Direct capture and amplification of nucleic acids using a universal, elution-free magnetic bead-based method for rapid pathogen detection in multiple types of biological samples

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Abstract
Nucleic acid amplification tests (NAATs) have become an attractive approach for pathogen detection, and obtaining high-quality nucleic acid extracts from biological samples plays a critical role in ensuring accurate NAATs. In this work, we established an elution-free magnetic bead (MB)-based method by introducing polyethylene-polypropylene glycol (PEPPG) F68 in lysis buffer and using NaOH solution instead of alcohols as the washing buffer for rapid nucleic acid extraction from multiple types of biological samples, including nasopharyngeal swabs, serum, milk, and pork, which bypassed the nucleic acid elution step and allowed the nucleic acid/MB composite to be directly used as the template for amplification reactions. The entire extraction process was able to be completed in approximately 7 min. Even though the nucleic acid/MB composite could not be used for quantitative real-time PCR (qPCR) assays, this elution-free MB-based method significantly improved the sensitivity of the loop-mediated isothermal amplification (LAMP) assay. The sensitivity of the quantitative real-time LAMP (qLAMP) assays combined with this elution-free MB-based method showed an improvement of one to three orders of magnitude compared with qLAMP or qPCR assays combined with the traditional MB-based method. In addition to manual operation, like the traditional MB-based method, this universal, rapid, and facile nucleic acid extraction method also has potential for integration into automated robotic processing, making it particularly suitable for the establishment of an analysis platform for ultrafast and sensitive pathogen detection in various biological samples both in centralized laboratories and at remote sites.

Keywords
Elution-free · Magnetic beads · Nucleic acid extraction · Pathogen detection · Nucleic acid amplification tests · Polyethylene-polypropylene glycol F68

Introduction
Nucleic acids, antigens, and antibodies are major biomarkers used for pathogen detection [1]. Although antigen-based tests can be easily carried out with test strips even by untrained users, their poor sensitivity makes it difficult for them to meet the demands of containing pathogen transmission [2, 3]. Additionally, antibody-based tests are not suitable for early and timely screening of pathogens, since antibodies are generally produced 2 weeks after infection [4]. In contrast, molecular detection of nucleic acids exhibits excellent sensitivity and specificity, while being effective in the earliest phase of the infection [5]. Thus, nucleic acid amplification tests (NAATs) based on polymerase chain reaction (PCR) and loop-mediated isothermal amplification (LAMP) have increasingly become the detection modality of choice in clinical diagnostics and food safety inspection [6, 7]. In particular, NAATs are the gold standard for...
COVID-19 diagnosis at present [8, 9]. Nucleic acid extraction is the key starting point for NAATs including two steps of nucleic acid isolation from samples and nucleic acid purification, which plays an extremely critical role in ensuring highly sensitive NAATs [10–12], as high concentrations of organic components that inhibit nucleic acid amplification are normally present in complex biological samples [13, 14]. In this context, the need for universal and efficient nucleic acid extraction methods for accurate disease diagnosis and foodborne pathogen detection cannot be overstated.

Magnetic beads (MBs) with a functionalized surface that enables nucleic acid capture have been widely used for nucleic acid extraction from biological samples, based on which a variety of formats have been developed, such as manual extraction by a magnetic rack or microfluidic chips [15, 16], as well as automated robotic processing [17, 18]. Compared with the traditional nucleic acid extraction approaches, such as spin column-based methods, thermal lysis, and alkaline lysis, MB-based methods combine the advantages of simple processing, short time consumption, high product purity, and easy integration for high-throughput detection, making them the preferred nucleic acid extraction methods for pathogen detection [19, 20]. In the process of using the traditional MB-based method, once the nucleic acids are captured on MBs, they are isolated from samples, followed by washing with alcohols and eluting into nuclease-free water, and then application as templates for NAATs. However, the elution process will lead to dilution and the loss of target nucleic acids, which may cause a reduction in sensitivity and false-negative results. In contrast, the approach in which nucleic acids together with MBs are directly used for downstream NAATs concentrates nucleic acid molecules and produces rapid and highly sensitive tests, thereby avoiding the elution step and simplifying the testing protocols. However, the commonly used MBs demonstrate nonspecific adsorption of organic components, inhibiting nucleic acid amplification in the biological samples and introducing them into the reaction systems, especially samples rich in organic ingredients, such as serum and meat [21, 22]. Therefore, elution-free MB-based methods are rarely reported, and those that have been reported have only been utilized for nucleic acid extraction from samples with low content of organic ingredients, such as nasopharyngeal swabs [23].

Herein, we developed a universal elution-free MB-based nucleic acid extraction method for multiple types of biological samples, which allowed the MBs to capture nucleic acids directly, adding to the available reaction systems for downstream NAATs. Polyethylene-polypropylene glycol (PEPPG) F68 was employed to reduce the nonspecific adsorption of organic interference components by the MBs. Moreover, sodium hydroxide solution instead of alcohol was utilized as the washing buffer to avoid the inhibition of downstream amplification reactions by alcohols, as well as their unpleasant smell. The ingredients of lysis buffers and extraction procedure were also optimized to improve the extraction efficiency and reduce residual organic interference components. Furthermore, the performance of this proposed elution-free MB-based method for pathogen detection was examined in four types of simulated samples, including severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2)-positive nasopharyngeal swabs, Streptococcus aureus-infected serum, Escherichia coli O157:H7-contaminated milk, and Salmonella typhimurium-contaminated pork, in comparison with a commercial kit based on the traditional MB-based method. Our work will provide a universal, rapid, and facile nucleic acid extraction method applicable for both automated robotic processing and manual operation, which can potentially be deployed for ultrafast and sensitive pathogen detection in various biological samples via NAATs both in the professional laboratory and at remote sites lacking instruments.

Materials and methods

Materials and reagents

SARS-CoV-2 pseudovirus was purchased from Fubio Biological Technology Co., Ltd (Shanghai, China). Staphylococcus aureus (ATCC 25923), E. coli O157:H7 (ATCC 35150), S. typhimurium (ATCC 14028), Vibrio parahaemolyticus (ATCC 17802), Pseudomonas aeruginosa (ATCC 27853), nonpathogenic E. coli (ATCC 25922) strains, inactivated Mycoplasma pneumoniae, inactivated influenza A (H1N1) virus, and inactivated influenza B (Victoria) virus, as well as colorimetric/fluorescence LAMP detection master mix, were provided by Navid Biotechnology Co., Ltd. (Qingdao, China). Hydroxyl (Si-OH) MBs with diameters of 100, 200, 500, and 1000 nm were purchased from BioMag Biotech Co., Ltd. (Wuxi China). PEPPG F68 and sodium hydroxide were purchased from Macklin Biochemical Co., Ltd. (Shanghai China). Guanidine hydrochloride, guanidine isothiocyanate, sodium chloride, EDTA-Na2, Tris, SDS, Triton X-100, Tween-20, and dNTPs were purchased from Sangon Biotech Co., Ltd. (Shanghai, China). PEPPG F68 and sodium hydroxide were purchased from Macklin Biochemical Co., Ltd. (Shanghai China). Guanidine hydrochloride, guanidine isothiocyanate, sodium chloride, EDTA-Na2, Tris, SDS, Triton X-100, Tween-20, and dNTPs were purchased from Sangon Biotech Co., Ltd. (Shanghai, China). 31000 EvaGreen dye (20× in water) was purchased from Biotium, Inc. (CA, USA). Pig serum was obtained from Solarbio Life Sciences Co., Ltd. (Beijing, China). The TIANamp DNA/RNA Kit was purchased from Tiangen Biotech Co., Ltd. (Beijing, China). The BCA Protein Assay Kit was purchased from Beyotime Biotechnology Co., Ltd. (Shanghai, China). All the other chemicals and reagents were of analytical grade.
Preparation of the biological samples

Simulated SARS-CoV-2-positive nasopharyngeal swabs were prepared by spiking 100 μL SARS-CoV-2 pseudovirus suspensions with the target gene at concentrations of 1.0×10⁴, 1.0×10³, 1.0×10², and 1.0×10¹ copies/mL on fresh swabs of the nasal cavity or throat of volunteers. Simulated S. aureus-infected serum was prepared by mixing 100 μL S. aureus cell suspensions at concentrations of 1.0×10⁸, 1.0×10⁷, 1.0×10⁶, 1.0×10⁵, 1.0×10⁴, 1.0×10³, and 1.0×10² CFU/mL into 900 μL pig serum. Simulated E. coli O157:H7-contaminated milk was prepared by mixing 100 μL E. coli O157:H7 cell suspensions at concentrations of 1.0×10⁶, 1.0×10⁵, 1.0×10⁴, 1.0×10³, 1.0×10², and 1.0×10¹ CFU/mL into 900 μL sterilized commercial milk. Simulated S. typhimurium contaminated pork was prepared by immersing 10 mg sterilized commercial pork in 1 mL S. typhimurium cell suspensions at concentrations of 1.0×10⁸, 1.0×10⁷, 1.0×10⁶, 1.0×10⁵, 1.0×10⁴, and 1.0×10³ CFU/mL for 15 min, which was then ground into tissue homogenate for subsequent use.

Elution-free nucleic acid extraction based on MBs

The protocol of the elution-free MB-based method was determined after optimization. Briefly, 200 μL of samples and 25 μL MB (diameter: 1000 nm) suspension (50 μg/μL) were thoroughly mixed with the lysis buffer (50 mM Tris, 10 mM EDTA, 0.7 M NaCl, 4 M guanidine isothiocyanate, 3 M guanidine hydrochloride, 0.1% [w/v] SDS, 5% [v/v] Tween-20, 3% [v/v] Triton X-100, 30% [w/v] PEPPG F68) in a 1.5-mL centrifuge tube by vortexing. Then the tube was placed at room temperature for 5 min, followed by placement on a magnetic rack and waiting 1 min until the MBs were concentrated. After removal of the supernatant, the MBs were washed with 600 μL NaOH solution (3 mM) for 15 s and collected again with the help of a magnetic rack. The MBs were directly transferred into reaction systems for downstream NAATs. Moreover, for comparison, the prepared biological samples were also treated with a TIANamp DNA/RNA Kit based on the traditional MB-based method according to the manufacturer’s instruction.

Primer design

Primer sets specific to the SARS-CoV-2 ORF1ab polyprotein (orf1ab) gene, S. aureus strain G9 thermonuclease precursor (nuc) gene, E. coli O157:H7 strain Shiga-toxin 1 (stx1) gene, and S. typhimurium strain invasion protein (invA) gene were designed and optimized by NCBI primer-BLAST (https://www.ncbi.nlm.nih.gov/tools/primer-blast) and NUPACK software (http://www.nupack.org/), and synthesized by Sangon Biotech (Shanghai, China). The sequences of the primer sets and fragments used in this work are shown in Table 1.

LAMP reactions

The collected MBs containing the captured nucleic acids were directly used for performing colorimetric and fluorescence quantitative real-time LAMP (qLAMP). In brief, all the collected MBs (approximately 1250 μg) were transferred into a 50-μL amplification mixture containing 40.6 μL colorimetric/fluorescence LAMP detection master mix, 1 μL F3 and B3 (10 μM), 0.8 μL FIP and BIP (100 μM), 0.4 μL LF and LB (100 μM), and 0.6 μL HiScript II Reverse Transcriptase (200 U/μL, for RNA templates only). Additionally, 5 μL of nucleic acid products prepared by the commercial kit were added into the same amplification mixture for comparison. Fluorescence qLAMP was performed using a CFX Connect™ Real-Time PCR System (Bio-Rad, CA, USA) at 65°C for 45 min, and monitored at 1-min intervals. The threshold was set at 10 times the standard deviation of the fluorescence values of the baseline. The time threshold (Tₙ) value was defined as the time point when the fluorescence values exceeded the threshold. Colorimetric LAMP assays were performed at 65°C for 50 min in a heating block, and the results were directly read by the naked eye after placing the tubes on white cardboard.

PCR reactions

Besides LAMP assays, the nucleic acid products prepared using the commercial kit were also employed as templates for fluorescence quantitative real-time PCR (qPCR) to simulate the most commonly used NAAT process at present. Briefly, 2 μL of the nucleic acid products prepared by the commercial kit were added into a 20-μL reaction mixture containing 0.4 μL dNTPs (10 mM), 0.8 μL each for forward and backward primer (10 μM), 0.2 μL Taq DNA polymerase (5 U/μL), 3 μL 10× FastTaq buffer, 0.5 μL EvaGreen dye, and 0.15 μL HiScript II Reverse Transcriptase (200 U/μL, for RNA templates only). The reaction procedure included denaturation at 95°C for 3 min and 40 cycles consisting of 95°C for 10 s, and 58°C for 30 s for amplification. For RNA templates, a reverse transcription step of incubating the reaction system at 55°C for 5 min was added before the denaturation step in the reaction procedure. Fluorescence real-time PCR was performed by a CFX Connect™ Real-Time PCR System (Bio-Rad, CA, USA) and monitored after each thermal cycling. The threshold was set at 10 times the standard deviation of the fluorescence values of the baseline. The cycle threshold (Cₜ) value was defined as the number of cycles required for the fluorescence values to exceed the threshold.
Protein and nucleic acid absorption capability assays

The influence of PEPPG F68 on the protein and nucleic acid absorption capability of the MBs was investigated. In brief, S. aureus cell suspensions with a concentration of 3.0×10^8 CFU/mL were mixed with the MBs and lysis buffer with/without PEPPG F68, respectively. After incubation for 5 min, the residual protein content of the mixtures was measured by a BCA protein assay kit. Moreover, the protein content of the mixtures without the MBs was also measured to evaluate the influence of PEPPG F68 on BCA protein assays. For nucleic acid absorption capability assay, a 77-nucleotide nucleic acid fragment containing the partial sequence of SARS-CoV-2 orf1ab gene was designed, and then the nucleic acid fragment solutions at concentrations of 1.0×10^{-12}, 1.0×10^{-11}, 1.0×10^{-10}, 1.0×10^{-9}, and 1.0×10^{-8} M were mixed with the MBs and lysis buffer with/without

| Name | Sequence (5′–3′) |
|------|------------------|
| SARS-CoV-2 orf1ab gene (aOP102293.1) | |
| LAMP primer set [24] | F3 CCGTGGAACAAAATTGTACAC |
| | B3 CTTCTCTGGAATTAAACACACTT |
| | FIP TACGACACAAAGCGCCAAAAATATTATTTCGCTGCAAGGAATTAAAGGAG |
| | BIP TAATGCTGGGACTAAATTAAAGCCATCTGTACAAATTCCCCTTTGAGTG |
| | LF TTACAGCTTAAAGAATGTCTGAACACT |
| | LB TTTGATTAAGGGAACATATTGTACAG |
| PCR primer set | F TCAACTAGGTTTCAAACTTTACTTT |
| | R AGCTGTCAACCACTTGAG |
| 77-nucleotide DNA fragment | TCACTAGGTTTCAAACTTTACTTGCTTACAGAAAGTATTTATGTACAG |
| S. aureus nuc gene (aDQ399678.1) | |
| LAMP primer set [25] | F3 ATGCAAAAGAAAAATTGAAGTCGA |
| | B3 GCCTTTCTTCTCGCTCCAAT |
| | FIP CTTTCTTACATTTTCATACAGCATGTGTGACAAAGGTCAAAAGA |
| | BIP TCAAGGCTCTGCTAAAGTTGCTTTATTTTCGCTGTACACTT |
| | LF TACGCTAAGGCAACGCATA |
| | LB CTTAACAATACACATAAGAACC |
| PCR primer set | F GTGCTGCGATATGATGCAG |
| | R CACTAAGCAACATGACTGAG |
| E. coli O157:H7 stx1 gene (aLC388498.1) | |
| LAMP primer set [26] | F3 TGTGGAAGAAAAATTGTACAC |
| | B3 GCCTTTCTTCTCGCTCCAAT |
| | FIP CTTTCTTACATTTTCATACAGCATGTGTGACAAAGGTCAAAAGA |
| | BIP TCAAGGCTCTGCTAAAGTTGCTTTATTTTCGCTGTACACTT |
| | LF TACGCTAAGGCAACGCATA |
| | LB CTTAACAATACACATAAGAACC |
| PCR primer set | F ATTCGCGGAGCTGCGGCAAT |
| | R CTCCCTCACCTACGAG |
| S. typhimurium invA gene (aM90846.1) | |
| LAMP primer set [27] | F3 CTGGACATTGTGTTAGTACAG |
| | B3 ACATACGGTACAGTACAGA |
| | FIP GAAGGTCGGGAGGAAATATGGGCAAGCAAGTCACACGAG |
| | BIP TTCGTGAAACCGCTGAAYGGAACAGCTGAGGGATTAG |
| | LF TCCGCTGTGTAACAGGAGTA |
| | LB TACACAGGAGATTACACAACATGG |
| PCR primer set | F CAAATCGGCGGCAATTCAGAG |
| | R AGGAAGGTACTGCGAG |

Table 1 Sequences of nucleic acids used in this work

*GenBank accession number*
PEPPG F68, respectively. The MBs were washed with the washing buffer and transferred into the PCR reaction system. The influence of PEPPG F68 on nucleic acid absorption was evaluated by comparing $C_t$ values for the MBs incubated in the lysis buffer with/without PEPPG F68.

Results and discussion

Optimization of the MB-based elution-free method

Before the protocol of the elution-free method was determined, it was optimized using fluorescence qLAMP with S. typhimurium (1.0×10^6 CFU/mL) as a template. First, the effects of particle size of the MBs on the nucleic acid extraction efficiency were explored. As shown in Fig. 1a, MBs with a diameter of 1000 nm exhibited the best performance with the lowest $T_t$ value, while the performance of smaller beads showed no evidence of differences. Despite the larger MBs possessing relatively smaller specific surface area, a higher amount of Si-OH was coated on a single magnetic bead, which may be beneficial for the macromolecules like nucleic acid to be more stably adsorbed on the MBs. The lower performance of the 100-nm, 200-nm, and 500-nm MBs was likely due to the lower amount of Si-OH on their surface. Besides appropriate particle size, a sufficient dosage of the MBs would also improve the extraction efficiency. The $T_t$ value decreased significantly with the increase in the dosage of MBs from 500 μg to 1250 μg, and failed to decrease further and increased slightly when the dosage of MBs was increased to 1500 μg (Fig. 1b). This was because the higher dosage of MBs introduced more organic interference components into the reaction system, leading to excessive absorption of LAMP components like primers and polymerase and resulting in reduced amplification efficiency. Thus, 1250 μg was determined as the optimal dosage of MBs.

In this work, PEPPG F68 was applied in lysis buffer to reduce the absorption of protein ingredients of the samples, the major organic interference components that inhibit the amplification reaction, such as immunoglobulin, protease, mucin, and hemoglobin [28–30]. Therefore, the concentration of PEPPG F68 in the lysis buffer was also optimized. As shown in Fig. 1c, the $T_t$ value of the nucleic acids isolated by the lysis buffer with 30% (w/v) PEPPG F68 was 22.55 min, whereas the values for nucleic acids isolated by the lysis buffer with 10% and 20% (w/v) PEPPG F68 were approximately 8 min greater, demonstrating that the application of PEPPG F68 indeed promoted NAAT efficiency. However, the lysis buffer with 40% (w/v) PEPPG F68 failed to further reduce the $T_t$ value, likely because, in addition to

![Fig. 1](image1.png)  

**Fig. 1** Optimization of the protocol for elution-free MB-based method. Effect of (a) the MB size, (b) the MB dosage, (c) the PEPPG F68 concentration in lysis buffer, (d) the incubation time for MBs and samples in lysis buffer, (e) the NaOH concentration in washing buffer, and (f) the sample volume applied for nucleic acid extraction on the efficiency of LAMP assays.

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proteins, this surfactant can also weaken the electrostatic interaction between nucleic acids and Si-OH on the surface of the MBs. Moreover, the application of PEPPG F68 would increase the viscosity of the lysis buffer, and thus excessive PEPPG F68 content would lead to inconvenient liquid handling and prolonged MB gathering time. Sufficient incubation time is of great importance for the complete lysis of cells and virions, and the release of genomic materials. The $T_1$ value of the samples decreased from 21.65 min to 18.53 min with the increase in the incubation time from 1 min to 5 min. However, no further decrease in the $T_1$ value was observed when the incubation time was extended to 7 min or 9 min (Fig. 1d). Accordingly, the 30% (w/v) PEPPG F68 was applied in the lysis buffer, and the optimal incubation time in lysis buffer was determined to be 5 min.

Besides the lysis buffer, the washing buffer was also optimized. The results illustrated that the $T_1$ value decreased sharply with the increase in the NaOH concentration when the MBs were washed with the NaOH solution of no more than 3 mM, and increased continuously with the increase in the NaOH concentration from 3 mM to 6 mM since a high-pH environment is not beneficial for the capture of nucleic acid by the Si-OH on MBs (Fig. 1e). Finally, considering that the MBs carrying nucleic acids are directly used for downstream amplification, we explored the influence of the volume of initial samples on extraction efficiency. As expected, the $T_1$ value decreased slightly with increased sample input from 200 μL to 1000 μL, revealing that the nucleic acids captured by the MBs increased. However, the $T_1$ value decreased only 1.74 min after the total input sample volume was increased fourfold, while the time consumption for MB collection by the magnetic rack was sharply increased from 1 min to 4.5 min due to the more dispersed MBs in larger-volume systems (Fig. 1f). Since both the total time and sample consumption were the lowest, the 200-μL protein content of the MBs treated by lysis buffer without PEPPG F68 showed no significant differences when no MBs were added to the mixtures, demonstrating that the presence of PEPPG F68 did not promote cell lysis or influence BCA protein assays (Fig. 3a). In contrast, the $C_1$ values for MBs incubated in lysis buffer with or without PEPPG F68 showed no significant differences, demonstrating that this surfactant at this dose would not affect the nucleic acid absorption on MBs (Fig. 3b). With the help of this low-cost surfactant (approximately $0.13/g$), this elution-free MB-based method involved no specially designed MBs and required only ordinary commercial Si-OH MBs (approximately $0.2/mg$), ensuring that the cost of this method was about the same as that of traditional MB-based methods.

Besides introducing PEPPG F68 in lysis buffer, the alternation of washing buffer from alcohol to NaOH solution also contributed to the improvement in LAMP assay efficiency, as the cells or virions in the samples could be further lysed by NaOH during the washing process.
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the interaction between nucleic acids and Si-OH of the MBs weakened in alkaline solution, allowing the nucleic acids to easily detach from MBs and participate in the LAMP reaction when they were transferred into the reaction system. Furthermore, since the LAMP reaction has better tolerance to 3 mM NaOH solution than alcohols, the MBs could be directly added into the reaction system without a drying step, which further simplified the protocol and reduced the probability of aerosol pollution and nonspecific amplification. As shown in Fig. 4, by combination with this elution-free MB-based method, the LAMP assays could specifically identify the target pathogen from the nucleic acid products of multiple types of biological samples spiked with the same amount of other pathogens that are frequently detected in corresponding biological samples, illustrating that the treatment of this method did not cause nonspecific amplification. In summary, by combination with the elution-free MB-based method, the entire pathogen detection process from sample to result could be completed within 30 min while exhibiting excellent specificity, which was approximately 40 min faster than the detection process involving traditional MB-based methods.

**Sensitivity of the LAMP assays combined with the elution-free MB-based method**

Apart from excellent efficiency and specificity, high sensitivity is another important property of NAATs to ensure successful diagnoses. Thus, the sensitivity of the LAMP assays combined with the elution-free MB-based method was explored next. Biological samples described in section 2.2 were treated with the elution-free MB-based method...
and the commercial kit for nucleic acid extraction, and the products were employed as templates for the following fluorescence qLAMP and colorimetric LAMP assays. As shown in Fig. 5a and e, the qLAMP assays with the nucleic acid products prepared by the elution-free MB-based method as templates showed higher sensitivity than those with the products prepared by the commercial kit. The products of the simulated SARS-CoV-2-positive nasopharyngeal swabs with 1.0×10^4, 1.0×10^3, and 1.0×10^2 copies/mL of target nucleic acid were detected as positive by the fluorescence qLAMP assays combined with the elution-free MB-based method, while only the samples with 1.0×10^4 and 1.0×10^3 copies/mL target nucleic acid were detected as positive by the fluorescence qLAMP assays combined with the commercial kit. The results demonstrated that the elution-free MB-based method could improve the sensitivity of fluorescence qLAMP assays to this RNA target by one order of magnitude. Moreover, fluorescence qLAMP assays combined with the elution-free MB-based method exhibited much better performances for DNA targets detection, as the limit of detection (LOD) for the *S. aureus* nuc gene, *E. coli* O157:H7 stx1 gene, and *S. typhimurium* invA gene in different simulated biological samples was 1.0×10^5 CFU/mL, 1.0×10^3 CFU/mL, and 1.0×10^5 CFU/mL (Fig. 5b–d), respectively, while the LOD of the fluorescence qLAMP assays combined with the commercial kit to the same targets was 1.0×10^5 CFU/mL, 1.0×10^3 CFU/mL, and 1.0×10^5 CFU/mL, respectively (Fig. 5f–h), suggesting the elution-free MB-based extraction method could improve the sensitivity to these DNA targets by two to three orders of magnitude as compared with the traditional MB-based method. Besides lower LOD, the *T*_i values of the fluorescence qLAMP assays combined with the elution-free MB-based method were significantly smaller than those combined with the commercial kit for the same targets, which was consistent with the results of extraction efficiency evaluation in section 3.2.

Unfortunately, the products prepared by the elution-free MB-based method could not be used as templates for qPCR assays, as almost no fluorescence signal accumulation was observed in the reactions, which might be attributed to the inhibition of the reaction by the Fe^{3+} released from the MBs during the high-temperature thermal cycling process, since this heavy metal ion can strongly inhibit qPCR at a concentration of 3 mg/L [31]. Therefore, only the nucleic acid products prepared by the commercial kit based on the traditional MB-based method were employed as templates for qPCR assays for the comparison, which is the most commonly used NAATs protocol at present [32]. As shown in Fig. 5i–l, by combination with the elution-free MB-based method, the sensitivity of the qLAMP assays to the target bacteria in the biological samples was even higher than the qPCR assays, whose LOD to SARS-CoV-2, *S. aureus*, *E. coli* O157:H7, and *S. typhimurium* was 1.0×10^3 copies/mL, 1.0×10^4 CFU/mL, 1.0×10^5 CFU/mL, and 1.0×10^4 CFU/mL, respectively. The results further indicated that the elution-free MB-based method was more effective for nucleic acid extraction from samples with high content of organic interference components, such as serum, milk, and pork, involved in this work than the traditional ones.

In addition to fluorescence qLAMP assays, the elution-free MB-based method improved the sensitivity of the colorimetric LAMP assays by one order of magnitude for the
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RNA targets and two to three orders of magnitude for DNA targets (Fig. 6). While the sensitivity of the colorimetric LAMP assays was still lower than the fluorescence qLAMP assays and qPCR in general, neither the colorimetric LAMP nor the elution-free MB-based method requires sophisticated bulky instruments or complicated protocols, and their combination provides a fast and facile approach suitable for on-site pathogen detection from sample to result in remote areas lacking instruments or professional staff, which is of great importance for the timely control of pathogen transmission. In all, the more concentrated nucleic acids and lower levels of organic interference components in the products prepared by the elution-free MB-based methods as compared with the traditional methods not only resulted in higher detection efficiency but also improved the sensitivity of the LAMP assays.

Conclusion

Herein, an elution-free MB-based nucleic acid extraction method was established by introducing PEPPG F68 in lysis buffer and using NaOH solution instead of alcohols as washing buffer, which was utilized for pathogen detection in multiple types of biological samples via NAATs. Following nucleic acid isolation and purification, the nucleic acid/Si-OH MB composite can be directly added to the reaction systems as templates for LAMP assays. Compared with the
traditional MB-based method, this proposed elution-free approach not only simplified the extraction protocol and reduced the time consumption, but also enhanced the efficiency and sensitivity of LAMP assays. The LOD of both fluorescence qLAMP and colorimetric LAMP assays combined with this elution-free MB-based method showed an improvement of one to three orders of magnitude compared with those combined with the traditional MB-based method.
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and the fluorescence qLAMP assays even exhibited higher sensitivity than the most widely used qPCR gold-standard assay for pathogen detection in samples with high content of organic interference components, despite the inability to use the products of this extraction method for qPCR assays. Moreover, the time consumption of the entire detection process from sample to result was less than 30 min, which was much shorter than the most commonly used approaches at present. In addition, this elution-free MB-based method was robust and reliable for pathogen detection in a variety of biological samples, in contrast with those that were only applicable for samples with low content of organic interference components, such as nasopharyngeal swabs, thus contributing to the progress of analytical chemistry applied for nucleic acid analysis in biological samples. Finally, the execution of nucleic acid extraction via this method did not require sophisticated bulky equipment or specialized instruments, making it easy to integrate into automated robotic processing, as well as for manual extraction by microfluidic chips and a magnetic rack, supporting its potential for pathogen detection from various biological samples via NAATs in both the centralized laboratory and remote sites.

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Authors’ contributions Qianqian Jiang, Jinling Guo, and Ailin Wang performed the experiments; Qianqian Jiang and Yang Li analyzed the data; Yang Li, Cuiping Ma, Chao Shi, and Lin Huang designed the study; Yang Li and Qianqian Jiang wrote the manuscript; and all authors contributed to the writing of the paper, had primary responsibility for the final content, and read and approved the final manuscript.

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Declarations

Ethics approval The authorized Human Health and Ethics Committee of the Affiliated Hospital of Qingdao University approved this study, and all the volunteers helping us prepare the simulated SARS-CoV-2-positive nasopharyngeal swabs provided informed consent. In addition, all methods were carried out in accordance with the relevant guidelines and regulations.

Conflicts of interest There are no conflicts of interest to declare.

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