Development and Evaluation of the Rapid and Sensitive RPA Assays for Specific Detection of Salmonella spp. in Food Samples

Liwei Zhao†, Jianchang Wang‡, Xiao Xia Sun‡, Jinfeng Wang‡, Zhimin Chen‡, Xiangdong Xu§, Mengyuan Dong†, Ya-nan Guo†, Yuanyuan Wang†, Pingping Chen†, Weijuan Gao†* and Yunyun Geng†*

1 Hebei Key Laboratory of Chinese Medicine Research on Cardiocerebrovascular Disease, Hebei University of Chinese Medicine, Shijiazhuang, China, 2 Food Microbiology and Animal Quarantine Laboratory, Technology Center of Shijiazhuang Customs, Shijiazhuang, China, 3 School of Public Health, Key Laboratory of Environment and Human Health, Hebei Medical University, Shijiazhuang, China

Salmonella spp. is among the main foodborne pathogens which cause serious foodborne diseases. An isothermal real-time recombinase polymerase amplification (RPA) and lateral flow strip detection (LFS RPA) were used to detect Salmonella spp. targeting the conserved sequence of invasion protein A (invA). The Real-time RPA was performed in a portable fluorescence scanner at 39°C for 20 min. The LFS RPA was performed in an incubator block at 39°C for 15 min, under the same condition that the amplifications could be inspected by the naked eyes on the LFS within 5 min. The detection limit of Salmonella spp. DNA using real-time RPA was $1.1 \times 10^1$ fg, which was the same with real-time PCR but 10 times higher than that of LFS RPA assay. Moreover, the practicality of discovering Salmonella spp. was validated with artificially contaminated lamb, chicken, and broccoli samples. The analyzing time dropped from 60 min to proximately 5–12 min on the basis of the real-time and LFS RPA assays compared with the real-time PCR assay. Real-time and LFS RPA assays’ results were equally reliable. There was no cross-reactivity with other pathogens in both assays. In addition, the assays had good stability. All of these helped to show that the developed RPA assays were simple, rapid, sensitive, credible, and could be a potential point-of-need (PON) test required mere resources.

Keywords: Salmonella, invA gene, real-time RPA, lateral flow strip (LFS), isothermal amplification

INTRODUCTION

Salmonella spp. is a Gram-negative bacterium belonging to the family of Enterobacteriaceae. Salmonella spp. is a major cause of the foodborne pathogen around the world (Nassib et al., 2003). It is widespread in nature and proliferates under ambient temperature with low nutritional demands (Li et al., 2013). Salmonella spp. infections attract much attention in public health especially in food safety. Salmonella spp. causes food poisoning, typhoid fever, gastrointestinal inflammation, and septicemia for both humans and animals (McGuinness et al., 2009; Rukambile et al., 2019).
Currently, there are over 80 million cases of foodborne salmonellosis in the world (Li et al., 2019). Additionally, reports revealed that outbreaks caused by *Salmonella* spp. were largely associated with animal derived products such as poultry, egg, and chicken, and contamination is common in retail raw meats (Nassib et al., 2003; Foley and Lynne, 2008; Foley et al., 2008). An accurate and fast diagnosis is needed in order to prevent *Salmonella* spp. infections.

*Salmonella* spp. is currently detected in foods primarily through traditional laboratory methods. These traditional laboratory methods are inconvenient, time-consuming and takes over 3 days to obtain the result following multiple analytical steps (Li et al., 2019; Hong et al., 2020). Moreover, the complexity of the samples had a great effect on the bacterial morphology colony. The cross-reactivity among bacteria in *Enterobacteriaceae* restricted the specificity and sensitivity of the test (Nassib et al., 2003; Li et al., 2019). Upgrading molecular diagnostics provides powerful means for detecting *Salmonella* spp. in light of sensitivity and specificity. Currently, many nucleic acid amplification-based assays have gained popularity such as polymerase chain reaction (PCR), real-time PCR, multiplex PCR, reverse transcriptase PCR (RT-PCR), and loop-mediated isothermal amplification (LAMP) (Hirose et al., 2002; Yeh et al., 2002; Techathuvanan et al., 2010; Domesle et al., 2018). Real-time PCR is extensively applied for the early field settings (Techathuvanan et al., 2011). PCR assays’ application is limited within the walls of the laboratories (Techathuvanan et al., 2011). In addition to the PCR assays, state-of-the-art isothermal amplification technologies such as LAMP, have been used for early and rapid detection of *Salmonella* spp. Simpler and more convenient techniques are required imperatively for the point-of-need (PON) diagnosis of *Salmonella* spp. in field conditions.

Recombinase polymerase amplification (RPA) acts as one of isothermal gene amplification techniques. RPA has the merit of amplification at a relatively low temperature (37–42°C) within 10–20 min (Piepenburg et al., 2006; Geng et al., 2018). The use of RPA-based methods has been proved to be a success in detecting pathogenic bacteria and viruses in clinical and food samples (Euler et al., 2012; Abd El Wahed et al., 2015; Wang et al., 2020). RPA-based methods have been designed to be a miniaturized diagnostic device that includes all the components for the RPA assay (Asiello and Baumann, 2011). RPA assay was a rapid, stable, and promising assay for the on-site detection. The objective of the study was to develop the real-time and LFS RPA assays using the exo probe and no probe combined with lateral flow strip respectively as a way of rapidly detecting *Salmonella* spp. in food samples.

### MATERIALS AND METHODS

#### Bacterial Strains and DNA Extraction

A total of 34 common pathogenic bacteria strains were used to validate the techniques employed in this study (Table 1). These pathogenic bacteria were purchased from the American Type Culture Collection (ATCC), China Center of Industrial Culture Collection (CICC), China Center for Medical Culture Collection (CMCC) or isolated in the lab. All strains were reserved in the lab. Stock cultures were stored at −80°C in 0.8 ml of Nutrient broth (Beijing Land Bridge Technology Co., Ltd., Beijing, China) and 0.2 ml of 80% glycerol. The DNA templates were extracted using the TIANamp Bacteria DNA Kit (Tiangen, Beijing, China). These DNA samples were stored at −20°C before the assays.

| Strain Name | Origin | Real-time RPA | LFS RPA | Real-time PCR |
|-------------|--------|---------------|---------|--------------|
| *Salmonella* CICC 22956 | + | + | + |
| *Salmonella*CMCC50786 | + | + | + |
| *Salmonella*CMCC50761 | + | + | + |
| *Salmonella*CMCC500359 | + | + | + |
| *Salmonella*Isolated by lab | + | + | + |
| *typhimurium* | | | | |
| *Salmonella enteritidis* Isolated by lab | + | + | + |
| *Salmonella* paratyphi Isolated by lab | + | + | + |
| *Salmonella enterica* Isolated by lab | + | + | + |
| *Enterobacter sakazakii* ATCC 29644 | − | − | − |
| *Staphylococcus aureus* ATCC 6538 | − | − | − |
| *Campylobacter jejuni* ATCC 33291 | − | − | − |
| *Vibrio parahaemolyticus* ATCC 9027 | − | − | − |
| *Pseudomonas aeruginosa* ATCC 27562 | − | − | − |
| *Vibrio vulnificus* ATCC 9027 | − | − | − |
| *Pseudomonas aeruginosa* CMCC 63301 | − | − | − |
| *Bacillus cereus* ATCC 19114 | − | − | − |
| *Listeria monocytogenes* ATCC 35659 | − | − | − |
| *Proteus mirabilis* ATCC 13048 | − | − | − |
| *Enterobacter aerogenes* ATCC 51592 | − | − | − |
| *Shigella sonnei* ATCC 14580 | − | − | − |
| *Proteus pneumonia* CMCC 49027 | − | − | − |
| *Shigella boydii* CMCC 51250 | − | − | − |
| *Shigella flexneri* CMCC51105 | − | − | − |
| *Shigella flexneri* CICC 21530 | − | − | − |
| *Escherichia coli* CICC 22956 + + + |
| *CMCC50761* CMCC50786 | + | + | + |
| *Salmonella dublin* CMCC50786 | + | + | + |
| *Salmonella* CICC 22956 | + | + | + |
| *CMCC51105* CICC 22956 | + | + | + |
| *CMCC50786* CMCC50786 | + | + | + |
| *Salmonella enteritidis* Isolated by lab | + | + | + |
| *Salmonella paratyphi* Isolated by lab | + | + | + |
| *Salmonella enterica* Isolated by lab | + | + | + |
| *Enterobacter sakazakii* ATCC 29644 | − | − | − |
| *Staphylococcus aureus* ATCC 6538 | − | − | − |
| *Campylobacter jejuni* ATCC 33291 | − | − | − |
| *Vibrio parahaemolyticus* ATCC 9027 | − | − | − |
| *Pseudomonas aeruginosa* ATCC 27562 | − | − | − |
| *Vibrio vulnificus* ATCC 9027 | − | − | − |
| *Pseudomonas aeruginosa* CMCC 63301 | − | − | − |
| *Bacillus cereus* ATCC 19114 | − | − | − |
| *Listeria monocytogenes* ATCC 35659 | − | − | − |
| *Proteus mirabilis* ATCC 13048 | − | − | − |
| *Enterobacter aerogenes* ATCC 51592 | − | − | − |
| *Shigella sonnei* ATCC 14580 | − | − | − |
| *Proteus pneumonia* CMCC 49027 | − | − | − |
| *Shigella boydii* CMCC 51250 | − | − | − |
| *Shigella flexneri* CMCC51105 | − | − | − |
| *Shigella flexneri* CICC 21530 | − | − | − |
| *Escherichia coli* CICC 21530 | − | − | − |
| *O157:H7* Mannheimia | Isolated by lab | − | − |
| *haemolytica* Eisenia | − | − | − |
| *Enterobacter cloacae* Isolated by lab | − | − | − |
| *Citrobacter freundii* Isolated by lab | − | − | − |
| *Streptococcus* Isolated by lab | − | − | − |
| *Bacillus subtilis* Isolated by lab | − | − | − |
| *Pseudomonas* aeruginosa | − | − | − |
| *Staphylococcus* pasteurianus | Isolated by lab | − | − |
| *Escherichia coli* Isolated by lab | − | − | − |

1. ATCC, American Type Culture Collection; CICC, China Center of Industrial Culture Collection; CMCC, China Center for Medical Culture Collection.
2. +, positive results; −, negative results.

### TABLE 1 | Bacterial strains and analytical specificity results for real-time RPA and LFS RPA assays.
RPA Primers and Probe

Nucleotide sequence data for Salmonella spp. strains from GenBank were aligned to identify conserved regions. Based on the reference sequences of different Salmonella spp. genotypes (accession numbers: AY594273, AY594271, DQ644632, DQ644633, EU348367, EU348368, JF951188, and JN982040), three pairs of primers targeting the conserved region of invA were designed (Rahn et al., 1992; Gonzalez-Escalona et al., 2009).

RPA, Real-time RPA primers, and probes were then selected through testing the combination that yielded the highest sensitivity (Table 2). Primers and exo probes were synthesized by Sangon (Sangon, Shanghai, China).

Real-Time RPA Assay

Real-time RPA was accomplished in the tube with 50 μl reaction volume, including 40.9 μl of Buffer A (rehydration buffer), 2.0 μl of each RPA primers (Sa-exo-F and Sa-exo-R, 10 μmol/L), 0.6 μl of exo probe (Sa-exo-P, 10 μmol/L), and 2.5 μl of Buffer B (magnesium acetate, 280 mmol/L). Furthermore, 1 μl of genomic DNA was used for the specificity and sensitivity analysis, or 2 μl of sample DNA was used for the clinical sample diagnosis. In the process, the Genie III scanner device (OptiGene Limited, West Sussex, UK) and TwistAmpTM exo kit (TwistDX, Cambridge, UK) were applied.

LFS RPA Assay

Moreover, the LFS RPA assay was performed according to the given instructions. The commercial TwistAmp™ nfo kit (TwistDX, Cambridge, UK) was used in the LFS RPA. The reactions were performed in a 50 μl volume with 29.5 μl of rehydration buffer with 2.5 μl of magnesium acetate (280 mM) included. Other components contained 420 nM RPA primer, 120 nM exo probe, and 1 μl of bacterial genomic DNA or 5 μl of sample DNA. The assay was performed in an incubator block at 39°C for 15 min and the lateral flow strips (Ustra Biotec GmbH, Germany) were employed to discover the RPA amplicons dual-labeled with FAM and biotin. Testing samples were considered positive when both the test line and the control line were visible. The testing sample was considered negative when the control line was visible. However, the sample was considered invalid when the control line was invisible.

Real-Time PCR

Real-time PCR was performed on the ABI 7500 instrument in which premix Ex Taq™ (Takara Co., Ltd., Dalian, China) was employed (Geng et al., 2019). The reaction was performed as follows: 95°C for 30 s, followed by 35 cycles of 95°C for 10 s and then 60°C for 34 s. The sequences of the primers and probes used for real-time PCR were listed in Table 2. The reporter and fluorescence quencher were marked with 6-FAM (6-CarboxyFluorescein) and BHQ1 (Black Hole Quencher 1) respectively.

Analytical Specificity and Analytical Sensitivity Analysis

For the food security, the real-time RPA and LFS RPA assays were evaluated to amplify the nucleic acid of some important pathogens. Five independent reactions were performed.

The genomic DNA of Salmonella spp. varying from 1.1 × 10^8 to 1.1 × 10^9 fg was diluted in nuclelease-free water for the analytical sensitivity analysis of the RPA. One microliter of each DNA dilution was amplified by both RPA assays to determine the limit of detection (LOD). The culture of Salmonella spp. was diluted in sterile water (ranging from 1.4 × 10^8 to 1.4 × 10^9 CFU) and counted by plate counting in 37°C overnight. The sensitivity of the real-time RPA and LFS RPA method were assessed with Salmonella spp. in pure culture. The analytical sensitivity analysis was repeated for five times.

Validation With Artificially Contaminated Samples

The pure colony of Salmonella spp. was picked into a tube containing 1 ml sterile saline. The solution was vortexed for 30 s and the turbidity was measured to 1.00 using a turbidimeter. Sterile saline was used for 10-fold gradient dilution until it attained 10^-8 dilution. With 10^-5, 10^-6, and 10^-7 diluents of 200 μl on the chromogenic medium of Salmonella spp., the initial concentration of the pure culture bacteria was calculated using three parallels.

Commercially available chicken/lamb/broccoli were purchased from a local supermarket free of Salmonella spp. to assess the potential use and suitability of the developed RPA assays. Testing of the samples was done according to the Chinese national standard (GB 4789.4-2016). A total of 4, 14, and 59 CFU/25g of Salmonella spp. with chicken, lamb, and broccoli respectively, were added into a sterile stomaching bag containing 225 ml Nutrient broth. These samples were mixed well to get homogenous samples and incubated for 6 or 8 h at 37°C to increase the bacterial concentrations to attain detectable levels. The Bacterial genomic DNA extraction, the real-time RPA, and

| Table 2 | Primer and probe sequences for Salmonella spp. real-time RPA, RPA, real-time RPA and LFS RPA assays. |
|---------|-------------------------------------------------------------------------------------------------|
| Method  | Name     | Sequence 5'-3' Amplication                                      | Size(bp) |
|---------|----------|----------------------------------------------------------------|----------|
| Real-time RPA | RPA-FP  | GTCATTCGATTACCTAATGCTGTGATTTCCT                                    | 200       |
| LFS-RPA | RPA-RP   | GGCAGCGCTCTCAAGATGTAAGAAGAAGGTG                                       |          |
|         | exo Probe| GTAGCTGCGATATGGTCTTTATGTGCGCTGT-(FAM-dT)-THF-(BHQ1-dT)-ACATGGCACAATCC-C3-spacer |          |
|         | nfo Probe| FAM-GTACTGGCAGATTTGCTTTAGTGGTCGTT-THF-T-ACATGGCACAGATCC-C3-spacer   |          |
| Real-time PCR | PCR-FP  | GAAGCTGGCTGGCCGGAAGG                                                                 | 68        |
|         | PCR-RP   | GGAGCTGGCTGGCCGGAAGG                                                                 |          |
|         | Probe    | JOE-GGAGCTGGCTGGCCGGAAGG                                                                 |          |

1FP, Forward primer; RP, Reverse primer.
LFS RPA reactions were performed, and each experiment was repeated for no less than three times to attain results.

RESULTS

Analytical Specificity and Sensitivity of the Real-Time RPA and LFS-RPA Assay

The invA gene coding of the invasion protein of Salmonella spp. is the most used specific gene for the discovery of many different Salmonella spp. serotypes. The RPA primers and probes were designed according to the invA gene of Salmonella spp. in this study. Both RPA assays provide excellent results at 39°C within 20 min. This was faster than any other common nucleic acid amplification method. The analytical sensitivity of RPA methods was evaluated by employing Salmonella spp. genomic DNA and bacterial pure culture as templates from $1.1 \times 10^6$ to $1.1 \times 10^9$ fg and from $1.4 \times 10^7$ to $1.4 \times 10^9$ CFU. The data on the analytical sensitivity of RPA methods was presented in Figures 1A, 2A. The detection limit (LOD) of real-time RPA was $1.1 \times 10^1$ fg similar to that of real-time PCR (the data was not shown) (Figure 1A). The limit of detection of the LFS RPA method was $1.1 \times 10^2$ fg for genomic DNA (Figure 2A), 10-times lower compared with the real-time PCR. The LOD of the real-time RPA and LFS RPA was $1.4 \times 10^2$ CFU for bacteria in pure culture (shown in Figures 1B, 2B).

![Figure 1A](Image)

**FIGURE 1 |** Analytical Sensitivity of the real-time RPA assay. The LOD of the real-time RPA was $1.1 \times 10^1$ fg/μl of Salmonella spp. standard DNA and $1.4 \times 10^2$ CFU/ml for bacteria in pure culture. Fluorescence development over time using a dilution range of $1.1 \times 10^6$–$1.1 \times 10^9$ fg of Salmonella spp. genomic DNA. For (A): Curve 1, $1.1 \times 10^6$ fg; Curve 2, $1.1 \times 10^5$ fg; Curve 3, $1.1 \times 10^4$ fg; Curve 4, $1.0 \times 10^3$ fg; Curve 5, $1.1 \times 10^2$ fg; Curve 6, $1.1 \times 10^1$ fg; Curve 7, $1.1 \times 10^0$ fg; Curve 8, ddH2O. For (B): Curve 1, $1.4 \times 10^6$ CFU; Curve 2, $1.4 \times 10^5$ CFU; Curve 3, $1.4 \times 10^4$ CFU; Curve 4, $1.4 \times 10^3$ CFU; Curve 5, $1.4 \times 10^2$ CFU; Curve 6, $1.4 \times 10^1$ CFU; Curve 7, $1.4 \times 10^0$ CFU.
Regarding specificity, only amplification signal was observed at the control line with *Salmonella* spp. and no cross-detection of other pathogens were shown in both real-time RPA and LFS-RPA assays (Table 1). Five independent reactions were repeated and similar results were obtained. This manifests the high specificity of the assays.

**Evaluation With Artificially Contaminated Samples After Enrichment**

The diagnostic performance of the developed RPA assays was compared with other detection approaches and was shown in Table 3. This was done while detecting *Salmonella* spp. in artificially contaminated food samples. The National Standard

![Diagram](image1.png)  
**FIGURE 2** | Analytical sensitivity of the LFS RPA assay. The LOD of the LFS RPA method was 1.1 × 10² fg for genomic DNA and 1.4 × 10² CFU/ml for bacteria in pure culture. For (A): Sample 1, 1.1 × 10⁸ fg; Sample 2, 1.1 × 10⁷ fg; Sample 3, 1.1 × 10⁶ fg; Sample 4, 1.1 × 10⁵ fg; Sample 5, 1.1 × 10⁴ fg; Sample 6, 1.1 × 10³ fg; Sample 7, 1.1 × 10² fg; Sample 8, 1.1 × 10¹ fg. For (B): Sample 1, 1.4 × 10⁷ CFU; Sample 2, 1.4 × 10⁶ CFU; Sample 3, 1.4 × 10⁵ CFU; Sample 4, 1.4 × 10⁴ CFU; Sample 5, 1.4 × 10³ CFU; Sample 6, 1.4 × 10² CFU; Sample 7, 1.4 × 10¹ CFU; Sample 8, 1.4 × 10⁰ CFU.

**TABLE 3** | The comparison of reaction time of different methods in contaminated foods.

| Food samples | Spiked samples¹ (CFU/25g) | Enrichment time(h) | Real-time RPA² (min:ss) | LFS-RPA² (Ct) | Real-timePCR² (Ct) | GB4789.4-2016 Viable cell counts (CFU/g) |
|--------------|--------------------------|-------------------|------------------------|--------------|-------------------|--------------------------------------|
| **Lamb**     |                          |                   |                        |              |                   |                                       |
| 4            | 6                        | −                 | −                      | −            | −                 | 0                                    |
| 14           | 6                        | 12:18             | −                      | 33.24        | +                 | 4.7 × 10³                            |
| 59           | 6                        | 11:44             | 15(+)                 | 33.38        | +                 | 5.1 × 10⁴                            |
| 8            | 6                        | 12:04             | 15(+)                 | 33.78        | +                 | 4.6 × 10³                            |
| 8            | 10:15                    | 15(+)             | 30.58                  | +            | 7.8 × 10³         |
| **Broccoli** |                          |                   |                        |              |                   |                                       |
| 4            | 6                        | −                 | −                      | −            | −                 | 0                                    |
| 14           | 6                        | 10:09             | 15(+)                 | 30.02        | +                 | 6.2 × 10³                            |
| 59           | 6                        | 6:55              | 15(+)                 | 28.47        | +                 | 1.3 × 10³                            |
| 8            | 5:16                     | 15(+)             | 19.34                  | +            | 5.3 × 10³         |
| 8            | 6:33                     | 15(+)             | 25.46                  | +            | 1.8 × 10⁴         |
| 8            | 5:39                     | 15(+)             | 20.58                  | +            | 1.8 × 10⁵         |
| **Chicken**  |                          |                   |                        |              |                   |                                       |
| 4            | 6                        | −                 | −                      | −            | −                 | 0                                    |
| 14           | 6                        | 11:54             | 15(+)                 | 33.49        | +                 | 3.8 × 10⁴                            |
| 59           | 6                        | 12:06             | 15(+)                 | 34.01        | +                 | 6.3 × 10³                            |
| 8            | 6                        | 6:14              | 15(+)                 | 25.18        | +                 | 2.8 × 10⁵                            |
| 8            | 11:22                    | 15(+)             | 33.28                  | +            | 3.9 × 10⁴         |
| 8            | 5:54                     | 15(+)             | 22.39                  | +            | 3.7 × 10⁵         |

¹CFU, colony-forming units; ²Ct, Cycle threshold; ³+, detected; −, not detected.

Total bacterial count, 1.5 × 10⁵ CFU/g and coliform group, 3.1 × 10⁴ CFU/g in Lamb;
Total bacterial count, <1.0 × 10⁵ CFU/g and coliform group, <1.0 × 10⁴ CFU/g in Broccoli;
Total bacterial count, 3.7 × 10⁵ CFU/g and coliform group, <1.0 × 10⁴ CFU/g in Chicken.
GB 4789.4-2016 method was used as a reference to guarantee that the food samples were successfully contaminated. However, after 6-h enrichment of food samples contaminated with 4 CFU/25 g of *Salmonella* spp., no *Salmonella* spp. was discovered through any of the detection methods. This signaled no false-positive results from samples containing low levels of *Salmonella* spp. All contaminated food samples were detected and showed increasing values of CFU/25 g of spiked samples and enrichment time. However, lamb contaminated with 14 CFU/25 g of *Salmonella* spp. was different from the other contaminated food samples and was enriched for 6 h. Furthermore, a diagnostic agreement of 100% with real-time PCR and the traditional method was indicated in the developed real-time RPA and LFS RPA assays. Moreover, the speed of the RPA assays outstripped. The time required to attain positive results with real-time RPA and LFS RPA was 5–12 min and 15 min respectively. Real-time PCR with CT values at 19.34–34.01 required approximately 20–35 min. Therefore, the results demonstrated that, with the equal sensitivity, the real-time and LFS RPA assays was faster than the real-time PCR.

**DISCUSSION**

The disease induced by foodborne pathogens remains a major public health issue worldwide despite ongoing measurements to ensuring food safety. *Salmonella* spp. frequently leads to infections and worldwide outbreaks accounting for huge economic costs and life loss every year (Foley and Lynne, 2008). Rapid and reliable diagnostic techniques play an important part to efficiently detect *Salmonella* spp. from contaminated specimens and make appropriate measures for preventing and controlling the risk of *Salmonella* spp. infection as early as possible.

The real-time RPA and LFS RPA assays are good choices for detecting *Salmonella* spp. as demonstrated in this report. These assays are specific, sensitive, and simple to perform. In the specificity analysis, both the real-time RPA and LFS RPA only amplified the genomic DNA of *Salmonella* spp. used in the study. This indicated high specificity of these assays. However, other more *Salmonella* strains are needed to further examine the cross-reactivity of these RPA assays. The real-time RPA had equal sensitivity (limit of detection) as real-time PCR in this study. This was 10 times higher than the LFS assay. However, it is possible for a varied reaction mechanism and enzyme kinetics between the different methods. The reaction time of RPA assays was much shorter than real-time PCR. The diagnostic performances of the developed real-time RPA and LFS RPA assays has been further assessed. These assays were proved to be a success in the detection of the artificially contaminated food samples, and performed better than the real-time PCR in light of the detecting speed. However, a pre-enrichment step was necessary when the level of pathogen contamination was low. A similar ideal result was obtained using the direct water boiling method to extract the bacterial DNA as the template of RPA reaction. Direct boiling method was used to extract the *Salmonella* spp. genomic DNA as the template of the RPA reaction. The LFS strip were combined to facilitate the detection of *Salmonella* spp. at quarantine stations, ports, or the site of outbreak by the PRA assay based on nfo-probe.

RPA was first introduced in 2006 and represented an innovative DNA isothermal detecting technology beyond the reach of PCR or traditional culture-based methods (Piepenburg et al., 2006; James and Macdonald, 2015). RPAAs have successfully been practiced in the discovery of pathogenic bacteria (Hong et al., 2020), fungus (Ahmed et al., 2014), and viruses (Boyle et al., 2013). The reagents in RPA are available in lyophilized form for long-term storage and are conveniently transported even without cold-chain (Wang et al., 2020). Moreover, under the prerequisite that the testing results were visible, the real-time RPA assay was accomplished on the user-friendly PON (point of need) detection platform (Genie III) with battery power. The developed LFS RPA assay only needed a simple incubator block. Therefore, these two pieces of equipment were portable, lightweight, and less expensive than the equipment for LAMP/PCR. As other isothermal DNA amplification methods, Loop-mediated isothermal amplification (LAMP) and the cross-priming amplification assay (CPA) have been adopted for rapidly and sensitively detecting *Salmonella* spp. Both RPA assays have the merits of amplification at a relatively lower temperature and within shorter time than that of LAMP and CPA assays. Both RPA reactions could be done at 37–42°C within 10–20 min. However, the optimum time and temperature were approximately 60 min and above 60°C respectively which were required for LAMP and CPA (Domesle et al., 2018; Wang et al., 2018). Moreover, several reports have shown that RPA is tolerant to mismatches, background DNA, and most of PCR inhibitors (Daher et al., 2016; Li et al., 2019; Liu et al., 2019). All of these outstanding characteristics make the assays readily suitable for the field, PON (point-of-need), or diagnosis of infectious diseases with poor environmental resources.

In conclusion, the current study proved that, the developed RPA assays with high specificity and sensitivity was convenient, rapid, and reliable for *Salmonella* spp. detection. In addition, the simple devices and easy operation protocol helped to improve the efficiency of detection. Among the isothermal amplification techniques, real-time RPA and LFS RPA assays play an outstanding role in preventing and controlling of *Salmonella* spp. especially in the settings with limited resources.

**DATA AVAILABILITY STATEMENT**

The datasets presented in this study can be found in online repositories. The names of the repository/repositories and accession number(s) can be found in the article/supplementary material.

**AUTHOR CONTRIBUTIONS**

YG, JCW, and WG designed and conducted the experiment. LZ, XS, JFW, XX, MD, Y-nG, YW, and PC performed the experiments and analyzed the data. YG drafted the manuscript. All authors contributed to the article and approved the submitted version.
FUNDING
This work was supported by the program of Traditional Chinese Medicine Scientific Research foundation in Hebei Administration of Traditional Chinese Medicine (2020142, Hebei, China), the Project of Excellent Young Teacher Fundamental Research (YQ2019003) and Doctoral Foundation (BSZ2019009) of Hebei University of Chinese Medicine.

REFERENCES
Abd El Wahed, A., Weidmann, M., and Hufert, F. T. (2015). 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. 69, 16–21. doi: 10.1016/j.jcv.2015.05.004
Ahmed, A., van der Linden, H., and Hartskeerl, R. A. (2014). Development of a recombinase polymerase amplification assay for the detection of pathogenic Leptospira. Int. J. Environ. Res. Public Health 11, 4953–4964. doi: 10.3390/ijerph110504953
Asiello, P. J., and Baeumner, A. J. (2011). Miniaturized isothermal nucleic acid amplification, a review. Lab. Chip. 11, 1420–1430. doi: 10.1039/c0lc00666a
Boyle, D. S., Lehman, D. A., Lillis, L., Peterson, D., Singhal, M., Armes, N., et al. (2013). Rapid detection of HIV-1 proviral DNA for early infant diagnosis using recombinase polymerase amplification. mBio. 4, e00135-13. doi: 10.1128/mBio.00135-13
Daher, R. K., Stewart, G., Boissinot, M., and Bergeron, M. G. (2016). Recombinase Polymerase Amplification for Diagnostic Applications. Clin. Chem. 62, 947–958. doi: 10.1373/chlm.2015.245829
Domesle, K. J., Yang, Q., Hammack, T. S., and Ge, B. (2018). Validation of a Salmonella loop-mediated isothermal amplification assay in animal food. Int. J. Food Microbiol. 264, 63–76. doi: 10.1016/j.ijfoodmicro.2017.10.020
Euler, M., Wang, Y., Otto, P., Tomaso, H., Escudero, R., Anda, P., et al. (2012). Recombinase polymerase amplification assay for rapid detection of Franciscella tularensis. J. Clin. Microbiol. 50, 2234–2238. doi: 10.1128/JCM.00504-11
Foley, S. L., and Lynne, A. M. (2008). Salmonella challenges: prevalence and antimicrobial resistance. J. Anim. Sci. 86, E173–E187. doi: 10.2527/2007.00157
Foley, S. L., Lynne, A. M., and Nayak, R. (2008). Salmonella challenges: prevalence in swine and poultry and potential pathogenicity of such isolates. J. Anim. Sci. 86, E149–E162. doi: 10.2527/2007.00146
Geng, Y., Liu, G., Liu, L., Deng, Q., Zhao, L., Sun, X. X., et al. (2019). Real-time detection of Salmonella enterica serovar Indiana by recombinase polymerase amplification assay. J. Microbiol. Methods 111, 285–290. doi: 10.1016/j.mimet.2018.12.017
Gonzalez-Escalona, N., Hammack, T. S., Russell, M., Jacobson, A. P., De Jesus, A. J., and Baeumner, A. J. (2011). Miniaturized isothermal nucleic acid amplification assay for the detection of Salmonella enterica serovars Typhi and Paratyphi A. J. Clin. Microbiol. 49, 633–636. doi: 10.1128/jcm.0063-3633.2002
Hirose, K., Itoh, K., Nakajima, H., Kurazono, T., Yamaguchi, M., Moriya, K., et al. (2002). Selective amplification of tyr (rfbE),prt (rfbS), viaB, and mY gene sequence to detect the Salmonella enterica serovars Typhi and Paratyphi A. J. Clin. Microbiol. 40, 633–636. doi: 10.1128/JCM.04330-02
Hong, H., Sun, C., Wei, S., Sun, X., Mutukumira, A., and Wu, X. (2020). Development of a real-time recombinase polymerase amplification assay for rapid detection of Salmonella in powdered infant formula. Int. Dairy J. 102. doi: 10.1016/j.idairyj.2019.104579
James, A., and Macdonald, J. (2015). Recombinase polymerase amplification: Emergence as a critical molecular technology for rapid, low-resource diagnostics. Expert Rev. Mol. Diagn. 15, 1475–1489. doi: 10.1586/14737519.2015.1090877
Li, R., Lai, J., Wang, Y., Liu, S., Li, Y., Liu, K., et al. (2013). Prevalence and characterization of Salmonella species isolated from pigs, ducks and chickens in Sichuan Province, China. Int. J. Food Microbiol. 163, 14–18. doi: 10.1016/j.ijfoodmicro.2013.01.020
Li, J., Ma, B., Fang, J., Zhi, A., Chen, E., Xu, Y., et al. (2019). Recombinase Polymerase Amplification (RPA) Combined with Lateral Flow Immunoassay for Rapid Detection of Salmonella in Food. Foods. 9, 27. doi: 10.3390/foods9010027
Liu, L., Li, R., Zhang, R., Wang, J., An, Q., Han, Q., et al. (2019). Rapid and sensitive detection of Mycoplasma hyopneumoniae by recombinase polymerase amplification assay. J. Microbiol. Methods 159, 56–61. doi: 10.1016/j.mimet.2019.02.015
McGuinness, S., McCabe, E., O’Regan, E., Dolan, A., Duffy, G., Burgess, C., et al. (2009). Development and validation of a rapid real-time PCR based method for the specific detection of Salmonella on fresh meat. Meat Sci. 83, 553–562. doi: 10.1016/j.meatsci.2009.07.004
Nassab, T. A., El-Din, M. Z., and El-Sharoud, W. M. (2013). Assessment of the presence of Salmonella spp. in Egyptian dairy products using various detection media. Lett. Appl. Microbiol. 37, 403–409. doi: 10.1111/j.1472-765X.2003.01420.x
Piepenburg, O., Williams, C. H., Stemple, D. L., and McEwen, S. A., Clarke, R. C., Galan, J. E., Ginocchio, C., et al. (1992). Amplification of an invA gene sequence of Salmonella typhimurium by polymerase chain reaction as a specific method of detection of Salmonella. Mol. Cell Probes. 6, 271–279. doi: 10.1016/0888-8808(92)90002-f
Rukambile, E., Sintchenko, V., Muscatello, G., Kock, R., and Alders, R. (2019). Infection, colonization and shedding of Campylobacter and Salmonella in animals and their contribution to human disease: A review. Zoonoses Public Health 66, 562–578. doi: 10.1111/zph.12611
Techathuvanan, C., Draughon, F. A., and D’Souza, D. H. (2010). Real-time reverse transcriptase PCR for the rapid and sensitive detection of Salmonella typhimurium from pork. J. Food Prot. 73, 507–514. doi: 10.4315/0362-028x-73.3.507
Techathuvanan, C., Draughon, F. A., and D’Souza, D. H. (2011). Comparison of reverse transcriptase PCR, reverse transcriptase loop-mediated isothermal amplification, and culture-based assays for Salmonella detection from pork processing environments. J. Food Prot. 74, 294–301. doi: 10.4315/0362-028x-JFP-10-306
Wang, Y. X., Zhang, A. Y., Yang, Y. Q., Lei, C. W., Cheng, G. Y., Zou, W. C., et al. (2018). Sensitive and rapid detection of Salmonella enterica serovar Indiana by cross-priming amplification. J. Microbiol. Methods 153, 24–30. doi: 10.1016/j.mimet.2018.08.003
Wang, J., Li, R., Sun, X., Liu, L., Hao, X., Wang, J., et al. (2020). Development and validation of the isothermal recombinase polymerase amplification assays for rapid detection of Mycoplasma ovipneumoniae in sheep. BMC Vet. Res. 16, 172. doi: 10.1186/s12917-020-02387-3
Yeh, K. S., Chen, T. H., Liao, C. W., Chang, C. S., and Lo, H. C. (2002). PCR amplification of the Salmonella typhimurium fimY gene sequence to detect the Salmonella species. Int. J. Food Microbiol. 78, 227–234. doi: 10.1016/S0168-1605(02)00115-0

Conflict of Interest: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.
Copyright © 2021 Zhao, Wang, Sun, Wang, Chen, Xu, Dong, Guo, Wang, Chen, Gao and Geng. This is an open-access article distributed under the terms of the Creative Commons Attribution License (CC BY). The use, distribution or reproduction in other forums is permitted, provided the original author(s) and the copyright owner(s) are credited and that the original publication in this journal is cited, in accordance with accepted academic practice. No use, distribution or reproduction is permitted which does not comply with these terms.