APPLICATION OF GRAPHENE OXIDE ON APTAMER-BASED BIOSENSOR DEVELOPMENT FOR AUTHENTICATION OF GELATIN

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ABSTRACT

Objective: The objective of this study was to perform aptamer selection using systematic evolution of ligands by exponential enrichment (SELEX) method which assisted by graphene oxide against target of porcine gelatin (non-halal gelatin).

Methods: The aptamer selection was carried out using SELEX method without target immobilization. Selection of aptamer capable of binding porcine gelatin by applying graphene oxide (GO) was known as GO-SELEX. The selection process was initially carried out by incubation of single-stranded DNA (ssDNA) libraries targeting on porcine gelatin with the addition of graphene oxide. The selected ssDNA was then purified by several stages namely, symmetric PCR amplification, purification of products with DNA purification kits, asymmetric PCR amplification, and continued purification of DNA with native PAGE. The analysis of each stage was done by agarose gel electrophoresis.

Results: The results showed that aptamer targeting porcine DNA could be selected. This was indicated by the results of DNA analysis using native polyacrylamide gel electrophoresis (PAGE) in which sharp separation band with a base length equivalent to the marker of the ssDNA library (about 80 base pair) was obtained.

Conclusion: Aptamer targeting on porcine gelatin has been successfully developed using GO-SELEX method. GO can increase selectivity in developing aptamer which will be used as a biosensor to detect porcine gelatin. The method could be proposed as a standard of aptamer based method for porcine gelatin detection on halal products authentication.

Keywords: Graphene oxide, Aptamer, Porcine gelatin, GO-SELEX

INTRODUCTION

Gelatine, obtained from partial hydrolysis of collagen of bones and skins of pig and cattle is commonly used in several industries including in food, pharmaceuticals, and cosmetics. In pharmaceutical application, gelatine is commonly exploited for preparing either hard or soft capsule shells [1] and components in pharmaceutical formulation [2, 3]. The encapsulated active pharmaceutical ingredient can be protected from moisture, heat or other extreme conditions in order to enhance its stability. It is reported that the main sources of gelatine used in pharmaceutical industries are cattle bones accounting of 23.1%, bovine skin accounting of 9.4% and pigskin accounting of 46.4%. Currently, gelatine is also prepared from bones accounting of 23.1%, bovine skin accounting of 9.4% and pigskin accounting of 46%. Generally, gelatine is derived mainly from bovine and porcine skins of pig and cattle is commonly used in several industries in pharmaceutical industries.

The moslem community, especially the follower of school of thought (madzhab) syafi'i is not allowed to consume food and pharmaceutical products containing porcine gelatin [6], as a consequence, some methods for identification of gelatin sources has been extensively proposed and applied. Physico-chemical methods based on spectroscopic such as Fourier transform infrared spectroscopy (FTIR) [7] and chromatographic like liquid chromatography [8], mass spectrometry [9] have been widely applied for differentiation among gelatines, however, this method is lack in selectivity, thus molecular biology-based methods were currently used for gelatin analysis.

DNA-based methods especially polymerase chain reaction is an excellent method to be applied for identification of gelatin sources. Real-time PCR [10], Restriction Fragment Length Polymorphism (RFLP) [11], PCR multiplex [12], and species-specific PCR [13] have been used for identification of DNA from pig species (porcine DNA). Aptamer, a short chain oligonucleotide of RNA or single-stranded DNA (ssDNA), can bind selectively ligand with high specificity [14]. Aptamer can bind several target molecules including DNA, peptide, nucleotide and other molecules with low molecular weight [15]. Aptamer could be obtained from multiple selection in vitro secara using the method known as SELEX (Systematic Evolution of Ligands by Exponential Enrichment). SELEX comprised 4 processes, namely binding, elution, amplification and separation. SELEX has emerged as a powerful method for isolation of nucleic acid aptamer with high sensitivity and selectivity and ease in its production [16].

Nanomaterial-based on graphene oxide has been widely used for making functional biosystem integrated with nucleic acids, peptides, protein and even with cells [16]. The capability of ssDNA to be absorbed on graphene and the nature difference between ssDNA and dsDNA in its retention has led the development of biosensor [17]. This study was aimed to perform a selection of aptamer using SELEX method assisted by graphene oxide (GO-SELEX) against analyte target (porcine gelatin). Aptamer obtained was used as biosensor using nanoparticle platform for detecting porcine (Sus scrofa domesticus) DNA.

MATERIALS AND METHODS

This research was intended to perform aptamer selection using SELEX method modified with the use of graphene oxide (GO). The selection was carried out by a series of processes and separation of aptamer using Sigma/Sartorius 3-18K andinkubator (Memmert In55). The separated aptamer was amplified using Bio-Rad T-100™ Thermal Cycler, analyzed using electrophoresis gel agarose and visualized with UV illuminator.

Primer design

Primer ssDNA libraries were designed based on gene Cyt-b babi with accession number: AF939170 [18]:

Forward primer: 5'-GGCTAAATCTCCCCCTCTCAGCTGTA-3'.
Reverse primer: 5'-ATGAAGAGCCATAATAGTTTTTCG-3'.

ssDNA library was synthesized randomly with the end of forward primer and its complement reverse primer as:
The designed primer and ssDNA was synthesized in Integrated DNA Technologies (IDT) through PT. Genetika Science, Jakarta, Indonesia. The primer sequence is 5’-GCCTAAATCTCCCTCAATGGTA—N40—CGAAATCTATTGGCCTCTT-3’.

Application of graphene oxide in aptamer selection

The library of ssDNA was reconstituted with binding buffer pH 7.4 and heated at 94 °C for 5 min. The mixture was cooled in ice bath for 15 min and allowed to stand at 25 °C for 10 min. Selection cycle 1: in the microtube, 1, 2, 3 mmol ssDNA incubated with 5, 10, 15 mg of porcine gelatin in 400, 300, 200 µl binding buffer was added with ethanol %, stirred for 2 hour at the temperature of 25 °C. The mixture was then added with 100, 200, 300 µl graphene oxide (2 mg/ml) into mixture, incubated at 25 °C for 40 min (mass ratio of graphene oxide/ssDNA = 40: 1). The mixture was centrifuged at 13000 rpm for 10 min. The supernatant containing ssDNA binded on gelatin was pooled, while deposite (ssDNA which was not binded and adsorbed on graphene oxide) was removed [19].

Purification of an aptamer using asymmetric PCR

The selection products were purified using asymmetric PCR which was started by amplification using PCR according to [20] with slight modification. The amplification step was: initial denaturation of 95 °C for 1 min, denaturation at 95 °C for 15 sec, annealing at 49.05 °C (from annealing temperature optimization) for 15 sec, extension at 72 °C for 10 sec, with PCR cycle of 25-35, and final extension at 72 °C for 5 min. PCR products were purified using purification kits of DNA Clean and Concentrator™ 5 from Zymo Research. The purified DNA was then amplified using asymmetric PCR with primer ration of 10/4, 12/2, 14/0, followed with the ratio of forward/reverse primers of: 5/0, 10/0, 15/0, 20/0. The products were analyzed using gel agarose 3% in buffer TAE (40 nM Tris-acetate and 1 mmol EDTA, pH 8.0) containing 0.5 µg/ml ethidium bromide and visualized with UV Transilluminator.

Purification of ssDNA gel

PCR product was precipitated with absolute ethanol until final volume of 20 µl. The product was purified with 10% native polyacrylamide gel electrophoresis, TBE buffer (89 mmol tris-Boric, 2 mmol EDTA, pH 8.3), and was stained with 0.5 µg/ml ethidium bromide in 1x buffer TBE for 7 min. ssDNA band was purified using crush and soak method. Band containing ssDNA was cut using the sterile cutter and was then placed into Effendorf. Gel was destroyed using steril pipette tip followed by addition of 400 µl eluting buffer (1 M Na-acetat; 0.01 M EDTA). The sample was incubated at 37 °C for 6-8 h with shaking at 900 rpm. The gel particles were precipitated with centrifugation for 2 min and the supernatant was precipitated with ethanol with final volume of 20 µl [21].

RESULTS AND DISCUSSION

The development of non-halal component detection becomes important to provide comfort for the ummah. Methods or detection tools that have high accuracy and sensitivity, inexpensive, and within easy reach will greatly assist the monitoring process. One of the potential methods to be applied in a non-halal analysis is based on a biosensor using aptamer, a short-stranded oligonucleotide in the form of RNA or single-stranded DNA (ssDNA) that can be selectively bound with ligands or target proteins with high affinity and specificity [22]. Graphene and graphene oxide (GO) are fairly wide-ranging two-dimensional nanoparticles used as platforms in biosensors. The difference in the affinity of graphene oxide to ssDNA and dsDNA is then adopted in the test to observe DNA.

The process was started by designing and synthesizing sequence library of nucleic acids (DNA/RNA) randomly, which were normally composed of aptamer with variation until ~10^15 capable of binding with analyze target (porcine gelatin) specifically. To obtain a variety ssDNA library, the bases of A: G: C: T with ratio of 1.5: 1.25: 1.15: 1.0 were used. The library diversity was determined by chain length among bases of ssDNA with the surface of GO. In order to separate ssDNA which do not bind porcine gelatin, the mixture was incubated at 25 °C for 40 min. The separation of ssDNA which was not bound to analyte target was carried out by addition of graphene oxide (GO). The binding capacity of GO toward ssDNA was enhanced by incubation at 25 °C for 40 min. ssDNA sequence which do not bind porcine gelatin could be adsorbed by GO via π-π interactions among bases of ssDNA with the surface of GO. In order to separate ssDNA bound and not bound with porcine gelatin, the centrifugation of mixture at 13000 rpm for 10 min was applied. The supernatant was ssDNA binding porcine gelatin, while precipitate was ssDNA which not bound and adsorbed by GO. The DNA profile could be separated and was analyzed by electrophoresis gel agarose 3% was shown in fig. 1.

Fig. 1: The electrophoresis of DNA on gel agarose; (a) using agarose 3%, (b) using agarose 0.8%. The arrow (-----) indicated band of ssDNA library as control
The DNA as result of selection was analyzed using electrophoresis gel agarose with 2 different agarose concentrations, namely 0.8\% and 3.0\% using buffer TAE 1x. The ssDNA library was used as marker to investigate whether products have the same length with the used library. The result showed that isolate has not appeared band clearly yet. This is due to low concentration of DNA target so that the isolate was continued with PCR amplification. To amplify ssDNA, symmetrical PCR was used, i.e. amplification of ssDNA target and its complementary DNA using forward and reverse primers with the same ratio. The results of symmetrical PCR was then purified using DNA purification kits using protocol set in the manufacture procedure. The purified products were then purified again using asymmetrical PCR, using primer ratio of x/0, i.e. forward primer was varied while reverse primer was made constant. The objective of ssDNA purification using asymmetrical PCR was to amplify template sequence which complements with ssDNA target.

The results of asymmetrical PCR using master mix KAPA Taq Ready Mix were then analyzed using electrophoresis gel agarose 3\% as shown in fig. 2. The result showed that all primer ratios of 5/1, 10/1 and 20/1 could obtain amplification products toward DNA target. This was indicated by band which parallel to marker band (ssDNA library). The next step was an amplification of ssDNA with primer ratio of x/0 (no reverse primer used during PCR analysis) so that the amplification only occurred in DNA template. The results could be seen in fig. 3.

![Fig. 2: Electrophoresis gel agarose 3\% of PCR products using primer ratio x/1. The ssDNA was marker. The arrow (→) indicated amplification band](image1)

Asymmetrical PCR with a decreased ratio of reverse primer (fig. 4) showed the different profiles of electrophoresis results. Reverse primer with ratio of 14/0 could yield higher amplification products, as indicated by clearer separation band under UV illuminator. In addition, using asymmetrical PCR with primer ratio x/o (5/0, 10/0, 15/0, and 20/0), the amplification products were shown in fig. 5. It can be stated that the higher the level of forward primer, the higher the amplification products.

![Fig. 3: Electrophoresis gel agarose 3\% of PCR products using primer ratio x/0; 5/0; 10/0; 20/0. The arrow (→) indicated amplification band](image2)
During purification of ssDNA, PCR product was precipitated until final volume of 20 μl with absolute ethanol. The product was purified on 10% native polyacrylamide electrophoresis (PAGE), TBE buffer (89 mmol tris-Borat, 2 mmol EDTA, pH 8.3), and stained with 0.5 μg/ml ethidium bromide in 1x bufer TBE for 7 min. ssDNA was purified using crush and soak. Band containing ssDNA was cut using the sterile cutter and placed into Eppendorf. Gel was destroyed using a sterile pipette tip followed by addition of 400 μl eluting buffer (1M Na-acetate; 0.01 M EDTA). A sample was incubated at 37 °C for 6-8 h with shaking at 900 rpm. Gel particles were precipitated using centrifugation at maximum speed for 2 min. The supernatant was precipitated carefully using ethanol until the final volume of 20 μl. The results of PAGE 10% of ssDNA and DNA ladder were shown in fig. 6.
CONCLUSION
Graphene oxide (GO) could be used to assist aptamer selection without immobilization process. The ability of GO could adsorb ssDNA specifically, not in dsDNA. In addition, the use of GO could assist the purification of selected aptamer because the residue of ssDNA which not bound to target (porcine gelatin) was bound into GO so that ssDNA target could be separated by centrifugation. Furthermore, the selected Aptamer can be used as a biosensor to detect porcine gelatin contamination for halal authentication purposes.

ACKNOWLEDGMENTS
The author thanks to the Ministry of research and higher education for financial support during this study through Penelitian Unggulan Perguruan Tinggi 2017-2018 awarded to Prof. Dr. Sismindari.

AUTHORS CONTRIBUTIONS
All the author have contributed equally

CONFLICT OF INTERESTS
The author declares that there is no conflict of interest

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