CODIS STR LOCI (CSF1PO, THO1, TPOX, vWA) GENETIC VARIATION ANALYSIS IN MADURESE

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ABSTRACT

Endogamy continues to occur among the Madurese people in rural areas of the island of Madura, especially those areas of the smallest islands around the mainland of Madura. Endogamy as seen from a genetic standpoint will increase the frequency of homozygote genotypes. With regard to genetic variations, STRs of nuclear DNA and polymorphisms in mtDNA are frequently examined. Genetic variations in humans undergo an evolutionary process through the accumulation of changes in DNA sequence, i.e. through the process of nucleotide substitutions that evolves in number with the directional development of lineage. So far, the genetic variations among the populations in Madura Island have not been known. The present study was an observational analytical research with the purpose of determining the genetic variations in STR CODIS in the populations of Madura Island. Results indicated that, based on loci allele: CSF1PO, THO1, TPOX, and vWA, there was homozygote genotypes. The allele variations is not specific for Madurese ethnic but this variations may represent married model in Madurese ethnic. According to Mustama (2007), a gene pool is not only a collection of genes but a dynamic system organized and containing the past history of a population. Any genetic information has certain historical, anthropological and statistical aspects necessitating an interdisciplinary coordination and collaboration. (FMI 2016;52:1-6)

Keywords: Genetic variations, CODIS STR, Madurese ethnic

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INTRODUCTION

Madurese communities have a distinctive, unique, stereotypical, and stigmatic culture. Madura island population group is an ethnic group that largely until today still retain indigenous ancestral traditions coexisting in harmony with their religion. Marriage in the community of remote areas in Madura island, especially in smallest island, still occur among their own ranks (endogamy). Actually, the reason they practically do endogamous marriages is that the treasure they have will not lost/moved to other family, as well as strong family ties. Endogamous marriage seen from a genetic genotype standpoint will increase the frequency of homozygote. Increased genetic homogeneity will appear if the endogamous marriage occurs continuously between generations up to the point where all alleles in one locus or all loci become homozygous (Bodmer & Cavalli-Sforza 1976).

Ease of transportation in the territory of Madura island with its Suramadu bridge will open wider opportunities
for the people of Madura island’s mobilization. Madurese may look for a life partner outside the island so that exogamous marriage becomes possible more than endogamous marriage. According to Bodmer & Cavalli-Sforza (1976) concept of marriage between populations is the basis of population genetics analysis observed from genes frequency and explains the process of change in gene frequency in the population.

In genetic variation, the frequently used examination is the Short Tandem Repeats (STR) on deoxyribonucleic acid (DNA) nucleus and polymorphisms in mitochondrial DNA. Regions in STR are generally area that do not code for proteins, but has a simple repetition of the nucleotide motif. Alleles at STR locus vary greatly depending on the number of units of its motives, so that it can be used to study population sub-structure and the short-term evolution as well as to measure the phylogenetic relationships between populations. Therefore STR is a marker for selection in observing population proximity from the point of ethnicity, language, culture and history of a population (Sahoo & Kashyap 2005).

Based on the above observations, genetic variation will vary across each population and has a history of occurrence of those variations, so that the study of genetic variation in a population is still relevant. This study was expected to find genetic variations that include STR alleles pattern in populations of Madura island and yield loci mapping that can be used as a reference in Forensic DNA profiling in forensic DNA laboratory in Indonesia.

MATERIALS AND METHODS

The type of this study was observational conducted to determine genetic variation through CODIS STR (CSF1PO, THO1, TPOX, VWA) in Madurese, with cross sectional design. Samples were Madurese ethnic groups residing on the island of Madura, parents or grandfather/grandmother married to fellow Madurese.

Material in this study were bloodstains-derived DNA. Materials for DNA extraction: DNAzol Reagent, 100% ethanol solution, 70% ethanol solution, and distilled water. Materials for PCR: PCR Mix, DW sigma. Primary Short Tandem Repeat (STR), which includes loci: THO1 (5'-CTTGGGACAGTGAGG CAGCGTCT-3' and 5'-TGCCGAAGTCATCTCTA CAGTC-3'), TPOX (5'-ACTGCGACAGAACAGGCT ACATAG-3' and 5'-GGAGGAACTGTGGAAGGAC CACACAGT-3'), CSF1PO: (5'-AACCCTGAGTCTGGCAAAGAGACTAGC-3' and 5'-TTCCACACACCACACTGCACATCTCTC-3'), VWA: (5'-CTTAGTGGATG AAAATAATCAGT ATG-3' and 5'-GGACAGAT GATAAATACATAG GATGAGATGG-3'). Materials acetaldehyde and loading dye, electrophoresis: Bis-acryl-amid, Agarose, Temed, Tris Boric EDTA (TBE) 0.5%, 100 bp marker, marker K562, and 0.03% bromphenol blue. Research instruments were cycle PCR, DNA sequencer ABI Prism 310, spectrophotometer, electrophoresis, whirli mixer, centrifuge, Eppendorf micropipette, micropipette tips, UV transluminator, Polaroid camera, Trassonnic 310, spinator, Eppendorf tubes, and microwave.

Bloodstain DNA extraction

DNA was extracted by piercing the FTA card that contains the sample by means of Harris punch of 2.0 mm, added 100 mL, FTA purification reagent, and incubated for 10 min at room temperature. It was stirred with pipette and exhausted, repeated 3 times. 100 mL TE-1 buffer was entered, incubated for 10 min at room temperature, stirred with a pipette and then the TE-1 was removed. It was repeated 2 times. Samples were dried and stored with the addition of 30 mL TE buffer.

PCR amplification

DNA amplification by PCR method was done with the protocol of Promega Corp, 2001. PCR amplification for THO1, CSF1FO, TPOX was as follows: Phase I: Initial denaturation 96°C for 2 minutes; Phase II: Cycle 1 (10 times), consisting of subsequent denaturation 94°C for 1 minute, annealing 64°C for 1 min, extension 70°C for 1 min 30 sec; 2 cycle (30 times), consisting of: denaturation 90°C for 1 min, annealing 64°C for 1 min, and extension 70°C for 1 minute 30 seconds. As for the VWA was follows: Phase I: Initial denaturation 96°C for 2 minutes, phase II: Cycle 1 (10 times), which consists of Subsequent denaturation 94°C for 1 min, Annealing 600°C for 1 min, Extension 700°C for 1 min 30 seconds.

Electrophoresis

This stage was done by using polyacrylamid composite agarose gel (PAGE) stained with silver staining. PAGE procedure was as follows: agarose gel was made from 30 cc EDTA 0.5x Tris Boric and Agarose (LE Analytical Grade) 0.15 grams, heated in a microwave until crisp then cooled to a temperature of 50°C. Then, it was added with acrylamid bis (N, N'-Methylenebis-acralade) 4.5 cc and Temed (N, N', N'-Tetramethyl-enediamine) 15 ul, ammonium persulfate was further added 100 ul, and then poured into the mold (Gel bed), waited until it froze, DNA produced from PCR 12.5 ul
with loading buffer 2 ul was included and run at voltage of 70 volts for 2 hours.

PAGE silver staining procedure was as follows: drying: (20% methanol + 2% glycerol) in 100 cc distilled water for 5 minutes, fixation (10% ethanol + 5% acetic acid glycerol) in 100 cc distilled water for 20 minutes, and washed/rinsed with distilled water 1x quickly. Staining: 0.1% AgN03 in 100 cc distilled water for 50-80 minutes. Developing: (NaOH + 100 ul 1.5% Formalin) in 100 cc of distilled water, then observed with a UV lamp until it looked clear.

RESULTS

Polymerase Chain Reaction (PCR) amplification process was begun with DNA template preparation through the process of cell lysis of the blood spots using FTA card kit. PCR results visualization on CSF1PO of 295-327 bp can be seen in Fig. 1.

**Fig. 1.** Visualization of PCR results at CODIS STR loci CSF1PO; 295-327 bp. (M) markers, (K +) positive control K562 allele 12 (11,12,13,21,22,23,31,32) samples.

Fig. 2 shows the emergence of the tape/band between 295-327 bp that CSF1PO locus primer pairs used are specific, only attaches to the expected position. PCR process was carried out to amplify CODIS STR loci using primer pair M1 and M2. PCR products were analyzed by PAGE with silver staining. As well as showing allele homozygotes (12.12) and heterozygous (12.11).

**Fig. 2.** Visualization of PCR results at CODIS STR loci CSF1PO; 295-327 bp. (M) markers, (K +) positive control K562 allele 12 (41,42,43,51,33) samples

**Fig. 3.** Results visualization CODIS STR loci PCR results: TPOX; 224-252 bp. (M) markers, (K +) positive control K562 allele 11 (11,12,13,21.22.23.31) samples

**Fig. 4.** Visualization of CODIS STR loci PCR results: ThoI; 179-203 bp. (M) markers, (K +) positive control K562 allele 9 (11,12,13,21,22) samples.

In Fig. 3 above shows the emergence of the tape/band between 224-252 bp that TPOX locus primer pairs used are specific only stick to the expected position. The PCR process is carried out to amplify TPOX CODIS STR loci using primer pair M1 and M2. PCR products
were analyzed by PAGE with silver staining showing homozygotes (11:11) and heterozygous (11,12) allele.

In the picture above shows the emergence of the tape/band between 179-203 bp that locus primer pairs used are specific Thoi just stick to the expected position. The PCR process is carried out to amplify Thoi CODIS STR loci using primer pair M1 and M2. PCR products were analyzed with PAGE silver staining, showing allele heterozygotes (9.11) (9.8) and homozygous (9.9).

In the picture above shows the emergence of the tape/band between 179-203 bp that locus primer pairs used are specific Thoi just stick to the expected position. The PCR process is carried out to amplify Thoi CODIS STR loci using primer pair M1 and M2. PCR products were analyzed with PAGE silver staining, showing allele heterozygotes (9.11) (9.8) and homozygous (9.9).
Table 1. Distribution of allelic variation CODIS STR loci (CSF1PO, Thoi, TPOX, VWA)

<table>
<thead>
<tr>
<th>Allele number</th>
<th>CSF1PO</th>
<th>THOI</th>
<th>TPOX</th>
<th>vWA</th>
</tr>
</thead>
<tbody>
<tr>
<td>12.12</td>
<td>15</td>
<td>15</td>
<td>15</td>
<td>15</td>
</tr>
<tr>
<td>(73.33%)</td>
<td>(20%)</td>
<td>(60%)</td>
<td>(66.67%)</td>
<td></td>
</tr>
<tr>
<td>12.11</td>
<td>9.9</td>
<td>11.11</td>
<td>17.19</td>
<td></td>
</tr>
<tr>
<td>(26.67%)</td>
<td>(46.67%)</td>
<td>(40%)</td>
<td>(26.67%)</td>
<td></td>
</tr>
<tr>
<td>9.11</td>
<td>18.19</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>(33.33%)</td>
<td>(6.67%)</td>
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</tbody>
</table>

n = 15

Table 1 shows distribution of allelic variation in the CODIS STR loci CSF1PO, Thoi, TPOX, VWA, demonstrating differences in allele frequencies. Loci CSF1PO, TPOX and VWA show that homozygotes allele have the highest frequency. The results of this study indicate that allele of loci CSF1PO, Thoi, TPOX and VWA of 4 examined families contained a pair of alleles (averagely homozygous) found in each locus of Madura tribe. A pair of allele of the variant is not specific to Madurese who settled in Madura island, so it needs careful calculation based on population samples. However, a pair of alleles locus of this study may illustrate the pattern model of marriage among Madurese.

DISCUSSION

Based on the highest allele frequencies, the results of this study had differences from similar studies on Indonesian population (Untoro 2009), populations of Java, Sumatra, Sulawesi (Linace 2001). The difference may be due to difference in samples where the research was on one population and traced up to 3 generations above/below and between samples were not mutually related. In this study, samples were those of native Madurese with endogamous marriage patterns, marrying the fellow Madurese. However, with the development of technology and adequate transportation facilities, endogamous marriage patterns began to decline. To study the patterns of inheritance at the population level is necessary to understand Mendelian population. Mendelian population is a group of individuals of a species that reproduce sexually, live in a certain place at the same time and marriage occur between themselves (interbreeding) so that each will provide genetic contribution to the gene pool, which is a set of genetic information carried by all individuals in the population.

According to Stinson et al (2000) there are four main mechanisms that can alter gene frequencies and genotype in the population, namely mutation, selection, gene flow and genetic drift. Mutations and gene flow will increase variations in population. Gene flow is the exchange of genetic material between populations caused by migration and marriage. According to Cavalli-Sforza et al (1994), genetic drift occurs in the population in area which is closed, isolated or slightly receiving migrants from other regions. There are two circumstances that can lead to genetic drift. They are the bottleneck, where the population size is reduced at a certain time, and founder effect in which all individuals in a population can be traced back to a small number of individuals of origin. Genetic diversity decreases with migration. When a small group of our ancestors migrated to new areas, they basically carry within them a smaller sample of genetic diversity of the original communities.

The results of this study recommend that research on population or other medical fields should not only focused on genetics alone, but also take into account culture and customs of a population. Gene pool is not only a collection of genes, but a dynamic system that is well organized and includes the past history of a population. Every aspect of genetic information has history, anthropology and certain statistics so that it necessitates coordination and collaboration of various disciplines.

CONCLUSION

Analysis of CODIS STR loci CSF1PO, Thoi, TPOX, VWA in Madurese shows high genotype/homozygous allele in their gene pool. Local endogamous marriage patterns is most prevalent although in a small percentage (48%-55%). From genetic variation analysis based on history, it is suspected that Madurese had experienced suspected genetic drift and founder effects exists. It is visible in the gene pool of the Madurese, which mostly is homozygous.

REFERENCES