Development of Tetra-primer Amplification Refractory Mutation System (ARMS) PCR for Detection of CHRNA3 rs8040868

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Received date: Nov 23, 2020; Revised date: Apr 19, 2021; Accepted date: Apr 21, 2021

Abstract

BACKGROUND: Single nucleotide variations (SNV) have been mapped to be associated with several human conditions and diseases. To validate the association between SNV to certain human traits or diseases, a large number of subjects must be included. Thus, in need of a fast, relatively economic, and reliable genotyping method. This can be achieved through the use of tetra-primer amplification refractory mutation system polymerase chain reaction (Tetra-primer ARMS PCR). This study reports strategy to develop Tetra-primer ARMS PCR-based genotyping of CHRNA3 rs8040868.

METHODS: The optimization of Tetra-primer ARMS PCR was done through these steps: identification of gene sequence and position of single mutation; designing outer and inner PCR primers; amplification of target gene fragments through PCR by using outer primer; confirming genotype of the PCR product by using sequencing; determining an optimum ratio of outer and inner primer; and determining optimum annealing temperature and cycles for the PCR program. The PCR products were run in 2% gel agarose electrophoresis and visualized under UV illumination.

RESULTS: Outer and inner primer ratio of 1:3 with annealing temperature of 64.4°C and 40x cycles was found to be the most optimum condition. Tetra-primer ARMS PCR was able to confirm the results of the DNA sequence of 2 samples, confirming wild-type variants (TT allele) and the heterozygous mutant (CT allele).

CONCLUSION: Tetra-primer ARMS PCR was able to genotype rs8040868 of the CHRNA3 gene.

KEYWORDS: tetra-primer ARMS PCR, CHRNA3, rs8040868, genotyping

Indones Biomed J. 2021; 13(2): 192-200

The variation of CHRNA3 rs8040868 is one of the known SNV in chromosome 15. This synonymous mutation (Valine) changes the allele from Thymine (T) to Cytosine (C) in region Exon 2. Detailed characteristics of this variant taken from the National Center for Biotechnology Information (NCBI) database can be found in Table 1.

Located in the 15q25.1 locus, the CHRNA3-CHRNAS-CHRNB4 gene cluster is known to be associated with smoking-associated cancer. Genetic variation of CHRNA3 rs8040868 is considered as a pathogenic mutation associated with a phenotype of non-small cell lung cancer.
Table 1. *CHRNA3* rs8040868 variant information.

| Gene Variant | Position | Region | Changes | Allele | Effect |
|--------------|----------|--------|---------|--------|--------|
| *CHRNA3* rs8040868 | chr15:78,618,839 Exon 2 | c.159 A>G | T/C | Synonymous variant (p. Val53=) |

All information was taken from the NCBI database (http://www.ncbi.nlm.nih.gov/).

in Iran.(4) This variant increased the risk of getting non-small cell lung cancer incidence among the Han Chinese population (OR = 2.50; 95% CI = 1.04-5.97).(5) This variant also had a high linkage disequilibrium with the *CHRNA5* rs16969968 variant, a pathogenic variant that changes amino acid from aspartic acid to asparagine. Mutant allele C from the *CHRNA3* rs8040868 is known to be associated with a decreased expression of the *CHRNA5* gene in lung tissue.(6)

Over the last years, genome-wide association studies and other candidate gene analyses have been done to map SNV and its association with various physiological and clinical traits in humans. To validate the association of SNV to a disease, a large number of subjects must be studied, thus making scientists in need of a feasible genotyping method without expensive tools or reagents.(7) The advanced laboratory uses DNA microarray genotyping/DNA biochip for mass genotyping analysis. This advancement is difficult to operate in general molecular laboratories commonly operated in developing countries such as Indonesia, due to expensive infrastructure and consumables. An example of genotyping methods that utilize common qualitative polymerase chain reaction (PCR) machines with more robust, more efficient, and yet valid is Tetra-primer Amplification Refractory Mutation Systems PCR (Tetra-primer ARMS PCR). This method uses two different sets of primers, outer and inner primer pairs. The outer primers were used to amplify the target gene sequence, while the inner primers were used to identify base variations that occur in the gene sequence.(8) Variations of alleles will be on different band sizes which can be differentiated by further visualization by using UV after gel electrophoresis. PCR efficiency and gene specificity will be ensured by outer primer pairs while allele specificity will be ensured by inner primer pairs.(2)

Tetra-primer ARMS PCR is an accurate, easy, practical, and inexpensive genotyping method. The required tools and materials are the same as conventional PCR. Further, the result of Tetra-primer ARMS PCR can be detected by gel electrophoresis to determine the genotype of the samples tested.(8) Besides, Tetra-primer ARMS PCR has advantages to detect variants present in low percentage in specific disease due to the ‘mutant enrichment’ method. Four low penetrance variants in breast cancer prediction had been successfully genotyped using this method.(9,10) Therefore, this method can be used to detect somatic mutation (i.e., tumor tissue, leukemia, etc.), not just germline variants from blood.(10)

This manuscript describes the strategy used to develop Tetra-primer ARMS PCR-based genotyping for rs8040868 (T/C) variation. Steps used within this report can also be used as a template to develop Tetra-primer ARMS PCR-based genotype detection.

### Methods

The experiment was performed under the approval of the ethics committee of the Faculty of Medicine, Public Health, and Nursing, Universitas Gadjah Mada (No. KE/0922/EC/2019).

### Samples

We used a total of 10 DNA samples isolated from the peripheral blood of volunteers. Two samples were used in the Tetra-primer ARMS PCR optimization step, while 8 were used in the validation step.

### DNA Extraction

Control genomic DNA used for this study was obtained from the buffy coat of peripheral blood of adult volunteers (n=2), coded as K1 and K2. DNA extraction was done by using QIAamp DNA Mini Kit No. 51304 (Qiagen, Germantown, MD, USA). The procedures were carried out according to the kit. DNA quantity DNA was examined by using spectrophotometer NanoDrop (Thermo Fisher Scientific, Waltham, MA, USA) and ran in 2% gel agarose electrophoresis (GeneDireX, Taoyuan, Taiwan) to check its quality.

### Primer Design

The minor allele frequency (MAF) of rs8040868 was checked in several genome projects based on its closure to the Indonesian population (Table 2). This step was done to make sure that the variant can be found in our tested samples. The DNA sequence of *CHRNA3* rs8040868 was
identified from an accessible NCBI database (http://www.ncbi.nlm.nih.gov/). Two specific sets of outer and inner PCR primers (later written as “primers”) were designed using web-based software Primer1 (http://primer1.soton.ac.uk/public_html/primer1.html)(8) Primer’s specificity to detect CHRNA3 gene was tested using NCBI BLAST program (http://www.ncbi.nlm.nih.gov/blast). The primers used for Tetra-primer ARMS PCR for detection CHRNA3 rs8040868 were provided in Table 3. DNA sequence used to design primers, the schematic illustration of the Tetra-primer ARMS PCR for CHRNA3 rs8040868, and its expected PCR product were started in Figure 1. Further, gene fragments of CHRNA3 from two DNA samples (K1 and K2) were amplified using the designed outer primers. Results were run in 2% agarose electrophoresis to observe the target-size DNA fragment.

**Confirmation of Genotype**

The genotype of the PCR product with the expected DNA size (Figure 2) was further confirmed by Sanger DNA sequencing (Figure 3). PCR product and the outer primer in a total volume of 30 µL, respectively, were sent to 1st Base Sequencing Laboratories (Selangor, Malaysia) for sample processing. The result was analyzed using free website software Poly Peak Parser (http://yosttools.genetics.utah.edu/PolyPeakParser/).

**Optimization Steps of Tetra-primer ARMS PCR**

Optimization of Tetra-primer ARMS PCR was done in three steps, which include: determining the ratio of outer and inner primers; searching for the optimum annealing cycle and temperature; and using the correct concentration of the PCR reagents to minimize nonspecific bands. The total volume of PCR reaction was 20 µL; containing 10 µL GoTaq® Green Master Mix (2X Green GoTaq® Reaction Buffer (pH 8.5), 400 µM dATP, 400 µM dGTP, 400 µM dCTP, 400 µM dTTP, and 3 mM MgCl₂ (Promega Corporation, Madison, WI, USA), an optimized ratio of outer and inner primers (each primer has working stock concentration 10 nM/µL), and genomic DNA (50-200 ng/µL). DNA amplification was done by using Thermocycler PCR ProFlex Applied Biosystem (Life Technologies Holdings Pte, Ltd, Singapore).

The optimization of Tetra-primer ARMS PCR was performed by applying different ratios of outer and inner primer concentration between 1:1, 1:2, 1:3, and 1:4. The annealing temperature was determined by lowering the average melting temperature by 5°C and gradually increased.
Figure 1. Tetra-primer ARMS PCR for *CHRNA3* rs8040868. A: DNA sequence used to design primers for *CHRNA3* rs8040868. The green-colored sequence indicates the specific exon where the variants present. The yellow-colored sequence indicates the changed codon. B: Schematic illustrations of the Tetra-primer ARMS PCR assay for genotyping. T and C indicate the allele changes between thymine to cytosine. C: Expected product from Tetra-primer ARMS PCR after ran on 2% agarose gel electrophoresis.

Figure 2. DNA quality of K1 and K2 samples, and conventional PCR product with outer primers under 2% agarose gel electrophoresis. Single K1 and K2 bands showed good and intact genomic DNA. The use of outer primers produced 353 bp of DNA band, as expected. (L: DNA ladder; K1: sample 1; K2: sample 2; KN: negative control).

Results

Table 3 shows the MAF of the reference population of the Malay, Chinese, Vietnamese, and East Asian population. We concluded it is possible to detect the *CHRNA3* rs8040868, as the MAF was spanned from 0.51-0.344.

Further, a set of outer primers were designed based on the *CHRNA3* DNA sequence that includes the rs8040868 locus. Following PCR using the outer primers, the product obtained confirmed that the outer primer set has successfully amplified targeted DNA fragments of *CHRNA3* at the correct size, 353 bp. The same PCR products were analyzed by the Sanger sequencing method, confirming the sequence...
Table 4. Reagent mix for Tetra-primer ARMS PCR CHRNA3 rs8040868.

| Reagent                        | Working Concentration | Volume |
|--------------------------------|-----------------------|--------|
| GoTaq® Green Master Mix        | 10 µL                 |        |
| Nuclease Free Water            | 7.4 µL                |        |
| Forward outer primer           | 10 nM                 | 0.2 µL |
| Reverse outer primer           | 10 nM                 | 0.2 µL |
| Forward inner primer           | 10 nM                 | 0.6 µL |
| Reverse inner primer           | 10 nM                 | 0.6 µL |
| DNA                            | 50-200 ng/ µL         | 1 µL   |
| **Total volume**               |                       | 20 µL  |

Figure 3. Sanger sequencing result of K1 (A) and K2 (B) samples. The red line shows the place where the “keywords” were found. The yellow-colored arrow shows that the K1 has G codon in forward sequence and C codon in reverse sequence; the blue line indicates that the changes are heterozygous, thus the K1 sample has CT allele. The K2 has A codon in forward sequence and T codon in reverse sequence, making the sample have the TT allele (wild-type). (Alt: alternative allele, means the sequence that is analyzed; Ref: Reference allele).
Figure 4. The optimization process of Tetra-primer ARMS PCR to detect CHRNA3 rs8040868. PCR products run in 2% of agarose gel. A: PCR with a variety of outer and inner primer ratios of 1:2, 1:3, and 1:4 in 64.4°C, 65.4°C, 66.4°C, and 67.4°C annealing temperatures (see the label in each picture), all done in 35x PCR cycles. Results showed an annealing temperature of 64.4°C as the optimum PCR condition, which showed the heterozygous band in K1 (yellow arrow). B: To get a clearer band, an additional cycle was added. Optimization with 1:3 outer and inner primer ratio by using 64.4°C annealing temperatures with 40x annealing cycle to increases specificity of the PCR process, shows the best result (L: ladder; K1: sample 1; K2: sample 2; KN: negative control).
Table 5. Tetra-primer ARMS PCR CHRNA3 rs8040868 program.

| Reagent          | Times    | Temperature | Cycle number |
|------------------|----------|-------------|--------------|
| Initial denaturation | 2 minutes | 95°C        | 1 cycle      |
| Denaturation      | 30 seconds | 95°C        |              |
| Annealing        | 30 seconds | 64.4°C      | 40 cycles    |
| Elongation       | 1 minute  | 72°C        |              |
| Termination      | 5 minutes | 72°C        | 1 cycle      |
| Cooldown         | ~        | 4°C         |              |

**Discussion**

SNV is a molecular marker which was widely used because of their ubiquitous distributions throughout the genome. The increasing number of SNV markers has led to increasing demand for numerous cost-effective genotyping platforms. (11) Genotyping to detect genetic susceptibility can be done by several methods, including sequencing, genotyping array, Restriction Fragment Length Polymorphism (RFLP), and RT-PCR with probes.(7) Several studies have reported the use of different genotyping methods to detect SNV. (12-15) Despite that, each method has different advantages and disadvantage to consider. Sequencing and genotyping array are a high-throughput genotyping technique but they need expensive machineries and reagents. Using qPCR will equally requires expensive reagents; the later technology the more expensive the technology. PCR-RFLP is more common to be used in less sophisticated research laboratory, as it needs only conventional PCR machine and generic PCR reagents, but it is limited as its analysis will be only possible if the SNV creates a restriction enzyme site.(7) Thus, this is a two-steps process prior to detection by using agarose gel electrophoresis, can not be used for mass analysis, and often not reproducible.

The application of Tetra-primer ARMS PCR to identify genetic variation has been widely reported. Tetra-primer ARMS PCR has been known to be the combination of Tetra-primer PCR and ARMS PCR, by using conventional PCR machine and generic PCR reagents. Tetra-primer ARMS PCR has a higher sensitivity and accuracy in comparison to ARMS PCR, specifically in distinguishing normal and heterozygotes allele.(16) Tetra-primer ARMS PCR is a reliable method to analyze point mutations. It delivers fast results and enables the identification of specific genotypes in a single PCR step. It is also less time-consuming, once the melting temperature is optimized, and more affordable than genotyping using restriction enzyme.(17) This method can detect insertion and deletions, such as variation of rs4343 A>G and rs4340 insertions/deletions in artery coronary disease study.(18) Also, this method can be used to detect variations in miRNA genes, such as miR-146a rs2910164 C>G gene in breast cancer study.(19) Not only in humans, but Tetra-primer ARMS PCR also has been reported to detect g.4140 A>G polymorphism in the SIRT2 gene of cattle.(20)

This study follows our previous optimization strategy to detect variation CYP2E1 gene.(1) First, we design the outer and inner primers using Primer 1.(21) Primer design is a critical step to ensure the success of the development of Tetra-primer ARMS PCR.(2) A successful primer set depends on the primer Tm, GC content percentage, and the selection of mismatch bases.(21) The interaction between the inner and outer primers may also affect the specificity of the reaction.(2)

Designing outer primer is important to amplify selected DNA target, that further becoming positive control once the genotype confirmation is done. Genotype confirmation by using sequencing is needed to validate whether the PCR product indeed amplifies the correct DNA sequence. Choosing the correct inner primer pairs is the key element in the development of Tetra-primer ARMS PCR, as this determines the genotype of DNA sample variant. To design the inner primers, the free software Primer1 equips the inner primer with the mismatch.(1,8,16,21) We designed the inner primer where each has 2 mismatches, at the 3’ end and position -2 from the 3’ end. The latter mismatch is supposed to increase the specificity of allele detection, as consequently DNA polymerase will only able to elongate when its 3’ end is complemented perfectly to the template.(7) It is also important to choose a set of inner primers which has a target band with a minimum difference of 50 bp, so it can be separately differentiated using 2% gel electrophoresis.
The second step was finding the optimum concentration ratio of the outer and inner primer.(1) The use of more than one pair of primers poses a challenge. Having two sets of primers will increase the chances of getting non-specific products that can be seen when they were visualized by showing multiple DNA bands. This result happens because dimers may be formed between the primers, and many unwanted reactions may increase during the PCR cycle.(7)

To overcome this, having the most suitable ratio between the outer and inner primer is a must. Previous studies reported that the ratios between the outer and inner primer differed depending on which codon was amplified. (22) Some studies reported different ratios one to another, but they always use less concentration of outer primer than inner primer.(8,23,24) Reducing the concentrations of the outer primer will enhance the amplification of the allele-specific product. Additionally, we also found that lowering the outer primers concentration truly gives the best result even though it may give a lesser clear DNA band under UV transillumination, similar to the previous finding.(8)

We were also searching for the optimum annealing temperature simultaneously with finding the best primer ratio. Both factors were solved by using a gradient PCR system which allows different annealing temperatures in one PCR process. For the optimum annealing temperature, 5°C lower than average melting temperature is suggested to show the most stable product.(8) This suggestion was confirmed in our study. However, other studies also showed that the optimum annealing temperature was closer to the average melting temperature.(1,7) We concluded that different variants may give different results; so, optimization still requires a variety of temperatures. It is also important to try adding PCR cycles if the expected DNA band cannot be seen clearly under the UV transilluminator.

This study posed a limitation, as we could only detect 2 of 3 expected variants, i.e., homozygous wild type and heterozygous mutant, despite our attempt to use DNA samples from 10 different subjects (part of data not shown). Additionally, the use of this method by using different DNA sources needs to be validated.

Conclusion

Tetra-primer ARMS PCR was successfully employed to accurately determine the genotype of the rs8040868 variant in the CHRNA3 gene. This method was simple, affordable, and can be easily adapted to detect other genetic variants.

Acknowledgements

This work was supported by “Hibah Rekognisi Tugas Akhir (RTA)”, Universitas Gadjah Mada, Indonesia year 2020 (PI: Fachiroh J). The authors would also thank the donors and technicians who participated in this research.

Authors Contribution

JF, ALD, DK designed the study. ALD carried out the laboratory work. ALD, DK, JF analyzed the data. ALD and JF wrote the manuscript. All authors read and approved the final version of the manuscript.

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