype AIV |
|-----------|------------------------------------------------------------------------------------------------------------------|
| 10^6      | 8/8                                                                                                               |
| 10^5      | 8/8                                                                                                               |
| 10^4      | 8/8                                                                                                               |
| 10^3      | 5/8                                                                                                               |
| 10^2      | 2/8                                                                                                               |
| 10^1      | 0/8                                                                                                               |

* Each dilution was tested in a total of eight replicates

Fig. 1 Analytical sensitivity of the H5 RT-RAA assay. A dilution range from 10^6 to 10^1 copies/μL of H5 subtype AIV cRNA was used to evaluate the detection limit of H5 RT-RAA assay. Negative represents negative control. 1–6: 10^6 copies/μL–10^1 copies/μL, 7: negative control
Analytical specificity of RT-RAA

The test group corresponding to the H5 subtype AIV RNA template showed standard fluorescence detection curves, and the other virus test groups and negative control groups did not show amplification curves. Thus, the RT-RAA assay did not crossreact with other subtype AIVs, NDVs and IBVs and demonstrated high specificity for the detection of H5 subtype AIV (Fig. 2).

Evaluation of the RT-RAA for clinical samples

The 420 conserved avian clinical samples collected from live poultry markets were tested by RT-RAA assay and compared with RT-qPCR. A threshold cycle value of 36 was used as the cut-off for a positive result in RT-qPCR. RT-qPCR detected 73 of the 420 samples as positive (17.38%), while RT-RAA correctly identified and differentiated 71 positive samples, with a sensitivity of 97.26% (95% CI, 89.56–99.52%) and 100% specificity (95% CI, 98.64–100%) (Table 4). The κ value for RT-RAA and RT-qPCR was 0.983 (p < 0.001).

Discussion

The frequent variation of AIV antigen increases the difficulty of AIV detection. Among all the AIV subtypes, H5 subtype HPAIV often leads to high morbidity and mortality in poultry. Nowadays, the H5 subtype AIV is widely prevalent and of significant concern to the poultry industry and public health in China. Therefore, early detection of AIV and H5 subtype is necessary in the surveillance and control of AIV outbreaks. Until now, there have been many methods used in AIV detection, such as gold immunochromatographic assay [16], microarray [17], immunosensor [18], immunofluorescence [19] and enzyme-linked immunosorbent assay [20]. However, many methods require complex and costly devices and are difficult to perform in the field. So far, many detection methods have been used for rapid detection of the AIV in the field, such as nucleic acid sequence amplification [21], RT-LAMP and RT-RPA [22]. Moreover, as the mutation rate of the H5 subtype AIV accelerates, the previous detection methods may not meet the actual detection needs. Some of them fail to subtype AIV, while others

![Fig. 2 Analytical specificity of the H5 RT-RAA assay. Detection signals were recorded by real-time fluorescence RT-RAA with four samples including H5 subtype AIVs (H5N1, H5N2 and H5N6), while no signals were detected from the 14 samples including other subtype AIVs, NDVs, IBVs and negative controls. 1: K144 (H5N1); 2: QD1 (H5N2); 3: G2324 (H5N6); 4: G2084 (H5N6)](image)

| Table 4 Detection of H5 subtype AIV in avian clinical samples |
|---------------------------------------------------------------|
| **RT-qPCR** | **Total** | **Kappa (κ)** | **p value of kappa** | **Sensitivity% (95% CI)** | **Specificity% (95% CI)** |
| Positive | 71 | 2 | 73 | 0.983 | <0.001 | 97.26 (89.56–99.52) | 100 (98.64–100) |
| Negative | 0 | 347 | 347 | | | | |
| Total | 71 | 349 | 420 | | | | |
cannot be applied in early diagnosis owing to inadequate sensitivity [23], so a rapid diagnostic method capable of detecting all avian influenza epidemic strains is needed.

RAA is a novel isothermal amplification and detection assay requiring only 20 min to complete, while RT-qPCR and conventional PCR need longer time. In PCR, thermocycling is required for double-stranded DNA separation, primer binding and amplification. RT-RAA could produce a positive signal in as little as 4 min and be completed in 30 min, at half the cost of RT-qPCR. RAA can be carried out using a portable device with no complicated processes, while the instruments for RT-qPCR and conventional PCR are more expensive. RAA can also use reverse transcriptase and a fluorescent probe system to detect RNA amplicons in real time [17]. Until now, RAA has widely been used for detecting human pathogens, including Salmonella, Listeria monocytogenes, respiratory syncytial virus, coxsackievirus, MERS-CoV, SARS-CoV-2, Mycobacterium Tuberculosis, hepatitis B virus and Schistosoma japonicum-specific gene fragments and there have been no previous reports of the use of RAA for detecting H5 subtype AIV [24–31]. RAA has also been used in the detection of Staphylococcus aureus in milk, African swine fever virus and animal derived ingredients in exported food, such as pigs, cows and chickens [32–36]. However, compared with fluorescence and ordinary PCR, the cost of RT-RAA is relatively high at present.

In this study, an H5 RT-RAA assay was created, which indicated the potential value of our method for early detection and rapid diagnosis of the H5 subtype AIV. In the detection of experimental and clinical samples, this method showed higher sensitivity along with high efficiency. RT-qPCR detected 73 of the 420 samples as positive (17.38%), while RT-RAA correctly identified 71 positive samples, with a sensitivity of 97.26% (95% CI, 89.56–99.52%) and 100% specificity (95% CI, 98.64–100%) (Table 4). The κ value for RT-RAA and RT-qPCR was 0.983 (p < 0.001). Moreover, the clinical swab samples detected as positive in RT-RAA were also certified as positive by PCR detection and sequencing. According to the primer/exo-probe design method and the sample detection results, the RT-RAA method was considered to be suitable for detecting almost all H5 subtypes. To the best of our knowledge, this is the first RT-RAA method for the detection of multiple H5 clades in AIV.

In conclusion, the RT-RAA method established in our study quickly and accurately identified H5 subtype AIV, including the current epidemic strains, which meets the need for H5 subtype AIV testing. This RT-RAA is expected to detect emerging H5 subtype AIV rapidly and provide a powerful and valuable tool for the control of H5 subtype AIV.

Author contributions
Conceptualization: SW, YL, FZ, KW, HL, BH. Data curation: SW, KW. Formal analysis: SW, YL, FZ. Funding acquisition: KW, BH. Investigation: SW, YL, FZ, NJ, QZ, GH. Methodology: SW, YL, FZ, KW. Project administration: KW. Resources: GH, QZ, LJ, JY, CY, LY. Supervision: KW. Validation: SW, YL, FZ, NJ. Visualization: SW, YL, FZ. Writing-original draft preparation: SW, YL, FZ. Writing-review and editing: SW, YL, FZ, KW, HL, BH. All authors read and approved the final manuscript.

Funding
This work was supported by the National Key Research and Development Program of China (2017YFC120050).

Availability of data and materials
All data generated or analyzed during this study are included in this published article [and its supplementary information files].

Declarations
Ethics approval and consent to participate
Not applicable.

Consent for publication
Not applicable.

Competing interests
The authors declare that they have no competing interests.

Author details
1 China Animal Health and Epidemiology Center, 369 Nanjing Road, Qingdao, Shandong Province, China. 2 Yanbian University, Agricultural College, Yanji, Jilin, China. 3 Shandong Vocational Animal Science and Veterinary College, Weifang, China. 4 Key Laboratory of Animal Biosafety Risk Prevention and Control (South), Ministry of Agriculture and Rural Affairs, Qingdao, China.

Received: 7 December 2021 Accepted: 6 May 2022
Published online: 30 July 2022

References
1. Dadonaite B, Gilbertson B, Knight ML, Trifkovic S, Rockman S, Laederach A, Brown LE, Fodor E, Bauer DL. The structure of the influenza A virus genome. Nat Microbiol. 2019;4(11):1781–9.
2. Nakajima K. Influenza virus genome structure and encoded proteins. Nihon Rinsho Jpn J Clin Med. 1997;55(10):2542–6.
3. Peiris JSM, de Jong MD, Guan Y. Avian influenza virus (H5N1): a threat to human health. Clin Microbiol Rev. 2007;20:243–67.
4. Drew TW. OIE manual of diagnostic test and vaccine for terrestrial animal. Paris: Avian Influenza; 2008. p. 465–81.
5. Capua I, Alenxander DJ. Avian influenza and Newcastle disease: a field and laboratory manual. Springer; 2009.
6. Sidoti F, Rizzo F, Costa C, Astegiano S, Curtoni A, Mandola ML, Rossana C, Massimiliano B. Development of real time RT-PCR assays for detection of type A influenza virus and for subtyping of Avian H5 and H7 hemagglutinin subtypes. Mol Biotechnol. 2010;44(1):41.
7. Notomi T, Okayama H, Masubuchi H, Yonekawa T, Watanabe K, Amino N, et al. Loop-mediated isothermal amplification of DNA. Nucleic Acids Res. 2000;28(12):E63.
8. Lutz S, Weber P, Focke M, Faltin B, Hoffmann J, Muller C, Mark D, Roth G, Munday P, Armes N, Piepenburg O, Zengerle R, Stetten FV. Microfluidic lab-on-a-foil for nucleic acid analysis based on isothermal recombinase polymerase amplification (RPA). Lab Chip. 2010;10:887–93.
9. Walker GT, Fraiser MS, Schram JL, Little MC, Nadeau JG, Malinowski DP. Strand displacement amplification—an isothermal, in vitro DNA amplification technique. Nucleic Acids Res. 1992;20:1691–6.
10. Bei L, Cheng HR, Yan QF, Huang ZJ, Shen GF, Zhang ZF, Li YN, Deng ZX, Lin M, Cheng Q. Recombinate-aid amplification, a novel technology of in vitro rapid nucleic acid amplification. Sci Sinica. 2010;40:983–8.
11. Tamura K, Stecher G, Peterson D, Filipski A, Kumar S. MEGA6: molecular evolutionary genetics analysis version 6.0. Mol Biol Evol. 2013;30:2725–9.
12. Rychlik W. Oligo 7 primer analysis software. Methods Mol Biol. 2007;402:35–60.
13. Stackman R, Senne DA, Myers TJ, Bulage LL, Suarez DL. Development of a real-time reverse transcriptase PCR assay for type A influenza virus and the avian H5 and H7 hemagglutinin subtypes. J Clin Microbiol. 2002;40(9):3256–60.
14. Cui D, Zhao D, Xie Y, Yang X, Hua Z, Zheng S, Fu C, Chen Y. Simultaneous detection of influenza A subtypes of H3N2 virus, pandemic (H1N1) 2009 virus and reassortant avian H7N9 virus in humans by multiplex one-step real-time RT-PCR assay. Springerplus. 2016;5:2045.
15. Chen C, Li XN, Li GX, Zhao L, Duan SX, Yan TF, Feng ZS, Ma XJ. Use of a rapid reverse-transcription recombinase aided amplification assay for respiratory syncytial virus detection. Diagn Microbiol Infect Dis. 2018;90:90–5.
16. Peng D, Hu S, Hua Y, Xiao L, Wang X, Bi DR. Comparison of a new gold-immunochromatographic assay for the detection of antibodies against avian influenza virus with hemagglutination inhibition and agar gel immunodiffusion assays. Vet Immunol Immunopathol. 2007;117(1–2):17–25.
17. Gall A, Hoffmann B, Harder T, Grund C, Höper D, Beer M. Design and validation of a microarray for detection, hemagglutinin subtyping, and pathotyping of avian influenza viruses. J Clin Microbiol. 2009;47(2):327–34.
18. Wang R, Wang Y, Lasster K, Li Y, Hargis B, Tung S, Berghman L, Botte J. Interdigitated array microelectrode based impedimcesensor for detection of avian influenza virus H5N1. Talanta. 2009;75(2):159–64.
19. Zhao GY, Shu-Chun WU, Wang LN, Chen GB, Cui SJ. A disposable amperometric enzyme immunosensor for rapid detection of avian influenza virus. Chin J Anim Vet Sci. 2008;39(10):1442–8.
20. Song JL, Zhang WD, Wang JP, Zuo-Sheng LI, Feng ZL, Yuan-Yuan Hu, Guo SH, Zhang YG, Fan QS, Song XL. Development of type and subtype-specific immunofluorescence techniques for detection of avian influenza virus. J Funan Univ. 2008;30(5):526–30.
21. Collins RA, Ko LS, Fung KY, Chan KY, Xing J, Lau LT. Rapid and sensitive detection of avian influenza virus subtype H7 using NASBA. Biochem Biophys Res Commun. 2003;303:507–15.
22. Wahed AEA, Weidmann M, Huffert FT. Diagnostics-in-a-Suitcase. Development of a portable and rapid assay for the detection of the emerging avian influenza A (H7N9) virus. J Clin Virol. 2015;69:16–21.
23. Zhou EM, Cantin MF. Evaluation of a competitive ELISA for detection of antibodies against avian influenza virus nucleoprotein. Avian Dis. 1998;42(3):517–22.
24. Yan TF, Li XN, Wang L, Chen C, Duan SX, Qi JJ, Li LX, Ma XJ. Development of a reverse transcription recombinase-aided amplification assay for the detection of coxsackievirus A10 and coxsackievirus A6 RNA. Adv Virol. 2018;2018:163:1455–61.
25. Qi J, Li X, Zhang Y, Shen X, Song G, Pan J, Fan T, Wang RH, Li LX, Ma XJ. Development of a duplex reverse transcription recombinase-aided amplification assay for respiratory syncytial virus incorporating an internal control. Adv Virol. 2019;164:1843–50.
26. Shen XX, Qiu FZ, She LP, Yan TF, Zhao MC, Qi JJ, Chen C, Zhao L, Wang L, Feng ZS, Ma XJ. A rapid and sensitive recombinase aided amplification assay to detect hepatitis B virus without DNA extraction. BMC Infect Dis. 2019;19:229.
27. Song Z, Ting L, Kun Y, Wei L, Jian-Feng Z, Li-Chuan G, Yan-Hong L, Yang D, Qing-Jie Y, Hai-Tao Y. Establishment of a recombinase-aided isothermal amplification technique to detect Schistosoma japonicum specific gene fragments. Chin J Schistosomiasis Control. 2018;20:273–7.
28. Zhou DG, Luo J, Chen JH, Yu XJ. Establishment and application of rt-rra rapid detection method for Middle East respiratory syndrome coronavi- rus. J Virol. 2018;34(01):45–51.
29. Qian CZ, Liu CY, Liu BY. Establishment and evaluation of New Coronavirus nucleic acid fluorescence RT-RAA detection method. J Anhui Med Univ. 2021;56(06):980–5.
30. Li XN. Detection of mutations in EV71, c1va16 and rp08 genes of Myco- bacterium tuberculosis by recombinant enzyme mediated amplification. China Center for Disease Control and prevention. 2020.
31. Huang XX, He YP, Yuan CG, Guo DH, Zhao Y, Sun XH, Zeng J, Zheng QY. Detection of Listeria monocytogenes in food by fluorescent recombinase mediated isothermal amplification. Bull Microbiol. 2021;48(12):4989–5000.
32. Zhao GJ. Study on rapid detection method of Salmonella and Staphylo- coccus aureus in dairy products and design of food microbiology labora- tory. Nanchang University. 2020.
33. Zhao KY, Zeng DX, Hu YX, Qian BX, Xue F, Qian YJ, Wu XD, Dai JJ. Establishment of real-time fluorescence RAA detection method for African classical swine fever virus. Chin Vet Sci. 2021;51(01):1–8.
34. Miao L, Wang JC, Wang XJ, Han JX, Shen H, Xu SF, Zheng QY, Wang L, Luo BZ, Wu S, Lin H, Kong FD. Rapid detection of pig derived components in export food, recombinant enzyme mediated strand replacement nucleic acid amplification (RAA). Zhengzhou Customs Technology Center, 2018.
35. Miao L, Zong K, Wang P, Shen H, Zhang JK, Wu S, Luo BZ, Kong FD, Wang L, Zheng QY, Xu SF. Rapid detection of sheep derived components in export food, recombinant enzyme mediated strand replacement nucleic acid amplification (RAA). Zhengzhou Customs Technology Center, 2018.
36. Miao L, Wang P, Wang JC, Zong K, Deng TT, Wang L, Luo BZ, Kong FD, Lin H, Zheng QY, Wu S. Rapid detection of chicken derived components in export food, recombinant enzyme mediated strand replacement nucleic acid amplification (RAA). Zhengzhou Customs Technology Center, 2018.

Publisher’s Note
Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.