Interactions of the G quartet forming semaphorin 3F RNA with the RGG box domain of the fragile X protein family

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ABSTRACT

Fragile X syndrome, the most common cause of inherited mental retardation, is caused by the transcriptional silencing of the fmr1 gene due to an unstable expansion of a CGG trinucleotide repeat and its subsequent hypermethylation in its 5' UTR. This gene encodes for the fragile X mental retardation protein (FMRP), an RNA-binding protein that has been shown to use its RGG box domain to bind to G quartet-forming RNA. In this study, we performed a detailed analysis of the interactions between the FMRP RGG box domain and one of its proposed RNA targets, human semaphorin 3F (S3F) RNA by using biophysical methods such as fluorescence, UV and circular dichroism spectroscopy. We show that this RNA forms a G quartet-containing structure, which is recognized with high affinity and specificity by the FMRP RGG box. In addition, we analyzed the interactions of human S3F RNA with the RGG box and RG cluster of the two FMRP autosomal paralogs, the FXR1P and FXR2P. We found that this RNA is bound with high affinity and specificity only by the FXR1P RGG box, but not by the FXR2P RG cluster. Both FMRP and FXR1P RGG box are able to unwind the G quartet structure of S3F RNA, however, the peptide concentrations required in this process are very different: a ratio of 1:6 RNA:FMRP RGG box versus 1:2 RNA:FXR1P RGG box.

INTRODUCTION

The fragile X mental retardation syndrome (FXMR/FXS), an X-linked disorder, is the most common cause of inherited mental retardation (1). At the molecular level, the progressive expansion of (CGG)ₙ repeats and the hypermethylation of the CpG island, in the 5'-untranslated region (5'-UTR) of the fmr1 gene causes its transcriptional inactivation. The resulting suppression of the encoded protein, named the fragile X mental retardation protein (FMRP), has been shown to be the underlying cause of this syndrome (2,3). FMRP is a putative nucleocytoplasmic shuttling protein (4), found abundantly expressed in the neurons and several studies suggested that this protein participates in the synaptic plasticity of neurons by acting on post-transcriptional control of gene expression (5–7). FMRP has been proposed to regulate the transport and translation of specific messenger RNA targets (mRNA) in a manner critical for neuronal development. It has also been shown that this protein has nucleic acid chaperone properties (8). The sequence analysis of FMRP revealed that the 632 amino acid protein contains two types of RNA-binding motifs: two K-homology (KH) domains and one arginine-glycine-glycine rich region (RGG box), suggesting that the protein exerts its function through RNA binding (9).

FMRP has two autosomal paralogs, the FXR1 and FXR2 proteins (FXR1P and FXR2P), with which it forms the fragile X-related protein family (10,11). Sequence analysis revealed that the two proteins have ~60% amino acid identity, with regions of 90% sequence identity to FMRP (11,12). FXR1P and FXR2P are also cytoplasmic RNA-binding proteins, each containing two KH domains. The FXR2P is divergent from FMRP and FXR1P in the C-terminal region, in that it has a RG cluster instead of an RGG box. The fact that the FXR proteins have been found to be associated predominantly with the ribosomal 60S subunit, and that they have similar RNA-binding domains lead to the suggestion that the FXR1P and FXR2P might compensate for the FMRP function (13). However, the comparison of the expression levels of each of these proteins in different tissues and cellular distributions suggests that each of the FXR proteins might have an independent function (14).

Biochemical studies conducted in vitro showed that FMRP uses its RGG box to bind with high affinity to target RNA sequences proposed to contain G quartet structures (15–19). A G quartet is formed from four guanine residues arranged in a planar configuration,
which is stabilized by Hoogsteen-type hydrogen bonds. Several such planar structures can stack and are stabilized by potassium or sodium cations, but they do not form in the presence of lithium cations (20–22). The specific mechanism by which FMRP interacts with its mRNA ligands and regulates their translation still remains poorly understood. Human semaphorin 3F (S3F) mRNA has been identified both in vivo and in vitro as a potential mRNA target of the FMRP (16,23) and it has been proposed that its interactions with the FMRP RGG box occur in a G quartet-dependent manner (16). S3F mRNA encodes for the SEMA 3F protein which belongs to the class 3 semaphorins, a family of secreted and transmembrane signaling molecules that play crucial roles in the nervous (neuronal migration and axon pathfinding), immune and cardiovascular systems. Every member of this family has the 500 amino acid signature, the semaphorin domain (24,25). SEMA 3F is a putative secreted protein that has been suggested to have chemotactic and repulsion functions. The expression of the SEMA 3F gene has also been reported to suppress tumor formation in nude mice and to cause the alteration of the cellular response to drugs inducing apoptosis (26).

The goal of this study is to contribute to our understanding of the principles of recognition between FMRP and its RNA target(s), by analyzing its interactions with S3F RNA. We show here that S3F RNA adopts a parallel intramolecular G quadruplex structure and we use thermodynamic methods to determine if the stability and the secondary structure of this RNA are altered by its interactions with the FMRP RGG box. We also investigate the binding of the RGG box and RG cluster of the FXR1P and FXR2P to S3F RNA, in an effort to determine if the recognition of the structural elements in this RNA is unique to the FMRP RGG box.

MATERIALS AND METHODS

RNA synthesis

The unlabeled RNA oligonucleotides (S3F-lg, S3F-M2 and Munc-13 site 1) were synthesized by in vitro transcription reactions using T7 RNA polymerase (produced in-house), following the procedure by Milligan and Uhlenbeck (27). The synthetic DNA templates were purchased from Trilink Biotechnologies, Inc. The RNA oligonucleotides were purified by denaturing gels and electrophoretic elution, followed by extensive dialysis against 10 mM Tris (pH 7.5) or 10 mM cacodylic acid (pH 6.5). S3F-M2 was constructed by introducing two point mutations in S3F-lg to stabilize the stem (Figure 2A).

The S3F-M2_15AP RNA used in the fluorescence spectroscopy studies was constructed by replacing the adenine at the 15th position in S3F-M2 with the highly fluorescent analog 2-aminopurine (2-AP) (Dharmacon, Inc.). The pure 2-aminopurine labeled RNA was re-suspended in sterile deionized water.

All experiments were performed with RNA samples that were annealed by heating at 95°C, followed by slow cooling at room temperature for 20 min.

Peptide synthesis

The FMRP, FXR1P RGG boxes and the FXR2P RG cluster were chemically synthesized and purified by the Peptide Synthesis Unit at the University of Pittsburgh, Center for Biotechnology & Bioengineering.

UV spectroscopy thermal denaturation experiments

The UV melting curves of the unlabeled S3F-lg, S3F-M2 and of the 2-AP labeled S3F-M2_15AP RNAs were collected using a Varian Cary 3E spectrophotometer equipped with a Peltier cell. The samples were annealed in the standard buffers 10 mM Tris, pH 7.5 or 10 mM cacodylic acid, pH 6.5, containing either 150 mM KCl or 150 mM LiCl. The RNA samples were heated from 20 to 99°C at a rate of 0.2°C/min, recording points every 1°C. Blank samples were treated in the same manner. Depending upon the RNA concentration, the spectral absorbance was measured either at 295 or 305 nm, wavelengths that have been previously identified to be sensitive to G quadruplex dissociation (28). To determine if S3F-M2 folds into an intermolecular or intramolecular conformation, the melting temperature of the G quadruplex structure was determined at different RNA concentrations in the range 10–80 μM.

RNA:RGG peptide complexes were formed by mixing the RGG boxes of the FMRP and FXR1P or the RG cluster of FXR2P with S3F-M2 RNA in a 1:1 ratio, and allowed to equilibrate for 30 min. The melting temperature of the complex of S3F-M2 with various RGG boxes was determined using the same parameters described above for the free RNA. All melting curves of the S3F-M2:RGG peptide complexes were corrected by subtracting the melting curves of the corresponding free RGG peptides.

The transition of the G quadruplex dissociation in S3F-M2 and S3F-M2_15AP was identified and fitted assuming an independent two state model:

\[
A(T) = \frac{A_U + A_F e^{\frac{-H}{RT}} e^{\frac{\Delta H}{kT}}}{e^{\frac{-\Delta H}{RT}} e^{\frac{\Delta S}{kT}} + 1}
\]

where \(A_U\) and \(A_F\) represent the absorbance of the unfolded and native G quadruplex RNA, respectively, and \(R\) is the universal gas constant.

Circular dichroism spectroscopy

The CD spectra were recorded on a Jasco J-810 spectropolarimeter at 25°C. The G quartet formation in the S3F-M2 RNA structure (at a 10 μM concentration in 10 mM Tris, pH 7.5) was monitored by titrating increasing amounts of KCl from a 4 M stock solution to a final concentration of 150 mM. The spectra were measured between 200 and 350 nm and corrected for solvent contributions and dilutions. Each spectrum was scanned three times with a 1 s response time and a 2 nm bandwidth. For the binding studies, increasing amounts of the RGG peptides (0–100 μM) were titrated into a fixed concentration of S3F-M2 RNA (10 μM) in 10 mM cacodylic acid, pH 6.5 and 150 mM KCl. The CD spectra were recorded after each addition of the peptide and the interactions...
were monitored by measuring the molar ellipticity of the RNA at 264 nm. All spectra were corrected by subtracting the free RGG peptide contributions at each peptide concentration.

To determine if the addition of a 1:10 RNA:RGG peptide ratio results in the degradation of the RNA, these samples were treated with proteinase K (1 μg) for 1 h at 25 °C, which degrades the RGG peptides. Subsequently, the CD spectra were re-recorded and corrected for the presence of proteinase K.

Fluorescence spectroscopy

Steady-state fluorescence spectroscopy measurements of S3F-M2_15AP RNA were performed on a J.Y. Horiba Fluoromax-3 equipped with variable temperature control. The excitation wavelength was at 310 nm and the emission spectrum was recorded in the range of 330–450 nm.

The binding of the FMRP RGG box to S3F-M2_15AP was measured by titrating increasing concentrations of the peptide to a fixed concentration of 150 nM S3F-M2_15AP. The same procedure was repeated for the FXR1P RGG box. The binding dissociation constant, $K_d$, was determined by fitting the binding curves to the equation:

$$F = 1 + \left(\frac{I_f}{I_B} - 1\right)$$

$$\frac{(K_d + [P])_i + [RNA]_i}{2} - \frac{(K_d + [P])_i + [RNA]_i}{2} - 4\sqrt{(K_d + [P])_i + [RNA]_i}$$

where $I_f$ and $I_B$ represent the steady-state fluorescence intensities of the free and bound S3F-M2_15AP, $[RNA]_i$ is the total concentration of S3F-M2_15AP and $[P]_i$ is the total RGG box peptide concentration.

Competition experiments were performed by monitoring the binding of the FMRP or FXR1P RGG peptides to S3F-M2_15AP in the presence of a 10-fold excess of unlabeled Munc-13 site 1 RNA or in the presence of a 6-fold excess of the FXR2P RG cluster peptide.

The thermodynamic parameters for the FMRP RGG box binding to S3F-M2_15AP were determined by measuring the $K_{obs} = 1/K_d$ at different temperatures in the range 20–45 °C. The standard enthalpy and entropy of binding were determined from the slope and intercept of the graph:

$$R \ln K_{obs} = \Delta S_b^0 - \frac{1}{T} \Delta H_b^0$$

NMR spectroscopy

The 1D $^1$H spectra of S3F-M2 RNA were acquired at 29 °C on a 500 MHz Varian Unity Plus spectrometer. The water suppression was accomplished using the jump-and-return pulse sequence (29) with the maximum of excitation set at 11 p.p.m. S3F-M2 RNA (387 μM) was prepared in 10 mM Tris (pH 7.5) at a 90% H$_2$O/10%D$_2$O ratio.

The melting of the S3F-M2 RNA stem structure was monitored by recording the 1D $^1$H NMR spectrum at different temperatures in the range 20–60 °C. These experiments were performed on a Bruker AVANCE™ 500 MHz NMR spectrometer.

Electrophoretic mobility gel shift assay (EMSA)

EMSA reactions were performed in a total volume of 15 μl. The RNA:peptide complexes were prepared by mixing the RGG peptides with S3F-M2 or S3F-M2_15AP in 1:1 or 1:2 ratios and resolved on 15% non-denaturing acrylamide gels that were run in the presence of 75 mM KCl, at 35 V. The electrophoretic mobilities of the free RNA and the S3F RNA:peptide complexes were visualized by UV-shadowing at 254 nm, using an AlphaImager HP (AlphaInnotech, Inc.).

RESULTS AND DISCUSSION

Semaphorin 3F mRNA adopts a G quartet structure

It has been proposed that the G quartet motif is important in the FMRP recognition of its RNA targets, and S3F mRNA has been identified as a potential in vivo target of the protein, based on the fact that its G-rich sequence could fold into this structural motif (16). The interactions of FMRP with the semaphorin mRNA fragment used in this study (Figure 1A) have been visualized in living mammalian cells; moreover, it has been shown that the mutation of the GG doublets proposed to be involved in the G quartet formation abolishes these interactions (23), supporting the idea that the S3F RNA recognition occurs in a G quartet-dependent manner.

We have expressed and purified a 38-nt RNA, containing the 34-nt G-rich fragment of human S3F mRNA proposed to interact with FMRP, to which four extra nucleotides (GGGA) were added at the 5'-end for transcription purposes (16) (named S3F-3g; Figure 1A). To determine if this RNA folds into a G quadruplex structure, we used a combination of CD, fluorescence, NMR and UV spectroscopy techniques. It is well known that the G-rich nucleic acid sequences fold in to G quartets in the presence of cations like K$,^+$, by forming cation–dipole interactions with the guanine residues (21). CD spectroscopy has been extensively used to analyze the G quadruplex structure in DNA and RNA. Typically, there are two types of CD spectra observed for G quadruplexes: type I, which exhibits a positive band around 265 nm and a negative band around 240 nm and type II, which exhibits a positive band around 295 nm and a negative band around 260 nm (30).

At least for ’intermolecular’ G quadruplexes, there is a strong correlation between the parallel quadruplex and type I CD spectrum (31,32) and between the antiparallel quadruplex and type II CD spectrum (33,34). In the case of intramolecular G quadruplexes, there are also examples of parallel type quadruplexes exhibiting type I of CD spectrum (35–38) and of antiparallel quadruplexes exhibiting type II of CD spectrum (35,39,40). However, this correlation is not as clearly established in the case of intramolecular G quadruplexes, since there are not enough high-resolution structures available. Generally, the CD spectroscopy results cannot be exclusively used to assign definitively a particular type of fold to a G quadruplex structure, as there are examples of more complex CD spectra exhibiting the features of both, type I and type II spectra (41).
The CD spectrum of S3F-lg RNA folded in the presence of K⁺ ions is of type I, with a positive band at 263 nm and a negative one at 238 nm (Figure 1B), confirming the presence of G quartet structural elements in this RNA. This result suggests that the fold of this G quadruplex is of parallel nature, however, as discussed above, this can only be confirmed by the high-resolution structure of this RNA.

The UV spectroscopy thermal denaturation profile of S3F-lg measured at 305 nm, shows a characteristic hypochromic transition between 52 and 72 °C, corresponding to G quadruplex dissociation (28) (Figure 1C), supporting the presence of a G quadruplex structure in this RNA.

G-rich sequences are notorious for forming alternate G quartet structures in vitro, and to determine if this is true for S3F-lg RNA as well, we used native gel electrophoresis. Two conformations are observed on a 15% native gel performed in the presence of 75 mM KCl at all RNA concentrations investigated (Figure 1D and data not shown).

To obtain higher-resolution information about the structure of S3F-lg RNA, we have used one-dimensional (1D) ¹H NMR spectroscopy. Resonances are present in the 10–12 ppm. proton region corresponding to imino protons involved in G quartets, however, they are very broad, suggesting that this RNA exchanges between different conformations. Surprisingly, we did not observe any imino proton resonances corresponding to Watson–Crick base pairs in the 12–14 ppm. region, indicating that the stem structure proposed in Figure 1A does not exist in S3F-lg (data not shown). One possible explanation for this finding is that the addition of the extra four nucleotides (GGGA) at the beginning of the S3F-lg sequence might actually promote the folding of this RNA into an alternate G quadruplex structure, since they contribute to the formation of an uninterrupted stretch of 10 purines.

In an effort to promote the folding of S3F RNA into a single conformer we have removed its first four GGGA
nucleotides [that were added only for transcription purposes (16)], and introduced specific point mutations at positions 3 (G to C) and 4 (G to U) and at the complementary positions 31 (U to A) and 32 (U to C), respectively (labeled in blue in Figure 2A). This mutated RNA, named S3F-M2 RNA, no longer contains a stretch of 10 purines at its 5' end and of six consecutive guanines in the region proposed to fold into a stem structure (Figure 2A). We first investigated whether S3F-M2 RNA maintained the ability to form a G quadruplex structure. Upon titration of increasing concentrations of KCl, the CD spectrum of S3F-M2 RNA showed the spectral features of a type I G quadruplex CD spectrum, with a strong positive band at 263 nm and a negative band at

Figure 2. (A) Secondary structure of the S3F-M2 RNA fragment with the mutated base pairs highlighted in blue (49). (B) Proposed G quadruplex parallel structure of S3F-M2 RNA. The 2-AP insertion in S3F-M2_15AP is highlighted in red. (C) CD spectra showing the G quadruplex formation in S3F-M2 RNA by titrating increasing concentrations of KCl. (D) Fifteen percent of native gel electrophoresis of S3F-M2 RNA at various RNA concentrations: 3 μM (lane 1), 6 μM (lane 2), 10 μM (lane 3), 20 μM (lane 4), 30 μM (lane 5), 50 μM (lane 6), 100 μM (lane 7). (E) 1D 1H NMR spectrum of S3F-M2 RNA (387 μM) in 10 mM Tris pH 7.5 and various KCl concentrations.
238 nm (Figure 2C), very similar to that of S3F-lg RNA. The native gel electrophoresis of S3F-M2 RNA indicates that at concentrations <10 μM this RNA exists in a single conformation (black arrow in Figure 2D, and data not shown), whereas at higher concentrations S3F-M2 adopts more conformations.

The 1D 1H NMR spectrum of S3F-M2 shows resonances corresponding to imino protons involved in G quartets, as well as resonances corresponding to Watson–Crick base pairs, indicating the presence of both a stem and a G quadruplex in the structure of this RNA (Figure 2E). However, the G quartet imino proton resonances are very broad, consistent with an exchange between the different S3F-M2 conformers formed at the high RNA concentration required when using this technique. The exchange between different RNA conformations is also supported by the Watson–Crick imino protons, whose resonances become much broader upon addition of increasing KCl concentrations. These findings hindered our efforts to pursue high-resolution NMR spectroscopy studies of the S3F-M2 RNA structure.

**Thermodynamics of G quartet formation in S3F-M2 RNA**

We have employed UV spectroscopy to obtain the thermodynamic parameters of G quartet formation in S3F-M2 RNA. The UV thermal melting profile of 10 μM S3F-M2 RNA folded in the presence of 150 mM KCl shows a hypochromic transition between 38 and 67°C (indicated in red in Figure 3A), and a hyperchromic transition starting around 75°C (indicated in red in Figure 3A). However, the G quartet imino proton resonances are very broad, consistent with an exchange between the different S3F-M2 conformers formed at the high RNA concentration required when using this technique. The exchange between different RNA conformations is also supported by the Watson–Crick imino protons, whose resonances become much broader upon addition of increasing KCl concentrations. These findings hindered our efforts to pursue high-resolution NMR spectroscopy studies of the S3F-M2 RNA structure.

For 'intramolecular' species with *n* number of strands, 1/*T*_m* depends linearly on the natural logarithm of the total RNA concentration (*c*_T*):

\[
\frac{1}{T_m} = \frac{R(n-1)}{\Delta H_{vH}^0} \ln c_T + \frac{\Delta S_{vH}^0 - (n-1)R \ln 2 + R \ln n}{\Delta H_{vH}^0} \tag{4}
\]

where *R* is the gas constant and \(\Delta H_{vH}^0\) and \(\Delta S_{vH}^0\) are the Van’t Hoff thermodynamic parameters. For 'intramolecular' species, *T*_m* is independent of the total RNA concentration *c*_T*:

\[
n = 1 \text{ and } \frac{1}{T_m} = \frac{\Delta S_{vH}^0}{\Delta H_{vH}^0}
\]

Lower concentrations of RNA (<10 μM) favor the formation of a single species, with a melting temperature *T*_m* of ~52°C (Figure 3C, blue trace). However, at higher RNA concentrations (>10 μM) a second hypochromic transition appears in the range 63–86°C, corresponding to a new S3F-M2 conformation with a melting temperature of ~79°C (Figure 3C, brown trace). These findings are consistent with the native gel electrophoresis results that
showed the presence of more conformations in S3F-M2 at RNA concentrations higher than 10 μM (Figure 2D). The Tm of the 38–67°C hypochromic transition is independent of the RNA concentration (Figure 3D), indicating that the G quartet conformation formed by S3F-M2 at low RNA concentrations is ‘intramolecular’.

The standard enthalpy, entropy and free energy of G quartet formation in S3F-M2 RNA, which were obtained by fitting the 38–67°C hypochromic transition to Equation (1) (Materials and Methods section), are summarized in Table 1. The values of the thermodynamic parameters for G quartet formation (ΔH° = -43.1 ± 0.1 kcal/mol and ΔG° = -3.6 ± 0.1 kcal/mol) are consistent with the presence of two G quartet planes in the structure of S3F-M2 RNA [the enthalpy of formation of a single G quartet plane in an intramolecular G quadruplex, measured in
Table 1. Thermodynamic parameters for the formation of the G quartet structure of S3F-M2 RNA

| Molecule | $T_m$ (°C) | $\Delta H_{VH}$ (kcal/mol) | $\Delta S_{VH}$ (cal/mol K) | $\Delta G_{VH}$ at 25°C (kcal/mol) |
|----------|------------|---------------------------|---------------------------|----------------------------------|
| S3F-M2   | 51.6±0.1   | -41.2±0.1                 | -135.4±0.4                | -3.2±0.1                         |
| S3F-M2   | 64.7±0.1   | -38.6±0.1                 | -114.3±0.4                | -4.5±0.1                         |
| + FMRP RGG |           |                           |                           |                                  |
| S3F-M2   | 60.6±0.1   | -50.7±0.1                 | -152.1±0.4                | -5.4±0.1                         |
| + FXR1P RGG |          |                           |                           |                                  |
| S3F-M2   | 53.0±0.2   | -39.7±0.1                 | -121.6±0.7                | -3.4±0.1                         |
| + FXR2P RGG |          |                           |                           |                                  |

similar experimental conditions, ranges from −18 to −25 kcal/mol (45). Figure 2B illustrates a possible S3F-M2 RNA structure consistent with these results, in which two G-tetrads are stacked in a parallel manner. It has been proposed that due to their sequence, the GGA-containing mRNA targets of FMRP might adopt a more complex structure containing a hexad formed by an G quartet flanked by two adenines (46), however, our results are not consistent with this proposal. First, the amino protons of two guanines involved in the hexad formation are hydrogen-bonded to adenines at the N7 position, giving rise to sharp resonances in the 1D $^1$H NMR spectrum. We do not observe any sharp guanine amino proton resonances in the $^1$H NMR spectrum of S3F-M2 RNA. Second, the CD spectrum of S3F-M2 lacks the small positive band at 305 nm that seems to be associated with the presence of these hexads (46) (Lipay, J. and M.-R.M., unpublished data).

Thermodynamics of FMRP RGG box binding to S3F-M2 RNA

To characterize the thermodynamics of FMRP RGG box binding to S3F-M2 RNA, we have employed fluorescence spectroscopy. The S3F-M2 RNA used in this study, named S3F-M2_15AP, was labeled at the 15th position by the highly fluorescent purine analog 2-AP (highlighted in red in Figure 2B). Based on previous studies in our laboratory, we anticipated that the 2-AP at the 15th position will be a sensitive reporter of the G quartet formation (42). The CD spectral features of S3F-M2_15AP indicate that it forms the same type of G quartet structure like S3F-M2 RNA. Moreover, the UV melting profiles of S3F-M2_15AP and S3F-M2 are very similar, indicating that the 2-AP insertion did not cause major perturbations in the structure and stability of S3F-M2_15AP RNA (data not shown). In addition, we determined by EMSA that the FMRP RGG box binds identically to S3F-M2 and S3F-M2_15AP RNAs (Figure 4A).

The steady-state fluorescence of 2-AP is sensitive to stacking interactions, and we expected to observe a change when S3F-M2_15AP is folded in the presence of K$^+$ versus Li$^+$, since the structures formed by the RNA in the presence of these ions are likely very different (compare Figure 2A and B). As shown in Figure 4B, the steady-state fluorescence of S3F-M2_15AP increases 5-fold when the RNA is folded in the presence of KCl (forming a G quartet structure in which the 2-AP reporter is located in a G quartet surrounding loop) as compared to the case when it is folded in the presence of LiCl (that does not promote G quartet formation). This result establishes that the 2-AP reporter in S3F-M2_15AP RNA is sensitive to the G quartet structure formation.

Next, we measured the binding of the FMRP RGG box to S3F-M2_15AP RNA by titrating increasing concentrations of the FMRP RGG peptide, and monitoring the steady-state fluorescence change of the 2-AP reporter (Figure 4C). Figure 4C also shows the results of two negative control experiments: in the first one increasing amounts of the FMRP RGG box were titrated into a solution of Sc1-sh RNA, an RNA previously shown by native gel electrophoresis not to be bound by the FMRP RGG box (16,42). Sc1-sh RNA forms a G quartet, but lacks a stem structure, and in our experiment we used a 2-AP labeled Sc1-sh RNA in which the 2-AP reporter is located in one of its G quartet surrounding loops (42). In the second negative control experiment, we have titrated increasing amounts of the FXR2P RG cluster, a nonbinding peptide (see subsequently) to S3F-M2_15AP RNA.

A dissociation constant, $K_d$, of (0.7±0.3) nM was obtained by fitting the FMRP RGG box binding curve to Equation (2) (Materials and Methods section), indicating this peptide binds with very high affinity to S3F-M2_15AP RNA. This corresponds to a free energy of binding $\Delta G_d$ of (−12.5±0.2) kcal/mol. We measured a $K_d$ value smaller by two orders of magnitude than the value of 75 nM reported by Darnell et al. (16) for the full-length FMRP or its RGG box binding to Sc1 RNA. This discrepancy could originate from the fact that Darnell et al. measured an average value for the FMRP RGG box binding to both S3F conformers that exist even at low RNA concentrations (our native gel electrophoresis results indicate that both S3F-Ig conformers are bound by the FMRP RGG box; data not shown). In the case of S3F-M2 RNA we measured the FMRP RGG box binding to the single conformer adopted by this RNA at nanomolar concentrations.

Another RNA for which the thermodynamics of FMRP binding has been determined is Sc1 (42), a model G quartet forming RNA identified by the SELEX method (16). It is interesting to note that the FMRP RGG box binds tighter to S3F-M2 RNA by approximately one order of magnitude: $K_d$ of 0.7 nM for S3F-M2 RNA versus 7 nM for Sc1 RNA, which translates to a difference in the free energy of binding of 1.5 kcal/mol. A comparison of the sequences of these two RNA molecules reveals differences in the G quartet surrounding loops and in the junction connecting the G quartet structure with the stem, which likely account for the difference in their binding by the FMRP RGG box.

To define the forces that drive the interactions between the FMRP RGG box and S3F-M2 RNA we have determined the enthalpy and entropy of binding by measuring the association binding constant, $K_{obs} = 1/ K_d$, as a function of temperature. The thermodynamic
parameters of binding (summarized in Table 2) were determined from the slope and intercept of the Van’t Hoff plot of \( \ln (K_{\text{obs}}) \) versus \( 1/T \), which is linear when the change in enthalpy (\( \Delta H^0 \)) is independent of temperature [Equation (3) and Figure 4D]. We found that the association of FMRP RGG box with S3F-M2_15AP RNA is enthalpically driven, \( \Delta H^0 = -41.4 \pm 3.9 \) kcal/mol, with an unfavorable entropic contribution \( T\Delta S^0 = -28.9 \pm 3.8 \) kcal/mol. Contributions from hydrogen bonds, van der Waal’s or electrostatic interactions are generally associated with the favorable negative enthalpy changes, whereas a decrease in the conformational flexibility or the exposure of hydrophobic residues to the complex surface are associated with unfavorable entropy changes. The association between Sc1 RNA and FMRP RGG box has also been reported to be enthalpically driven, with an unfavorable entropic change (42). However, this finding cannot be generalized for all mRNA targets of FMRP since we determined that the FMRP RGG box binding to the microtubule-associated protein, 1B RNA (another proposed G quartet forming RNA target of FMRP) is.
enthalpically driven only at temperatures higher than 30°C (Menon, L., et al., manuscript in preparation).

To evaluate the role played by electrostatic interactions in the S3F-M2_15AP RNA:FMRP RGG box recognition, we have determined the association constant $K_{\text{obs}} = 1/K_d$ in the presence of increasing salt concentrations in the range 150–1000 mM KCl. The dependence of $K_{\text{obs}}$ on the concentration of monovalent salt concentrations is known as the salt dependence $\Delta S_\text{b}$. We found that within experimental error $K_{\text{obs}}$ does not change in the presence of increasing KCl concentrations in the range 150–1000 mM KCl (Figure 4E and Table 3), indicating that electrostatic contributions do not play a dominant role in the binding of FMRP RGG box to S3F-M2_15AP RNA.

### The FMRP RGG box stabilizes the G quartet structure of S3F-M2 RNA

To determine if the FMRP RGG box binding has any influence upon the stability of the G quartet structure of S3F-M2 RNA, we measured its melting temperature when the RNA is in complex with the RGG peptide. The UV melting curve of the S3F-M2 RNA:FMRP RGG box complex shown in Figure 3E has been corrected by subtracting the UV melting curve of the free FMRP RGG box peptide. The 38–67°C UV hypochromic transition corresponding to G quartet dissociation in the free RNA is now shifted in the range 42–78°C, corresponding to an increase of the G quartet structure $T_m$ from $\sim$52°C to $\sim$65°C (Table 1). Thus, in a 1:1 ratio, the FMRP RGG box increases the stability of S3F-M2 G quartet structure. It is very interesting to note that upon binding the RGG peptide, a second hypochromic transition appears in the range 79–91°C, indicating that the peptide promotes the formation of an alternate more stable structure of S3F-M2 RNA.

The FMRP RGG box binding has also been shown to stabilize the G quartet structure of Scl RNA (42), however to a different extent: the difference in melting temperatures of the G quartet structure in the free RNA and in the RNA in complex with the RGG peptide is $\sim$20°C for Sc1 RNA and $\sim$13°C for S3F-M2 RNA. Thus, in the case of S3F-M2, only a small fraction of the FMRP RGG box binding free energy is used to stabilize the RNA G quartet structure (Table 1 and Figure 3E). In contrast, in the case of Sc1 RNA, a significant fraction of the binding free energy is used to stabilize its G quartet structure (42).

### Interactions of S3F-M2 RNA with the FXR1P and FXR2P

We inquired next if the recognition of the G quartet structure in S3F-M2 RNA is unique to the FMRP RGG box, or if this RNA is also recognized by the two FMRP autosomal paralogs, the FXR1P and FXR2P. FXR1P has an RGG box different in sequence from that of the FMRP RGG box, whereas the FXR2P has a RG cluster, but not RGG repeats per se (Figure 5A). Using EMSA we determined that S3F-M2 RNA is bound by the FXR1P RGG box (Figure 5A, lanes 5 and 6), but not by the FXR2P RG cluster (Figure 5A, lanes 3 and 4). To get a quantitative measure of the FXR1P RGG box binding we determined the binding curve by titrating increasing amounts of the peptide into a solution of 150 nM S3F-M2_15AP RNA and monitoring the steady-state fluorescence change of the 2-AP reporter (Figure 5B, green trace). The $K_d$ of (55.0±3.8) nM, which was determined by fitting the binding curve with Equation (2), indicates that the FXR1P RGG box binds with high affinity to S3F-M2_15AP RNA.

To determine if the binding of the FMRP and FXR1P RGG boxes to S3F-M2_15AP is specific, we measured their binding either in the presence of a 10-fold excess of a non-specific RNA, Munc13 site1 (16) or in the presence of 6-fold excess of the FXR2P RG cluster. As shown in Figure 5B and C, both the FMRP and FXR1P RGG boxes bind specifically to S3F-M2_15AP RNA, since their binding curves are identical in the presence or absence of a large excess of the Munc13 site1 RNA or of the FXR2 RG cluster (the $K_d$ values are reported in the Figure 5 legend).

This result is in contrast to what has been reported for the G quartet-forming Sc1 RNA (42), since this RNA is bound specifically only by the FMRP RGG box, but not by the FXR1P RGG box. This difference in specificity of the FXR1P RGG box binding to Sc1 and S3F-M2 RNAs is likely due to structural differences in their G quartet and/or junction regions. Thus, the ability of FXR1P RGG box to bind specifically and with high affinity to the G quartet-forming mRNA targets of FMRP might be modulated by subtle differences in the particular G quartet structure adopted by the RNA. Another level of complexity becomes apparent if one considers the G quartet RNA-binding activity of the full-length FXR1P, as opposed to just of its RGG box domain. Recently, it has been reported that of three different FXR1P isoforms, only one is able to bind specifically to N19 RNA, a G quartet-forming segment of the FMRP mRNA (47). Since all these three FXR1P isoforms contain the RGG box domain, it has been suggested that the FXR1P RGG box domain is not sufficient per se to bind to the G quartet structure and

### Table 2. Thermodynamic parameters for the binding of FMRP RGG box to S3F-M2_15AP RNA

| $\Delta H^\circ$ (kcal/mol) | $\Delta S^\circ$ (cal/mol K) | $\Delta G^\circ$(kcal/mol) |
|---------------------------|---------------------------|---------------------------|
| $-41.4 \pm 3.9$          | $-96.9 \pm 12.7$         | $-12.5 \pm 0.4$           |

### Table 3. The dissociation constants for the S3F M2 RNA: FMRP RGG box complex measured at different salt concentrations

| KCl (mM) | $K_d$ (nM) |
|----------|------------|
| 150      | 0.7±0.3    |
| 250      | 0.8±0.2    |
| 400      | 0.7±0.2    |
| 1000     | 1.1±0.3    |
that a 27 amino acid stretch, present only on the FXR1P isoform that binds specifically to N19 RNA, might directly assist the FXR1P RGG box in binding the G quartet, or it might alter the structure of C-terminal portion of FXR1P, thereby allowing binding. We show here that the FXR1P RGG box domain can per se bind specifically to G quartet forming RNA, so it is more likely that the differences observed in the binding activity of the
full-length FXR1P isoforms originate from differences in the C-terminal structure that might have an impact on the accessibility of the RGG box domain.

Since the FXR1P RGG box binds with high affinity and specificity to S3F-M2_15AP RNA, we assessed also if this peptide has any effect upon the stability of the RNA G quadruplex structure. Figure 5D shows that the 38–67°C UV hypochromic transition corresponding to G quartet dissociation in the free RNA is shifted in the range 50–72°C when the RNA is complexed with the FXR1P RGG box, corresponding to a $T_m$ of 54°C (Table 1). Thus, the FXR1P RGG box binding is also slightly stabilizing the G quadruplex structure of S3F-M2 RNA. Moreover, like the FMRP RGG box, the FXR1 RGG box binding induces the formation of an alternate G quadruplex structure in S3F-M2 RNA, whose dissociation has a hypochromic transition in the range 73–93°C.

The FXR2 RG cluster has no effect upon the stability of the S3F-M2 G quadruplex structure (Table 1) and it does not promote the formation of a secondary alternate structure (Figure 5E).

At high concentrations the FMRP and FXR1P RGG boxes induce the unwinding of the G quadruplex of S3F-M2 RNA

To investigate the effect of the FMRP and FXR1P RGG boxes on the intramolecular G quartet structure of S3F-M2 RNA, we compared the CD spectra of the free RNA with those of the RNA in complex with the RGG peptides. At a 1:1 ratio of S3F-M2 RNA:RGG peptide, the intensity of the CD band at 263–238 nm was almost unchanged, for both the FMRP and FXR1P RGG boxes (Figure 6A and B). However, at higher ratios of the RNA:RGG box, both peptides induced the unstacking of the G quadruplex structure of S3F-M2, as reflected by a decrease of intensity of the 263 nm CD band as well as a shift to 265 nm. As a negative control we
have also performed the same experiment in the presence of the non-binding FXR2 RG cluster (Supplementary Figure 3) It is interesting to note that the actions of the FMRP and FXR1P RG peptides are very different: the FXR1P RG peptide starts to unwind the G quartet RNA structure at only an 1:2 RNA:peptide ratio, whereas this effect occurs for the FMRP RG box at an 1:6 ratio. At RNA:RGG peptide ratios higher than 1:4 the solutions become turbid, similar to what has been reported for the RG box of nucleolin protein interactions with MS2 phage RNA (48). To rule out the possibility that the RNA is degraded in the presence of the large excess of the RGG peptides, we have treated the solution containing the 1:10 RNA:peptide complexes with proteinase K, which degrades the RGG peptides, and re-acquired its CD spectrum (corrected for the proteinase K contribution). As shown in Figure 6C and D, the removal of the RGG peptides, allows the free RNA to refold into a G quadruplex structure, indicating that the spectral changes we observed in the presence of a large excess of the RGG peptides are not due to the RNA degradation, but due to the G quadruplex structure unstacking.

We cannot easily reconcile the findings that in an 1:1 RNA:peptide ratio, both the FMRP and FXR1P RGG boxes stabilize the S3F-M2 G quadruplex structure, whereas at higher ratios, they induce the G quadruplex unwinding. The unwinding of the S3F-M2 RNA G quartet structure by the FMRP RG G box occurs at high RNA:peptide ratios and it is accompanied by an increase in the solution turbidity; thus it is not clear if this event is biologically significant. However, the finding that the FXR1P RG box starts to unwind the G quartet structure of S3F-M2 RNA at only an 1:2 RNA:peptide ratio might be relevant considering that the FXR1P has been shown to exist in living cells as a homo-multimeric complex (12).

It is tempting to speculate that the FMRP and FXR1P might compete for, and act differently on the G quartet RNA structure; however, it remains to be seen if this G quadruplex unwinding effect is observed for other FMRP G quartet forming RNA targets besides S3F-M2 and also for the full-length FMRP and FXR1P.

**SUPPLEMENTARY DATA**

Supplementary Data are available at NAR Online.

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