Casein Kinase II Phosphorylates the Fragile X Mental Retardation Protein and Modulates Its Biological Properties

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Fragile X syndrome is caused by loss of FMR1 protein expression. FMR1 binds RNA and associates with polysomes in the cytoplasm; thus, it has been proposed to function as a regulator of gene expression at the posttranscriptional level. Posttranslational modification of FMR1 had previously been suggested to regulate its activity, but no experimental support for this model has been reported to date. Here we report that FMR1 in Drosophila melanogaster (dFMR1) is phosphorylated in vivo and that the homomer formation and the RNA-binding activities of dFMR1 are modulated by phosphorylation in vitro. Identification of a protein phosphorylating dFMR1 showed it to be Drosophila casein kinase II (dCKII). dCKII directly interacts with and phosphorylates dFMR1 in vitro. The phosphorylation site in dFMR1 was identified as Ser406, which is highly conserved among FMR1 family members from several species. Using mass spectrometry, we established that Ser406 of dFMR1 is indeed phosphorylated in vivo. Furthermore, human FMR1 (hFMR1) is also phosphorylated in vivo, and alteration of the conserved Ser500 in hFMR1 abolishes phosphorylation by CKII in vitro. These studies support the model that the biological functions of FMR1, such as regulation of gene expression, are likely regulated by its phosphorylation.

Fragile X syndrome is the most frequent cause of inheritable mental retardation and is also one of the most common single-gene disorders (14). Cognitive deficits reported for fragile X children range from mild to severe, and behavioral disturbances include social and attention deficits, autistic-like behaviors, unusual responses to sensory stimuli, hyperactivity, and sleep problems (12, 14). The gene directly responsible for these conditions, FRMS, was identified in 1991 (47). In most cases, the syndrome is caused by a trinucleotide repeat expansion in the 5′ untranslated region of the FMR1 gene (49). The expansion of the CGG repeat results in an absence of the encoded protein. It is therefore clear that the pathophysiological mechanisms leading to symptoms in fragile X syndrome can be elucidated by studying the functions of the FMR1 gene.

The fact that the FMR1 protein is a cytoplasmic protein with RNA-binding motifs (two KH domains and an RGG box) (2, 38, 39) and the fact that it associates with ribosomes (6, 9, 21, 41) suggest that this protein functions in the posttranscriptional regulation of some specific mRNAs (16). Indeed, FMR1 has recently been proposed to be a translational repressor in a cell-free system and in frog oocytes (26, 27). Interestingly, although the inhibitory effect was demonstrated not to be influenced by the features of the mRNAs used in those studies, FMR1 seemed to specifically inhibit its own translation (36). It has been known that FMR1 is found at postsynaptic sites enriched in ribosomes, where some specific mRNAs, including FMR1, are translated in response to synaptic activation (13). It has also been observed that in fragile X patients the numbers of dendritic spines are different and the morphology is also changed from that of those cells in normal individuals (18). These data suggest that FMR1 takes part in normal synaptic formation during development by regulating the expression of specific mRNAs at the postsynaptic sites.

What kind of mRNAs does FMR1 specifically bind to in vivo? It has been reported that FMR1 in vitro binds specifically to poly(G) and poly(U), but not to poly(A) and poly(C) (38), and it was estimated that FMR1 interacts with about 4% of human fetal brain mRNAs (2). These results indicate that FMR1 has an intrinsic ability to distinguish its target molecules from among all the native RNA molecules and to bind specific sequences. Recently, mRNAs to which FMR1 preferentially binds have been identified by several groups (5, 7, 36, 53). In the mRNAs identified, the nucleotide sequence through which FMR1 binds have been identified by several groups (2, 36, 53). In the mRNAs identified, the nucleotide sequence through which FMR1 interacts with about 4% of human fetal brain mRNAs (2). These results indicate that FMR1 has an intrinsic ability to distinguish its target molecules from among all the native RNA molecules and to bind specific sequences. Recently, mRNAs to which FMR1 preferentially binds have been identified by several groups (5, 7, 36, 53). In the mRNAs identified, the nucleotide sequence through which FMR1 interacts with about 4% of human fetal brain mRNAs (2). These results indicate that FMR1 has an intrinsic ability to distinguish its target molecules from among all the native RNA molecules and to bind specific sequences.

The discovery of the existence of two FMR1-related genes, FXR1 and FXR2, has revealed an additional level of complexity in the study of FMR1 functions in vertebrates (40, 52). For many human disorders, redundancy of gene function due to gene duplication in the same species makes it difficult to interpret clearly the effects of loss-of-function mutations on cellular physiology and biochemistry. Recently, a Drosophila homolog of FMR1 (dFMR1) has been identified (48). In agreement with the statement of Wan et al. (48), the recently completed Drosophila melanogaster genome sequence (1) revealed that dFMR1 is a unique, single gene with no homologs in the fly genome. Drosophila casein kinase II (dCKII) interacts with and phosphorylates dFMR1 in vitro. The phosphorylation site in dFMR1 was identified as Ser406, which is highly conserved among FMR1 family members from several species. Using mass spectrometry, we established that Ser406 of dFMR1 is indeed phosphorylated in vivo. Furthermore, human FMR1 (hFMR1) is also phosphorylated in vivo, and alteration of the conserved Ser500 in hFMR1 abolishes phosphorylation by CKII in vitro. These studies support the model that the biological functions of FMR1, such as regulation of gene expression, are likely regulated by its phosphorylation.

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genome. The *Drosophila* and vertebrate FMR1 proteins share a number of topographical landmarks, including RNA-binding motifs (two KH domains and an RGG box), and show similar biological properties such as a binding preference in vitro for certain RNA sequences and ribosomal association in vivo (19, 48; M. C. Siomi and H. Siomi, unpublished data). Moreover, a number of genes related to complex behaviors such as learning and memory are conserved between flies and humans (30, 34). Therefore, *Drosophila* could be used to define the molecular pathways leading to human neurological diseases such as fragile X syndrome and to identify the genes involved. In fact, several groups have individually produced fly models of fragile X syndrome and have shown that dFMR1 is necessary for normal synaptic activities (53) and for rhythmic circadian output regulating locomotor activities (8, 17, 31).

For a number of proteins, posttranslational modifications have been well studied and are known to be critical for regulating physiological and biochemical functions. So far, however, very little is known about the posttranslational modifications of FMR1 protein and their effects at the functional level. Here we provide the first evidence that FMR1 protein, both in flies and in humans, is phosphorylated in vivo. By purification from the cytoplasmic lysate of a *Drosophila* culture cell line, Schneider 2 (S2), and by mass spectrometric analysis, we identified *Drosophila* casein kinase II (dCKII) (35) as a protein phosphorylating dFMR1. Indeed, recombinant dCKII was able to interact with and phosphorylate dFMR1 in vitro. Systematic mutation analyses revealed that the phosphorylation site in dFMR1 is Ser406, located just upstream of the RGG box. This amino acid residue is conserved among FMR1/FXR family members across many species, and mutation to Ala of the Ser residues in dFMR1 and hFMR1 proteins abolishes the phosphorylation by CKII in vitro. Finally, we demonstrate that the efficiency of dFMR1 in forming homomers and binding RNA is profoundly altered by phosphorylation in vitro. Taken together, our data strongly support the hypothesis that phosphorylation by CKII could play an important role in regulating physiological functions of FMR1, such as regulation of gene expression for the normal formation of synapses during development.

**MATERIALS AND METHODS**

**Cell culture.** S2 cells were grown in Schneider’s *Drosophila* medium (Gibco BRL) supplemented with 10% fetal bovine serum (Sigma), 1% l-glutamine 50 U of penicillin/ml, and 50 μg of streptomycin/ml (Gibco BRL) at 26°C. HeLa cells were grown in Dulbecco’s modified Eagle medium (Sigma) supplemented with 10% fetal bovine serum, 1% l-glutamine, 50 U of penicillin/ml, and 50 μg of streptomycin/ml (Gibco BRL) at 37°C. Subcellular fractionation, CIP treatment, and Western blot analysis. A cytoplasmic lysate of S2 cells was prepared in a lysis buffer and proteins were separated on an SDS-polyacrylamide gel after heat denaturation. Western blotting was carried out essentially as described previously (41) by using an anti-dFMR1 monoclonal antibody (48). Protein bands were visualized with an ECL Western blotting detection kit (Amersham Bioscience).

In vitro phosphorylation assay. Either full-length or truncated dFMR1 cDNA (48) was subcloned into the pGEX-5X (Amersham Bioscience) bacterial expression vector to produce glutathione S-transferase (GST) fusion proteins for assay. GST fusion proteins and GST itself were produced and purified as described by the manufacturer. For Fig. 6C, full-length hFMR1 cDNA (38, 47) was used and GST-hFMR1 was produced as well. All the point mutants were constructed by performing PCR with primers specifically designed to alter Ser to Ala. The final products were inserted into the pGEX-5X expression vector, and GST fusion proteins were produced as well. The assay was carried out in P buffer containing 20 mM Tris-HCl (pH 8.0), 200 mM NaCl, 2 mM ATP, and 1 mM dithiothreitol (DTT). In the presence of [γ-32P]ATP, a cytoplasmic lysate of S2 cells after centrifugation at 100,000 × g for 30 min was used for the experiments for which results are shown in Fig. 1B and 2. Purified mammalian CKII (Fig. 6D) was a kind gift from U. Kikkawa (22). After the reaction, proteins were separated on SDS-polyacrylamide gels and autoradiography was carried out to visualize phosphorylated proteins.

**Mass spectrometry for identification of in vivo phosphorylation sites.** Coomassie-stained dFMR1 bands were digested with trypsin (37), and the digest solution was desalted with a ZipTip C18 tip (Millipore). The peptide mixture eluted with 70% acetonitrile and 1% formic acid was analyzed by using a Q-ToF mass spectrometer (Micromass, Manchester, United Kingdom) equipped with a nanoelectrospray ionization source, and the phosphorylation sequence was obtained by tandem mass spectrometry (MS/MS). MS/MS data were analyzed by the Mascot Search program (Matrix Science, London, United Kingdom). MS/MS data were analyzed by the PepSeq program in the BioLyx software package (Micromass) for determination of peptide sequences.

**Purification and identification of dCKII.** A cytoplasmic lysate of S2 cells prepared as described above was resolved on a linear sucrose gradient (5 to 30%). The gradient was centrifuged at 4°C in a Beckman MLA-130 rotor at 40,000 rpm for 90 min. Following centrifugation, fractions were collected and assayed for kinase activity toward GST-dFC285. The active fractions from the gradient were pooled, applied to a gel filtration PC-10 column (Amersham Bioscience), and eluted with 100 mM NaCl prior to being applied to a UnoQ ion exchange column (Bio-Rad). The column was washed extensively with buffer A, and a 100 mM to 1 M NaCl gradient of buffer A was applied to the column. Fractions (0.5 ml) were collected and assayed for kinase activity. The phosphorylation-active fractions were pooled and diluted with 10 mM Tris-HCl (pH 7.5) to a final 100 mM NaCl concentration prior to being applied to a UnoQ ion exchange column (Bio-Rad). The column was washed extensively with buffer A, and a 100 mM to 1 M NaCl gradient of buffer A was applied to the column. Fractions (0.5 ml) were collected and assayed for activity. The active fractions were pooled and diluted with 10 mM Tris-HCl (pH 7.5) to a final 100 mM NaCl concentration prior to being applied to a heparin column. The column was washed and extensively with buffer A, and proteins were eluted stepwise with 10 mM Tris-HCl (pH 7.5) containing NaCl at the concentrations indicated in Fig. 4C. Each fraction was assayed for kinase activity and for protein content by silver staining after being resolved on an SDS-polyacrylamide gel. Both protein bands (as indicated in Fig. 4C, lane 1 M NaCl) were sent to the Genomic Research Center, Shimadzu Biotech, and mass spectrometric analysis was carried out to identify them.

**Production of recombinant alpha and beta subunits of dCKII.** In order to obtain the cDNAs encoding the alpha and beta subunits of dCKII, poly(A) + RNA was purified from S2 cells and reverse transcription-PCR (RT-PCR) was carried out using primers specifically designed for each clone (35). The cDNAs of the alpha and beta subunits of dCKII were cloned into pGEX-5X (Amersham Bioscience) bacterial expression vector to produce GST-dCKII alpha and His-dCKII beta, respectively. Both fusion proteins were produced and purified as described by the manufacturers.

**Phosphorylation of hFMR1 in vivo.** HeLa cells were grown as described previously (42). HeLa cells transiently expressing myc-tagged hFMR1 were labeled with [32P]orthophosphate (1 mCi/ml) (Amersham Bioscience) in phosphate-free medium supplemented with 10% fetal calf serum. After incubation for 3 h, the cells were lysed in phosphate-buffered saline buffer containing 1% Empigen, 1 mM EDTA, 0.1 mM DTT, 2 μg of leupeptin/ml, 2 μg of pepstatin/ml, and 0.5% aprotinin. After brief sonication, the lysate was centrifuged at 16,000 × g for 5 min and the supernatant was immunoprecipitated with either an anti-myc antibody (9E10) or a nonimmune control antibody (HP60). After extensive washing, the immunoprecipitated samples were resolved on an SDS-polyacrylamide gel and proteins labeled with [32P] were visualized by using BAS-2500 (Fujifilm).
**RESULTS**

**dFMR1 protein is phosphorylated in vivo and in vitro.** In the course of experiments to elucidate the function of dFMR1, we observed that dFMR1 in S2 cells migrates as several bands when probed with an anti-dFMR1 antibody. This suggested that dFMR1 is posttranslationally modified in vivo and that one such modification is likely phosphorylation. In order to test this hypothesis, the cytoplasmic lysate from S2 cells was treated with CIP, an enzyme known to dephosphorylate both nucleic acids and proteins, and Western blot analysis was performed. In the lysate without CIP, dFMR1 was detected as a strong doublet (~75 kDa) and a weak upper band (~80 kDa) on an SDS–7.5% polyacrylamide gel (Fig. 1A, CIP lane in S2 panel). After CIP treatment, however, the doublet became a discrete single band, apparently comigrating with the lower band of the doublet in the untreated lanes (Fig. 1A, +CIP lane in S2 panel), demonstrating that dFMR1 is phosphorylated in vivo and that the phosphorylation state of dFMR1 is normally heterogeneous in the cells. After CIP treatment, the weak upper band also migrated slightly faster, but did not disappear, indicating that this weak band also represents dFMR1 in a phosphorylated form but that its larger size suggests other modifications and/or alternative splicing for dFMR1. The same phenomenon was observed with fly ovaries (Fig. 1A, right panel), demonstrating that phosphorylation of dFMR1 is not restricted to cultured cells but also occurs in animals. We next investigated if dFMR1 could be phosphorylated in vitro. Reconstituted full-length dFMR1 tagged with GST was purified from *Escherichia coli* and mixed with a cytoplasmic lysate prepared from S2 cells in the presence of Mg^{2+} and [γ-^32P]ATP. The phosphorylated proteins were visualized by autoradiography. The full-length GST fusion protein is clearly phosphorylated, whereas GST alone and the truncated by-product of GST-dFMR1 (~65 kDa in the CBB panel) (see the text) are not, suggesting that a protein with a dFMR1-phosphorylating activity exists in the cytoplasm of S2 cells and that the activity is sequence specific. It is noted that EDTA abolishes the activity. The left panel shows the protein substrates stained with Commassie brilliant blue (CBB), and the positions of molecular mass markers are indicated on the left of the gel.

**FIG. 1.** dFMR1 is phosphorylated in vivo and in vitro. (A) dFMR1 proteins in cytoplasmic lysates of S2 cells and of fly ovaries before and after CIP treatment were visualized by Western blotting using an anti-dFMR1 antibody (50). Migration changes of dFMR1 proteins are observed before and after CIP treatment on SDS–7.5% polyacrylamide gels, indicating that dFMR1 proteins both in cultured cells and in animals are phosphorylated in vivo. (B) After production of GST-dFMR1, an in vitro phosphorylation assay was carried out using the cytoplasmic lysate of S2 cells in the presence of [γ-^32P]ATP, and the phosphorylated proteins were visualized by autoradiography. The full-length GST fusion protein is clearly phosphorylated, whereas GST alone and the truncated by-product of GST-dFMR1 (~65 kDa in the CBB panel) (see the text) are not, suggesting that a protein with a dFMR1-phosphorylating activity exists in the cytoplasm of S2 cells and that the activity is sequence specific. It is noted that EDTA abolishes the activity. The left panel shows the protein substrates stained with Commassie brilliant blue (CBB), and the positions of molecular mass markers are indicated on the left of the gel.

**Protein-protein interaction assays.** A fusion construct encoding dFMR1 and a tandem affinity purification (TAP) tag (33) under the control of the *Drosophila* metallothionein promoter in the pRHA3-3 vector (25) (a kind gift of F. Lafont) was transfected into S2 cells. Expression of the dFMR1-TAP fusion protein was induced by adding copper ions to the medium, and the fusion protein and associated components were recovered from the cytoplasmic lysates of the S2 cells by affinity selection on an immunoglobulin G (IgG) Sepharose 6 Fast Flow column (Amersham Bioscience). After extensive washing with the binding buffer (20 mM Tris-HCl [pH 7.5], 0.5% Triton X-100, 150 mM NaCl, 2 μg of leupeptin/ml, 2 μg of pepstatin/ml, and 0.5% aprotinin), the bound materials were separated on an SDS-acrylamide gel and analyzed by Western blotting using an anti-dFMR1 antibody. For the GST pulldown assay, GST-dCKII alpha, GST-dFMR1, and GST alone were bound to 30 μl of glutathione-Sepharose 4B resin (Amersham Bioscience) in 1 ml of binding buffer (20 mM Tris-HCl [pH 7.5], 0.5% Triton X-100, and 150 mM NaCl). Following incubation at 4°C for 1 h, the resin was pelleted and washed with the binding buffer. Either His-dCKII beta or His-dFMR1 was then added to the resins and bound in the absence or in the presence of MgCl₂ at 10 mM and ATP at 1 mM at 4°C for 2 h. After extensive washing with the binding buffer, the bound proteins were analyzed by Western blotting to visualize His-dCKII beta and His-dFMR1 by using an anti-His antibody (Santa Cruz). A cytoplasmic lysate of S2 cells prepared as described above was used for the in vivo interaction of dCKII with dFMR1. The antibodies against rat CKII cross-reacting to dCKII (a kind gift from U. Kikkawa) were bound on GammaBind G Sepharose resin (Amersham Bioscience) and incubated with the S2 cytoplasmic lysate at 4°C for 1 h. After extensive washing of the resins, the bound proteins were analyzed by Western blotting using an anti-dFMR1 antibody. For Fig. 7, in vitro-phosphorylated GST-dFMR1 and His-dFMR1 were obtained as described above with a minor modification, namely, that [γ-^32P]ATP was replaced with additional 2 mM ATP. In addition, the 1 mM DTT in the buffer was replaced with 0.1 mM tris(2-carboxyethylphosphine)hydrochloride for HIS-dFMR1 phosphorylation. The enzymatic source used in the experiments was the recombinant dCKII shown in Fig. 5A. Phosphorylated GST-dFMR1 was bound to 30 μl of glutathione-Sepharose 4B resin in 1 ml of binding buffer (20 mM Tris-HCl [pH 7.5], 0.5% Triton X-100, and NaCl at the concentration indicated). Following incubation at 4°C for 1 h, the resin was pelleted and washed with the binding buffer. His-dFMR1, either in a phosphorylated state or in a unphosphorylated state, was then added to the resin with 1 ml of the binding buffer and incubated at 4°C for 2 h. After extensive washing with the binding buffer, the bound proteins were detected on Western blots by using an anti-His antibody.

**RNA-binding assay.** Binding of His-dFMR1 in phosphorylated and unphosphorylated states was carried out by using 30 μl of either AGP[0.15±0.35] (Amersham Bioscience) or poly(G) immobilized on polyacrylamidoagarose (Sigma) in 1 ml of binding buffer (20 mM Tris-HCl [pH 7.5], 0.5% Triton X-100, and NaCl at the concentration indicated). Following incubation at 4°C for 1 h, the resin was pelleted and washed with the binding buffer. Bound proteins were then eluted by addition of SDS-PAGE sample buffer to the beads and heat denaturation. Western blotting was performed to visualize His-dFMR1 by using an anti-His antibody.
blotting using an anti-GST antibody (data not shown), was not phosphorylated under the same circumstances. These results suggest that the phosphorylation is sequence-specific and that the phosphorylation site is likely located towards the C terminus of dFMR1.

The phosphorylation site in dFMR1 is Ser406 preceding the RGG box. To determine the phosphorylation site in dFMR1, we decided to perform systematic mutation analyses. First, dFMR1 was divided into four individual fragments, each of which was fused to GST. After these proteins were purified from E. coli, an in vitro phosphorylation assay was carried out. We found that only one substrate, termed GST-dFC285, which contains 285 amino acid residues at the C terminus of dFMR1 (amino acids 397 to the end), was phosphorylated in this assay (Fig. 2A). Since GST-dFC277 was phosphorylated efficiently, whereas GST-dFC240 was not, indicating that the region covering amino acid residues 405 to 441 in dFMR1 contains the phosphorylation site. (C) In the region identified in panel B, each of four Ser residues was mutagenized to Ala in the context of GST-dFC285, and mutants were assayed to determine the phosphorylation site in dFMR1. The phosphorylation level of the S406A mutant was detectable but was markedly reduced compared to those of the others, demonstrating that Ser406 is the major phosphorylation site in dFMR1. (D) The amino acid sequence of the region containing the phosphorylation site, Ser406 (circled and shaded), and the RGG box of dFMR1 (48) is shown. Five RGG sequences found in the RGG box are indicated by boxes.

FIG. 2. Determination of the phosphorylation site in dFMR1 by systematic mutation analyses. (A) The four parts of dFMR1 (shown in the diagram at the bottom) were fused individually to GST and assayed for in vitro phosphorylation. It may be observed that only GST-dFC285, containing 285 amino acids from the end of dFMR1, becomes phosphorylated, indicating that the phosphorylated region includes the RGG box. In the schematic drawings, the locations of the first and the second KH domains, the Leu-rich region, and the RGG box are indicated as KH1, KH2, Leu, and RGG, respectively. (B) To narrow down the phosphorylation region of dFMR1, the dFC285 fragment was further delineated from its N terminus (shown in the diagram at the bottom), fused to GST, and assayed. GST-dFC277 was phosphorylated efficiently, whereas GST-dFC240 was not, indicating that the region covering amino acid residues 405 to 441 in dFMR1 contains the phosphorylation site. (C) In the region identified in panel B, each of four Ser residues was mutagenized to Ala in the context of GST-dFC285, and mutants were assayed to determine the phosphorylation site in dFMR1. The phosphorylation level of the S406A mutant was detectable but was markedly reduced compared to those of the others, demonstrating that Ser406 is the major phosphorylation site in dFMR1. (D) The amino acid sequence of the region containing the phosphorylation site, Ser406 (circled and shaded), and the RGG box of dFMR1 (48) is shown. Five RGG sequences found in the RGG box are indicated by boxes.

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To purify a protein with dFMR1-phosphorylating activity, a cytoplasmic lysate of S2 cells was first fractionated on a linear density sucrose gradient and assayed for phosphorylation activity toward GST-dFC285 (Fig. 4A). Fractions showing the highest activity (F3, F4, and F5 in Fig. 4A) were pooled, and further purification was performed using various chromatog-
raphy columns (Fig. 4B). Individual fractions eluted from the last column (a heparin column) were assayed and separated on an SDS–7.5% PAGE gel, followed by silver staining to visualize the protein pattern (Fig. 4C). The fraction containing kinase activity eluted at 1 M NaCl and contained two predominant proteins with apparent molecular masses of 35 and 28 kDa (Fig. 4C). Interestingly, these two proteins were observed to exist in an approximately equal molar ratio, suggesting that they might comprise the enzyme that phosphorylates dFMR1. The 35- and 28-kDa proteins in the fraction at 1 M NaCl were isolated and subjected to MS analysis. We found that they were identical to the alpha and beta subunits of dCKII (35), respectively (Fig. 4C).

Confirmation of dCKII as the protein phosphorylating dFMR1 in vitro. To confirm that dCKII is indeed the protein phosphorylating dFMR1, we obtained by RT-PCR the cDNA clones encoding both the alpha and beta subunits of dCKII (35) and inserted them into vectors to produce the GST-dCKII alpha and His-dCKII beta proteins (Fig. 5A). We first examined whether these two recombinant proteins can interact with each other, as might be expected (see Discussion) (11, 35). We found that GST-dCKII alpha interacted with His-dCKII beta and that the affinity of the two proteins was stronger in the presence of Mg2+ and ATP (Fig. 5B). Interaction between dFMR1 and each subunit of dCKII was also examined. Each subunit interacted with dFMR1 independently of the other, and the affinity of the alpha subunit for dFMR1 was again increased by addition of Mg2+ and ATP to the buffer (Fig. 5B). We also examined if dCKII interacts with dFMR1 in vivo. Immunoprecipitation from an S2 cytoplasmic lysate was first performed with anti-rat CKII (rCKII) antibodies cross-reacting with dCKII (data not shown), and then Western blotting was performed using an anti-dFMR1 antibody. A band corresponding to dFMR1 was observed in the lane with anti-rCKII antibodies, but not with the nonimmune control antibody (SP2/0), demonstrating that dFMR1 indeed interacts with dCKII in vivo (Fig. 5C). We next examined the phosphorylation activity of recombinant dCKII toward GST-dFC285. As expected, the
alpha subunit efficiently phosphorylated the substrate (see Discussion), whereas the beta subunit did not (Fig. 5D). When GST-dFMR1 was substituted for GST-dFC285, the same data were obtained (data not shown). Interestingly, addition of the beta subunit to the reaction mixture together with the alpha subunit did not appear to alter the kinase activity of the alpha subunit (Fig. 5D). The contribution of the beta subunit to dFMR1 phosphorylation remains unknown.

The Ser406 phosphorylation site in dFMR1 is highly conserved among FMR1/FXR family members across species. Determining whether FMR1/FXR family members from other species are also phosphorylated by CKII was of great interest. First, we inspected the amino acid sequences of those proteins (40, 47, 48, 52) and found that the region surrounding Ser406 in dFMR1 is highly conserved (Fig. 6A). To analyze whether hFMR1 is phosphorylated in cultured cells, in vivo $^{32}$P labeling of HeLa cells expressing myc-tagged hFMR1 was performed, and myc-tagged hFMR1 was immunoprecipitated with antmyc antibodies. Autoradiography of the immunoprecipitates separated by SDS-PAGE revealed that predominantly one protein band was labeled with $^{32}$P and that it migrated at the expected size of myc-tagged hFMR1 (∼80 kDa) (Fig. 6B); this suggests that hFMR1 is a phosphoprotein in vivo. To see if the residue (Ser500) corresponding to Ser406 of dFMR1 (Fig. 6A) is the phosphorylation site in hFMR1, the residue was altered to Ala in the context of full-length hFMR1 and an in vitro phosphorylation assay was carried out using recombinant dCKII. As shown in Fig. 6C, a GST-hFMR1 mutant (S500A) was not phosphorylated, whereas wild-type hFMR1 was.

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Ser500 in hFMR1 was, therefore, determined to be the phosphorylation site of hFMR1, and there seem to be no other phosphorylation sites in this case. We also tried purified rCKII in the assay and found that the mammalian enzyme phosphorylated wild-type hFMR1, but not the S500A mutant (Fig. 6C). Moreover, we also observed that the mammalian CKII phosphorylated dFMR1 (data not shown).

Phosphorylation of dFMR1 modulates the efficiency of forming homomers and of binding RNA in vitro. It has been reported that dFMR1 forms homomers in vitro (48), just as mammalian FMR1/FXR proteins do (52). To address the question of whether dFMR1 also forms homomers in vivo, we used the TAP tag method (33). The TAP-tagged dFMR1 protein was expressed in S2 cells, and the fusion protein and associated components were recovered from cell extracts by affinity selection on an IgG matrix and were then analyzed by SDS-PAGE and Western blotting. As shown in Fig. 7A, not only the TAP-tagged dFMR1 but also endogenous dFMR1 Ser500 in dFMR1 was, therefore, determined to be the phosphorylation site of dFMR1, and there seem to be no other phosphorylation sites in this case. We also tried purified rCKII in the assay and found that the mammalian enzyme phosphorylated wild-type hFMR1, but not the S500A mutant (Fig. 6C). Moreover, we also observed that the mammalian CKII phosphorylated dFMR1 (data not shown).

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was found in the IgG-bound material, indicating that dFMR1 forms homomers in vivo. We then tried to ascertain if the phosphorylation affects the oligomerization of dFMR1. For this experiment, GST- and His-tagged dFMR1s were purified from E. coli were phosphorylated by recombinant dCKII. That most of the population of GST- and His-dFMR1 proteins were phosphorylated was confirmed by observing the mobility change on Western blots (see Fig. 7B, input lanes, for His-dFMR1). GST-dFMR1 in a phosphorylated state (data not shown) was first bound to glutathione-Sepharose resins, and after extensive washing, a GST pulldown assay was carried out at 150 and 450 mM NaCl. By performing Western blotting using an anti-His antibody, it was clearly observed that the phosphorylated His-dFMR1 bound much more efficiently than the unphosphorylated His-dFMR1 (Fig. 7B), demonstrating that the oligomerization of dFMR1 is upregulated by dCKII phosphorylation. We next investigated whether the RNA-binding activity of dFMR1 was also influenced by dCKII modification. When the assay was carried out with poly(U) beads, phosphorylated dFMR1 showed a higher affinity for them than did unphosphorylated dFMR1 (Fig. 7C). Considering the result shown in Fig. 7B, the phosphorylated dFMR1 likely forms homomers more efficiently than the unphosphorylated dFMR1 at the NaCl concentrations used in this experiment. Thus, it is speculated that phosphorylation followed by oligomerization confers a higher RNA-binding activity on dFMR1 protein. We also examined dFMR1 binding to poly(G) beads. Recently, it was reported that dFMR1 preferentially binds G quartets in RNA (7, 36). The G quartet is a secondary structure formed in G-rich sequences and is known to be stabilized by certain cations such as K\(^+\). Poly(G) is known to form G quartets (23). Thus, this particular experiment was performed in a buffer containing KCl. In this assay, phosphorylated and unphosphorylated dFMR1s showed approximately the same affinity for poly(G) at any given concentration of KCl (Fig. 7D). Interestingly, it was clearly observed that the unphosphorylated form of dFMR1 coexisting with the phosphorylated form in the input (see the right input lane in Fig. 7D) was not detected in the bound fraction but was observed in the unbound fraction (Fig. 7D). When the unphosphorylated form was allowed to bind to the beads first and then the phosphorylated form was added, both forms were detected in the bound fraction (data not shown), suggesting that phosphorylated dFMR1 does not have the ability to displace unphosphorylated dFMR1 from poly(G) beads. It is known that G quartets are stabilized more by K\(^+\) than by Li\(^+\) and that hFMR1 shows a higher affinity for a G quartet formed in the presence of K\(^+\) (36). Thus, the poly(G)-binding activities of phosphorylated and unphosphorylated dFMR1s were compared in the presence of K\(^+\) and Li\(^+\). Interestingly, the unphosphorylated form of dFMR1 showed no significant difference between K\(^+\) and Li\(^+\), whereas the phosphorylated form showed much higher affinity for poly(G) in a buffer containing K\(^+\) (Fig. 7D), suggesting that in binding to a G quartet, the phosphorylated dFMR1 behaves as does hFMR1.

**DISCUSSION**

In this study we have shown that dFMR1 protein is phosphorylated in vivo, both in cultured cells and in animals, and that bacterially produced GST-dFMR1 is phosphorylated with a cytoplasmic lysate from S2 cells in vitro. We have also shown that the protein with dFMR1-phosphorylating activity purified from the lysate was dCKII (35) and that recombinant dCKII does indeed phosphorylate dFMR1 in vitro.

CKII is known to phosphorylate both nuclear and non-nuclear proteins (46, 50) and is highly conserved from yeasts to humans. The high degree of conservation suggests that CKII has an essential role in organisms. Indeed, genetic analyses in yeast have demonstrated that the enzyme is essential for viability (32, 44). We found in this study that at least in vitro, both dCKII and mammalian CKII phosphorylate both dFMR1 and hFMR1 at the sites we identified by performing mutation analyses. The cross-species activity implies that the phosphorylation of FMR1 proteins is highly conserved through evolution and is important, perhaps even required, for function of the FMR1/FXR proteins in divergent organisms.

CKII is composed of alpha and beta subunits that are associated as a heterotetramer (α2β2) (22). The alpha subunit contains a protein kinase domain that is homologous to the catalytic domain of other protein kinases (15). The beta subunit has been proposed to play a complex role in regulating the basal activity of the alpha subunit (4). Recently, the physical interaction of the CKII enzyme and its substrates was shown to be mediated by the beta, but not the alpha, subunit (10). In the present study, we observed that both the beta and the alpha subunits are able to interact directly with dFMR1 in vitro and that the affinity of the alpha subunit for dFMR1 is increased in the presence of Mg\(^{2+}\) and ATP. We also found that the alpha subunit is necessary and sufficient for efficient phosphorylation of dFMR1 in vitro; that is, the beta subunit seemed not to influence that activity. The role of the beta subunit of CKII in the phosphorylation of FMR1 remains undefined.

CKII phosphorylates Ser/Thr residues of many protein substrates in vivo, and the consensus sequence is known to be (Ser/Thr)-(Asp/Glu)-X-(Asp/Glu) (24). We determined that the peptide sequence in dFMR1 containing the phosphorylation site (Ser406) was Ser-Asp-Ile-Glu, which perfectly fits the consensus. The phosphorylation site is highly conserved within FMR1/FXR family members from flies to humans. The sequences of these other family members also match the CKII consensus. We have mentioned that there might be minor phosphorylation sites in dFMR1, since the S406A mutation did not result in a total loss of phosphorylation. However, there are no other sequences in dFMR1 that perfectly fit the consensus sequence for CKII phosphorylation. In the case of hFMR1, we could find two other sites that fit the consensus (namely, Thr455 and Thr518). But neither of these is likely to be a site of modification by CKII, at least in vitro, since the GST-hFMR1 mutant (S500A) that contains intact Thr455 and Thr518 residues is not phosphorylated by either dCKII or mammalian CKII. It remains possible, however, that other, unidentified kinases or even CKII itself contributes phosphorylation of the other sites of hFMR1 in vivo. It is known that the negative-charge requirement within the CKII consensus can be supplied by phosphorylated residues (46). If so, once Ser500 in hFMR1 is phosphorylated, Ser497 and consequently Ser494 become appropriate targets for CKII. This sequential phosphorylation may have occurred even in vitro (see Fig. 6C), but at this time we do not have data to support this scenario.
Although it is understood that the phosphorylation activity of CKII appears to be constitutive, there are studies available showing the regulation of CKII phosphorylation activity in vivo. Recently, it was demonstrated that CKII phosphorylates the human p53 protein in response to UV irradiation and that the chromatin transcriptional elongation factor, which forms a complex with CKII, alters the specificity of CKII to selectively phosphorylate p53 over other substrates including casein (20). In the case of Dishevelled (Dsh), another example of a substrate for dCKII, phosphorylation is regulated in response to Dsh and Dfz2 (a Wingless receptor) expression in the signal transduction pathway of the Wingless protein (50). When we used a linear density sucrose gradient as a first means to purify dCKII, a relatively high activity was obtained in fractions E3 to F5 (see Fig. 4A). This indicated that it was likely that dCKII associates with other molecules, constituting a relatively heavy complex. However, the complex may not be very stable and probably is dissociated through a series of purification procedures, since no other proteins eluted with dCKII from the last heparin column (see Fig. 4C). At this point, we have no data identifying molecules that associate with dCKII. It is possible, as in the case of p53, for example, that the dCKII-interacting proteins and/or nucleic acids regulate the dCKII activity in the phosphorylation of dFMR1. We are currently attempting to isolate the dCKII interactors and understand their involvement in the phosphorylation of dFMR1.

It has been reported that the DNA-binding activity of some nuclear proteins is regulated by CKII phosphorylation (3, 28, 29, 45). In this study we have shown that the RNA-binding activity of dFMR1 to poly(U) is upregulated by phosphorylation by dCKII in vitro. This property may be a result of the favorable homotypic interactions of dFMR1 caused by phosphorylation (see Fig. 7B). We speculate that phosphorylation of dFMR1 induces a conformational change that results in a stronger association with itself and perhaps also with poly(U). Stable oligomerization of dFMR1 simply means that the KH domains and the RGG boxes are also multiplied and are brought close enough together to cooperate, which we think confers the stronger ability to bind poly(U) beads. Surprisingly, different results were obtained when we used poly(G) beads. In such experiments, dFMR1, whether phosphorylated or unphosphorylated, showed approximately the same ability to bind poly(G). On the other hand, when both proteins were added to the beads together, only phosphorylated dFMR1 was detected in the bound fraction. In different experiments, however, where unphosphorylated dFMR1 was allowed to bind poly(G) first, and phosphorylated dFMR1 was added later, both proteins were observed in the bound fraction. One explanation for these observations is that the on-rate and off-rate of phosphorylated or unphosphorylated dFMR1 for poly(G) binding may differ from each other. Data to support this speculation are currently not available, and further detailed studies will be necessary.

These studies constitute the first demonstration that CKII is phosphorylatable in vivo and that the phosphorylation alters its biochemical properties, such as its interactions with itself and with RNA in vitro. Although the in vivo relevance of these observations is currently unknown, our results should be the starting point for a better understanding of the effects of phosphorylation on the function of FMR1.

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