ABSTRACT

A fluorescence assay combined with PCR, catalytic hairpin assembly (CHA), and graphene oxide (GO) was established to detect emetic *Bacillus cereus* in milk samples. The processes of the assay are not new, but components of the processes make the assay useful. Two partially complementary hairpin probes (H1 and FAM-H2) were designed according to the target single-strand DNA (ssDNA). The CHA reaction could be initiated only by the target ssDNA, which was generated by the denaturation of PCR amplicons. In the absence of the target ssDNA, CHA reaction could not be triggered, which caused the H1 and FAM-H2 adsorbing on the surface of GO and exhibiting a low fluorescence intensity. Addition of the target ssDNA resulted in opening of the hairpin H1 that subsequently hybridized with H2. Then, target ssDNA would be replaced from the H1 and recycled to promote another CHA reaction. Through the CHA reaction, multiple H1-H2 duplexes were generated that could not adsorb on the surface of GO. Thus, a strong fluorescence signal would be obtained. The assay showed a limit of detection for emetic *B. cereus* of $6.2 \times 10^1$ cfu/mL in pure culture and $5.9 \times 10^2$ cfu/mL in spiked milk without enrichment. By changing the PCR primer, the assay developed in this study had potential to detect other bacteria.

**Key words:** catalytic hairpin assembly, graphene oxide, fluorescence, emetic *Bacillus cereus*

INTRODUCTION

*Bacillus cereus* is a gram-positive, rod-shaped, and spore-forming bacillus that can pose significant threats to public health (Priha et al., 2004; Dierick et al., 2005; Fricker et al., 2007). *Bacillus cereus* can cause 2 different gastrointestinal diseases of food poisoning, namely diarrhea and emesis (Arslan et al., 2014; Du et al., 2018; Li et al., 2018). The diarrhea type is caused by heat-labile enterotoxins whereas the emesis type is the result of heat-stable cereulide (Martínez-Blanch et al., 2011). It has been reported that *B. cereus* can be found in food such as pasta, rice, meat, noodles, milk, and other dairy products (Martinez-Blanch et al., 2010; Shiota et al., 2010). Moreover, emetic *B. cereus* can cause spore germination when foods are boiled, cooled, and stored at room temperature. Considering emetic *B. cereus* poses significant threats to public health (Delbrassinne et al., 2015), a specific and accurate method for detection of emetic *B. cereus* in food is in demand.

Conventional detection methods for *B. cereus* are time consuming (~7 d; Madiyar et al., 2015; Porcellato et al., 2016). Polymerase chain reaction has been developed for the rapid and accurate detection of foodborne pathogens (Mao et al., 2016; Luo et al., 2017). However, conventional PCR has been limited by the low resolution of gel electrophoresis. Even worse, the use of ethidium bromide could affect the operator’s health (Yu et al., 2018). Real-time PCR provides a higher sensitivity with no additional postamplification; however, real-time PCR needs expensive equipment and fluorescence dyes, and even worse, it is difficult to design the primer and requires trained analysts to operate (Zhou et al., 2017; Hiitiö et al., 2018). Therefore, the development of low-cost and simple methods to make up for these shortcomings is vitally important.

In recent years, a series of detection methods based on fluorescence was established. However, most of the fluorescence strategies required a fluorophore-labeled DNA and the quencher-labeled cDNA or a dual-labeled beacon at each end of the DNA (Zheng et al., 2012). On the one hand, it was expensive to synthesize fluorophore-labeled DNA, and on the other hand, the labeled DNA had weak affinity to the target DNA (Liu et al., 2013; Xing et al., 2016). To make up for the disadvantages, researchers have been working on new approaches.
Graphene oxide (GO) is an important derivative of graphene and is well-known for its extraordinary electronic, optical, and thermal properties as well as large surface area, good water dispersibility, and biocompatibility (Jhaveri et al., 2000). Graphene oxide exhibits unique fluorescence-quenching capability when organic fluorescent molecules adsorb on it (Ho and Leclerc, 2004). Also, it has been reported that single-strand DNA (ssDNA) can adsorb onto the surface of GO by π–π stacking and electrostatic interactions, but hardly interacts with rigid double-stranded DNA (dsDNA) or well-folded structures (Feng et al., 2013). Based on the fluorescence-quenching capability and the affinity difference, a novel bioassay platform for DNA detection through GO fluorescence has been reported. However, a single target DNA usually conjugates only to one fluorescence probe, which limits the sensitivity. Recently, a series of signal amplification methods was developed; catalytic hairpin assembly (CHA) amplification technique is an enzyme-free amplification technique that relies on hybridization and strand-exchange reactions to achieve amplification (Chen et al., 2016; Li et al., 2016; Dai et al., 2018). The reaction mechanism can be explained as follows: the initiator strands can trigger a pair hairpin to form a duplex, then the initiator strands can be released to the reaction system for the subsequent CHA reaction. Thus, by introducing only one ssDNA, the reaction could produce multiple dsDNA (Zhang et al., 2011; Luan et al., 2017). Furthermore, CHA is a sequence-dependent reaction, and thus, it ensures specificity.

Based on the above considerations, we developed a novel and enzyme-free assay based on catalyzed hairpin DNA assembly combined with GO to detect emetic B. cereus. In this assay, the target ssDNA was generated by PCR amplicons and the hairpin probes opened only after binding with the target ssDNA. In the absence of the target ssDNA, hairpin probes were adsorbed on the surface of GO, resulting in fluorescence quenching. Upon the addition of the target ssDNA, a cascade of dsDNA between the 2 hairpin probes was formed, which had low affinity with GO, and then the hairpin probes would separate from the GO and a strong fluorescence signal would be obtained. The proposed method could be a cost-effective and label-free fluorescence system. By changing the PCR primer, the assay developed in this study had potential to detect other bacteria.

### MATERIALS AND METHODS

#### Reagents and Materials

Luria-Bertani medium was purchased from Land Bridge Technology Co. Ltd. (Beijing, China). The GO was bought from XFNANO Materials Technology Co. Ltd. (Nanjing, China). The TaKaRa MiniBest DNA Fragment Purification Kit was purchased from TaKaRa Biotech Co. Ltd. (Dalian, China). Two × Taq master mix was purchased from Tiangen Biotech Co., Ltd. (Beijing, China). HEPES buffer (25 mM HEPES; 200 mM KCl; 200 mM NaCl; 1% DMSO; pH 7.4) was used in the experiments. The oligonucleotides (Table 1), including specific primers, hairpin structure H1, FAM modified hairpin structure H2, and the target ssDNA (93 nt), were synthesized by TsingKe Biotech. Co. Ltd. (Beijing, China). To ensure the formation of hairpin structure, the hairpin structure oligonucleotides were heated at 95°C for 5 min and then cooled to room temperature.

#### Bacterial Culture

The bacteria used in this experiment were listed in Table 2. All bacterial strains were cultured in Luria-Bertani broth overnight at 37°C with continuous shaking (180 rpm). Sterile PBS (0.01 M, pH 7.4) was applied to suspend fresh target bacteria to obtain 10 serial dilutions. The counts of bacterial colonies were determined using the conventional plate counting method at 37°C for 18 h.

#### DNA Extraction and PCR Conditions

The genomic DNA for all the bacteria were harvested using the boiling method. One milliliter of the bacteria

<table>
<thead>
<tr>
<th>Nucleic acid</th>
<th>Sequence (5′–3′)</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>cesB-Forward</td>
<td>CTCACGAGAGATGCCGATTGC</td>
<td>Yu et al. (2018)</td>
</tr>
<tr>
<td>cesB-Reverse</td>
<td>CGTGGTTACCATTCCGGAAGG</td>
<td>Yu et al. (2018)</td>
</tr>
<tr>
<td>H1</td>
<td>ATGGAAGAGAGATGGCCCAAGAGAGTGGTGGG</td>
<td>This study</td>
</tr>
<tr>
<td>FAM-H2</td>
<td>FAM-ATGGCCAAACCCTTCTCTTTTGGGCTTCCATCTCTCTAGAAGAGTTTAAGTA</td>
<td>This study</td>
</tr>
<tr>
<td>Target ssDNA1 (93 nt)</td>
<td>CGTGGTACATTCCGGAGGCGAAATACCTCTCTCTAAGAATTTGCGCAATCTCTCTTTCATGGGCTCAG</td>
<td>This study</td>
</tr>
</tbody>
</table>

1ssDNA = single-strand DNA.
was washed twice in an equal volume of bacteria-free PBS and resuspended in 50 μL of sterilized water, then boiled in a water bath for 25 min. After centrifugation at 12,000 × g for 5 min at room temperature, the DNA in supernatant was used as template for PCR.

The specific primers used in this study were listed in Table 1. The PCR amplification was carried out in a total reaction volume of 50 μL containing 25 μL of 2 × Taq Master Mix, 2 μL of 10 μM each forward and reverse cesB primer, 2 μL of genomic DNA, and 19 μL of dH2O. The PCR conditions were as follows: 94°C for 5 min, followed by 30 cycles of denaturation at 94°C for 30 s, annealing at 60°C for 30 s, and elongation at 72°C for 30 s, with a final elongation at 72°C for 10 min.

**Purification and Denaturation**

After the PCR, the PCR amplicons were subjected to purification and denaturation. The PCR amplicons were purified according to the operation steps of the DNA Fragment Purification Kit. After purification, the PCR amplicons were put into the PCR apparatus and denatured at 94°C for 5 min. Then the denatured amplicons were quickly transferred to ice water to obtain the target ssDNA.

**Agarose Gel Electrophoresis**

Agarose gel electrophoresis was used to test feasibility of the catalyzed hairpin assembly. The H1, H2, mixture 1 (target ssDNA and H1), mixture 2 (H1 and H2), and mixture 3 (H1, H2, and target ssDNA) were analyzed and the final concentration of all the DNA was 1 μM. After incubation at 37°C for 30 min, the test samples were added into 4% agarose gel. The gel was run under a constant voltage of 80 V for 45 min followed by imaging under the UV trans illuminator (Bio-Rad, Hercules, CA).

**GO-Based CHA Reaction and Fluorescence Detection**

The H1, FAM-H2, and denatured PCR amplicons were mixed in HEPES buffer, followed by incubation at 37°C for 30 min. Subsequently, GO was added to the mixture and incubated 10 min at room temperature. Finally, the reaction mixture was transferred to 96-well plate for fluorescence detection. The fluorescence intensity was monitored using a multifunctional microplate reader (Thermo Fisher Scientific, Waltham, MA) by exciting the sample at 488 nm and measuring the emission at 520 nm. The slits for excitation and emission were both set at 5 nm. All the experiments were repeated in triplicate.

**Detection of Emetic B. cereus Using Conventional PCR**

The practicability of the conventional PCR assay was evaluated using the emetic *B. cereus* in pure culture. One milliliter of emetic *B. cereus* in pure culture was centrifuged at 12,000 × g for 5 min at room temperature and resuspended in an equal volume of sterile water. Then the emetic *B. cereus* was diluted with sterile water ranging from 6.2 × 10⁶ to 6.2 × 10⁹ cfu/mL. Next, DNA templates were obtained using the boiling method as mentioned above, followed by the PCR amplification. Finally, the PCR amplicons were added into a 2% agarose gel electrophoresis, run under a constant voltage of 150 V for 20 min, followed by imaging under the UV trans illuminator (Bio-Rad).

**Process of Emetic B. cereus Detection in Artificially Inoculated Milk Samples**

The milk purchased from a local supermarket used in this study was verified for the absence of *B. cereus* by standard culture assays. The milk matrix was obtained by mixing 1 mL of milk with 9 mL of sterile PBS. Then a 900-μL milk matrix was contaminated with the different concentrations of emetic *B. cereus* to obtain a final concentration of 5.9 × 10⁶ to 5.9 × 10⁹ cfu/mL. These were centrifuged at 8,000 × g for 3 min at room temperature to remove the large debris. After extracting genomic DNA for PCR, the purified and denatured PCR amplicons were used in a fluorescence detection method. All of the experiments were repeated in triplicate.

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**Table 2.** Target and nontarget bacteria strains used in this study

<table>
<thead>
<tr>
<th>Strain of bacteria</th>
<th>Strain ID</th>
<th>Source</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>Bacillus cereus</em> (n = 5)</td>
<td>JDZ0102Y</td>
<td>JX-CDC</td>
</tr>
<tr>
<td></td>
<td>PZ0063L</td>
<td>JX-CDC</td>
</tr>
<tr>
<td></td>
<td>JA0061</td>
<td>JX-CDC</td>
</tr>
<tr>
<td></td>
<td>JX0121LY</td>
<td>JX-CDC</td>
</tr>
<tr>
<td></td>
<td>FX0106Y</td>
<td>JX-CDC</td>
</tr>
<tr>
<td><em>Staphylococcus aureus</em></td>
<td>26001</td>
<td>CMCC</td>
</tr>
<tr>
<td><em>Bacillus subtilis</em></td>
<td>65501</td>
<td>CMCC</td>
</tr>
<tr>
<td><em>Cronobacter sakazakii</em></td>
<td>45401</td>
<td>CMCC</td>
</tr>
<tr>
<td><em>Salmonella Paratyphi B</em></td>
<td>50049</td>
<td>JX-CDC</td>
</tr>
<tr>
<td><em>Listeria monocytogenes</em></td>
<td>13932</td>
<td>ATCC</td>
</tr>
</tbody>
</table>

1ATCC = American Type Culture Collection; CMCC = China Medical Culture Collection; JX-CDC = Jiang Xi Province Center for Disease Control and Prevention.
RESULTS AND DISCUSSION

Principle of Emetic B. cereus Detection

The principle of the fluorescence assay is shown in Figure 1. The 2 partially complementary hairpin probes were designed according to the target ssDNA. The H1 included recognition sequence of the target ssDNA and the FAM-H2. The FAM-H2 included the recognition sequence of H1 and partly the same sequence as the target ssDNA. First, we used the boiling method to harvest the genomic DNA. Then PCR was used to obtain the target DNA. After PCR amplification, the PCR amplicons were subjected to purification. Then the target ssDNA was obtained by denaturing the PCR amplicons at 94°C for 5 min and quickly transferring them to ice water. Finally, the target ssDNA was used for the fluorescence detection. Part A shows the fluorescence assays with target ssDNA. When the system contained target ssDNA, the H1 would hybridize with target ssDNA and expose the recognition sequence of H2. Subsequently, H2 would hybridize with the recognition sequence of H1 and replace the target ssDNA to generate the H1-H2 duplexes. The released target ssDNA would recycle to promote another CHA reaction. Thus, by introducing the target ssDNA, multiple H1-H2 duplexes could be generated that could not absorb on the surface of GO, then a strong fluorescence signal would be obtained. However, in the absence of the target ssDNA (part B), 2 hairpin probes could not hybridize spontaneously because the complementary sequences were locked in their stem. Thus, the sticky ends of the hairpin probe would adsorb on the GO surface by π–π interaction, which resulted in low fluorescence intensity. Therefore, the concentration of the bacteria could be quantified based on the intensity of the fluorescence signal. Thus, the complete time for the detection of emetic B. cereus including the time for sample preparation, PCR amplification, purification, and denaturation, and CHA amplification would be about 3.5 h.

Feasibility of This Method

To demonstrate the viability of the proposed strategy, the fluorescence signals of the system containing differ-
ent conditions were shown in Figure 2A. The system without target ssDNA exhibited very weak fluorescence intensity. Upon the addition of the target ssDNA, the fluorescence intensity increased significantly due to the formation of the H1-H2 duplexes though the CHA reaction. To determine the occurrence of CHA reaction, 4% agarose gel electrophoresis analysis was performed. As shown in Figure 2B, when H1 mixed with target ssDNA in lane 4, a new brighter band appeared. This indicated the target ssDNA could open the H1 and produced target ssDNA-H1 duplexes. When the H1 and H2 were mixed, there were no new bands in lane 5, indicating that the H1 and H2 did not react spontaneously with each other. When the H1, H2, and target ssDNA were mixed, there were new bands in lane 6. This suggested that the target ssDNA initiated a CHA reaction and generated the H1-H2 duplexes. These 2 results demonstrated the target ssDNA could catalyze the hybridization of H1, H2 and subsequently generated the H1-H2 duplexes. All of the experiments were repeated in triplicate.

Optimization of Experimental Parameters

To obtain better performance of emetic B. cereus detection, the dosage of GO, incubation time, and CHA reaction time were carefully optimized. As GO acted as a quencher, it was important to optimize the concentration of GO and the incubation time. If the GO concentration was low, it could cause a strong background signal. Also, if the GO concentration was too high, it could cause a negative signal. Thus, a suitable concentration of GO was vitally important to the assay. In this study, we used 3 nM H1/H2 and 50 μg/mL of GO to study the quenching effect. The signal to noise ratio F/F0 (F and F0 represented the reaction system with and without the target ssDNA, respectively) was used to investigate the quenching effect. As shown in Figure 3A, as the volume of GO increased from 0 to 10 μL, the F/F0 also increased. When the volume reached 10 μL, it recorded the highest signal. Therefore, the optimized volume of 50 μg/mL of GO was 10 μL.

The effects of CHA reaction time and GO incubation time were also investigated. As shown in Figure 3B, the F/F0 increased with the CHA reaction time from 0 to 30 min, then decreased after 30 min; thus, the optimal CHA reaction time was 30 min. Figure 3C showed the fluorescence intensity changed with the incubation time from 0 to 22 min after the addition of GO. It also showed the fluorescence intensity of the negative control group decreased rapidly and then stabilized. However, the fluorescence intensity of the positive group after the addition of GO decreased slightly and then stabilized. Thus, 12 min was adopted as the optimal incubation time of the GO for subsequent studies. All of the experiments were repeated in triplicate.

Application of the CHA-GO Method for Detection of Emetic B. cereus in Pure Culture

To demonstrate the sensitivity of the developed method, different concentrations of emetic B. cereus were measured under the optimal experimental conditions. Figure 4 shown the fluorescence intensity changed with the concentrations of emetic B. cereus. As the concentration of emetic B. cereus increased, the fluorescence intensity was also increasing. A linear regression equation between the concentration of the emetic B. cereus and the fluorescence intensity was given by y = 0.3427x + 0.1534 (R² = 0.9857), where y was the fluorescence intensity at 520 nm and x was the concentration of emetic B. cereus that ranged from 6.2 × 10⁰ to 6.2 × 10⁷ cfu/mL. Based on the cut-off [negative control (NC) + 3 SD; Xu et al., 2017], the sensitivity of the method in pure culture was 6.2 × 10¹ cfu/mL. All of the experiments were repeated in triplicate.
Specificity of the CHA-GO Assay in Pure Culture

The specificity of the proposed assay was evaluated by selecting 8 strains of nontarget bacteria (Table 2). As shown in Figure 5, the nontarget bacteria (approximately 10^7 cfu/mL) containing *Bacillus subtilis*, *Listeria monocytogenes*, *Salmonella Paratyphi*, *Staphylococcus aureus*, *Cronobacter sakazakii*, and non-emetic *B. cereus* showed low fluorescence intensity. However, a high fluorescence intensity was obtained using the emetic *B. cereus*. This suggested that this fluorescence method had a high specificity to distinguish between target bacteria and other bacteria. All of the experiments were repeated in triplicate.

Detection of Emetic *B. cereus* Using Conventional PCR

The ability of the conventional PCR assay was also evaluated using the emetic *B. cereus* in pure culture and the limit of detection (LOD) was established. The concentrations of emetic *B. cereus* ranged from 6.2 × 10^7 to 6.2 × 10^0 cfu/mL. As shown in Figure 6, the NC lane (the sterile water was used as the template for PCR) had no target DNA band, and when the concentration of emetic *B. cereus* was below the 10^4 cfu/mL, no target DNA band was exhibited on the gel. Thus, the LOD of the traditional PCR assay for the detection of emetic *B. cereus* in pure culture was 6.2 × 10^4 cfu/mL. All of the experiments were repeated in triplicate.

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**Figure 3.** (A) Optimization of graphene oxide (GO) volume, (B) optimization of catalytic hairpin assembly (CHA) reaction time, and (C) fluorescence intensity (a.u. = arbitrary units) under different incubation times of GO and CHA products. All of the tests were repeated in triplicate; each error bar represents SD. The F/F₀ represent the fluorescence intensity by CHA reaction with target single-strand DNA (ssDNA)/the fluorescence intensity by CHA reaction without target ssDNA.

**Figure 4.** The limit of detection (LOD) of the catalytic hairpin assembly (CHA)-graphene oxide (GO) assay for emetic *Bacillus cereus* in pure culture. All of the tests were repeated in triplicate. a.u. = arbitrary units. Each error bar represents SD, and the dashed line represents the cut-off [negative control (NC) + 3 SD].
DETECTION OF EMETIC BACILLUS CEREUS IN MILK

Application of the CHA-GO Method for Detection of Emetic B. cereus in Artificially Inoculated Milk Samples

To verify the practical application, the proposed method was assessed in artificially inoculated milk samples. The results showed (Figure 7) that the fluorescence intensity increased with the concentration of emetic B. cereus from $5.9 \times 10^1$ to $5.9 \times 10^8$ cfu/mL. A linear relationship between spiked emetic B. cereus from $10^2$ to $10^8$ cfu/mL and fluorescence intensity was given by $y = 0.2986x - 0.196$ ($R^2 = 0.9684$), where $y$ was the fluorescence intensity at 520 nm and $x$ was the concentrations of spiked emetic B. cereus. Based on NC + 3 SD, the LOD in spiked milk samples was $5.9 \times 10^2$ cfu/mL. All of the experiments were repeated in triplicate.

Specificity of the CHA-GO in Artificially Inoculated Milk Samples

The specificity of the proposed assay was also evaluated in artificially inoculated milk samples. The groups of bacteria ($10^7$ cfu/mL in the milk sample) were used to evaluate the specificity of the assay. Figure 8 showed that only in the presence of emetic B. cereus, a high fluorescence intensity would be obtained. This result indicated that the CHA-GO method had a high specificity to distinguish target bacteria or other bacteria in an artificially inoculated milk sample. All of the experiments were repeated in triplicate.

CONCLUSIONS

Recently, a series of methods for detection of foodborne pathogens has been established. However, these methods are time-consuming and have a high detection limit. In this study, an enzyme-free fluorescence platform for emetic B. cereus detection was developed. Fluorescence intensity was restored though CHA reaction, which was initiated by denatured PCR amplicons.
hairpin probes H1/H2 designed in this study achieved the specific detection of emetic B. cereus. Under optimal conditions, the CHA-GO method resulted in an LOD of 6.2 \times 10^2 cfu/mL in pure culture and 5.9 \times 10^3 cfu/mL in spiked milk, respectively. Although the method described in this manuscript is relatively time-consuming and costly, it provides an effective solution to apply CHA to the detection of pathogens, and a relatively sensitive result has been obtained. This work would be interesting to readers of the analysis of other food/dairy products, and by changing the PCR primer, the work has the potential to detect other bacteria.

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