ACMG Standards and Guidelines for fragile X testing: a revision to the disease-specific supplements to the Standards and Guidelines for Clinical Genetics Laboratories of the American College of Medical Genetics and Genomics

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Molecular genetic testing of the \(FMR1\) gene is commonly performed in clinical laboratories. Mutations in the \(FMR1\) gene are associated with fragile X syndrome, fragile X tremor ataxia syndrome, and premature ovarian insufficiency. This document provides updated information regarding \(FMR1\) gene mutations, including prevalence, genotype–phenotype correlation, and mutation nomenclature. Methodological considerations are provided for Southern blot analysis and polymerase chain reaction amplification of the \(FMR1\) gene, including triplet repeat–primed and methylation-specific polymerase chain reaction. In addition to report elements, examples of laboratory reports for various genotypes are also included.

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**Key Words:** \(FMR1\); fragile X syndrome; Southern blot analysis; triplet repeat–primed PCR

The American College of Medical Genetics and Genomics Laboratory Quality Assurance Committee has the mission of maintaining high technical standards for the performance and interpretation of genetic tests. In part, this is accomplished by the publication of the document “ACMG Standards and Guidelines for Clinical Genetics Laboratories,” which was published in its second edition in 1999 and is now maintained online (http://www.acmg.net/StaticContent/SGs/Section_G_2010.pdf). This subcommittee also reviews the outcome of national proficiency testing in the genetics area and may choose to focus on specific diseases or methodologies in response to those results. Accordingly, the subcommittee selected fragile X syndrome to be the first topic in a series of supplemental sections, recognizing that it is one of the most frequently ordered genetic tests and that it has many alternative methods with different strengths and weaknesses. This document is the second update to the original Standards and Guidelines for Fragile X testing that were published in 2001. It is designed for genetic testing professionals who are already familiar with the disease and the methods of analysis.

**FX 1: INTRODUCTION**

Disease-specific statements are intended to augment the current general *ACMG Standards and Guidelines for Clinical Genetics Laboratories*. Individual laboratories are responsible for meeting the Clinical Laboratory Improvement Amendments/College of American Pathologists (CAP) quality assurance standards with respect to appropriate sample documentation, assay validation, general proficiency, and quality control measures.
FX 2: BACKGROUND ON FRAGILE X SYNDROME

FX 2.1: Gene symbol/chromosome locus

*FMR1* is the gene symbol recognized by the HUGO gene nomenclature committee (http://www.genenames.org/data/hgnc_data.php?hgnc_id=3775). The gene was referred to as *FX* in the past. The chromosome locus is Xq27.3.

FX 2.2: OMIM number

The OMIM numbers are as follows: *309550 (FMR1), 300624 for fragile X mental retardation syndrome (includes fragile X-associated primary ovarian insufficiency), and 300623 for fragile X tremor/ataxia syndrome (FXTAS).

FX 2.3: Brief clinical description

The features of fragile X syndrome include specific cognitive deficits and certain characteristic, but nonspecific, physical features and behaviors. Most individuals with the premutation do not show fragile X syndrome–related features; however, some with high repeat sizes (>100 repeats) have been identified with learning difficulties, emotional problems, or even intellectual disability. Females with premutations (usually >80 CGG repeats) are at ~20% risk for premature ovarian insufficiency (POI). This condition is now referred to as FXPOI. There is no evidence to support an association between high normal and intermediate range *FMR1* alleles with a risk of POI. Older males and females with premutations are at risk for FXTAS. FXTAS is a late-onset, progressive development of intention tremor and ataxia often accompanied by progressive cognitive and behavioral difficulties including memory loss, anxiety, reclusive behavior, deficits of executive function, and dementia. The risk for FXTAS is higher in males who carry a premutation as compared with females. The penetrance of FXTAS increases with age and with premutation repeat length. Guidelines to identify patients for whom FXTAS testing is indicated are available. For more information on these disorders, see the online GeneReviews profile for *FMR1*-related disorders at http://www.ncbi.nlm.nih.gov/sites/GeneTests/ and the National Fragile X Foundation at http://www.nfxf.org.

FX 2.4: Mode of inheritance

Inheritance of the *FMR1* mutation is X-linked, although the pattern of fragile X syndrome is complicated due to the characteristics of the unstable repeat sequence mutation. In typical fragile X families, the mutation is a multistep expansion occurring over one or more generations in a region of CGG repeats in the 5′ untranslated region of the gene. Small expansions (premutations) are not generally associated with cognitive deficits in males and females. Large expansions (full mutations) are penetrant in all males and many females. With extremely rare exceptions, the parent of origin of the expansion to the full mutation is female.

FX 2.5: Gene description/normal gene product

The gene product is fragile X mental retardation protein (FMRP), a widely expressed RNA-binding protein. The fragile X syndrome is caused by a loss of the FMRP. FMRP is a selective RNA-binding protein that can form a messenger ribonucleoprotein complex that can associate with polysomes. FMRP has been shown to behave in *vitro* as an inhibitor of protein translation. At the neuroanatomic level, the fragile X brain differs from normal brain due to the presence of unusually long and thin dendritic spines in the cortical regions. The dendritic spines are the location where excitatory synaptic transmission occurs. FMRP appears to be associated with polyribosomes within dendritic spines of “wild-type” neurons. From these data, it has been proposed that FMRP is a translation suppressor that regulates protein synthesis locally in dendrites in response to synaptic stimulation signals. In the fragile X brain, translation of certain messages may be exaggerated because the normal inhibition provided by FMRP is absent.

Studies of *FMR1* mRNA expression provide evidence that expansion in the premutation range perturbs gene expression and may have pathophysiological consequences, particularly those related to FXTAS and ovarian failure (see section 2.3). Reductions in the amount of FMRP have been found in both lymphocytes and transformed lymphocytes of premutation carriers. Using a highly sensitive fluorescent assay, Kenneson et al. demonstrated a decrement in FMRP in individuals with expansions only slightly larger than the upper edge of the normal range. The reduction in FMRP is associated with an increase in *FMR1* mRNA in individuals with premutations. Expansion of the CGG repeats into the premutation range can shift transcription of *FMR1* mRNA from the usual, downstream-most start site to upstream sites. The utilization of alternative start sites may be correlated with increased *FMR1* mRNA transcription levels. RNA-mediated toxicity may result in the FXTAS phenotype.

FX 2.6: Mutational mechanism/abnormal gene product

Fragile X syndrome is caused by the deficiency or absence of FMRP. Theoretically, this can occur through any type of deletion or inactivating mutation, but in more than 99% of cases, there is an expansion of a segment of CGG repeats in the 5′ untranslated region of *FMR1*. Large CGG expansions in this region are associated with hypermethylation and inhibition of transcription.

FX 2.7: Listing of mutations

Mutations at locations other than the CGG repeat have been described. A listing can be found in the Human Gene Mutation Database at http://www.hgmd.cf.ac.uk/ac/index.php. Guidelines for detecting these relatively rare mutations are beyond the scope of this document.

FX 2.8: Prevalence and ethnic association of common mutations

FX 2.8.1: Full mutations. The general prevalence of males with a full mutation is estimated as ~1 in 4,000. The female prevalence rate is presumed to be approximately one-half of the male rate, or ~1 in 5,000–8,000. Two large North American studies using
anonymized newborns identified a prevalence of 1 in 5,161 US males and 1 in 6,209 Canadian males.

All major ethnic groups and races appear to be susceptible to expansion of the FMR1 CGG region. An extensive literature review indicated a prevalence of fragile X syndrome ranging from 1 in 3,717 to 1 in 8,198 among Caucasian males in the general US population. In another study carried out over 4 years in metropolitan Atlanta, the prevalence of fragile X syndrome was estimated to be 1 in 2,545 among African-American males and 1 in 3,717 among Caucasian males. However, the prevalence estimate for Caucasian males, determined from this and from other studies, fell within the 95% confidence interval for African-American males. The prevalence of the fragile X mutation in an Afro-Caribbean population in the French West Indies was similar (1 in 2,539) to that in the African-American population in Atlanta. Falik-Zaccai et al. have suggested that the Tunisian Jewish population is the only other ethnic group to have a higher prevalence of the fragile X syndrome than the Caucasian population. However, these studies have not been supported by other data. A blinded Taiwanese study estimated the prevalence of fragile X as 1 in 10,046 males. Further studies are required to determine if the frequency of the fragile X syndrome differs in ethnic populations.

**FX 2.8.2: Premutations.** A study of 2,300 US women identified 1 in 382 as carriers. In the largest US study to date (of over 119,000 tested), the overall carrier frequency among US females was 1.3% (0.61% for full mutation and 1.7% for premutation). A recent large-scale study of 21,411 anonymous Canadian females (mothers of newborns) identified 1 in 549 as carriers. Previous screens for the prevalence of premutations (with 55–101 repeats) in French-Canadian women estimated the carrier frequency to be 1 in 259. A subsequent study revealed a 1 in 1,760 prevalence of premutation alleles among Canadian males.

A study from Israel of 36,483 women who requested screening identified 1 in 157 as carriers. This is consistent with an earlier study of 9,459 women in Israel that found 1 in 152 with alleles having >54 repeats. In the individuals with no family history of the fragile X syndrome, 1 in 166 women were determined to have premutations with a CGG-repeat range of 55–101. This estimate of the premutation carrier frequency is approximately twofold higher than that reported in the studies performed in Canada. Toledano-Alhadef et al. obtained similar values when studying 14,334 preconceptual or pregnant women in Israel, namely, 1 in 113 women with >54 CGG repeats. This study excluded women with a family history of mental retardation. In addition, they found that the premutation carriers were well distributed among all the Jewish ethnic groups, in contrast to a previous study.

An Italian study of ~2,000 mothers and their newborns in the general population found a premutation carrier frequency of 1 in 109 females and 1 in 225 newborn males (56–70 CGG repeats). A fragile X screen of 10,000 newborn males in Taiwan showed a prevalence of 1 in 1,674. Therefore, the carrier frequencies vary widely among populations and may be higher than those determined in the French-Canadian population.

Among females with POI and simplex cases of adult males with cerebellar ataxia, the FMR1 premutation is identified in 4–6% and 2%, respectively. FX 2.9: Special testing considerations

**FX 2.9.1: Sensitivity and specificity.** CGG-repeat-expansion full mutations account for >99% of cases of fragile X syndrome. Therefore, tests that effectively detect and measure the CGG-repeat region of the FMR1 gene are >99% sensitive. Positive results are 100% specific. There are no known forms of FMRP deficiency that do not map to the FMR1 gene. Fragile X syndrome should not be confused with the unrelated syndrome associated with the FRAXE locus.

**FX 2.9.2: Indications for testing.** The ACMG and American Congress of Obstetricians and Gynecologists have published recommendations regarding fragile X testing for diagnostic testing and carrier detection. The identification of a full mutation in a male is considered diagnostic rather than predictive, inasmuch as penetrance of fragile X syndrome is virtually 100% in males and the age of onset is not variable.

The identification of a full mutation in a female may be diagnostic, but <50% of females with full mutations have intellectual disability. They may, however, have some manifestations of the disease such as avoidance personality, mood, or stereotypic disorders. Nonrandom X inactivation may explain the milder phenotype in females, although the extent of symptoms cannot be determined by X-inactivation patterns from diagnostic tests that determine the expansion and methylation in blood.

The identification of a premutation in an asymptomatic male or female undergoing carrier testing (e.g., due to a family history of intellectual disability) is predictive because FXPOI and FXTAS are not fully penetrant and are dependent on both age and allele size.

Population carrier screening and newborn screening for fragile X syndrome are not recommended at this time and should occur only under a research protocol.

**FX 2.9.3: Prenatal testing.** This test can be used for prenatal diagnosis in cells obtained from amniocentesis and chorionic villus sampling (CVS). Because methylation is not fully established at the time of CVS, the appearance of full mutations examined by a methylation-specific method may vary in CVS as compared with blood and amniocytes. Laboratories offering testing of chorionic villi must be aware of this tissue’s unique properties:

- Methylation associated with lyonization is usually not present, and methylation associated with full mutations may or may not be present. In the past, the hemimethylated status of this locus in this tissue had been thought of as a limitation or possible source of confusion. However, because it is unwarranted to use methylation status or
X-inactivation for phenotypic prediction of a full mutation, the possible hypomethylation of this tissue is no disadvantage, provided that the tissue-specific basis of the hypomethylation is understood.\textsuperscript{56,57} It is an acceptable option to omit methylation analysis entirely when testing CVS specimens. In the minor fraction of CVS cases with a result that is ambiguous between a large premutation and a small full mutation by size criteria alone, a follow-up amniocentesis may be required;

- The degree of somatic variation in a full mutation “smear” has a wider range of possibilities than is typically seen in blood specimens, from very limited to extraordinarily diffuse;
- Mosaicism between trophoblasts and somatic cells is theoretically possible. For this reason, when CVS results indicate a premutation, follow-up amniocentesis has been suggested to rule out mosaicism for a full mutation. However, there has been no known occurrence of this type of mosaicism.

FX 2.10: Nomenclature
The use of standard nomenclature is important for the accurate communication of results to health-care providers and is recommended by the ACMG\textsuperscript{58} and CAP.\textsuperscript{59} According to Human Genome Variation Society recommendations regarding the nomenclature for short sequence repeats, the designation for the \textit{FMR1} triplet repeat (based on the coding DNA reference sequence, NM\_002024.5) is c.-129CGG[X]. The start of the variable repeat is specified by -129, and CGG indicates the sequence of the repeat unit. The number of triplet repeats present is specified by X. If the exact size of the repeat cannot be determined (e.g., full mutations sized by Southern blot analysis), then the square brackets are replaced by parenthesis, (X), to signify uncertainties.\textsuperscript{60} Current Human Genome Variation Society recommendations do not address nomenclature for nucleotide repeat mutations with size mosaicism.

Standard nomenclature is recommended, although laboratorians and clinicians may continue to use common mutation nomenclature. Therefore, to avoid confusion, it is acceptable to describe a mutation using standard nomenclature followed by the common name in parentheses or to use the common name in one section of the report (e.g., results) and the standard name in another section (e.g., description of methods or interpretation).

FX 3: GUIDELINES

FX 3.1: Definition of normal and mutation categories
There are four allelic forms of the gene: normal, intermediate, premutation, and full mutation. The associated number of CGGs for each can be defined on the basis of our current information up to date. It must, however, be recognized that the borders of each definition may change with increased empirical data and research.

\textbf{FX 3.1.1: Normal alleles.} Normal alleles have a range of ~5 to ~44 repeats. The most common repeat length is 29 or 30 CGG repeats. Normal alleles have no meiotic or mitotic instability.

\textbf{FX 3.1.1.1:} In stable, normal alleles, the CGG region is interrupted by an AGG triplet after every 9 or 10 CGG repeats. The AGG triplets are thought to anchor the region during replication and prevent strand slippage. Direct testing for the AGG triplets is available clinically, although it is not routinely performed and its clinical usefulness is yet to be determined. It may help predict risk of expansions from premutations of <100 repeats.\textsuperscript{61}

\textbf{FX 3.1.2: Intermediate (gray zone, inconclusive, borderline).} The range from ~45 to ~54 repeats is intermediate (also referred to gray zone, inconclusive, or borderline). Alleles in this range can be considered normal in the sense that such alleles are not associated with fragile X syndrome and have not been observed to expand to a full mutation in one generation. Although earlier studies suggested an association between alleles in this size range and FXPOI, larger subsequent studies did not support these initial findings.\textsuperscript{6,8} A small number of patients meeting the criteria for FXAS with \textit{FMR1} intermediate alleles have been described,\textsuperscript{62} although larger studies are needed to determine the significance of this finding.

Minor increases and decreases in repeat number can occur when alleles of this size are passed on, but there is no measurable risk of a child with fragile X syndrome in the next generation. Alleles of this size may be associated with fragile X syndrome in future generations or in distant relatives. Alleles in this range can be referred to as premutations if they are confirmed by family studies to be traceable to a known full mutation or unambiguous premutation. A gray-zone allele of 52 repeats was reported to expand to a premutation allele of 56 repeats in one generation, which subsequently expanded to a full mutation allele in the next generation.\textsuperscript{63} Testing at-risk relatives of individuals with an intermediate allele may determine the stability of the allele in the family. However, the rate of expansions of intermediate alleles is not well understood.

\textbf{FX 3.1.3: Premutation.} Premutation alleles range from ~55 to ~200 repeats. These alleles are long repeat tracks that are unstably transmitted from parent to child. Expansions from the premutation size range to the full mutation typically occur during maternal transmission. Due to the possibility of somatic mosaicism, careful examination for mosaicism into the full mutation range is recommended when a premutation is detected. \textit{FMR1} alleles in the premutation size range are not hypermethylated and are not associated with fragile X syndrome. Although males and females with premutations and manifestations of some symptoms of fragile X syndrome have been reported, further studies are needed.\textsuperscript{64} Women with alleles in this range are considered to be at risk for having affected children.\textsuperscript{65-67} The smallest \textit{FMR1} premutation allele reported to expand to a full mutation in a single generation is 56 repeats.\textsuperscript{63} Females who carry an \textit{FMR1} premutation should be offered prenatal diagnosis for all pregnancies. Some clinicians may offer prenatal diagnosis to women whose reproductive partner carries a premutation as well. All at-risk family members of
known carriers should be offered testing to determine their status.

FX 3.1.3.1: The upper limit of premutations is sometimes said to be ~230. In fact, both numbers (200 and 230) are rough estimates derived from Southern blots in which large premutations were measured with increases of 0.5 to 0.6 kb, implying ~170 to 200 more triplet repeats than normal.

FX 3.1.3.2: Traditional PCR. The lower limit of premutations is ~170 repeats, however, due to diffusion artifacts in Southern blots. This number is derived from a restriction fragment length polymorphism (RFLP) analysis where size mosaics and methylation mosaics have been observed.68–71

FX 3.1.3.3: When methylation is present, tissue-specific differences can be seen. Individuals with size or methylation mosaicism may be higher functioning than individuals with completely methylated full mutations.

FX 3.1.4: Southern blot analysis

FX 3.2.1: Southern blot analysis

FX 3.2.1.1: Probe and restriction site combinations. Table 1 describes several single- and double-enzyme options that are commonly used and several probes that are available. Other restriction enzymes and probes can be used, if equivalence is demonstrated. Probes can be sustained in plasmid DNA and isolated using a plasmid preparation procedure, they can be prepared by PCR amplification, or they can be purchased commercially.

FX 3.2.1.2: In general, when using the StB12.3 probe, small premutations are more easily detected when the normal fragment is small and/or electrophoretic migration is long, whereas large/diffuse full mutations are more easily detected when the normal fragment is large and/or electrophoretic migration is short.

FX 3.2.1.3: Controls should be included to confirm the proper choice and activity of restriction enzymes and probe. They should ideally represent the more difficult-to-recognize genotypes. To verify digestion and hybridization parameters, a normal control will suffice. However, in fragile X blot analyses, the abnormal controls are extremely important because they provide quality control on the resolution of small premutations and the detectability of diffuse smears. Characterized reference material possessing specific FMR1 premutation CGG-repeat sizes may be obtained from the CDC Genetic Testing Reference Materials Coordination Program through the Coriell Institute for Medical Research.76

FX 3.2.1.4: For female patients, it should be noted that the degree of separation between two differently sized normal alleles could appear identical with that between a normal and a premutation allele (e.g., 20 and 44 repeats vs. 35 and 59 repeats). A Southern blot analysis with superior resolution and appropriate size standards or controls is required to distinguish between these possibilities (Figure 1). Alternatively, most PCR-based methods can provide the required resolution. Similar considerations apply to detection of premutation alleles in normal transmitting males.

FX 3.2.1.5: Because full mutations can be extremely diffuse and faint, signal to noise ratios must be very good. Laboratories are advised to be aware of the many different

Table 1 Probe and restriction site combinations

<table>
<thead>
<tr>
<th>Probe and restriction site combinations</th>
<th>Primary restriction sites and normal length</th>
<th>Optional internal methyl-sensitive sites</th>
<th>Probes and references</th>
</tr>
</thead>
<tbody>
<tr>
<td>EcoRI, 5.2 kb</td>
<td>Eagl, BssHII, NruI, etc.</td>
<td>StB12.3 (ref. 72), pE5.1 (ref. 73)</td>
<td></td>
</tr>
<tr>
<td>HindIII, 5.4 kb</td>
<td>Oxl1.9 (ref. 74)</td>
<td>BglII, 12 kb</td>
<td></td>
</tr>
<tr>
<td>PstI, 1.0 kb</td>
<td>NA</td>
<td>pfxa3 (ref. 75), Oxo0.55 (ref. 74)</td>
<td></td>
</tr>
</tbody>
</table>

NA, not appropriate.
Help discriminate between premutations and full mutations for the rare alleles that fall near the boundary (i.e., around 200 repeats); and

- Detect rare individuals who are methylation mosaics.

FX 3.2.1.7.2: In DNA extracted from tissues other than chorionic villi, the results of routine methylation analysis and PCR are sometimes confounded by an abnormal karyotype such as 45,X or 47,XXY. Individuals with testicular feminization (XY females) will have a male methylation pattern. In these cases, gender should be confirmed. Interpretation of results should take the individual’s karyotype into account, when it is available to the molecular laboratory.

FX 3.2.1.7.3: In DNA extracted from tissues other than chorionic villi, methylation analysis increases the difficulty of detecting females with small premutations who have highly skewed X-inactivation. Double digestion with a methyl-sensitive restriction enzyme causes the signal from each allele in a female to be split into active and inactive bands, forming four bands in a carrier female. When X-inactivation is balanced in a carrier, the two active bands are readily seen, although the two inactive bands may comigrate. However, if X-inactivation is heavily skewed, there will be only two visible bands. This is particularly challenging when the premutation is predominantly inactive because then it appears only in the upper region of the gel, where resolution is considerably poorer. For an example of a carrier with extremely skewed X-inactivation, see lane 13 of Figure 1. Lanes 3 and 4 show two females with oppositely skewed X-inactivation. The above data are true for the use of the StB 12.1 probe. Use of other probes such as pE5.1 will yield an additional small control band.

FX 3.2.1.7.4: *FMR1* methylation status should not be used to predict severity in fetal or newborn cases, regardless of whether the DNA was extracted from amniocytes, chorionic villi, or blood.

FX 3.2.1.7.5: In DNA extracted from chorionic villi, the *FMR1* region usually does not have methylation associated with X-inactivation, and it may or may not have hypermethylation associated with full mutations if the CVS procedure was performed before 12.5 weeks gestation. When testing DNA extracted from chorionic villi, methylation analysis is optional. Incidentally, methylation analysis before 12.5 weeks gestation can serendipitously alert a laboratory to maternal cell contamination in chorionic villus specimens inasmuch as methylation associated with X-inactivation is usually not present at this locus in tissue obtained via CVS; a strong normal inactive band can be a sign of possible maternal cell contamination. Other explanations for such a band include X-inactivation in some fetal cells or incomplete digestion. Further investigation would be called for.

**FX 3.2.2: PCR methods**

FX 3.2.2.1: Several sets of primers, PCR conditions, and methods of separation and detection have been published. Other primers and methods can be used if equivalence is
demonstrated. A particular region to be aware of in primer design is the deletional hotspot.\textsuperscript{82} PCR can be performed using \textsuperscript{32}P and electrophoresis on an acrylamide gel, or more commonly incorporating a fluorescently labeled primer followed by capillary electrophoresis (CE).

**FX 3.2.2.2:** Regardless of the locus, any PCR can theoretically fail to detect an allele if there is a polymorphism at the primer-binding site. There are no known polymorphisms that would affect any of the commonly used primers.

**FX 3.2.2.3:** Patient amplicon sizes should be determined using a size standard. For CE, a standard fluorescent-labeled size marker can be used. For acrylamide gels, a standard ladder, such as an M13 sequencing reaction or a set of carefully chosen, independently tested human references, can be used.

**FX 3.2.2.4:** Controls representing the genotypes to be distinguished should be used for each run. Refer to section 3.2.1.3 for a discussion of reference materials. The upper limit of allele size that can be successfully detected should be known, and a control corresponding to that size should be included in each run. Laboratories should confirm the size of their control DNA by sequencing (if possible) or by using verified reference materials.

**FX 3.2.2.5:** Amplification of CG-rich regions is difficult, and special conditions are required. The difficulty increases with increasing numbers of CGG repeats; therefore, many PCR strategies do not attempt to detect large alleles. In such a system, it is not possible to tell the difference between a female who is homozygous for a normal allele and one who has a large nonamplifiable second allele. Similarly, patients who are mosaics for premutations and full mutations will appear to have only premutations.

**FX 3.2.2.6:** When a PCR strategy is capable of detecting large alleles, amplification nevertheless may favor the smaller allele in any specimen with multiple alleles, i.e., females and mosaics. Such methods should be validated with carrier females and mosaics, in addition to males. Because of disproportinate amplification, PCR is not reliable for determining the ratio of different species in a mosaic individual.

**FX 3.2.2.7:** In PCR amplification of samples from females and mosaics, heteroduplexes can form. If denaturing polyacrylamide gel electrophoresis is used, conditions must be sufficiently denaturing to avoid heteroduplex artifact. If nondenaturing polyacrylamide gel electrophoresis is used, steps must be taken to distinguish between heteroduplexes and true abnormal alleles.

**FX 3.2.2.8:** Basic PCR amplification is not affected by methylation. Although PCR tests specifically modified to detect methylation status have been described,\textsuperscript{83,84} the original PCR strategies that have been in use for many years are completely independent of methylation.

**FX 3.2.2.9:** When a PCR strategy is used to detect full mutations, the presence of a deletion hotspot in the CGG-repeat region should be noted.\textsuperscript{82} Primers located within the deletion hotspot may result in failure to detect the expanded allele. Primers located upstream of the deletion hotspot may result in apparent size mosaicism.

**FX 3.2.2.10:** Triplet repeat–primed PCR

**FX 3.2.2.10.1:** Triplet repeat–primed PCR (TRP PCR) allows rapid detection of PCR products formed by a chimeric primer binding inside a triplet-repeat region. In TRP PCR for fragile X, one primer is anchored completely outside of the CGG-repeat region, whereas the other overlaps the CGG repeat and the adjacent nonrepeated sequence. A third primer can be anchored outside of the CGG region that, when paired with the opposite anchored primer, will amplify “over” the CGG repeat. This will increase the amount of full-length product from the largest CGG-repeat allele and in some assays enables accurate sizing of alleles up to 200 CGG repeats. From the chimeric primer annealing at each CGG repeat, multiple amplicons are made, forming products each with a length differing by three bases. A number of reports have been published describing TRP PCR for fragile X.\textsuperscript{85–89}

**FX 3.2.2.10.2:** Although products can be separated by ethidium-stained agarose gels to detect “smears,” combining TRP PCR with single-base-resolution fragment analysis, the “smear” on a lower-resolution agarose gel becomes characteristic “stutters” or “ladders” that are easily visualized. The stuttering will end at the allele with the greatest number of CGG repeats. An increase of this product can be seen, particularly if the third primer is used. For alleles with >200 repeats, a “compression” product can be seen, which can be used as a marker for an allele with >200 repeats, although the fragment cannot be sized (Figure 2). If a third primer is used, full mutations appear as a compression (or compacted) product. These are seen by CE as a compacted product of ~200 CGG repeats, thereby indicating a full mutation (Figure 3).

**FX 3.2.2.10.3:** PCR followed by CE at a single-base resolution has a high analytical sensitivity and specificity for detecting expanded alleles. A threshold can be set to distinguish premutation and full mutation (and intermediate, if desired) alleles from normal alleles. If no expansion is detected, no further testing is necessary. Alleles with a stuttering pattern past a threshold and consistent with an expansion can be tested further to determine methylation (by Southern blot analysis or methylation-sensitive TRP PCR) or determine size by traditional PCR and/or Southern blot analysis. Therefore, the simple yes/no answer for the presence of expansions can eliminate the need for Southern blot analysis in samples with a normal-sized and intermediate-sized \textit{FMR1} allele(s).

**FX 3.2.2.10.4:** TRP PCR resolves the challenges associated with apparent homozygous females, because the normal allele will not outcompete the expanded allele.

The increased sensitivity of the TRP PCR assay also resolves the difficulty of detecting mosaic males because mosaicism can be detected to ~10%. Laboratories using TRP PCR are encouraged to define the sensitivity of their assay using DNA with a normal \textit{FMR1} allele titrated with serial dilutions of full-mutation \textit{FMR1} DNA.

**FX 3.2.2.10.5:** Methylation PCR methods. Several methods besides Southern blot analysis have been described to determine methylation. Methylation-specific PCR involves...
the differential treatment of DNA with methylation-specific restriction enzymes followed by allele-specific PCR and resolution of the PCR products with CE. This method not only determines methylation status but also \textit{FMR1} allele size up to 250 repeats. However, to accurately size alleles >250 repeats, Southern blot analysis is needed.

\textbf{FX 3.2.2.11: Other methods.} Multiplexed ligation probe amplification has been described to identify males with methylated fragile X alleles. In this method, sequence-specific probes are hybridized to methylated and unmethylated alleles. Probes are simultaneously ligated and digested with a methylation-sensitive restriction endonuclease. A universal PCR primer set will amplify only probes that are ligated and undigested, indicating methylated alleles.

Real-time PCR has also been described with TaqMan chemistry and by melt-curve analysis, using methylation-specific PCR. TaqMan chemistry amplifies separately methylated and unmethylated specific alleles and provides a ratio based on amplification cycle thresholds. Using melt analysis, methylated and unmethylated alleles are amplified simultaneously but due to differences of GC content can be resolved by differences in melting temperature between methylated and unmethylated alleles. These methods have high analytical sensitivity and specificity for detecting methylation in males but are less sensitive and specific in females.

\textbf{FX 3.3: Interpretations}

\textbf{FX 3.3.1:} Example reports are included in the Supplementary Appendix online. In addition to the items described in the current general ACMG Standards and Guidelines for Clinical Genetics Laboratories (http://www.acmg.net/StaticContent/SGs/Section_G_2010.pdf), the following elements should be included in the report:

\textbf{FX 3.3.1.1:} State whether the method used was PCR, Southern blot analysis, or both. If Southern blot analysis, state the restriction enzymes and probes used. If PCR, describe the PCR method used (e.g., TRP PCR) and method used for separation and detection (e.g., CE).

\textbf{FX 3.3.1.2:} State the definitions used for normal, intermediate (gray-zone, borderline, inconclusive), premutation, and full mutation.

\textbf{FX 3.3.1.2.1:} Note that it is not necessarily obvious that the borderline category refers to the border between normal and premutation and not to the border between premutation and full mutation. Similarly, note that the term instability, which is often used with regard to borderline alleles to describe minor inter-generational or mitotic changes, may unintentionally suggest a risk of having an affected child or personal late-onset symptoms.

\textbf{FX 3.3.1.3:} Classify the patient’s result using the defined categories and Human Genome Variation Society nomenclature (refer to section 2.9.4). Common nomenclature can be included for clarity. The term size mosaic should be used for alleles that have significant subpopulations in both the premutation and full mutation range.

\textbf{FX 3.3.1.4:} All positive results should state that genetic counseling is recommended and testing is available for at-risk family members.

\textbf{FX 3.3.2:} The following descriptive elements may appear, with caution:

\textbf{FX 3.3.2.1:} The size of the alleles may be reported and could be of clinical use for premutation carriers. The premutation allele
ACMG STANDARDS AND GUIDELINES

size may be used for risk assessment in determining the chance of expansion in the offspring of carriers and also in determining the chance of FXTAS or FXPOI. If so, the precision used in quoting the size must be supportable by the precision of the size marker used, the sharpness of the bands or peaks, degree of stutter, and so on. It may be appropriate to state a range or use qualifying terms such as “approximately.” Descriptions such as “positive for an allele with 55–200 repeats” are ambiguous and should not be included in a laboratory report.

It has been the intent of the CAP/ACMG Biochemical and Molecular Genetics Resource Committee to standardize the accuracy of CGG-repeat quantitation. The acceptable range for sizing CGG repeats is based on the analysis of CAP/ACMG proficiency testing survey results for fragile X. Acknowledging the technical limitations of size analysis, the ACMG supports the following acceptable ranges for FMR1 clinical testing and/or as grading criteria for the CAP/ACMG proficiency testing survey: consensus size ± 5 repeats for alleles with <55 repeats; consensus size ± 10 repeats for alleles with 56–100 repeats; and consensus size ± 2 SDs for alleles with >100 repeats.

FX 3.3.2.2: Description of methylation may be provided. The two kinds of methylation must be clearly distinguished: methylation due to X-inactivation and hypermethylation of full mutations. The term methylation mosaic or incomplete methylation may be used if not all molecules in a full mutation are hypermethylated.

FX 3.3.3: The following helpful points on alternative diagnoses may be included:

FX 3.3.3.1: There are rare forms of FMRP deficiency not caused by CGG expansion, which may not be detected by this test.

FX 3.3.3.2: Intellectual disability associated with other fragile X sites, in particular FRAXE, or other gene mutations will not be detected with this test.

FX 3.3.3.3: DNA analysis for fragile X syndrome should be performed as part of a comprehensive genetic evaluation that includes routine cytogenetic analysis and more recently chromosomal microarray analysis as recommended by ACMG.

FX 3.3.4: Comments on phenotype, if included, should be abstract rather than case specific. The following concepts apply:

FX 3.3.4.1: All males with full mutations have fragile X syndrome to some degree. The severity cannot be predicted from the size of the full mutation, but if premutations are also present or if the majority of the full-mutation molecules are unmethylated, the phenotype may be less severe.

FX 3.3.4.2: Females with full mutations exhibit a wide spectrum of phenotypes. They may be as severely affected as a male with an expanded fragile X allele (which is itself a range of phenotypes). Females with full mutations may also exhibit very mild learning disabilities or have no detectable deficits. The severity cannot be predicted from the size of the full mutation, nor can it be predicted from the pattern of X-inactivation.

FX 3.3.4.3: Individuals with premutations should not be interpreted as unaffected carriers. Females who carry a premutation are at risk for FXPOI and FXTAS. Males with the premutation are at risk for FXTAS. If an individual referred for diagnostic testing due to intellectual disability, autism, or learning disability is found to carry a premutation, no association can be stated at this point in time. It should be considered as a coincident finding unless FMRP deficiency or mosaicism for a full mutation is detected.

FX 3.3.4.4: Individuals with intermediate alleles should be interpreted as unaffected. Even more so than a premutation, an intermediate allele is considered a coincidence when found in an individual referred for diagnostic testing due to intellectual disability, learning disability, or autism. FMRP deficiency or mosaicism for a full mutation can be investigated by methylation-sensitive Southern blot analysis but with less
likelihood of success because intermediate alleles are common in the general population.

FX 3.3.5: Comments on reproductive risk, if included, should be abstract rather than case specific. The following concepts apply:

FX 3.3.5.1: All affected males and the overwhelming majority of affected females inherit their mutations from their mothers. The mothers carry either a premutation or full mutation.

FX 3.3.5.2: Women with full mutations have a theoretical 50% chance of passing on the full mutation with each pregnancy.

FX 3.3.5.3: Women with premutations have a theoretical 50% chance of passing on the fragile X mutation with each pregnancy. If it is passed on, the chance the allele will increase to a full mutation depends on its size in the mother. Probabilities range from 3% for maternal alleles with CGG repeats from 55 to 59 (1/23 transmissions) to ~100% for maternal alleles with 90 CGGs and above. The smallest allele known to expand to the full mutation is 56 repeats. Laboratories should be familiar with publications on this topic, including any current publications.

FX 3.3.5.4: Men with premutations will almost always pass premutations to all of their daughters. An extremely rare phenomenon involves unaffected males with premutations who have had affected daughters, apparently by gonadal mosaicism for full mutations. The sons of men with premutations are not at risk for developing the fragile X syndrome or FXTAS.

FX 3.3.5.5: To date, there have been no reports of male or female carriers of intermediate alleles having offspring with an FMR1 allele in the full mutation range. Instability may be identified if the allele can be traced through the family to a known full mutation or unambiguous premutation. In the absence of such a connection, it may be possible to show meiotic instability or a specific repeat sequence pattern (absence of AGG interruptions) that is at higher risk for instability. Testing for AGG status is available, although the clinical usefulness of this information is not known at this time.

FX 4: ALTERNATIVE TESTING METHODS

FX 4.1: Cytogenetic analysis
Testing for the fragile site FRAXA at Xq27 is no longer an acceptable diagnostic method. Clinical and analytical specificity and sensitivity are both insufficient.

FX 4.2: Protein analysis
Immunohistochemical staining for FMRP is a valid diagnostic method in lymphocytes. Willemsen et al. demonstrated that staining for the FMRP protein in chorionic villus samples could be used as an alternative prenatal diagnostic method for detection of full mutations in male fetuses. The situation is more complicated in female fetuses for which some chorionic villi may be completely positive and others from the same sample may be completely negative for FMRP staining. The authors’ data shed light on the timing of X-inactivation in chorionic villus cells of the female fetus. The diagnostic application of this method is not recommended at this time for the prenatal diagnosis of females carrying FMR1 full mutations.

FX 5: POLICY STATEMENTS

FX 5.1
The American College of Medical Genetics and Genomics issued a policy statement titled “Fragile X Syndrome: Diagnosis and Carrier Testing” in 1994 (Am J Med Genet 53:380–381), which was updated in October 2005 (Genet Med 7:584–587). This document is also available online at http://www.acmg.net/resources/policy-list.asp. These Standards and Guidelines are in general agreement with that statement.

FX 5.2
In 2010, the American College of Obstetricians and Gynecologists issued a Committee Opinion, No. 469, on carrier screening for fragile X syndrome. These Standards and Guidelines are in general agreement with that opinion.

SUPPLEMENTARY MATERIAL

Supplementary material is linked to the online version of the paper at http://www.nature.com/gim

DISCLOSURE

The authors direct laboratories that offer clinical molecular genetic testing for fragile X syndrome.

REFERENCES


Example report 1: Diagnostic test for female with normal results

The Laboratory
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www.laboratory.com

<table>
<thead>
<tr>
<th>Patient Name:</th>
<th>Jane D.</th>
</tr>
</thead>
<tbody>
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<td>ID number:</td>
<td>057398468</td>
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<tr>
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<td>Gender:</td>
<td>Female</td>
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<tr>
<td>Ordering Physician:</td>
<td>Dr. Good</td>
</tr>
<tr>
<td>Date of Report:</td>
<td>June 15, 2012</td>
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<th>Test Performed:</th>
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<tr>
<td>Specimen Type:</td>
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</table>

Result: PCR analysis using DNA obtained from a blood sample detected 29 CGG repeats on one X chromosome and 32 CGG repeats on the other X chromosome. Using standard nomenclature, this genotype is described as c.-129CGG[29];[32].

Interpretation: These results do not provide evidence of the common trinucleotide repeat expansion observed in almost all patients with fragile X syndrome, indicating that a diagnosis of fragile X syndrome is highly unlikely in this patient. Due to the nature of the assay, we estimate that the number of CGG repeats is correct to within 5 repeats.
**Description of Methods:** Direct DNA analysis of the CGG repeat in the fragile X mental retardation 1 gene (*FMR1*) was performed by Triplet Repeat Primed PCR followed by capillary electrophoresis. Southern blot analysis, when performed, uses the StB12.3 probe and genomic DNA digested with EagI/EcoRI. Up to 44 copies of the CGG repeat within FMR-1 is considered normal. Premutation alleles are usually between 55-200 repeats. A gray zone exists for alleles between 45-54 repeats. The full mutation is characterized by the presence of approximately 200-230 repeats and greater.

Rare *FMR1* mutations, unrelated to trinucleotide expansion may not be detected by this assay. Possible diagnostic errors include sample mix-ups and genotyping errors resulting from trace contamination of PCRs, maternal cell contamination of fetal samples and from rare polymorphisms, which interfere with analysis. Low level mosaicism may not be detectable by this method. Mistaken paternity may be inadvertently identified. Mutation analysis should be combined with phenotypic, cytogenetic and pedigree data for the most accurate interpretation.

**Comment:** This test was developed and its performance characteristics determined by The Laboratory. It has not been cleared or approved by the U.S. Food and Drug Administration. The FDA has determined that such clearance or approval is not necessary. This test is used for clinical purposes. It should not be regarded as investigational or for research. This laboratory is certified under the Clinical Laboratory Improvement Amendments of 1988 (CLIA-1988) as qualified to perform high complexity clinical laboratory testing.

---

**This report was reviewed and approved by:**

________________________________________  ____________
John Doe, M.D., Ph.D.                  Date

Director, The Laboratory
Example report 2: Carrier test for male with an intermediate allele

The Laboratory
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www.laboratory.com

<table>
<thead>
<tr>
<th>Patient Name:</th>
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<td>Personal History:</td>
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<tr>
<td>Family History:</td>
<td>Intellectual disability</td>
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<tr>
<td>Ordering Physician:</td>
<td>Dr. Good</td>
</tr>
<tr>
<td>Date of Report:</td>
<td>June 15, 2012</td>
</tr>
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| Test Performed: | Fragile X |
| Indication for testing: | Carrier testing |
| Specimen Type: | Blood |

**Result:** PCR analysis using DNA obtained from a blood sample detected 48 CGG repeats. Using standard nomenclature, this genotype is described as c.-129CGG[48].

**Interpretation:** These studies indicate that this patient is not a carrier of a premutation or full mutation for the fragile X gene. If the family history is suggestive of fragile X syndrome, DNA studies and clinical evaluation of potential affected family members should be considered. Although CGG repeats of 45-54 are considered normal, the stability of this number of repeats over multiple generations has not been clearly established. In a male, the number of repeats would not be expected to increase significantly when passed to his daughters. However, the risk of expansion from his daughters is not known for certain. Studies of other family members could help determine the stability of the FMR-1 gene within this family. Due to the nature of the assay, we estimate that the number of CGG repeats is correct to within 5 repeats.
Description of Methods: Direct DNA analysis of the CGG repeat in the fragile X mental retardation 1 gene (FMR1) was performed by Triplet Repeat Primed PCR followed by capillary electrophoresis. Southern blot analysis, when performed, uses the StB12.3 probe and genomic DNA digested with Eagl/EcoRI. Up to 44 copies of the CGG repeat within FMR-1 is considered normal. Premutation alleles are usually between 55-200 repeats. A gray zone exists for alleles between 45-54 repeats. The full mutation is characterized by the presence of approximately 200-230 repeats and greater.

Rare FMR1 mutations, unrelated to trinucleotide expansion may not be detected by this assay. Possible diagnostic errors include sample mix-ups and genotyping errors resulting from trace contamination of PCRs, maternal cell contamination of fetal samples and from rare polymorphisms, which interfere with analysis. Low level mosaicism may not be detectable by this method. Mistaken paternity may be inadvertently identified. Mutation analysis should be combined with phenotypic, cytogenetic and pedigree data for the most accurate interpretation.

Comment: This test was developed and its performance characteristics determined by The Laboratory. It has not been cleared or approved by the U.S. Food and Drug Administration. The FDA has determined that such clearance or approval is not necessary. This test is used for clinical purposes. It should not be regarded as investigational or for research. This laboratory is certified under the Clinical Laboratory Improvement Amendments of 1988 (CLIA-1988) as qualified to perform high complexity clinical laboratory testing.

This report was reviewed and approved by:

_________________________    __________
John Doe, M.D., Ph.D.         Date
Director, The Laboratory
Example report 3: Carrier test for female with a premutation allele

The Laboratory
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www.laboratory.com

<table>
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<tr>
<th>Patient Name:</th>
<th>Jane D.</th>
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<tbody>
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<td>Dr. Good</td>
</tr>
<tr>
<td>Date of Report:</td>
<td>June 15, 2012</td>
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| Test Performed:       | Fragile X                   |
| Indication for testing: | Carrier testing            |
| Specimen Type:        | Blood                       |

**Result:** PCR analysis using DNA obtained from a blood sample detected 29 CGG repeats on one X chromosome and 95 CGG repeats on the other X chromosome. Using standard nomenclature, this genotype is described as c.-129CGG[29];[95].

**Interpretation:** These results indicate that this patient is a carrier of a premutation FMR1 gene. Males with a fragile X premutation are at increased risk of developing the fragile X-associated tremor/ataxia syndrome (FXTAS). Females who carry a fragile X premutation may also develop tremor and ataxia. In addition, females heterozygous for a premutation allele are at increased risk for premature ovarian insufficiency (FXPOI). Fragile X DNA studies are recommended for other at-risk family members. Prenatal diagnosis is available for all future pregnancies, since expansion to a full mutation can occur. Genetic counseling is recommended. Due to the nature of the assay, we estimate that the number of CGG repeats is correct to within 10 repeats.
Description of Methods: Direct DNA analysis of the CGG repeat in the fragile X mental retardation 1 gene (FMR1) was performed by Triplet Repeat Primed PCR followed by capillary electrophoresis. Southern blot analysis, when performed, uses the StB12.3 probe and genomic DNA digested with Eagl/EcoRI. Up to 44 copies of the CGG repeat within FMR-1 is considered normal. Premutation alleles are usually between 55-200 repeats. A gray zone exists for alleles between 45-54 repeats. The full mutation is characterized by the presence of approximately 200-230 repeats and greater.

Rare FMR1 mutations, unrelated to trinucleotide expansion may not be detected by this assay. Possible diagnostic errors include sample mix-ups and genotyping errors resulting from trace contamination of PCRs, maternal cell contamination of fetal samples and from rare polymorphisms, which interfere with analysis. Low level mosaicism may not be detectable by this method. Mistaken paternity may be inadvertently identified. Mutation analysis should be combined with phenotypic, cytogenetic and pedigree data for the most accurate interpretation.

Comment: This test was developed and its performance characteristics determined by The Laboratory. It has not been cleared or approved by the U.S. Food and Drug Administration. The FDA has determined that such clearance or approval is not necessary. This test is used for clinical purposes. It should not be regarded as investigational or for research. This laboratory is certified under the Clinical Laboratory Improvement Amendments of 1988 (CLIA-1988) as qualified to perform high complexity clinical laboratory testing.

This report was reviewed and approved by:

___________________                   __________
John Doe, M.D., Ph.D.             Date
Director, The Laboratory
Example report 4: Diagnostic test for male with a full mutation

The Laboratory
1111 Laboratory Avenue
City, State 00839
(555) 920-3333
www.laboratory.com

Patient Name: Tom D.
ID number: 057398468
DOB: 04/05/2009
Gender: Male

Patient Information
Personal History: Intellectual disability
Family History: Intellectual disability

Ordering Physician: Dr. Good
Date of Report: June 15, 2012

Test Performed: Fragile X
Indication for testing: Diagnostic testing
Specimen Type: Blood

Result: PCR analysis detected a full mutation \textit{FMR1} allele. Southern blot analysis revealed an expanded \textit{FMR1} allele in the full mutation size range of approximately 500 repeats. Methylation studies were consistent with the presence of a completely methylated \textit{FMR1} allele. Using standard nomenclature, this genotype is described as c.-129CGG(500).

Interpretation: These results confirm the diagnosis of fragile X syndrome. The severity of the clinical phenotype cannot be determined by these studies. Fragile X DNA studies are recommended for all at-risk family members. Genetic counseling is recommended.

Description of Methods: Direct DNA analysis of the CGG repeat in the fragile X mental retardation 1 gene (\textit{FMR1}) was performed by Triplet Repeat Primed PCR followed by capillary electrophoresis.
Southern blot analysis, when performed, uses the StB12.3 probe and genomic DNA digested with EagI/EcoRI. Up to 44 copies of the CGG repeat within FMR-1 is considered normal. Premutation alleles are usually between 55-200 repeats. A gray zone exists for alleles between 45-54 repeats. The full mutation is characterized by the presence of approximately 200-230 repeats and greater.

Rare FMR1 mutations, unrelated to trinucleotide expansion may not be detected by this assay. Possible diagnostic errors include sample mix-ups and genotyping errors resulting from trace contamination of PCRs, maternal cell contamination of fetal samples and from rare polymorphisms, which interfere with analysis. Low level mosaicism may not be detectable by this method. Mistaken paternity may be inadvertently identified. Mutation analysis should be combined with phenotypic, cytogenetic and pedigree data for the most accurate interpretation.

Comment: This test was developed and its performance characteristics determined by The Laboratory. It has not been cleared or approved by the U.S. Food and Drug Administration. The FDA has determined that such clearance or approval is not necessary. This test is used for clinical purposes. It should not be regarded as investigational or for research. This laboratory is certified under the Clinical Laboratory Improvement Amendments of 1988 (CLIA-1988) as qualified to perform high complexity clinical laboratory testing.

This report was reviewed and approved by:

John Doe, M.D., Ph.D. Date

Director, The Laboratory
Example report 5: Diagnostic test for male with a full mutation and methylation mosaicism

The Laboratory
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www.laboratory.com

<table>
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<th>Patient Name:</th>
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<td>Specimen Type:</td>
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Result: PCR analysis detected a full mutation \( FMRI \) allele. Southern blot analysis revealed an expanded \( FMRI \) allele in the full mutation size range of approximately 500 repeats with methylation mosaicism (also referred to as incomplete methylation). Using standard nomenclature, this genotype is described as \( c.-129CGG(500) \).

Interpretation: These results confirm the diagnosis of fragile X syndrome. The severity of the clinical phenotype cannot be determined by these studies. Fragile X DNA studies are recommended for all at-risk family members. Genetic counseling is recommended.

Description of Methods: Direct DNA analysis of the CGG repeat in the fragile X mental retardation 1 gene (\( FMRI \)) was performed by Triplet Repeat Primed PCR followed by capillary electrophoresis.
Southern blot analysis, when performed, uses the StB12.3 probe and genomic DNA digested with EaeI/EcoRI. Up to 44 copies of the CGG repeat within FMR-1 is considered normal. Premutation alleles are usually between 55-200 repeats. A gray zone exists for alleles between 45-54 repeats. The full mutation is characterized by the presence of approximately 200-230 repeats and greater.

Rare \textit{FMR1} mutations, unrelated to trinucleotide expansion may not be detected by this assay. Possible diagnostic errors include sample mix-ups and genotyping errors resulting from trace contamination of PCRs, maternal cell contamination of fetal samples and from rare polymorphisms, which interfere with analysis. Low level mosaicism may not be detectable by this method. Mistaken paternity may be inadvertently identified. Mutation analysis should be combined with phenotypic, cytogenetic and pedigree data for the most accurate interpretation.

\textbf{Comment:} This test was developed and its performance characteristics determined by The Laboratory. It has not been cleared or approved by the U.S. Food and Drug Administration. The FDA has determined that such clearance or approval is not necessary. This test is used for clinical purposes. It should not be regarded as investigational or for research. This laboratory is certified under the Clinical Laboratory Improvement Amendments of 1988 (CLIA-1988) as qualified to perform high complexity clinical laboratory testing.

\textbf{This report was reviewed and approved by:}

___________________                   __________
John Doe, M.D., Ph.D.             Date
Director, The Laboratory
Example report 6: Diagnostic test for male with a full mutation with size mosaicism

The Laboratory
1111 Laboratory Avenue
City, State 00839
(555) 920-3333
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Patient Name: Tom D.
ID number: 057398468
DOB: 04/05/2009
Gender: Male

Ordering Physician: Dr. Good
Date of Report: June 15, 2012

Test Performed: Fragile X
Indication for testing: Diagnostic testing
Specimen Type: Blood

Result: PCR analysis detected a full mutation \textit{FMR1} allele with mosaicism for a premutation of approximately 150 repeats. Southern blot analysis revealed an expanded \textit{FMR1} allele in the full mutation size range of approximately 500 repeats with mosaicism for a premutation of approximately 150 repeats. Methylation studies were consistent with the presence of a completely methylated \textit{FMR1} full mutation allele and unmethylated premutation.

Interpretation: These results confirm the diagnosis of fragile X syndrome. The severity of the clinical phenotype cannot be determined by these studies. Fragile X DNA studies are recommended for all at-risk family members. Genetic counseling is recommended.

Description of Methods: Direct DNA analysis of the CGG repeat in the fragile X mental retardation 1 gene (\textit{FMR1}) was performed by Triplet Repeat Primed PCR followed by capillary electrophoresis. Southern blot analysis, when performed, uses the StB12.3 probe and genomic DNA digested with
EagI/EcoRI. Up to 44 copies of the CGG repeat within FMR-1 is considered normal. Premutation alleles are usually between 55-200 repeats. A gray zone exists for alleles between 45-54 repeats. The full mutation is characterized by the presence of approximately 200-230 repeats and greater.

Rare FMR1 mutations, unrelated to trinucleotide expansion may not be detected by this assay. Possible diagnostic errors include sample mix-ups and genotyping errors resulting from trace contamination of PCRs, maternal cell contamination of fetal samples and from rare polymorphisms, which interfere with analysis. Low level mosaicism may not be detectable by this method. Mistaken paternity may be inadvertently identified. Mutation analysis should be combined with phenotypic, cytogenetic and pedigree data for the most accurate interpretation.

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**This report was reviewed and approved by:**

____________________  __________
John Doe, M.D., Ph.D.     Date
Director, The Laboratory