Multiplex PCR for detection of MCR genes in clinical fecal samples

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Abstract. Plasmid-mediated colistin-resistance genes have been reported worldwide in recent years. A multiplex polymerase chain reaction (Multi-PCR) protocol was developed to detect transferable colistin-resistance genes (mcr-1 to mcr-6) in Enterobacteria for clinical laboratory purposes. The authors first designed six new primer pairs to amplify mcr-1 to mcr-6 gene products to achieve stepwise separation of amplicons between 87 to 216 bp, then divided these primers into two subgroups with the assistance of a pair of universal primers for the detection of currently described mcr genes and their variants in Enterobacteria. The protocol was validated by testing 29 clinical isolates of Escherichia coli of human origin, each well characterised and prospectively validated. The Multi-PCR assay showed full concordance with whole-genome sequence data and displayed higher sensitivity and 100% specificity. The assay could detect all variants of the various mcr alleles described. It was able to detect mcr-3 and mcr-4 as singletons or in combination. This type of test is critical for the epidemiological surveillance of plasmid-encoded resistance in limited resources conditions, and this method allows rapid identification of mcr-positive bacteria and overcomes the challenges of phenotypic detection of colistin resistance.

1 Introduction

Colistin (polymyxin E) has been used in veterinary medicine worldwide, but human use is restricted in terms of its nephrotoxicity and neurotoxicity [1-3]. The increasing emergence of multidrug-resistant gram-negative bacteria in the past decade has led colistin administration be in a dilemma as colistin sometimes act as the last-resort antibiotic for human infections [1]. Since the first report of mcr-1 [2], mobile colistin-resistance (MCR) genes have been found in many clinical and environmental samples. Hitherto, ten types of MCR genes have been reported worldwide [4-14], and MCR genes ranging from mcr-1 to mcr-6 are frequently found in enterobacteria [15]. Thus the regular surveillance of these human isolated MCR genes also help the appropriate use of colistin in human and animals.

The transferability, high carrying rate and wide distribution of the mcr-1 gene have aroused great concern and anxiety in society. The ongoing introduction of new MCR genes and its variants also is a disadvantage for correct global surveillance. Since the positive growth on selective enrichment media was not equivalent to the presence of mcr genes, neither phenotypic resistance nor growth on colistin selective media are perfect indicators for the presence of mcr genes [16], molecular detection methods that targets specific genes are needed. The current routine molecular procedures for MCR gene detection and confirmation depend largely upon technologies such as singleplex polymerase chain reaction (PCR) [17], multilocus sequence typing (MLST) [18] or whole-genome sequencing (WGS) [19] or sequencing approach. Considering the global concerns, we sought to develop and validate a rapid, efficient and inexpensive method that enables screening of relevant isolates and identification of those clinical samples that may harbour plasmid-mediated colistin-resistance genes.

The discovery of MCR genes has raised international concern because the spread of those genes might hamper the efficacy of colistin in clinical use. Therefore the carriage of mcr-1n health individuals was much more associated with the living next to a farm, the production of sheep and freshwater aquaculture, annual consumption of total meat, pork and mutton, and daily intake of aquaculture products[20,21]. It is urgent to develop a reliable molecular which can rapidly detect variable MCR genes popular in China. Here, we describe a multiplex PCR protocol for rapid and reliable detection of mcr-1, mcr-2, mcr-3, mcr-4, mcr-5 and mcr-6 genes which are frequently found in China for surveillance and epidemiological research, particularly in clinical faecal samples.

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2 Materials and methods

2.1 Bacterial strain and growth condition

Table 1. Bacterial strains used in this study

| Bacterial strain | Strains ID/Source | qPCR | Shen, et al.(2018) |
|------------------|-------------------|------|---------------------|
| MCR-positive strain (n=14) | | | |
| Klebsiella (n=3) | isolation 1* | + | + |
| Klebsiella pneumoniaispp. rhinoscleromatis | isolation 2 | + | + |
| Klebsiella pneumoniaispp. rhinoscleromatis | isolation 3 | + | + |
| Escherichia coli (n=11) | isolation 1 | + | + |
| Escherichia coli | isolation 2 | + | + |
| Escherichia coli | isolation 3 | + | + |
| Escherichia coli | isolation 4 | + | + |
| Escherichia coli | isolation 5 | + | + |
| Escherichia coli | isolation 6 | + | + |
| Escherichia coli | isolation 7 | + | + |
| Escherichia coli | isolation 8 | + | + |
| Escherichia coli | isolation 9 | + | + |
| Escherichia coli | isolation 10 | + | + |
| Escherichia coli | isolation 11 | + | + |
| None MCR-positive strain (n=19) | | | |
| Enterobacter (n=4) | | | |
| Enterobacter cloacae | CICC 21539b | - | - |
| Enterobacter sakazakii | ATCC 29544 | - | - |
| Enterobacter aerogenes | CICC 10293 | - | - |
| Enterobacter aerogenes | ATCC13408 | - | - |
| Escherichia coli (n=12) | | | |
| E. coli O157:H7 | CICC 21530 | - | - |
| E. coli O157:H7 | SZCIQ 13813 | - | - |
| E. coli O157:H7 | ADCPC 931 | - | - |
| Escherichia coli | ATCC 9637 | - | - |
| Escherichia coli | SZCIQ eco3 | - | - |
| Escherichia coli | SZCIQ eco5 | - | - |
| Escherichia coli | SZCIQ jm109 | - | - |
| Escherichia coli | CMCC 44104 | - | - |
| Escherichia coli | CMCC 44105 | - | - |
| Escherichia coli | CMCC 44106 | - | - |
| Escherichia coli | CMCC 44111 | - | - |
| Escherichia coli | CGMCC 1.129 | - | - |
| Klebsiella (n=2) | | | |
| Klebsiella pneumonia | CMCC 46102 | - | - |
| Klebsiella pneumonia | CICC 10781 | - | - |
| Salmonella (n=1) | | | |
| Salmonella dublin | GZCDCdb1a | - | - |

Table 1 lists the 33 bacterial strains used in this study. In addition to MCR-positive Enterobacteriaceae (MCRPE), most MCR gene host bacteria are associated with E. coli, Salmonella, K. pneumoniae and Moraxella [22-25]. Strains with strain identification were purchased from culture collections. Isolates of MCRPE from fecal samples were obtained as described by Shen et al. [17]. All strains listed in Table 1 can be isolated from fecal samples and were genetically identified and characterised by 16S RNA sequencing, multilocus sequence typing (MLST) or conventional methods. The strains were grown on appropriate media before DNA extraction. Specifically, Salmonella spp., E. coli and Shigella spp. were aerobically incubated at 37 °C for 24 h in Luria-Bertani (LB) broth (Gangdong Huankai Microbial Sci & Tech Co, Ltd, Guangzhou, China), and K. pneumoniae was aerobically incubated at 37 °C for 24 h in nutrient broth (Gangdong Huankai Microbial Sci. & Tech. Co., Ltd.).

2.2 Primer design and DNA extraction

Fig. 1. The nucleotide sequences of the mcr alleles obtained from the GenBank.

The nucleotide sequences of the mcr alleles were obtained from the GenBank database (http://www.ncbi.nlm.nih.gov/) and a comparison was made. Figure 1 lists the accession numbers of these obtained from the
database. The primers were based on the presumptively conserved sequences of these six genes and their variants using Primer Express 3.0 [26]. To minimise the putative interactions among primers, primer combinations were evaluated for the formation of primer-dimer structures.BLAST analysis was then used to assess the primers’ specificity (http://www.ncbi.nlm.nih.gov/). Cell pellets from 1-mL bacterial cultures were used to isolate bacterial DNA using the QIAamp DNA mini kit (Qiagen, USA) or the Tiangen Bacteria DNA Kit (Tiangen Biotech Beijing Co., Ltd., Beijing, China) according to the manufacturer’s instructions. The concentration and purity of the extracted DNA samples were spectrophotometrically determined.

### 2.3 Real-time PCR analysis

Resolution melting real-time PCR amplification of small regions of the MCR genes was performed with a total volume of 20 μL, which contained 2 μL DNA template of each bacterium (0.5 μmol/L), 10 μL of Premix Taq (Ex Taq Version 2.0) (Takara Bio Group, Dalian, Japan), 2 μL (10 μmol/L) of primers, 1.25 μL of Eva-Green (Takara) and 0.2 μL of bovine serum albumin (20 mg/mL); nuclease-free water was then added to bring the total reaction volume to 20 μL. Nuclease-free water was used as the no-template control. The optimised experimental protocol for the ABI 7500 sequencer (Life Technologies, Inc., USA) was as follows: 95 °C for 10 min, 40 cycles at 95 °C for 10 s and 60 °C for 50 s. A melting curve temperature profile was finally undertaken from 60 °C to 95 °C at a melting rate of 0.2 °C/s. EvaGreen was used as the intercalating dye, and the data were analysed with the manufacturer’s software (ABI 7500 software 2.3). For Multiplex PCR, the universal bacteria primers were added at the concentration of 0.4 μmol/L, and other primers were added equally to achieve a system containing primers at concentration of 1 μmol/L.

### 2.4 Analytical specificity of real-time PCR assay

Thirty-three DNA templates were included in this assay, including 14 target bacteria and 19 non-target bacteria (as shown in Table 1). The analytical specificity was expressed as the percentage of samples with known identity of the target genes that returned a positive outcome in the assay, whilst the diagnostic specificity was expressed as the percentage of non-target samples with known identity that returned a negative outcome in the assay. All available samples were used to determine the analytical and diagnostic specificity of identification of MCR-positive types.

### 2.5 Analytical sensitivity of high-resolution melting real-time PCR assay

To evaluate the analytical sensitivity of the multiplex PCR method, each strain of mcr-positive bacteria was cultured for 16 h at 37 °C in tryptic soy broth and used for DNA extraction as described above. DNA from each strain was purified from liquid culture using the Genomic DNA Purification System (Promega, Madison, WI, USA). The DNA concentrations were spectrophotometrically determined as described above. A series of 10-fold dilutions for each purified MCR DNA was used for real-time PCR. All samples were processed in triplicate.

### 2.6 Detection of mcr variants in faecal samples

One hundred twelve fresh faecal samples were collected from healthy people with the MGI Easy collection suite for intestinal microbial genetic testing; these samples were suspended in buffer solution (version 3.0, for research only; MGI) and stored at 4 °C for less than 1 week. After no less than 10 hours of enrichment in colistin-chromagar media, each sample was submitted to the detection methods described by Shen et al. [17] and to our newly developed multiplex PCR.

### 2.7 Statistical analysis

All statistical data analyses were performed using Microsoft Excel. Means and standard deviations of the Tm values were calculated.

### 3 Results

#### 3.1 Primer design and real-time PCR analysis

Using the DNA from the six bacteria strains positive for mcr variants (Table 1) as the template, we amplified products ranging from 87 to 216 bp with the specific primer pairs. The differences between the amplified fragments, which were amplified by different primers, resulted in various melting temperature (Tm) values in the six mcr variants. We firstly designed six pairs of primers targeting on six mcr gene types (for example primers for mcr-3/mcr-4 shown in table 2), and then artificially synthesized new primers by adding individual nucleotides as the universal bacteria primers (as shown in table 2) to the 5’ end of the chain while do not change the targeting sites (for example primers for mcr-1/mcr-2/mcr-5/mcr-6 shown in table 2). Then we separately test the efficiency of each primer pairs via PCR procedure described in section 2.3. Specifically, for the primers without bacterial universal primers, the Tm peak of mcr-I was found at 84.8±0.01 °C, the Tm peak of mcr-2 was detected at 85.2±0.01 °C, the Tm peak of mcr-5 was observed at 84.1±0.01 °C, the Tm peak of mcr-6 was found at 83.3±0.02 °C, the Tm peak of mcr-3 was detected at 80.7±0.01 °C, and the Tm peak of mcr-4 was observed at 74.8±0.06 °C. For the primers adding by bacterial universal primers, the Tm peak of mcr-1 was found at 84.8±0.02 °C, the Tm peak of mcr-2 was detected at 85.2±0.01 °C, the Tm peak of mcr-5 was observed at 84.1±0.02 °C, the Tm peak of mcr-6 was found at 83.3±0.02 °C, the Tm peak of mcr-3 was detected at 80.7±0.01 °C and the Tm peak of mcr-4 was observed at 74.8±0.06 °C.
### Table 2. Primers and amplification fragment.

| Group   | Primers match | Primers sequence | Fragment length (bp) |
|---------|----------------|------------------|----------------------|
| A       | Universal primer | F: 5’ TAGCACACGCAGAGTA CTAGCT-3’ | mcr-1: 216bp |
|         |                 | R: 5’ CGAGACAGCAGTCAAT ACCGTC-3’ |               |
| mcr-1   |                 | F: 5’ TAGCACACGCAGAGTAC GTAGCTCGTGTAGCTAGCTCTCT-3’ | mcr-2: 193bp |
|         |                 | R: 5’ CGAGACAGCAGTCAAT ACCGTCGAGTCTAGTCTAGCT-3’ |           |
| mcr-2   |                 | F: 5’ TAGCACACGCAGAGTAC GTAGCTCGTGTAGCTAGCTCTCT-3’ | mcr-5: 140bp |
|         |                 | R: 5’ CGAGACAGCAGTCAAT ACCGTCGAGTCTAGTCTAGCT-3’ |           |
| mcr-5   |                 | F: 5’ TAGCACACGCAGAGTAC GTAGCTCGTGTAGCTAGCTCTCT-3’ | mcr-6: 152bp |
|         |                 | R: 5’ CGAGACAGCAGTCAAT ACCGTCGAGTCTAGTCTAGCT-3’ |           |
| B       | mcr-3           | F: 5’ TGCTCATCGTACGTTCA GGT-3’ | mcr-4: 87bp |
|         |                 | R: 5’ ATGGTGTGCTATAGTT CTTAAGT-3’ |               |
| mcr-4   |                 | F: 5’ TTAAACACCTACGACAC ACT-3’ | mcr-5: 90bp |
|         |                 | R: 5’ CGAACATATCTCTGGTTA CTT-3’ |               |

#### 3.2 Multiplex PCR for detection of homogeneous mcr variants

To achieve simultaneous detection of the currently described mcr genes and their variants, we developed several multiplex PCR strategies that involves testing the efficiency of random combinations of primers. Finally, a multiplex PCR strategy consisting of two PCR reactions was screened. Figure 2 presents the melting curves of the six mcr variants’ gene sequences when the homogeneous mcr variant was applied to the newly developed multiplex PCR. Using the DNA of the six bacteria strains positive for mcr variants (Table 1) as the template, products between 87 and 216 bp (Table 2) were amplified with specific primer pairs. The resolution melting curve profiles with distinct Tm peaks were consistently obtained, and each variant was represented by a single peak ranging from 74.8 to 85.2 °C (Figure 2). Specifically, in group A, the Tm peak of mcr-1 was found at 84.8±0.02 °C, the Tm peak of mcr-2 was detected at 85.2±0.01 °C, the Tm peak of mcr-3 was observed at 84.1±0.02 °C and the Tm peak of mcr-4 was found at 83.3±0.05 °C. In group B, the Tm peak of mcr-3 was detected at 80.7±0.02 °C, and the Tm peak of mcr-4 was observed at 74.8±0.05 °C (Figure 2).

#### 3.3 Multiplex PCR for the detection of mixed mcr variants

Since we have developed a multiplex PCR strategy that could identify mcr-1 to mcr-6 when they are separately existing, we following try to test its efficiency when several different types of mcr gene variants co-existing in one sample. According to the results of our pre-tests, primers that target mcr-1 and mcr-3 react as templates for each other, and the Tm peak of the by-product was 79.9 °C. In reality, mcr variants scarcely co-exist in a single sample. But to test the efficiency of this multiplex PCR, we mixed six mcr variants proportionally and tested the melting resolution curve profile. When mcr-1 and mcr-3 co-existed (Fig. 3a), in group A, the Tm peak of the amplification fragment was found at 84.8±0.02 °C, which was identical to the result of the single mcr-1 test; in group B, the Tm peak of the amplification fragment was detected at 79.6±0.05 °C, which was identical to the by-product. These findings indicate that multiplex PCR could detect only mcr-1, rather than both, when mcr-1 co-existed with mcr-3. When mcr-2, mcr-5 and mcr-6 co-existed (Fig. 3b), in group A, the Tm peak of the amplification fragment was found at 84.1±0.02 °C, which was identical to the result of the single mcr-5 test; in group B, there was no positive amplification, which indicates that multiplex PCR can only detect mcr-5 when mcr-2, mcr-5 and mcr-6 co-exist. When mcr-3 and mcr-4 co-existed (Fig. 3c), in group A, the Tm peak of the
amplification fragment was found at 79.6±0.05 °C, which was inferred as the by-product; in group B, the Tm peak of the amplification fragment was found both at 74.8±0.05 °C and at 80.7±0.01 °C, which was identical to the findings for singleton mcr-3 and mcr-4, respectively. This indicates that multiplex PCR can successfully detect mcr-3 and mcr-4 when they co-exist. In summary, the newly developed multiplex PCR not only detects all MCRs (mcr-1 to mcr-6) effectively when present in the sample as singletons, it can also detect specific MCR genes when they co-exist.

Fig. 3. Multiplex PCR for the detection of mixed mcr variants

3.4 Analytical specificity of multiplex PCR assay

In this study, we assessed the analytical specificity of multiplex PCR analysis with DNA from 33 strains (Table 1) that potentially hosted mcr variants. After multiplex PCR, positive amplification curves were observed only if the MCR genes (mcr-1 to mcr-6) existed in the culture, and the peaks of the dissolution curves were consistent with those of any standard dissolution curves of mcr-1 to mcr-6. If the MCR genes did not exist in the culture medium to be examined, no amplification signal was found. These primers have good sensitivity and specificity. Fourteen MCR-positive bacteria strains (3 K. pneumoniae and 11 E. coli) were correctly identified, whilst no positive fluorescent signal was observed from the 19 non-target controls, which indicates that the method has good specificity.

3.5 Analytical sensitivity of multiplex PCR assay

In this study, we determined the analytical sensitivity of the multiplex PCR analysis by analysing various dilutions of purified DNA for each mcr-positive bacteria with 10-fold diluted genomic DNA from 1 ng to 0.01 pg. Figure 4 shows that the detection limit was 1 pg per reaction for mcr-5 and mcr-6, 10 pg per reaction for mcr-1 and mcr-2, and 100 pg per reaction for mcr-3 and mcr-4. As shown in Figure 4a to 4d, when the template concentration was lower than the detection limit, the curve shows a peak of the by-product around 79.9 °C, which was identical to that seen in Figure 2a to 2d.

Fig. 4. Analytical sensitivity of the Multiplex PCR assay. The resolution melting curves of 10-fold diluted genomic DNA of mcr-positive E. coli: dilutions 1 ng to 0.01 pg, respectively

3.6 Detection of mcr variants in faecal sample

One hundred twelve fresh faecal samples were collected from healthy people with the MGI Easy collection suite for intestinal microbial genetic testing; the samples were suspended in buffer solution (v3.0, for research only, MGI) and stored at 4 °C for less than 1 week. The results show that the positive rate in our study was 13.4% (15/112). While the that positive rate of the same samples was 12.5% (14/112) when there were screened by the method of Shen et al. [17]. This method of extracting DNA from the sample to be measured as a template with good detection rates by boiling method positive MCR genes could be detected both with multiplex PCR and with traditional methods. However, the entire newly developed multiplex PCR procedure can be performed within 1 day.

4 Discussion

Plasmid-mediated resistance genes have the potential for further transmission in regions with high levels of antimicrobial resistance and global spread to multidrug-resistant bacteria [27]. Colistin has become a last-resort antibiotic for infection of some multi-drug-resistant gram-negative strains [28]. Early detection of MCR-positive bacteria would help setting policy in preventing further spread and providing appropriate antimicrobial therapy. Despite of traditional loop-mediated isothermal amplification, multiplex-LAMP assay has been developed and successfully detection ofmcr-1 to mcr-5in colistin-resistant bacteria [29,30]. MLST was used to detect and characterise ESBL-producing E. coli that express mcr-1 obtained from dairy cows in China [18]. In addition to traditional molecular detection, novel methods have also been recently developed, including matrix-assisted laser desorption/ionisation—time of flight mass spectrometry [31], Raman spectroscopy combined
with hierarchical cluster analysis [32] and liquid chromatography–dual mass spectrometry [33]. Other methods such as the DPA-based BMD test (colistin-MAC test) could be implemented by clinical laboratories that were unable to perform molecular tests and was able to screen large collections of isolates to reveal the expression of new mcr-like genes that were not targeted by current molecular assays [34], whilst the accuracy of the colistin-MAC test still requires improvement. These methods are limited by their sensitivity and requirement for instruments. For example, the detection sensitivity and specificity of detection methods based on Raman spectroscopy were found to be 90.9% and 91.1%, respectively, compared with the reference broth micro-dilution method [32]. In our study, in total of fourteen MCR-positive bacteria strains (3 K. pneumoniae and 11 E. coli) were correctly identified, whilst no positive fluorescent signal was observed from the 19 non-target controls. These primers were proofed to have good sensitivity and specificity. Furthermore, the entire newly developed multiplex PCR procedure can be performed within 1 day. In the present study, there solution melting analysis was applied to pure cultures and clinical samples to evaluate its use in detection and species identification of MCR genes in a single real-time PCR assay. The sensitivity and specificity of the mcr real-time PCR assay were both 100% in a method validation performed with a set of 112 previously well-characterised bacterial isolates that contain mcr-positive or mcr-negative bacteria. This newly developed assay also led to a significant reduction in the assay volume and reagent costs. This method has potential to be as a low-cost approach of high-throughput screening method for Enterobacteriaceae isolates in which a low prevalence of colistin resistance is expected with minimal hands-on time and costs, and it will assist in combating the spread of colistin resistance in bacteria for its shorter reaction times, simpler instruments and greater sensitivity while showing the same high specificity.

For genotyping of MCR variants, DNA is usually screened firstly with PCR and followed by sequencing of the amplicons [35,16]. To distinguish colistin-resistant isolates that carry the plasmid-encoded mcr genes, many researchers have developed multiplex PCR or other high-throughput detection methods, including in silico tested primers for PCR assay (mcr-1 to mcr-7) [36], multiplex PCR (mcr-1 to mcr-5) [37-39] and multiplex PCR (mcr-1 and mcr-2) [40]. While methods discussing the detection efficiency and distinguish ability when two or more types of MCR genes exist in one clinical sample are scarce. The multiplex PCR system developed by Lescat et al. can identify a single isolate with both mcr-1 and mcr-5 genes [38]. Although mcr-1 and mcr-5 were not found together, 4 of the 152 samples (2.7%) were positive for both mcr-1 and mcr-3, which indicates that mcr-1 and mcr-3 did appear at the same time [41]. Our results show that multiplex PCR can successfully detect mcr-3 and mcr-4 when they co-exist. With the supplement of PCR-A and PCR-B, the kit can successfully detect mcr-1 and mcr-3 when they co-exist. Furthermore, the PCR-A reaction system of this kit uses a combination of a large number of universal primer pairs and a very small number of specific primers. Specific primers specifically bind to target fragments and amplify them with universal primers using the initial amplified fragments as templates. This combination design can minimise the interaction between primers and improve the efficiency of multiple PCR.

5 Conclusions
Colistin is one of the last-resort antibiotics for treatment of multidrug-resistant infections in humans, but transmissible colistin-resistance genes have emerged in bacteria obtained from animals, humans and even food chain. The rapid and sensitive detection bacteria that carry these genes in clinical samples is critical to help control further spread. Here we describe a method for broth enrichment of colistin-resistant E. coli from human faecal samples followed by multiplex-PCR for simultaneous detection of two of the main colistin-resistance genes, mcr-1 to mcr-6. Real-time monitoring of the PCR products will greatly reduce the monitoring mobility of colistin tolerance. The purpose of the newly developed multiplex PCR is to provide a method to detect the MCR gene, which according to the existing technology rarely covers all MCR genes (ranging from mcr-1 to mcr-6). Based on all MCR gene sequences found to date, we designed two multiplex PCR systems for detection of all MCRs. This method is especially suitable for medical, clinical or environmental monitoring scenarios. It can detect any MCR genes in faecal samples quickly, simply and effectively. The real-time monitoring of PCR products will greatly reduce the resource consumption in our daily monitoring task. Furthermore, the group A system consists of a large portion of universal primer pairs and a very small portion of specific primers. These specific primers specifically bind to target fragments and amplify them by universal primers using the initial amplified fragments as templates. This combination strategy can minimise the interaction between primers and improve the efficiency of multiple PCR. The method adopts PCR technology as the fluorescence detection method, and the reaction was carried out in a closed-tube reactor to prevent PCR electrophoresis from forming aerosol pollution caused by false-positive results.

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