Role of Replication and CpG Methylation in Fragile X Syndrome CGG Deletions in Primate Cells

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Instability of the fragile X CGG repeat involves both maternally derived expansions and deletions in the gametes of full-mutation males. It has also been suggested that the absence of aberrant CpG methylation may enhance repeat deletions through an unknown process. The effect of CGG tract length, DNA replication direction, location of replication initiation, and CpG methylation upon CGG stability were investigated using an SV40 primate replication system. Replication-dependant deletions with 53 CGG repeats were observed when replication was initiated proximal to the repeat, with CGG as the lagging-strand template. When we initiated replication further from the repeat, while maintaining CGG as the lagging-strand template or using CCG as the lagging-strand template, significant instability was not observed. CpG methylation of the unstable template stabilized the repeat, decreasing both the frequency and the magnitude of deletion events. Furthermore, CpG methylation slowed the efficiency of replication for all templates. Interestingly, replication forks displayed no evidence of a block at the CGG repeat tract, regardless of replication direction or CpG methylation status. Templates with 20 CGG repeats were stable under all circumstances. These results reveal that CGG deletions occur during replication and are sensitive to replication-fork dynamics, tract length, and CpG methylation.

Introduction

Fragile X syndrome (FRAXA [MIM 309550]), the leading cause of inherited mental retardation, is caused by an expansion of the CGG trinucleotide repeat (TNR) located within the 5′ UTR of the FMR1 gene (Verkerk et al. 1991). TNR diseases, such as FRAXA, bring to the forefront important issues regarding DNA replication and its effect on the dynamic nature of repeat stability. Experiments on multiple transgenic mice have been unable to reproduce the parent-of-origin effects observed in humans, with the most instability observed for knock-in and human-YAC mice (Lavedan et al. 1998; Bontekoe et al. 2001; Peier and Nelson 2002; Pearson 2003). Furthermore, the majority of investigations into the underlying mechanism(s) of TNR instability are based on CAG/CTG model systems. However, there are striking differences between CGG- and CAG/CTG-associated diseases, including parent of origin, tissue specificity, somatic instability, aberrant CpG methylation, and the formation of unusual DNA structures. Given the numerous differences between CGG and CAG/CTG genetics, conclusions based on CAG/CTG experiments may not be applicable to the mechanisms of CGG instability; therefore, parallel studies of CGG instability should be performed.

Somatic and gametic instability play a crucial role in the etiology of most TNR diseases. Unlike the extraordinarily high level of somatic CTG instability—both within and between tissues—displayed by patients with myotonic dystrophy (DM1 [MIM 160900]) (Martorell et al. 1995; Wohrle et al. 1995), patients with FRAXA do not exhibit postnatal somatic CGG instability (Reyniers et al. 1999). Distinct patterns of CGG repeat–length heterogeneity formed during fetal development persist postnatally in patients with FRAXA (Wohrle et al. 1995), which contrasts DM1, in which the CTG length may vary over the patient’s life (Martorell et al. 1995; Wohrle et al. 1995). This lack of postnatal somatic instability, along with data from studies published elsewhere (Wohrle et al. 1993; Malter et al. 1997; Moutou et al. 1997), supports the suggestion that FRAXA expansions are the result of either meiotic events in oocytes or mitotic processes restricted to the earliest postzygotic cell divisions (i.e., before days 7–12). It is important to note, however, that only unmethylated premutation repeat lengths (50–200 repeats) exist in the sperm of males with full-mutation FRAXA, even though all other tissues display disease-length alleles (>200 repeats) (Reyniers et al. 1993; Tassone et al. 1999). The
Figure 1  Replication templates. A, Templates containing human FRAXA CGG repeats (20 or 53 repeats) (blackened box), with genomic nonrepeating sequences (gray boxes) flanking the repeat. The SV40-ori fragment encompassed the unique origin of bidirectional replication (OBR) (Hay and DePamphilis 1982) and was inserted at the BglI, EcoRI, NheI, or AflIII site. The location of the SV40-ori, relative to the repeat, defined the direction of replication, as well as which strand (CGG or CCG) would serve as the leading- or lagging-strand template. B, Diagram of templates. The distance to the repeat tract was measured from the SV40 OBR (circled) to the nucleotide preceding either the 5′ or the 3′ end of the CGG repeat.

absence of full-mutation alleles in the sperm of males with FRAXA suggests that deletions in the male germ-line arise later in fetal development, after the initial expansion event (Wohrle et al. 1993; Malter et al. 1997; Moutou et al. 1997; reviewed in Pearson 2003).

Unlike other TNR diseases, the FRAXA phenotype is modulated not only by repeat tract length (Nolin et al. 2003; Rife et al. 2004) but also by aberrant CpG methylation. Patients with FRAXA exhibit complete methylation of their expanded CGG tract, whereas rare individuals classified as “high-functioning” males or “methylation mosaics” possess unmethylated CGG repeats within the disease-length range and display considerable levels of postnatal somatic CGG instability (mostly deletions), highlighting the increased instability of the unmethylated repeats (Wohrle et al. 1998; Taylor et al. 1999). Interestingly, not only does CpG methylation modulate phenotype and influence CGG instability (Wohrle et al. 1998; Nichol and Pearson 2002), but it also has also been documented to delay the already slowed replication of the FRAXA chromosomal locus (Hansen et al. 1993; Subramanian et al. 1996).

Previous work of our lab, performed using an SV40 system of primate replication, revealed that the location of replication initiation and fork direction, relative to the repeat, can lead to different types and frequencies of repeat instability in CTG/CAG repeat tracts (Cleary et al. 2002). By taking advantage of this SV40 system, it was possible to examine not only the influence of replication-fork dynamics on CGG stability but also how CpG methylation may modulate this stability. Our results support a role for replication-mediated events in CGG instability, with the frequency of deletions depending on length, sequence type (i.e., CGG or CCG), and location of replication initiation. Furthermore, the occurrence and magnitude of deletions, as well as the efficiency of replication, could be modulated by CpG methylation. Interestingly, replication-induced deletions arose independently of a replication-fork block at the CGG repeat.

Material and Methods

Replication Templates

Genomic FRAXA sequence (nucleotides 2523–2911 of pE5.1 [GenBank accession number X61378]) was cloned into pPCR-Script Amp (Stratagene), with the repeat in the stable orientation, relative to the unidirectional bacterial ColE1 origin of replication (with CCG as the lagging-strand template) (Pearson et al. 1998). SV40 viral sequence (nucleotides 5171–128 [GenBank accession number NC_001669]), containing the origin of bidirectional replication (nucleotide 5210/5211), was inserted into the FRAXA vector at various locations (fig. 1A).

CpG Methylation

pFX/SV40 clones were methylated with SssI methylase (New England Biolabs), in accordance with the manufacturer’s protocol, by use of a buffer without MgCl2, to avoid topoisomerase activity (Matsuo et al. 1994). A
small amount of each plasmid was digested with AcI to confirm the presence of CpG methylation (data not shown).

Cell Transfections

COS-1 cells (American Type Culture Collection) (Gluzman 1981) were grown in Dulbecco modified Eagle medium (Invitrogen) with 10% fetal bovine serum (HyClone). Cells were incubated with 5 μg of each pFX/SV40 template and 50 μl of LipoTAXI (Stratagene), in accordance with the manufacturer’s specifications. Episomal DNAs were extracted 48 h posttransfection with 100 mM Tris-HCl (pH 7.5), 1 mM EDTA, and 0.6% SDS (Hirt 1967).

Mutation Analysis

Mutations were assessed using the stability of trinucleotide repeats by individual product (STRIP) assay (Cleary et al. 2002). Episomal DNAs were digested with DpnI (New England Biolabs) to eliminate unreplicated parental templates. DpnI-resistant material was transformed, and plasmid DNA from individual bacterial colonies, each representing an individual product of primate replication, was prepared. To minimize repeat instability due to propagation in Escherichia coli, the XL1-Blue MR cell strain (Stratagene) was selected for analysis of all pFX/SV40 replication products except pFX53/pFX20A, for which the XLmutS cell strain (Stratagene) was used.

Repeat-containing restriction fragments were resolved on 4% polyacrylamide gels, with tract lengths scored relative to starting material and a 123-bp ladder (Invitrogen). As described in detail elsewhere (Cleary et al. 2002), corrected mutation frequencies were calculated only for repeat-length distributions that were significantly different from the parental template (by use of χ² analysis) (fig. 2), by subtracting the background-length heterogeneity within the parental plasmid from the frequencies generated by primate replication (table 1). Repeat-length distributions that were not significantly different from the parental background (P > .05) were classified as either stable or having no primate cell–induced events (“0”) (table 1). Statistical analysis of the magnitude data was performed using t-test analysis (fig. 3). To confirm that the deletions or expansions observed were repeat-length changes, samples were electrotitransferred and hybridized with a 32P-(CGG)₁₀ oligonucleotide. Selected expansion and deletion products were also sequenced.

Replication Efficiency

Replication efficiency (fig. 4) was calculated by cotransfecting the pFX53 plasmids, with and without CpG methylation, with unmethylated, repeat-free pSV40 (fig. 1). Posttransfection, episomal DNA was digested with AluNI and DpnI, resolved on a 1% agarose gel, transferred to Biodyne B membrane (Pall Corporation), and probed with the 32P-SV40 origin of replication (SV40-ori). An aliquot of the starting mixture used for the cotransfection was linearized with AluNI and was analyzed on the same gel. ImageQuant version 1.2 (Molecular Dynamics) was used to quantify the ratio of starting materials between pFX53 and pSV40 and the ratio of completely replicated materials. These ratios were used to determine the replication efficiency of each template in relation to unmethylated pSV40.

Two-Dimensional (2D) Gel Electrophoresis

Replication intermediates of pFX53E and pFX53N were digested with StuI/XmnI and StuI/AluNI, respectively (fig. 5B and 5C). Samples were run in the first dimension for 24 h at 1 V/cm on a 0.4% agarose gel. Lanes were then cut out, embedded in 1% agarose gel with 0.5 μg/ml ethidium bromide, and run in the second dimension for 8 h at 5 V/cm at 4°C, with buffer recirculation (Cordeiro-Stone et al. 1997; Krasilnikova and Mirkin 2004). Gels were transferred as described above and were probed with the 32P-XmnI-AluNI fragment of pPCR-Script Amp (Stratagene).

Results

Primate Replication Assay for CGG Stability

The role of primate DNA replication in FRAXA CGG repeat stability was investigated using an established SV40 replication system (Cleary et al. 2002). The SV40-ori was inserted once within a construct containing both the CGG tract and FRAXA human genomic flanking sequence (pFX). By positioning the SV40-ori on either side of the repeat, it was possible to change which strand (CGG or CCG) was used as the lagging-strand template in a series of replication constructs comprising either 20 or 53 pure CGG repeats (fig. 1).

pFX/SV40 replication templates were transfected into SV40 T-antigen (T-Ag)–expressing COS-1 cells and were replicated by primate proteins. The episomally replicated DNAs were isolated and assayed for repeat stability by use of STRIP analysis (Cleary et al. 2002) (fig. A1 [online only]). The STRIP assay allowed for length analysis of individual products of primate replication, with the unreplicated starting material removed by DpnI digestion prior to transformation into E. coli.

Primate-Replication–Induced Deletions of a (CGG)₅₃ Template

In pFX53E, the SV40-ori location permitted replication to initiate 74 nt 5′ of the CGG tract, with CGG as the template for lagging-strand synthesis (fig. 1). Rep-
mutation analysis. As shown elsewhere (Cleary et al. 2002), bacterial preparations of the starting parental templates contained a distribution of repeat lengths that were classified into one of three categories: <53 repeats, 53 repeats, and >53 repeats (unblackened bars). After primate replication, the repeat-length distribution of the primate-replicated material for each template (unmethylated template [blackened bars] and CpG premethylated template [gray bars]) was similarly determined. The bar height (Y-axis) reflects the percentage of replication events within each length category. For each length category, the number of replication events observed within that length category, over the total number of replication events analyzed, is indicated inside the bar. After primate replication, templates with a statistically significant (P < .05) increase in the number of molecules with lengths >53 repeats were marked with an asterisk (*). All constructs were transfected a minimum of three times.

Replication Direction and the Location of Replication Initiation

Data from bacterial and yeast models suggest a role for direction of replication in CGG instability (Shimizu et al. 1996; Hirst and White 1998; White et al. 1999). To determine whether this phenomenon extends to primate replication, pFX53N was constructed such that CCG was lagging-strand template (fig. 1). In contrast to pFX53E, the frequency of deletions did not differ significantly from background (fig. 2 and table 1) (P = .637; χ² = 0.223) compared to the significant increase in deletion events observed for pFX53E, which was replicated in the same orientation. This shift—from high levels of deletions to levels that do not differ from background—results from a change in the distance between replication initiation and the repeat without changing the direction of replication, highlighting the contribution of cis-elements to CGG instability.

To further address the influence of the location of
Table 1
Summary of pFX Data

<table>
<thead>
<tr>
<th>TEMPLATE</th>
<th>CORRECTED FREQUENCY* (%) OF EXPANSION</th>
<th>EXPANSION LENGTH(S)*</th>
<th>REPLICATION EFFICIENCY* NORMALIZED TO pFX53E (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>pFX53A + CH3</td>
<td>0</td>
<td>68 (67)</td>
<td>43</td>
</tr>
<tr>
<td>pFX53B</td>
<td>0</td>
<td>NA</td>
<td>80</td>
</tr>
<tr>
<td>pFX53E + CH3</td>
<td>0</td>
<td>NA</td>
<td>71</td>
</tr>
<tr>
<td>pFX53E</td>
<td>0</td>
<td>20 NA (58)</td>
<td>100</td>
</tr>
<tr>
<td>pFX53N + CH3</td>
<td>0</td>
<td>69</td>
<td>54</td>
</tr>
<tr>
<td>pFX53N</td>
<td>0</td>
<td>54, 55 (NA)</td>
<td>37</td>
</tr>
<tr>
<td>pFX53A + CH3</td>
<td>0</td>
<td>60, 74</td>
<td>27</td>
</tr>
<tr>
<td>pFX53A</td>
<td>0</td>
<td>57 (62)</td>
<td>45</td>
</tr>
<tr>
<td>pFX20E/pFX20N +/− CH3</td>
<td>0</td>
<td>NA</td>
<td>ND</td>
</tr>
<tr>
<td>pFX53E in CV-1d</td>
<td>0</td>
<td>NA</td>
<td>ND</td>
</tr>
<tr>
<td>pFX53E + CH3 CV-1d</td>
<td>0</td>
<td>NA</td>
<td>ND</td>
</tr>
</tbody>
</table>

NOTE.—As described elsewhere (Cleary et al. 2002) (see the “Material and Methods” section), corrected mutation frequencies were calculated only for repeat-length distributions that were significantly different from the parental template background (fig. 2). NA = not applicable; ND = not determined.

* Expansion and deletion frequencies mediated by primate-cell replication were corrected for background by subtracting the frequencies of >53 or <53 events, respectively, for the parental template molecules from the frequencies of the same category for the replicated products. Repeat-length distributions that were not significantly different from the parental background (P > .05) were classified as stable or as having no primate-replication-induced events (“0”).

b In cases in which expansions were observed, the length of the primate-replicated expanded CGG repeat is noted in parentheses.

c Replication efficiencies were determined as in figure 4 (see “Material and Methods” section).

d Replication not permitted.

replication initiation on instability, pFX53A was analyzed. In pFX53A (fig. 1), replication proceeded in the same direction as pFX53N (with CGG as the lagging-strand template), but it was initiated further from the repeat tract (613 nt vs. 93 nt). Both pFX53N and pFX53A (fig. 2 and table 1) (P = .182; χ² = 1.781) did not show significant levels of deletions. Therefore, the effect of initiation location on instability was evident only when CGG, not CCG, served as the lagging-strand template.

Effect of CpG Premethylation on Stability

To examine the possible role of CpG methylation in CGG stability, the pFX/SV40 constructs were premethylation with SssI methylase prior to transfection. CpG premethylation of pFX53E decreased not only the frequency of deletion events incurred during primate replication (from 20% to 0%) (see table 1 and the “Material and Methods” section) but also significantly reduced the magnitude of CGG length changes within the observed deletions (fig. 3) (P = .0152). CpG premethylation did not modulate the frequency of deletions for pFX53B, pFX53N, or pFX53A (fig. 2). However, CpG premethylation of pFX53N significantly decreased the magnitude of CGG length change of deletions observed when compared with unmethylated pFX53N (fig. 3) (P = .0467). Only CpG methylation was changed in this experiment, providing strong support for the hypothesis that CpG methylation stabilizes the CGG repeat during replication.

Length Dependence on Instability

To assess whether the stability of shorter CGG repeat tracts was affected by replication, an identical series of pFX/SV40 templates with 20 CGG repeats (fig. 1B) was replicated in COS-1 cells. None of the pFX20 templates displayed significant levels of instability, regardless of replication direction and location or CpG methylation status (tables 1 and 2), suggesting an association between repeat length and instability; longer pure repeat tracts were more prone to mutation.

Rare Expansions of the (CGG)₅₃ Templates

One hallmark of FRAXA is the ability of premutation alleles containing 50–200 CGG repeats to expand upon transmission (Nolin et al. 2003; Rife et al. 2004).
Figure 3  Length distributions of observed deletion events. All deletion events were sized and graphed as the number of events within each CGG size category. Unmethylated primate-replicated deletions (blackened bars) and CpG premethylated primate-replicated deletions (gray bars) are noted. Statistical analysis comparing the difference in distribution between the two groups was performed using t-test analysis, with P values noted when significant (P < .05).

Somatic replication of templates with 53 CGG repeats resulted in rare expansion events, which were not statistically significant, as their numbers were not greater in magnitude or frequency than those observed in the bacterial background (table 1). However, it was interesting to note that most of the expansions occurred when CGG was used as the lagging-strand template (six of the seven expansion events; pFX53N and pFX53A) (table 1), perhaps suggesting a role for sequence specificity not only in deletion events but in expansion events as well.

Replication Efficiency Is Correlated with CGG Stability

The timing of replication through the FRAXA chromosomal locus occurs late in the cell cycle (Howell and McDermott 1982) and is further slowed by CpG methylation of the expanded CGG repeat (Hansen et al. 1993). To address whether the replication rate is correlated to the observed deletion frequency, the replication efficiency of each template (with and without CpG premethylation) was measured relative to an unmethylated internal control SV40-ori template containing no CGG repeats (fig. 1B). pFX53E, which displayed the highest frequency of deletions, was also the most efficiently replicated (fig. 4 and table 1). Interestingly, CpG premethylation of pFX53E decreased not only its levels of instability but also its replication efficiency. The effect of CpG premethylation on replication efficiency was observed for all templates when compared with their unmethylated replication efficiencies (fig. 4 and table 1).

Replication Efficiency Is Not Associated with a Block in the Replication Fork at the CGG Tract

A decrease in replication efficiency may be due to a block in replication-fork progression in or around the CGG repeat tract (Samadashwily et al. 1997). By use of 2D gel electrophoresis (fig. 5), replication forks derived from replication intermediates of pFX53E and pFX53N were assessed for the presence of a replication block.
Figure 4  Replication efficiency. An aliquot of the starting mixture, composed of each pFX53 construct with unmethylated pSV40, was linearized with AlwNY. Primate-replicated DNA resulting from the cotransfection of this same starting mixture was also linearized with AlwNI, was further digested with DpnI to remove the unrepli-
cated parental template, and was then probed with the SV40-ori
fragment. Replication efficiency was calculated for each template in re-
lacion to pSV40 (as described in the “Material and Methods” section).

Neither pFX53E nor pFX53N, regardless of CpG meth-
ylation, displayed a block in replication that was local-
ized to the CGG tract within their respective replication
forks, by analysis of either y arcs (fig. 5D–5G) or bubble
arcs (data not shown). The lack of a replication block
may indicate that, during primate-mediated replication,
instead of forming complete replication stalls, the CGG
tract may actually incur pauses that are rapidly pro-
cessed and overcome.

Discussion

Our data suggest that the location of replication initi-
ation, in terms of proximity to the CGG tract, as well
as the sequence of the lagging-strand template, is a major
determinant of CGG deletions. Furthermore, CpG meth-
ylation stabilizes the repeat by decreasing the frequency
and magnitude of replication-mediated CGG deletions.
We also demonstrate, using a sensitive assay, that a block
in replication-fork progression does not occur at the re-
ppeat tract and, therefore, is not related to CGG deletions.
The lack of a distinct block in replication-fork progres-
sion (regardless of CpG methylation status) is consistent
with previous observations in cells from patients with
FRAXA (Hansen et al. 1993). Our observation that the
context of the CGG tract, relative to its location within
the replication fork, influences repeat stability may re-
flex the increased instability observed in knock-in and
YAC transgenic mice—supporting the contribution of
cis-elements to CGG instability (Bontekoe et al. 2001;
Peier and Nelson 2002).

Replication Initiation

Studies focusing on stability of the FRAXA CGG re-
peat within fetal tissues (Devys et al. 1992; Wohrle et
al. 1993; Malter et al. 1997) have provided evidence
suggesting that maternally inherited CGG expansions
either are meiotic in origin or occur early in postzygotic
development and are then stably maintained. The ability
to incur deletions within a construct containing 53 pure
CGG repeats, simply by moving the position of repli-
cation initiation, highlights the potential impact of the
location of replication initiation, with respect to the lo-
cation of the repeat within the replication fork (Cleary
et al. 2002; J. D. Cleary and C. E. Pearson, unpublished
data), on early postzygotic CGG instability. This im-
portance is further demonstrated by the rare occurrence
of expansions, in our system, mainly when CCG is used
as the lagging-strand template.

During early embryogenesis, replication initiation oc-
curs randomly throughout the genome before becoming
localized to specific replication origins (Hyrien et al.
1995; Sasaki et al. 1999). If localized replication initi-
ation becomes set at a very early time point, the window
for random replication initiation within human embryo-
genesis, and, therefore, the window for postzygotic CGG
instability, may be limited to a very early stage in de-

development. This limited window for instability can ex-
plain fetal-tissue data displaying minimal CGG heter-
ogeneity within and between various tissues, a pattern
that does not change throughout life.

CpG Methylation Stabilizes the CGG Repeat and
Prevents Deletions

For most TNR diseases, the parent-of-origin effect
plays an important role in repeat instability (reviewed
in Pearson 2003). What is unique to FRAXA, however,
is that expansion to the disease range occurs only upon

Table 2
Analysis of Templates Containing 20 CGG Repeats

<table>
<thead>
<tr>
<th>Template</th>
<th>No. of Expansions</th>
<th>No. of Deletions</th>
<th>Level of Stability (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>pFX20E + CH3</td>
<td>0</td>
<td>1</td>
<td>32 (97)</td>
</tr>
<tr>
<td>pFX20E</td>
<td>0</td>
<td>0</td>
<td>38 (100)</td>
</tr>
<tr>
<td>pFX20N + CH3</td>
<td>0</td>
<td>1</td>
<td>36 (97)</td>
</tr>
<tr>
<td>pFX20N</td>
<td>0</td>
<td>0</td>
<td>38 (100)</td>
</tr>
</tbody>
</table>
Figure 5  Replication-fork progression, 2D gel electrophoresis. A, Schematic representation of predicted Y arcs, as well as expected position of a block (if resulting from the CGG repeat tract) with respect to the restriction digests used for analysis. B and C, Maps of CGG repeat (blackened box), FRAXA genomic flanking sequences (dark gray lines), SV40-ori (blackened arrows), and the restriction digests used for 2D gel analysis of pFX53E (B) and pFX53N (C). The black rectangle below the line denotes the Xmn I/AlwNI fragment used as a probe. D and F, Replication forks of pFX53E and pFX53N. E and G, Replication forks of CpG premethylated pFX53E and pFX53N.

maternal transmission. If expansions were occurring postzygotically and were modulated only by repeat length, then premutation alleles should behave similarly (i.e., show a tendency toward expansion), regardless of the transmitting parent; yet, they do not. This disparity in CGG instability may be driven by the differing CpG methylation status of maternally and paternally inherited DNA during zygote formation. In murine zygotes, the paternally derived alleles undergo rapid demethylation before the very first round of replication (Tamanini et al. 1997). This rapid demethylation contrasts with the maintenance of CpG methylation found for the maternally inherited alleles. Our data show that replication-mediated deletion events can be stabilized by the presence of CpG methylation, which perhaps hints at why paternally inherited premutation alleles, void of CpG methylation and potentially prone to deletions, do not expand during early fetal development.

The presence of only unmethylated premutation-length alleles in the sperm of full-mutation males diverges from the stable maintenance, within the rest of their tissues, of aberrantly methylated full-mutation-length CGG repeats. Whereas sperm contain only premutation-length alleles, CGG length analysis of the adult testis revealed the presence of both full-mutation and unmethylated premutation lengths (Reyniers et al. 1993; Tassone et al. 1999). Furthermore, expression studies of the fragile X mental retardation protein (FMRP) in fetal primordial germ cells (PGCs) from male fetuses with FRAXA demonstrate that FMRP is being expressed in some PGCs by 17 wk of development (Malter et al. 1997; Tamanini et al. 1997). This would suggest that, within the PGCs expressing FMRP, a CGG deletion event occurred after the original expansion event, resulting in unmethylated premutation-length CGG alleles.

Early in embryogenesis, male PGCs develop, migrate to the extraembryonic mesoderm, proliferate, and then return to the developing fetus to form the gonadal tissue (Sutton 2000). While in the extraembryonic mesoderm, paternal—not maternal—X chromosomes become preferentially inactivated by CpG methylation (Goto et al. 1997). The lack of CpG methylation on the maternally inherited X chromosome, as inherited by a male with FRAXA, may provide an opportunity for CGG deletions to occur before the PGCs return to the fetus and become mitotically arrested. As development continues, the cells that have incurred deletions to premutation lengths are perhaps selectively maintained by their expression of FMRP, which is thought to be necessary for gonadal development (Tamanini et al. 1997).

Absence of a Block in Replication at the CGG Repeat

Similar to the chromosomes of patients with FRAXA, which show delayed and/or slowed replication but no distinct pause (Hansen et al. 1993), primate replication of the pFX/SV40 constructs displayed reduced replication efficiency when they were CpG methylated but did not display a replication block at the CGG tract, regardless of replication direction or CpG methylation.
The reduced replication efficiency we observed may reflect the generally slowed replication over the FRAXA locus, reported by Hansen et al. (1993), which may suggest delayed initiation or slowed fork progression through the region. The latter may permit enhanced replication fidelity due to an increase in the time available for replication-fork progression. The absence of a pause contrasts the strong replication-fork blocks observed at the CGG tract in both yeast and bacterial systems (Samadashwily et al. 1997; Pelletier et al. 2003). The difference in the ability to replicate through CGG repeats (primate replication vs. yeast or bacteria replication) may result from the fact that neither bacterial nor yeast genomes contain endogenous CGG/CGG repeats (Astolfi et al. 2003) and therefore may not have the mechanisms in place to properly replicate through the repeat tract without experiencing blockages. However, other explanations for the differences are possible, including the unidirectional replication versus bidirectional replication or the location of the initiation relative to the repeat tract (the yeast study used distances $\geq 2.5$ kbp, greater than the 74–613 bp distances used herein). Supporting this, replication stalls were recently observed for (CGG)$_n$ repeats in a primate SV40-derived episomal system, while positioned further apart from the origin (Sergei Mirkin, personal communication). However, the CGG deletions or their stabilization by CpG methylation reported here do not appear to correlate with a detectable propensity to pause replication-fork progression.

CGG deletions, similar to CAG/CTG instability, are influenced by tract length and replication-fork dynamics (J. D. Cleary and C. E. Pearson, unpublished data). However, CGG deletions are further influenced by CpG methylation, which highlights the need to study dynamic DNA mutations, not only in the proper sequence context, but also with as many epigenetic modifiers as allowed by the experimental system.

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Electronic-Database Information

Accession numbers and URLs for data presented herein are as follows:


References