Single step for *Neisseria gonorrhoeae* genomic DNA detection by using gold nanoparticle probe.

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**Abstract**

*Neisseria gonorrhoeae* is a cause of gonorrhea which is the most common sexually transmitted diseases (STDs). Bacterial culture is a gold standard method for this bacterial detection, but it is time-consuming and skilled technicians are needed. Therefore, the development of a rapid, easy and sensitive detection technique is still required. In the present study, we developed a specific gold nanoparticle probe for the detection of *N. gonorrhoeae* genomic DNA compared to a conventional PCR method. The average size of the synthesized gold nanoparticles (GNPs) was approximately 13 nanometers. GNP probes could bind specifically with complementary sequences of DNA. Due to their salt-tolerance, GNPs solution remained in red color, which could be detected by naked eyes, after adding 100 mM MgSO₄ solution. This method is easy and rapid with the detection limit of 20 ng/µL genomic DNA of *N. gonorrhoeae*. Twelve strains of bacteria were tested by this detection method and the results showed high specificity to *N. gonorrhoeae*. In conclusion, this developed method has the potential to apply for medical diagnosis.

**Keywords:** *Neisseria gonorrhoeae*, Medical diagnosis, Gold nanoparticles, DNA detection, DNA probe.

**Abbreviations**


**Introduction**

Gonorrhea is a sexually transmitted disease caused by *Neisseria gonorrhoeae* [1]. The rate of gonococcal infection is still one of the most common worldwide, even using the condom [2], because of untreatable multidrug-resistance [1]. Those who have asymptomatic infections with *N. gonorrhoeae*, with no treatment, may get many complications in future such as prostatitis, orchitis, ectopic pregnancy, abortion and conjunctivitis in foetus.

Methods for *N. gonorrhoeae* detection include bacterial culture, gram stain, conventional Polymerase Chain Reaction (PCR), hybridization, nucleic acid amplification test (NAAT) and serological test. Bacterial culture is a gold standard method for *N. gonorrhoeae* detection with great specificity, but also low sensitivity and time-consuming [3,4]. Gram-stained method has high sensitivity and specificity for the diagnosis of symptomatic men with urethral discharge but it is not appropriate for diagnosis of cervical, rectal and pharyngea gonorrhea, or for asymptomatic patients because of its low sensitivity [5]. PCR sensitivity and specificity depend on primer design [6]. NAAT is more sensitive than culture technique, especially for pharyngeal and rectal specimens, but inhibitory substances in patient specimens may cause the false-negative results [7]. Hybridization method is a rapid and specific diagnosis of pathogens that are fastidious or cannot be cultured [6].

Taken together, we aimed to develop an accurate and rapid detection of *N. gonorrhoeae* by modification of hybridization technique to obtain higher sensitivity, cheap and rapid compared to culture technique. Specific DNA probe for *N. gonorrhoeae* conjugated with GNPs was designed for the specific sequence in the *porB* gene which codes for the major outer membrane protein (*PorB*) of *N. gonorrhoeae*. *PorB* can
act as an anion-selective porin, interact with host cells, induce or inhibit apoptotic signaling in eukaryotic cells and has a relationship with human host pathogenesis [8,9]. The sensitivity and specificity of this method were also studied.

Materials and Methods

Bacterial strains and reagents

*N. gonorrhoeae* were obtained from the Clinical Microbiology Laboratory of Srinagarind Hospital, Khon Kaen University, Thailand. Reference strains of *Klebsiella pneumonia*, *Corynebacterium* spp., *Pseudomonas aeruginosa*, *Staphylococcus epidermidis*, *Proteus mirabilis*, *Escherichia coli*, *Enterococcus* spp., and *Staphylococcus aureus*, Group B *streptococcus*, *Listeria monocytogenes* were obtained from the Department of Microbiology, Faculty of Associated Medical Sciences, Khon Kaen University. HAuCl$_4$$\cdot$4H$_2$O and sodium citrate dihydrate were purchased from Sigma-Aldrich Co., LLC (Missouri). The other chemicals were all of analytical grade.

Probe and primers design

The sequences of a DNA probe and two primers were designed from the *porB* gene. All oligonucleotide probes were generated as sequences using clustalW under the high homology among the *N. gonorrhoeae*. The NCBI BLAST (Basic Local Alignment Search Tool) search of the base target in *porB* gene showed highly specific for *N. gonorrhoeae* and no exact match against non-gonorrhoeae family members. Probes were labeled with thiol group (-SH) at 5’ end to enable conjugation with the GNPs surface. The probe and primer set were synthesized and then purified by Pacific Science Co., Ltd., Thailand.

*N. gonorrhoeae* detection by PCR method

The reaction mixtures of PCR were carried out in a final volume of 25 µL containing 50 mM KCl, 10 mM Tris-HCl pH 8.3, a 0.2 mM concentration of each dNTP, 2 mM MgCl$_2$, 3 unit of thermostable DNA polymerase and 0.3 µM of each primer. The primary PCR amplification was performed using primers which listed in Table 1. The primary PCR program was preceded by a 4-min denaturation step at 94°C, followed by 40 cycles of 94°C for 1 min, 40°C for 1 min, and 72°C for 2 min, and the last cycle was followed by an additional 10-min elongation step at 72°C. PCR amplicons were analyzed by electrophoresis at 100 V in 1.5% agarose gels staining with ethidium bromide and visualized under UV light.

Preparation of gold nanoparticles

Gold nanoparticles (GNPs) were synthesized by the slight modification of the citrate reduction [10]. Briefly, 100 mL of 1 mM H-AuCl$_4$$\cdot$4H$_2$O solution was mixed with 10 mL of a 38.8 mM trisodium citrate solution and refluxed for 15 min and then cooled down to room temperature. The morphology and size of the GNPs were characterized by using a field emission high-resolution transmission electron microscope (TEM) (TECNAI G2 20 Model, Hillsboro, OR) operating at 200 kV. GNPs sample was prepared for TEM analysis by dropping them onto a carbon film copper grid and drying at room temperature. The result of TEM was analyzed with Image J (NIH, Bethesda, Maryland) for measuring the average size of GNPs. The absorption spectra (400-700 nm) were also measured using a SpectraMax M5 fluorescence microplate reader (Molecular Devices LLC, Sunnyvale, CA, USA).

Preparation of oligo probe-conjugated GNPs

The conditions for labeling the GNPs with the *N. gonorrhoeae* oligonucleotide probes were modified from a previous study [10]. The reaction mixture of GNP probes was carried out at a final concentration of 0.50 and 1 µM oligonucleotide probes containing 1 mL of GNPs solution (GNP-0.5 µM probe and GNP-1 µM probe, respectively) and incubated on an orbital shaker overnight at 50°C. The phosphate buffer, SDS, and NaCl solution were added to obtain a final concentration of 10 mM phosphate buffer (pH 7.4), 0.01% (W/V) SDS, and 0.1 M NaCl. The mixture was incubated on an orbital shaker for 48 hours at 50°C, followed by centrifugation at 15,000 rpm 4°C for 30 min to remove unconjugated oligonucleotides. The GNP probes were washed with washing buffer (100 mM PBS containing 100 mM NaCl and 0.01% SDS) twice, then suspended in 200 µL of washing buffer and stored in dark at 4°C.

Estimation of the percentage of GNP probes

The concentration of unconjugated oligonucleotide probes after conjugation with GNPs was estimated by high-performance liquid chromatographic technique (HPLC) using an indirect assay [11]. After the reaction mixture of 1 mL of GNPs, probe (a final concentration of 0.50 µM oligonucleotide probe containing 1 mL of GNPs solution), phosphate buffer, SDS, and NaCl solution were performed; a final concentration of probe was 0.43 µM. The supernatant from the GNP probe conjugation step was injected into a Luna C18 (15 cm × 3.0 mm) HPLC column (Phenomenex, California). Gradient elution was conducted at a flow rate of 1.5 mL/min using 0.1% trifluoroacetic acid and 60% acetonitrile in 0.1% trifluoroacetic acid at ratios of 100:0, 80:20, and 0:100 at 0-5, 5-10, 15-25, and 30 min, respectively. The concentration of 0.43 µM of the oligo probe was used as a standard. Peak area corresponded to unbound probe in the supernatant was calculated with an oligonucleotide probe standard curve.

Genomic DNA extraction

The bacteria were cultured in chocolate agar at 37°C for 18-24 hours. Colonies were transferred to new tubes in 200 µL of deionized water and then boiled in boiling water for 10 min, then centrifuged at 13,000 rpm for 10 min and the supernatant was collected. The concentration of genomic DNA was quantified by measuring the optical density at 260 and 280 nm by using a NanoDrop 2000 spectrophotometer (NanoDrop Technologies, Wilmington, DE, USA). The supernatant was kept at -20°C until use.
GNP probes hybridization assay

Optimization of salt concentration: The 3 µL of 50 mM of N. gonorrhoeae genomic DNA were added to 3 µL GNP probes. The mixtures were incubated at 95°C for 5 min, followed by 63°C for 5 min. After that MgSO₄ at various concentration (3, 15, 30, 60, 80, 100 mM) was added. The positive result was red or pink and the negative result was blue solution which could be observed by naked eye. Deionized water was used as a negative control.

The sensitivity of GNP probes: N. gonorrhoeae genomic DNA was serially diluted to be 2.5, 5, 10, 20, 40, 60 ng/µL. Each genomic DNA concentration was hybridized with GNP probes. After the hybridization step, MgSO₄ solution was added.

The specificity of GNP probes: Genomic DNA of N. gonorrhoeae and other 11 bacterial strains were diluted to final concentration of 50 ng/µL. All bacterial strains were hybridized with GNP probes. After the hybridization step, MgSO₄ solution was added.

Results

Characterization of GNPs

These GNPs, prepared by sodium citrate reduction method, were spherical shape and size distribution about 13 nm diameter by TEM study. The maximum absorption of GNPs was 520 nm (Figure 1).

Estimation of the percentage of GNP probes

To evaluate the percentage of the GNP probes, the supernatant from the conjugation step of GNP probes was analyzed by HPLC and compared with a standard DNA probe (Figure 2). Peak area of unbound oligo probes and 0.43 µM of the standard oligo probes were 966 and 3647. The percentage of unbound oligo probes was 26.49%. Thus, the percentage of GNP probes was approximate 73.51%.

Figure 2. HPLC chromatogram of the 0.43 µM of the standard oligo probe (blue line) and the supernatant which was collected from the conjugation step of GNP probes (black line).

Optimization of salt concentration

After hybridization step, each of MgSO₄ concentrations was added into mixtures containing GNP-0.5 µM probe and N. gonorrhoeae reaction mixtures. The results showed that 30, 60, 80, and 100 mM MgSO₄ can distinguish N. gonorrhoeae from negative control (Figure 3). The concentration of MgSO₄ at 100 mM was selected as an optimum salt concentration.

Figure 3. Optimization of MgSO₄ concentrations in GNP-0.5 µM probe and Neisseria gonorrhoeae reaction mixtures. (NC) = negative control, (PC) = positive control, (+) = positive result, (-) = negative result.

The sensitivity of GNP probe

GNP probes were hybridized with 2.5, 5, 10, 20, 40, 60 ng/µL of N. gonorrhoeae genomic DNA by using sterile deionized water as a negative control (Figure 4). The solution containing 20, 40 and 60 ng/µL of N. gonorrhoeae genomic DNA remained red solution after MgSO₄ was added, whereas the solution containing genomic DNA concentration less than 20 ng/µL and negative control solution turned into purple. The sensitivity in between GNP-0.5 µM probe and GNP-1 µM probe was not different. Therefore, GNP-0.5 µM probe was selected to evaluate the specificity of this method. The detection limit was 20 ng/µL genomic DNA.

Figure 4. Characterization of GNPs by using UV–vis absorption spectroscopy and transmission electron microscopic (TEM) techniques.
Figure 4. Comparison of sensitivity in detection among various probe concentrations (A) GNP-0.5 µM probe (B) GNP-1 µM probe. (NC) = negative control, (+) = Positive result, (-) = Negative result.

The specificity of GNP probes

To assess the specificity of GNP-0.5 µM probe, the probes were hybridized with 12 bacterial strains including *N. gonorrhoeae*. Only reaction containing *N. gonorrhoeae* genomic DNA turned into red solution, whereas other reactions with 11 bacterial strains turned into purple solution after addition of 100 mM MgSO$_4$ (Figure 5).

Figure 5. Specificity of the GNP probes at probe concentration of 0.5 µM (A) The probes were hybridized with 1: Neisseria gonorrhoeae, 2: Escherichia coli, 3: Staphylococcus aureus, 4: Staphylococcus epidermidis, 5: Group B streptococcus, 6: Enterococcus spp., 7: Listeria monocytogenes, 8: Corynebacterium spp. (B) The probes were hybridized with 1: Neisseria gonorrhoeae, 2: Klebsiella pneumoniae ATCC 700603, 3: Klebsiella pneumonia ATCC 27736, 4: Pseudomonas aeruginosa, 5: Proteus mirabilis, (+) = positive result, (-) = negative result.

*N. gonorrhoeae* detection by PCR

In this study, the PCR technique was used as a reference method. We used specific primers to *porB* gene of *N. gonorrhoeae* and probe sequence was between the DNA sequence of PCR product. PCR products contained a band of 284 bp in length. The *N. gonorrhoeae* and other bacteria were examined by PCR assay to confirm the presence of the *porB* gene, which was the target gene of the GNP-probe assay. *N. gonorrhoeae* exhibited a band in gel electrophoresis at about 300 bp, whereas the others showed negative results (Figure 6). The result of PCR method was similar to the result of specificity by using GNP-probe.


Discussion

The incidence of gonorrhea are about 106 million adults each year worldwide, and infection rates in the developing countries are about 10-20 times more than in the developed countries [12]. Although bacterial culture is a gold standard method to identify *N. gonorrhoeae* infection but it has low sensitivity and time-consuming [12]. The other methods also have both advantages and disadvantages [5-7]. Therefore, we develop a novel detection method for diagnosis of *N. gonorrhoeae*. A previous study reported that colorimetric assay using GNP conjugated with thiol-modified oligonucleotide probes are rapid and the product result from this reaction can be observed with the naked eye without complicated and expensive instrumentation [13]. The colorimetric using GNPs-probe assay is faster than culture method and requires only a thermal block. Forming the linkage of thiol-ssDNA and GNPs for gold nanoprobe assay is easy, inexpensive in common molecular assays, and this solution can be kept at room temperature for 1 month [14,15]. In the present study, we used ~13 nm diameter GNPs for the colorimetric detection of DNA because the result of hybridization can be easily detected visually [11,16]. The hybridization of GNPs-probe with the target DNA formed double-stranded DNA helix that made GNPs more salt-tolerant than single-stranded DNA (No target DNA). Therefore, the hybridization of GNPs-probe with the target DNA showed red solution. The concentration of MgSO$_4$ was optimized after hybridization. A 100 mM MgSO$_4$ was used in this study due to the specificity test of GNPs-probe colorimetric assay, genomic DNA of *N. gonorrhoeae* and some other bacteria could not be discriminated by naked eye at less than 100 mM MgSO$_4$ (data not shown). Therefore, some concentration of MgSO$_4$ can distinguish between negative control (deionized water) and *N. gonorrhoeae* but not between bacterial strains. The detection limit of our study was 20 ng/µL unamplified genomic DNA.
which is better than a previous study that the limit of detection was ~54 ng/µL [17]. The different detection limit may arise from the different preparation step of GNPs-probe. The developed GNPs-probe assay had specificity to *N. gonorrhoeae* DNA. The result of PCR method was related to the result of GNP probe colorimetric assay in term of specificity and can confirm that positive control is true.

**Conclusion**

The present study reports a novel method for *Neisseria gonorrhoeae* genomic DNA detection based on GNP probe colorimetric assay. This developed method can distinguish positive and negative control by naked eye. The detection limit of this method was 20 ng/µL genomic DNA. It is a rapid and easy method with high sensitivity and specificity. This method also has a potential method to apply for medical diagnosis.

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**Conflict of Interest**

The authors declare that there are no conflicts of interest.

**References**


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