Carrier screening for (CGG)n repeat expansion of FMR1 gene in Korean women

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Purpose: We examined the prevalence and CGG/AGG repeat structure of expanded alleles of the FMR1 gene in preconceptional and pregnant Korean women.

Materials and Methods: The CGG repeats in the FMR1 genes of 1,408 women were analyzed by polymerase chain reaction and Southern blot analysis. To estimate the prevalence of expansion alleles, the individuals were divided into low risk and high risk group.

Results: Within this population, 98.4% had normal alleles and 1.6% had abnormal alleles including intermediate (0.6%), premutation (0.5%), full mutation (0.1%), and hemizygous (0.4%) alleles. There were 2 premutation alleles (1:666, 95% confidence interval [CI] 1:250-1,776) in the low risk group and 5 premutation alleles (1:15, 95% 1:6-36) in the high risk group. There were 8 intermediate alleles (1:167, 95% CI 1:130-213) in the low risk group and 1 intermediate alleles (1:76, 95% CI 1:11-533) in the high group. Six of the 7 premutation alleles did not contain AGG interruptions within the repeats and 1 had a single AGG interruption. Four of the 9 intermediate alleles contained 2-3 AGG, 4 had a single AGG, and 1 had no AGG interruptions.

Conclusion: Our study demonstrates the prevalence and CGG/AGG structure of expansion alleles in Korean women. The identified premutation prevalence is higher than that of other Asian populations and lower than that of Caucasian populations. Although our study is limited by size and population bias, our findings could prove useful for genetic counseling of preconceptional or pregnant women.

Key words: Fragile X syndrome, Trinucleotide repeat expansion, Primary ovarian failure, Gene frequency, Carrier state.

Introduction

Fragile X syndrome (FXS) is one of the most common mental retardation diseases and a major risk factor of inherited intellectual disabilities. FXS is caused by an expansion of CGG repeats in the FMR1 gene, which are hypermethylated within CpG islands at the promoter region, leading to transcriptional silencing of the FMR1 gene and the absence of fragile X mental
According to the American College of Medical Genetics and Genomics guidelines, expanded alleles can be classified as follows; normal (6-44 CGG repeats), intermediate or gray zone (GZ; 45–54 CGG repeats), premutation (55–200 repeats), or full mutation (>200 repeats). Most normal individuals have 6–44 CGG repeats including 2–3 interrupting AGG trinucleotides, found at positions 10 or 11, and 20 or 21 of the CGG repeat tract [2,3]. Premutation carriers have 55–200 CGG repeats with 1–2 AGG interruptions or tend to have fewer AGG interruptions resulting in an increasing length of the CGG repeat tract. The loss of AGG interruption appears to increase repeat instability and the risk of full mutation expansion during transmission of maternal premutation alleles [2,4]. Intermediate or GZ alleles (45–54 repeats) are potentially unstable and thus their clinical implications are currently under investigation [5]. Nolin et al. [5] defined the role of the AGG interruptions in the instability of 45–54 repeat alleles and identified a 19-fold difference in alleles with no AGG interruptions compared to those with 2 AGG interruptions. In addition, they found that full mutation expansions were transmitted from alleles with no AGG interruptions.

The aim of this study was to determine the prevalence and CGG/AGG structure of expanded alleles as well as the distribution of the FMR1 alleles in Korean women.

Materials and Methods

1. Subjects
A total of 1,408 preconceptional or pregnant women were tested for fragile X carrier screening at CHA Gangnam Medical Center (Seoul, Korea) between March 2010 and December 2015. The individuals applied for testing on their own initiative or on the advice of their physician, on a self-pay basis. Each of the women completed a questionnaire to ascertain any history of FXS, mental retardation, or premature ovarian failure (POF). All women identified as carrier with a premutation allele were offered genetic counseling. Those women who were already pregnant and preconceptional women who became pregnant were provided with information regarding prenatal diagnosis by amniotic-fluid analysis or chorionic villus sampling. This study was approved by the Institutional Review Board of Cha Gangnam Medical Center (IRB No. GCI-13-35).

2. DNA extraction and FMR1 region-specific CGG PCR
Genomic DNA was extracted from peripheral blood samples using the QuickGene DNA kit (Fujifilm Co., Tokyo, Japan) according to the manufacturer’s instructions. The (CGG)n repeats of the FMR1 gene were analyzed by polymerase chain reaction (PCR) using the GC rich PCR system kit (Roche Diagnostics, Mannheim, Germany).

3. FMR1 triplet repeat–primed (TP)–PCR and genetic analyzer
To confirm the number and structure of CGG/AGG repeats in intermediate, premutation and full mutation samples, we used the AmplideX FMR1 kit (Asuragen Inc., Austin, TX, USA) according to the manufacturer’s instructions. The size of the PCR products was converted into the number of repeats using an Microsoft Excel–based data analysis macro (FMR1 analysis macro version 2.1.2; Asuragen Inc.). A mixed internal standard DNA sample was tested in the same plate for each experiment to provide a process control [5,6]. The PCR products were loaded on an ABI Prism 3130xl Genetic Analyzer (Applied Biosystems, Foster City, CA, USA) and analyzed with GeneMapper version 3.2 (Applied Biosystems); 1 μL of the PCR products mixed with 9 μL formamide (Applied Biosystems) and 0.3 μL GeneScan™-500LIZ™ internal size standard (Applied Biosystems) internal size standard were loaded into the genetic analyzer.

4. Southern blot analysis
Southern blots were performed to identify premutation or full mutation when the PCR results were inconclusive (only one allele within normal range of [CGG]n repeats or amplification failed). Genomic DNA (10 μg) was digested with EcoRI and EagI (New England Biolabs, Ipswich, MA, USA). Hybridization and detection were performed using a DIG–labeled PCR probe (forward; TGAAGAGAAGATGGAGGAGCT/reverse; TCTCATTICGATAGGGAGCTAG) according to Gold et al. [7].

Results

Of 1,408 individuals 1,385 samples (98.4%) were within the

<table>
<thead>
<tr>
<th>CGG repeat range</th>
<th>Sample</th>
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<tbody>
<tr>
<td>5–45 (normal)</td>
<td>1,385 (98.4)</td>
</tr>
<tr>
<td>46–54 (intermediate)</td>
<td>9 (0.6)</td>
</tr>
<tr>
<td>55–200 (premutation)</td>
<td>7 (0.5)</td>
</tr>
<tr>
<td>&gt;200 (full mutation)</td>
<td>1 (0.1)</td>
</tr>
<tr>
<td>Only one normal allele (hemizygote)</td>
<td>6 (0.4)</td>
</tr>
<tr>
<td>Total</td>
<td>1,408 (100)</td>
</tr>
</tbody>
</table>

Values are presented as number (%).
normal range; 2 normal CGG repeat (16–44 repeats) alleles. There were 17 samples with one normal allele and one expanded allele; 9 had an intermediate (45–54 repeats) allele (0.6%), 7 had a premutation (55–200 repeats) allele (0.5%) and 1 had a full mutation (>200 repeats) allele (0.1%). The remaining 6 samples (0.4%) were hemizygous, meaning they had only a single normal X chromosome or one normal and a partially deleted X chromosome (Table 1).

The most prevalent alleles were 29 (45.0%), 30 (31.3%) and 36 repeats (10.1%) and these three alleles accounted for 86.4% of the total (Fig. 1).

The allele patterns identified in this study was shown in Fig. 2. Nine hundred and eighteen samples (65.2%) were heterozygous and 490 (34.8%) were homozygous. Of the heterozygous samples, 903 had two normal alleles, 9 (1.0%) had one normal allele and one intermediate allele and 6 were premutation carriers (0.7%). For homozygous samples, Southern blot analyses were performed to identify whether the homozygous samples had a large expansion allele (premutation or full mutation). One case (0.2%) was identified as a premutation carrier and one case (0.2%) had a full mutation allele. Interestingly, 6 cases (1.2%) were hemizygous. The CGG repeat length in these premutation, full mutation and hemizygous samples was then re-examined using TP-PCR. The premutation sample was shown to contain 29/93 repeats and the full mutation sample had >200 repeats (Fig. 3A). The hemizygous samples were shown to contain only the 2.8 kb unmethylated X chromosome allele; the 5.2 kb methylated allele

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**Fig. 1.** Distribution of triplet repeats in normal FMR1 alleles.

**Fig. 2.** The results for alleles pattern of FMR1 gene in total samples.
was not detected in these samples (Fig. 3B).

The CGG/AGG interspersion patterns were examined in 17 samples with an expanded allele (intermediate, premutation, and full mutation allele). Six of the 7 premutation alleles did not contain AGG interruptions within the repeats and 1 had a single AGG interruption (data not shown). In contrast, 4 of the 9 intermediate alleles contained 2-3 AGG interruptions, 4 had a single AGG with an expanded 3' pure CGG repeat stretch (41-43 repeats) and 1 had no AGG interruptions. Intermediate range alleles with 2-3 AGG interruptions exhibited a similar pattern to that of normal alleles, while alleles with 0-1 AGG interruptions demonstrated a similar pattern to that of premutation alleles with 0-1 AGG (Table 2).

### Discussion

In this study, we screened 1,408 Korean women for (CGG)n repeat expansion of FMR1 gene. Of total samples, abnormal samples containing intermediate, premutation, full mutation and hemizygous alleles accounted for 1.6% (n=23). The distribution of normal FMR1 alleles was 29, 30 and 36 repeats (86.4%), similar to that of other Korean population [8,9].

For the more details, the samples were divided into two groups a low risk group without indication of FXS and a high risk group with an indication of FXS history, mental retardation or gynecological diseases such as POF. we identified 8 intermediate alleles in the low risk group (1:167, 95% CI 1:130-213), 1 intermediate allele in the high risk group (1:76, 95% CI 1:11-533), 2 premutation alleles in the low risk group (1:666, 95% CI 1:250-1,776), 5 premutation alleles in the high risk group (1:15, 95% CI 1:6-36), 1 full mutation allele in the high risk group (1:76, 95% CI 1:11-533), 1 hemizygous allele in the low risk group (1:1332, 95% CI 1:188-9456) and 5 hemizygous alleles in the
high risk group (1:15, 95% CI 1:6-36). As mentioned just earlier, the prevalence of premutation carriers in our study was 1:666, similar to Kim et al. (1:781) in Korean population [8]. This finding is higher the prevalence of other Asian populations and lower than that of Caucasian populations [8,9]. Our population-based screening for FXS may not directly reflect the overall prevalence in Korean women because the study was limited to women who requested the test themselves or were referred by a doctor and had to pay for it.

Of the 6 hemizygous samples, 4 had an indication of POF, 1 was infertile for unknown reasons and 1 was detected from the prenatal test of a pregnant woman. Table 3 details the cytogenetic analyses of the hemizygous samples. The 6 hemizygous samples contained the 2.8 kb unmethylated allele of the active X chromosome and 4 of the hemizygous samples had an indication of POF. In 3-15% of women premutation aberrations of the FMR1 gene can manifest as a POF/POI phenotype [10]; 10-15% of the cases are X chromosome abnormalities, such as numerical and structural aberrations (deletions, inversions, and X/autosome translocations) [11,12]. Haploinsufficiency of genes located in the missing region of the X chromosome could provide a promising explanation for the POF disease background, especially when it involves Xq28. A haploinsufficient gene is defined as a gene that requires 2 functional alleles for wild type expression. Lack of gene expression because of X inactivation may affect ovarian function [10].

Inherited patterns of three cases with a premutation allele were examined. The premutation alleles of patient I (29/129 repeats) was transmitted to her daughter and underwent expansion to full mutation. Patient II, a pregnant woman with premutation alleles, has a son with FXS and full mutation alleles were detected in her prenatal testing. Patient III was found to have premutation alleles (29/93); the sister of patient III also has premutation alleles, however, her alleles are further expanded (29/96; data not shown).

Our results revealed dynamic mutation events in the (CGG)n repeat structure. One patient with intermediate allele (29/53 repeats, no AGG interruptions) had the POF disease background (Anti-Müllerian hormone: >0.14). Bodega et al. [13] demonstrated a significant prevalence of intermediate alleles in the POF population with respect to control group. Rybak et al. [14] reported that sisters with POF had intermediate alleles (45 repeats). In contrast, in Bennett et al. [15] and Guo et al. [16], they showed that the prevalence of intermediate FMR1 was not increased significantly in sporadic POF than that in controls. So, further study should be performed about association of intermediate allele with POF.

Although an allele is within the intermediate range, if it does not contain AGG interruptions or the length of the 3’ pure CGG repeat is greater than the appropriate level, the instability of the CGG repeat region might be increased and it could then expand into a premutation allele [5,6]. Thus, it is important that this information be conveyed not only to women with premutation alleles but also to those with intermediate alleles.

Since the identification of the FMR1 gene in the 1990s, expansion risk estimates have been based solely on maternal repeat length alone. However, data pointing to the role of interspersed AGGs in the stability of the CGG repeat region has accumulated over the past 20 years [17–19]. Nolin et al. [5,20] demonstrated a 19-fold difference for alleles with no AGGs compared to those with 2 AGG disruptions. Yrigollen et al. [18] showed that the combination of total CGG repeat tract length and the number of AGG interruptions was significantly associated with the risk of expansion to a full mutation. It is important that this more accurate risk information be provided to families through genetic counseling, which will serve to further facilitate their decision making process. Additional studies with larger cohorts and various populations will be required to confirm our findings because these data are limited by study size and population bias. Studies involving the children of patients with intermediate alleles containing 0–1 AGG interruptions might also provide more accurate risk information.

Although our study is limited by size and population bias, our
findings will be useful in terms of genetic counseling offered to preconceptional or pregnant women.

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References