Screening for \textit{FMR1} expanded alleles in patients with Autism Spectrum Disorders in Manaus, Northern Brazil

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\textbf{Abstract:} Fragile X Syndrome (FXS) is a neurodevelopmental disorder caused by dynamic mutations of a CGG repetition segment in an X chromosome’s single gene. It is considered the leading hereditary cause of both Autism Spectrum Disorders and Intellectual Disability. Some authors suggest that all individuals diagnosed with some of these latter conditions to be clinically and molecularly trialled for FXS due to the high levels of comorbidity between both conditions and also due to the variable expressiveness of this syndrome. This study has focused on verifying the presence of \textit{FMR1} expanded alleles since there is a lack of information about this kind of mutation in autism patients from the northern region of Brazil. The presence of large alleles for this gene could offer new therapeutic or pharmacological methods for the treatment of these patients. Both the presence and the frequency of CGG expansions were verified in 90 autism males by molecular analysis. Four of them had intermediate alleles and four others presented premutated alleles. Premutation carriers are on the propensity of developing the late onset Fragile X-associated tremor/ataxia syndrome. No full mutation alleles were found. Further studies are necessary to obtain more accurate statistical data about this kind of dynamic mutation.

\textbf{Key words:} autism spectrum disorders, dynamic mutation, fragile X syndrome, molecular screening, pre-mutation, trinucleotide repeats.

\section*{INTRODUCTION}

Fragile X-Syndrome (FXS), previously named Martin-Bell syndrome, was first identified by Martin and Bell (1943) when they observed eleven patients with intellectual deficiency and facial dysmophia in a six-generation family. The authors noticed that this neurological-morphological condition was only present in males, but not in their mothers and sisters. FXS is a monogenic recessive neurodevelopmental condition linked to the X-chromosome, with low penetrance, variable expressiveness, caused by the dynamic expansion of non-coding unstable repetitions compose by cytosine-guanine-guanine (Jin and Warren 2000, Bagni et al. 2012). This syndrome is a neurodevelopmental disorder considered the leading hereditary cause of both Autism Spectrum Disorders.
Disorders (ASD) and Intellectual Disability (ID) (Choi et al. 2015, Saldarriaga et al. 2016).

The primary cause of more than 98% cases of FXS is a dynamic mutation of CGG expansion at the 5'-UTR of FMRI gene (Fragile X Mental Retardation 1 gene), located at the FRAXA locus in Xq27.3 (Sherman et al. 2005, Peprah 2014, Winarni et al. 2013). This gene is classified into four different forms, according to their CGG-repeat size: healthy individuals have normal alleles ranging from 5 to 44 copies (Jang et al. 2014, Sethna et al. 2014). Affected patients usually exhibit mutated alleles longer than 200 CGGs (De Esch et al. 2014, Latham et al. 2014, Myrick et al. 2015). This significant expansion, known as the full mutation, leads to hypermethylation of the promoter region and transcriptional silencing (Sethna et al. 2014). The deficiency or lack of expression of a protein called FMRP (Fragile X Mental Retardation Protein) is then related to the classical symptoms of FXS, such as behavioral disorders, cognitive impairments and facial dysmorphia (Heulens et al. 2013, Winarni et al. 2013). The others two remaining classes for FMRI gene are called intermediate or grey-zone alleles (45–54 triples) and premutation alleles (55–200 repeats), whose carriers are not affected but have increased risks for expansion to premutation or full mutation, respectively, in the next generation (Fernandez-Carvajal et al. 2009, Jang et al. 2014).

Some authors (Mandel and Biancalana 2004, Hagerman et al. 2010) suggest that clinical and molecular trials for FXS be performed in all individuals diagnosed with ASD and or ID due to the high levels of comorbidity between ASD and FXS and to the variable expressiveness of FXS. The co-occurrence of ASD and FXS has been estimated at 30 to 50% (Kaufmann et al. 2004, Harris et al. 2008, Abbeduto et al. 2014) and both neurological conditions exhibit common etiological relationship. Also, the clinical characteristics of affected FXS patients are nonspecific and are quite subtle, varying from mild to severe (Mandel and Biancalana 2004, Bagni et al. 2012). Individuals with FXS may present with anything from learning problems and a normal IQ to severe mental retardation and autistic behaviors (Garber et al. 2008). Therefore, the presence of FMRI mutations in ASD patients could offer new therapeutic or pharmacological methods for their treatment.

The aim of this study is to verify the presence of expanded alleles for the FMRI gene in ASD patients since there is a lack of information about this kind of mutation in the northern region of Brazil.

**MATERIALS AND METHODS**

This study was approved by the Committee of Ethics on Human Research of the Universidade do Estado do Amazonas (No CEP-UEA 363.912/2013). Genomic DNA of 90 male patients (ranging in age from 3 to 23 years old) was analysed for the polymorphism of locus FRAXA, located at the 5’UTR of FMRI gene. All subjects included in the research were patients diagnosed with non-syndromic ASD, assisted by two specialized multidisciplinary institutions for the care of autistic people in the city of Manaus-Brazil (Centro de Educação Especial André Vidal de Araújo and Espaço de Atendimento Multidisciplinar ao Autista Amigo Ruy). All parents or legal guardians signed a free and informed consent form.

Cells from the jugal mucosa of each patient were collected using a SWAB stick and stored in Tris-EDTA solution (10mM / 0.1mM). Genomic DNA was extracted using CTAB 2% (Doyle and Doyle 1990) and then quantified in Eppendorf® BioSpectrometer instrument. Target locus was amplified by PCR using fluorescent primers developed by Fu et al. (1991): forward 5’-gtcagctcgctttaggggattactggt-3’ and reverse 6-FAM-5’-ageccgacccacccaccagctctctca-3’. Amplifications were performed in a SimpliAmp™ Applied Biosystems thermocycler programmed with the following conditions: initial denaturation at 94°C for 5’, 32 cycles of 94°C for 45”, 65°C
for 1’30” and 72°C for 2”, and a final extension at 72°C for 10’. Reaction total volume was 24µL, with 2.4µL of DNA amplification buffer (Biotech Amazonia, Manaus, BRA); 0.85mM of each dNTP; 0.33µM of both forward and reverse primer; 0.63% DMSO (dimethylsulfoxide – Sigma-Aldrich, St Louis, MO, EUA); 1.65µM de BSA (bovine serum albumin acetylated) (Promega, Madison, WI, EUA); and ~60ng of gDNA. Amplified DNA fragments were separated by electrophoresis on a 1% agarose gel, which was then photographed by a transilluminator UV L-PIX HE. Allele sizes were determined by capillary electrophoresis at an automatic sequencer ABI-3130XL using the polymer 3130 POP-7 (Applied Biosystems, Foster City, CA, EUA).

Allele sizes were determined by capillary electrophoresis at an automatic sequencer with an internal marker pUC19 ROX-labelled size standard. GeneMarker v.2.6.0 computer program performed the analysis of electropherograms. Expected allele size of the amplicon was 221bp, excluding the CGG repetition region itself. A formula developed by Hamdan et al. (1997) was used to calculate the number of CGG repetitions in the \(FMR1\) gene. Allele frequency was calculated according to the ratio between the quantities of alleles of a particular allelic form and the total sample number. Modal value and arithmetic mean for CGG copies were obtained on Excel 2016® program. For the purposes of this paper, we will henceforward refer to the names of the four classes of alleles (normal, grey zone, premutation and full mutation) as NM, GZ, PM and FM, respectively.

**RESULTS**

Of 90 male patients analysed in this study, 75 presented NM alleles; four showed GZ alleles, and four showed PM alleles. FM alleles were not observed. Seven sample could not be amplified due to PCR issues. The frequency of each allele is shown in Table I.

Both the arithmetic mean and the modal value for CGG repeats in the normal range were 27. The amplitude of CGG codons and the major and minor alleles for each allelic class are presented in Figure 1.

**DISCUSSION**

This study is the first one aiming the prevalence of dynamic mutations in the \(FMR1\) gene among ASD patients in the northern region of Brazil. No full mutation alleles were found in this population. However, four patients showed GZ alleles and four others showed PM alleles.

Grey-zone (or intermediate) allele is believed to be a potential precursor for premutation alleles and both are unstably transmitted from mother to children (Nolin et al. 2015). Intermediate-to-premutation expansions are rare and have not been observed in a single generation (Cronister et al. 2008). Although, Fernandes-Carvajal et al. (2009) reported an intermediate-to-full mutation increasing in the CGG region within two generations when studying a family where a 52 CGGs grandfather transmitted to his grandson a ~538 repeats allele.

Premutation carriers (55-200 CGGs) are relatively common in the general population. About one in 260 females and one in 813 males are in this range (Leehey 2009). Premutation-to-full mutation expansion occurs exclusively from PM carriers mothers, given that, during maternal meiosis, premutation alleles are highly unstable (Sherman et al. 2005), which depends on their CGG length (Fernandez-Carvajal et al. 2009) and

<table>
<thead>
<tr>
<th>Allele class</th>
<th>Nº of alleles found</th>
<th>Allele frequency (f)</th>
</tr>
</thead>
<tbody>
<tr>
<td>NM (5-44 CGGs)</td>
<td>75</td>
<td>(f): 0.90</td>
</tr>
<tr>
<td>GZ (45-54 CGGs)</td>
<td>04</td>
<td>(f): 0.05</td>
</tr>
<tr>
<td>PM (55-200 CGGs)</td>
<td>04</td>
<td>(f): 0.05</td>
</tr>
<tr>
<td>FM (&gt;200 CGGs)</td>
<td>0</td>
<td>(f): 0.0</td>
</tr>
</tbody>
</table>

**TABLE I**

Frequency of the four classes of alleles.
Figure 1 - The higher CGG repeat size found for the normal range was 42. For the grey-zone and premutation range were 53 and 90, respectively. The arithmetic mean and the modal value were both only calculated for the normal allele type.

<table>
<thead>
<tr>
<th>Minor allele found</th>
<th>Major allele found</th>
<th>Arithmetic mean</th>
<th>Modal value</th>
</tr>
</thead>
<tbody>
<tr>
<td>NM (5-44)</td>
<td>16</td>
<td>27</td>
<td>27</td>
</tr>
<tr>
<td>GZ (45-54)</td>
<td>42</td>
<td>53</td>
<td>90</td>
</tr>
<tr>
<td>PM (55-200)</td>
<td>55</td>
<td></td>
<td></td>
</tr>
<tr>
<td>FM (&gt;200)</td>
<td></td>
<td></td>
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on the number and position of AGG (adenine-guanine-guanine) inside the CGG repeat (Latham et al. 2014). According to Nolin et al. (2015), the chances of a >90 CGGs allele undergo to full mutation exceed 94%. Paternal premutation alleles are stably transmitted to daughters and do not result in FM (Jang et al. 2014), although one case of such enlargement of the FMR1 gene has been related by Alvarez-Mora et al. (2017) when they showed a 16-year-old girl who inherited mosaic PM-FM alleles (175 repeats and >200 repeats) from her 88 CGG father.

The four patients with PM alleles found in this study were from 6 to 13 years old. According to Hagerman and Hagerman (2013), boys with premutation in the early childhood present at higher rates of attention deficit hyperactivity disorder (ADHD), shyness, social deficits and intellectual disability. Thus, there should be given particular importance to these symptoms in these patients, given that they already have an ASD diagnosis. Male PM carriers, depending on their premutation range, are also on the propensity of developing Fragile X-associated tremor/ataxia syndrome (FXTAS) (Jacquemont et al. 2014), which is a late-onset neurodegenerative disorder that affects mostly men in the early seventh decade, causing cerebellar gait ataxia and intention tremor (Leehey 2009). Another point to bear in mind is that, due to the mechanisms of trinucleotide repeats expansion during maternatal transmission, the mothers of the eight patients we found with FMR1 grey-zone and premutated alleles may also have enlarged regions in this gene, smaller ones, but still in the grey-zone or the premutation range or even in the border of a previous allele class. Premutation alleles, for example, undergo an expansion in almost all cases when transmitted by a female (Barasoain et al. 2016).

Those carries mothers of GZ or PM alleles have a 20% chance of experiencing Fragile X-associated premature ovarian insufficiency, which is a cessation of menses prior to 40 years old (Allingham-Hawkins et al. 1999). They also have a 10% to 15% chance of suffering seizures (Tassone et al. 2014), and anxiety symptoms (Jacquemont et al. 2014).

The arithmetic mean of CGG triplets observed within those patients who showed NM alleles was 27. Others studies performed in Indonesia (Faradz et al. 2000), China (Faradz et al. 2000, Zhou et al.
and Mexico (Rosales-Reynoso et al. 2005) reported mean values of 29, 29 and 32 CGGs, respectively. The modal value we observed for CGG repetitions was also 27. Others studies performed with the Brazilian population found a modal value of 20 CGG repetitions (Sucharov et al. 1999, Mingroni-Netto et al. 2002). Although it is well-known that normal population present polymorphic alleles ranging from 4 to 55 repeats (Sethna et al. 2014), the importance of calculating the allelic frequency for FMR1 gene is that it is possible to verify the genetic instability at the FRAXA site and to identify the potential risks of allelic expansion, which can be attractive to medical geneticists and carriers of pre-mutated alleles at risk of increase for the full mutation (Peprah 2014).

There are several institutions for the care of autism children in the city of Manaus, Brazil. Here, we proposed a simple and free molecular test to only a small portion of the many parents who signed up to conduct the study with their children. Most parents informally reported that sometimes they blame themselves for their children neurological conditions, based on local beliefs. According to them, they are not properly informed about the possible causes of ASD. Only a small fraction of these parents are aware of the multifactors of autism spectrum disorders, including the genetic ones, but they relate that could not afford genetic tests for FXS requested by therapists. Such tests are quite expensive and often inaccessible and time-consuming in this region. Discovering genetic mutations in ASD patients is an important step not only to establish appropriate intervention and treatment but also to carry out genetic counselling that will inform the recurrent risk of FXS and Fragile X-associated disorders in the family. Once few molecular analysis have been carried out with the Brazilian population in order to verify the incidence of expanded alleles among the autistic population, more in-depth studies are necessary to obtain more accurate statistical data about this kind of dynamic mutation.

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AUTHOR CONTRIBUTIONS

All three authors conceived the original idea and planned the experiment. Jorge F.B. Ferreira carried out the experiments (obtaining of samples, DNA extraction, PCR reaction and electropherograms analysis) and wrote the manuscript with support from Jacqueline S. Batista and Cleiton Fantin. Jacqueline S. Batista performed the genotyping reaction. Cleiton Fantin was in charge of overall direction. All authors discussed the results and contributed to the final version of the manuscript.

REFERENCES


