NEUROSCIENCE

Augmented noncanonical BMP type II receptor signaling mediates the synaptic abnormality of fragile X syndrome

Risa Kashima,¹ Sougata Roy,¹ Manuel Ascano,² Veronica Martinez-Cerdeno,^{3,4} Jeanelle Ariza-Torres,³ Sunghwan Kim,¹ Justin Louie,¹ Yao Lu,¹ Patricio Leyton,⁵ Kenneth D. Bloch,⁵* Thomas B. Kornberg,¹ Paul J. Hagerman,⁶ Randi Hagerman,⁴ Giorgio Lagna,¹ Akiko Hata^{1†}

Epigenetic silencing of *fragile X mental retardation 1 (FMR1*) causes fragile X syndrome (FXS), a common inherited form of intellectual disability and autism. FXS correlates with abnormal synapse and dendritic spine development, but the molecular link between the absence of the *FMR1* product FMRP, an RNA binding protein, and the neuropathology is unclear. We found that the messenger RNA encoding bone morphogenetic protein type II receptor (BMPR2) is a target of FMRP. Depletion of FMRP increased BMPR2 abundance, especially that of the full-length isoform that bound and activated LIM domain kinase 1 (LIMK1), a component of the noncanonical BMP signal transduction pathway that stimulates actin reorganization to promote neurite outgrowth and synapse formation. Heterozygosity for *BMPR2* rescued the morphological abnormalities in neurons both in *Drosophila* and in mouse models of FXS, as did the postnatal pharmacological inhibition of LIMK1 activity. Compared with postmortem prefrontal cortex tissue from healthy subjects, the amount of full-length BMPR2 and of a marker of LIMK1 activity was increased in this brain region from FXS patients. These findings suggest that increased BMPR2 signal transduction is linked to FXS and that the BMPR2-LIMK1 pathway is a putative therapeutic target in patients with FXS and possibly other forms of autism.

INTRODUCTION

Fragile X syndrome (FXS) is the most common heritable form of intellectual disability and autism spectrum disorder (ASD) (1-3). FXS is caused by an expansion of trinucleotide (CGG) repeats in the fragile X mental retardation 1 (FMR1) gene that leads to transcriptional silencing and loss of the FMR1 protein (FMRP) (3-7). FMRP is an RNA binding protein that contains three RNA binding domains: two K homology (KH) domains (KH1 and KH2) and one arginine-glycine-glycinerich domain (8-11). FMRP binds mRNAs directly, modulating transport and stability and negatively regulating translation (12-15). A missense mutation in the KH2 domain (I304N) that abolishes RNA binding activity has been identified in a patient with FXS (16, 17). This suggests that the pathogenesis of FXS might be caused by the absence of the RNA binding function of FMRP. Both postmortem brain tissues from FXS patients (18, 19) and from young adult mice lacking FMR1 (20, 21) exhibit widespread defects in synaptic plasticity and development, including an increased number of long and thin dendritic spines instead of the mature and strong mushroom-shaped spines. Transcripts of 40 genes have been identified as FMRP targets (22-25). Most encode proteins that are implicated in synaptic growth, plasticity, cell adhesion, or cytoskeletal structure

*Deceased

+Corresponding author. Email: akiko.hata@ucsf.edu

and remodeling (15, 26). However, it is not known whether an increase in the abundance of these targets is responsible for the FXS phenotype or whether there are additional FMRP targets that contribute to the disease.

Bone morphogenetic proteins (BMPs) are members of the transforming growth factor- β (TGF β) superfamily of secreted proteins (27, 28). Their receptors consist of type I and type II subunits, which are both serine/threonine kinases (28). BMPR2 is a type II subunit that, in addition to the kinase domain (KD) in its C-terminal domain (CTD) that is typical of type II subunits, has a unique, evolutionarily conserved ~500-amino acid region. BMP binding leads to activation of the type I receptor and activation of the canonical pathway through phosphorylation of SMAD1, SMAD5, and SMAD8 (28); it also leads to a "noncanonical" signal that is mediated by proteins, such as LIM (Lin-11 Isl-1 Mec-3) domain kinase 1 (LIMK1), that interacts with the CTD (29). Once activated by BMPR2, LIMK1 phosphorylates and inhibits cofilin to promote neurite outgrowth and dendritogenesis (29). In cortical neurons, activation of the noncanonical pathway also involves the p21-activated kinase 1 (PAK1), which facilitates the recruitment of the type I BMP receptor to BMPR2 and the subsequent activation of LIMK1 (30).

The BMPR2-CTD-LIMK1 signaling pathway is conserved between human BMPR2 and the *Drosophila* BMPR2 ortholog wishful thinking (Wit). Although the CTD sequence homology is only ~30%, both CTDs interact with LIMK1, promote cofilin phosphorylation, and induce actin remodeling (29–36). At the *Drosophila* neuromuscular junction (NMJ), Glass bottom boat, a *Drosophila* BMP ortholog that is released from the postsynaptic muscle, binds Wit and regulates neuromuscular synapse formation, stability, and function (31, 32, 37, 38). Mutant *wit* larvae exhibit a reduced number of mature boutons, and this phenotype can be rescued by Wit but not by Wit that lacks the CTD (32), supporting an essential role of the CTD-LIMK1 interaction and noncanonical signaling in synaptic

¹Cardiovascular Research Institute, University of California, San Francisco, San Francisco, CA 94143, USA ²Department of Biochemistry, Vanderbilt University, Nashville, TN 37232, USA ³Institute for Pediatric Regenerative Medicine, Department of Pathology, University of California, Davis, Davis, CA 95817, USA ⁴MIND (Medical Investigation of Neurodevelopmental Disorders) Institute, University of California, Davis, Davis, CA 95817, USA ⁵Anesthesia and Critical Care, Massachusetts General Hospital, Boston, MA 02114, USA ⁶Department of Biochemistry and Molecular Medicine, University of California, Davis, Davis, CA 95817, USA ⁵Anesthesia, CA 95817, USA ⁵Anesthesia,

stability and growth at the NMJ. *BMPR2* mRNA is not among the previously known targets of FMRP. However, because published genome-wide analysis data imply that *BMPR2* mRNA could be an FMRP binding partner and because BMP signaling is involved in synaptic formation in *Drosophila*, we investigated the role of BMPR2 and noncanonical BMP signaling in FXS.

RESULTS

Two isoforms of BMPR2 are differentially expressed

In mammals, alternative splicing of BMPR2 produces two mRNA forms: one "full length" (FL) that contains 13 exons and encodes a 1038-amino acid protein, and a shorter one (Δ CTD) that encodes a 530-amino acid protein. Δ CTD lacks exon 12 (ex12), which encodes 80% of the entire CTD (Fig. 1A). Immunoblot analysis of various tissues from adult mice reveals that the two isoforms are differentially expressed (Fig. 1B). However, although the ratio between BMPR2 protein isoforms varied from tissue to tissue, it did not correlate with the amount of the corresponding mRNA isoform (for example, Δ CTD in liver and heart; Fig. 1B). To investigate the cause of this discrepancy, we expressed the two BMPR2 mRNA isoforms from cytomegalovirus promoter complementary DNA (cDNA) constructs in COS-7 cells. Despite similar mRNA amounts, the Δ CTD protein was ~7 times more abundant than the FL protein in immunoblots probed with the BMPR2 antibody (Fig. 1C, lanes 2 and 3), or with a FLAG antibody and FLAG-tagged BMPR2 protein constructs (fig. S1, lanes 2 and 3). The different amounts of the two protein isoforms could not be explained by the stability of the correspondent transcripts because the ΔCTD mRNA was slightly less stable than the FL mRNA in an experiment in which de novo mRNA synthesis was blocked by actinomycin D, an inhibitor of RNA polymerase II (Fig. 1D). Furthermore, the stability of FL and \triangle CTD proteins was similar over a period of 24 hours when new protein synthesis was inhibited by cycloheximide (Fig. 1E). These results strongly suggest that the BMPR2 isoforms are regulated posttranscriptionally.

BMPR2 translation is inhibited by the mRNA sequence encoding the CTD

To test whether this posttranscriptional regulation is caused by protein size or by the presence of the CTD coding sequence (hereinafter CTD_{seq}), we introduced a nonsense mutation at amino acid 530 in the FL construct (530X; Fig. 2A). The 530X transcript matches the FL transcript in length, but the 530X-encoded protein has the size of the Δ CTD deletion mutant (Fig. 2A). The difference between the transcripts encoding Δ CTD and 530X is the presence of CTD_{seq} (Fig. 2A). Upon transfection in COS-7 cells, we observed that the Δ CTD protein (Fig. 2A) was about three times more abundant than the FL protein (Fig. 2A), which is consistent with the expression data above (Fig. 1C). On the contrary, the expression of 530X and FL (Fig. 2A) was similar. Similar results were obtained by in vitro transcription and translation, suggesting that a cis-acting sequence (or sequences) within the CTD_{seq} affects translational efficiency.

To test the effect of the CTD_{seq} on transcripts originating from the endogenous genomic locus, we used two in vivo systems expressing a Δ CTD protein, one in which the CTD_{seq} is deleted and one in which a frameshift mutation truncates the BMPR2 protein but preserves the CTD_{seq} in the mRNA. We established two independent lines of primary pulmonary artery smooth muscle cells (PASMCs) from a heterozygous knock-in mouse in which ex12 of the *BMPR2* gene has been replaced by a cassette encoding the enhanced green fluorescent protein (eGFP) with a stop codon (39). In these cells (Δ ex12), one *BMPR2* allele generates transcripts that lack ex12 and encode a Δ CTD (amino acids 1 to 529)–eGFP fusion protein (Δ CTD- eGFP); the other allele is intact (Fig. 2B, top). Independent cell lines were established from two wild-type (Fig. 2B, 1 lanes and 3) and two Δ ex12 mice (Fig. 2B, lanes 2 and 4). In both wild-type PASMCs lines (Fig. 2B, lanes 1 and 3), only the FL protein is detected by immunoblot, indicating that FL is the dominant form of BMPR2 protein in PASMCs (Fig. 2B, right). In the two Δ ex12 cells lines (Fig. 2B, lanes 2 and 4), the amount of FL protein is reduced to ~50% (Fig. 2B, lanes 2 and 4) compared to wild-type cells



Fig. 1. Different abundance of BMPR2 isoforms. (A) Schematic of the exon/ intron structure (left) and the protein structure (right) of two isoforms of BMPR2: the FL and the CTD-truncated isoform (Δ CTD). ECD, extracellular domain; TM, transmembrane domain; aa, amino acid. (B) Immunoblot analysis and quantitative reverse transcription polymerase chain reaction (gRT-PCR) for FL and ∆CTD BMPR2 in brain, heart, liver, and kidney isolated from 12-week-old wild-type (WT) mice. Data are means ± SD of five experiments. (C) Immunoblot analysis and qRT-PCR in lysates from COS-7 cells transfected with vectors encoding FL or Δ CTD BMPR2 normalized to β -actin and GAPDH, respectively. Data are means ± SD of five experiments. **P < 0.05 by Student's t test. GAPDH, glyceraldehyde-3phosphate dehydrogenase. (D) qRT-PCR analysis for BMPR2 expression in COS-7 cells transfected with vectors encoding MBP-BMPR2 [in which the BMPR2 ECD was replaced by maltose-binding protein (MBP)], FL or MBP-ACTD (ACTD) BMPR2, and treatment with actinomycin D (ActD) (5 µg/ml) 24 hours afterward for the period of time indicated. Expression was normalized to GAPDH. Data are means ± SD of five experiments. *P < 0.05 by Student's t test. (E) Immunoblot analysis for BMPR2 in COS-7 cells transfected with optimized amount of MBP-BMPR2-FLAG (FL) or MBP-ΔCTD-FLAG (ΔCTD) expression vectors (to adjust for similar abundance of expressed protein) and treated with cycloheximide (CHX) (10 µg/ml) 48 hours afterward for the period of time indicated. Protein was normalized to β -actin. Data are means \pm SD of four experiments.

(Fig. 2B, lanes 1 and 3), consistent with the gene dosage of wild-type *BMPR2*, but the Δ CTD-eGFP protein accumulates ~7 times more than the FL protein (Fig. 2B, right). This result demonstrates a more efficient translation of the Δ ex12 transcripts compared to wild-type transcripts, presumably because of the lack of *CTD_{seq}* (Fig. 2B). We used our second in vivo approach to test this hypothesis. Using TALEN (transcription activator-like effector nucleases)-mediated genome editing in rat PAC1 pulmonary arterial smooth muscle cells, we engineered a frameshift mutation at Leu⁵⁰⁴ in ex11 of *BMPR2* (L504fs; Fig. 2C), which results in a premature termination at Met⁵⁰⁵. *L504fs* transcripts encode the Δ CTD protein but still contain the *CTD_{seq}* (Fig. 2C). Therefore, if the *CTD_{seq}* mediates translation inhibition, the L504fs protein should be expressed less efficiently than the Δ CTD-eGFP protein in Δ ex12 cells, compared to the respective wild-type proteins were



Fig. 2. Translational regulation of BMPR2 through the mRNA sequence encoding the CTD. (A) Top: Schematic of the expected mRNA and protein structures of C-terminal FLAG-tagged FL, △CTD, and 530X BMPR2. CTD_{seq} is the mRNA sequence encoding CTD. Bottom: Immunoblotting and qRT-PCR of FLAG or BMPR2, respectively, in transfected COS-7 cells. BMRP2 mRNA was normalized to GAPDH. Data are means ± SD of three experiments. *P < 0.05 by analysis of variance (ANOVA) with post hoc Tukey's test. (B) Top: schematic of WT and BMPR2 ex12 deletion mutant (Δ ex12) allele. In the Δ ex12/WT mice, the ex12 of one allele of the BMPR2 gene was replaced with eGFP. Right: Immunoblot for BMPR2 abundance in two independent primary vascular smooth muscle cell cultures generated from WT (WT/WT) and heterozygous (Δ ex12/WT) mice. (C) Top: Schematic of WT and BMPR2 L504fs allele. A frameshift mutation was introduced in at Leu⁵⁰⁴ (ex11) of BMPR2 gene using TALEN-mediated genome editing technology (L504fs) in rat vascular smooth muscle PAC1 cells. Consequently, L504fs allele produces mutant BMPR2 lacking 504 to 1038 amino acids; however, the transcripts contain CTD_{sea}. Right: Immunoblots for FL BMPR2 and L504fs in WT and heterozygous cells. Blots in (B) and (C) are representative of two experiments.

expressed at about a 1:1 ratio (Fig. 2C), in marked contrast to the Δ ex12 cells (Fig. 2B). Thus, these genome-driven expression results confirm a role of the *CTD*_{seq} in the *BMPR2* mRNA as a negative translational regulator.

FMRP associates with *BMPR2* mRNA through CTD_{seq} and controls its translation

FMRP binds a small subset of mRNAs and inhibits their translation (10, 14, 22, 40, 41). On the basis of the results of two independent high-throughput RNA immunoprecipitation (RIP) analyses (11, 22) in which *BMPR2* appeared in the list of potentially FMRP-bound mRNAs, we hypothesized that FMRP might be involved in the translational inhibition of FL BMPR2. To first test whether FMRP associates with the CTD_{seq} through putative FMRP binding motifs (FBMs; ACUK, WGGA, or a combination of both) (22), we transfected human embryonic kidney (HEK) 293 cells with the FL or Δ CTD expression constructs and performed an RIP assay. The FMRP-bound mRNAs were sheared to an average length of ~200 nucleotides to compare the enrichment of different regions of the FL or Δ CTD mRNAs (Fig. 3A). We investigated eight putative FBMs



Fig. 3. FMRP binds BMPR2-*CTD*_{seq} and negatively regulates translation. (A) RIP assay for the relative enrichment of *FL BMPR2* mRNA regions [schematic (top) marked with FMRP binding regions (black or red) and corresponding primers (orange; see table S2)] in FMRP immunoprecipitates from HEK293 cells transfected with FLAG-tagged FL or Δ CTD BMPR2 constructs. Data are representative of two experiments. Pulldown with non-specific immunoprecipitation. (B) RIP assay as in (A) and immunoblotting in HEK293 cells transfected with FL or mutant (Δ 3240 or Δ 4125) BMPR2. Data are means \pm SD of three experiments. **P* < 0.05; ***P* < 0.01 by ANOVA with post hoc Tukey's test.

(22) that span the entire length of the BMPR2 transcript (Fig. 3A and Table 1). Primer set #3468, which amplifies a region of the CTD_{seq} that does not contain FBMs and was not enriched in a high-throughput RIP study (22), was added as negative control (Fig. 3A). The result indicated that all the five tested FBMs located in the CTD_{seq} (#2736, #3116, #3240, #3630, and #4125; Fig. 3A) were highly enriched, demonstrating that FMRP binds to the CTD_{seq} (Fig. 3A). No enrichment of RNAs was observed for the negative control primer #3468 (Fig. 3A), suggesting that an FBM is required for FMRP binding within the CTD_{seq}. The three FBMs found in the KD were not enriched. Thus, it appears that the presence of short FBMs is not sufficient to bind FMRP to the 5' region of the BMPR2 transcript and suggests that additional determinants are required for FMRP recruitment. This result is consistent with our finding that the CTD_{seq} in FL transcripts [as well as in 530X (Fig. 2A) and L504fs (Fig. 2C)] inhibits translation, whereas translation of Δ CTD and Δ ex12 (Fig. 2B) transcripts was not inhibited because they lack the CTD_{seq}. To examine whether the binding of FMRP to FBMs localized in the CTD_{seq} is required for translational control, we generated deletion mutants, $\Delta 3240$ and $\Delta 4125$, in which sets of ACUK/WGGA combination FBMs, which are reported to be potent FBMs (22), were deleted (Fig. 3B) and subjected them to FMRP-RIP, as well as protein and mRNA expression analyses (Fig. 3B

and fig. S2). Each mutant produces a three–amino acid deletion but preserves the translation reading frame in FL BMPR2. Either mutant decreased the association with FMRP by 75 to 90% (Fig. 3B), suggesting that FMRP binds synergistically to RNA and both clusters of FBMs are required for FMRP binding. Concurrently, the amount of both Δ 3240 and Δ 4125 mutant proteins was increased by 1.3- and 1.8-fold, respectively, in comparison with wild type (Fig. 3B). Thus, we conclude that the binding of FMRP to FBMs in the *CTD*_{seq} results in translational inhibition of FL transcripts.

Modulation of FMRP alters FL-BMPR2 protein abundance

Next, we examined whether loss or gain of FMRP dictates the BMPR2 protein amount and the signal downstream of BMPR2. FMRP was depleted in COS-7 cells (Fig. 4A) by small inhibitory RNAs (siRNAs) (si*FMR1*). Whereas si-*FMR1* did not affect FL mRNA abundance, FL protein was increased ~1.5-fold over controls (Fig. 4A). Conversely, expressing exogenous *FMR1* under doxycycline control (Tet-On system) in a stable HEK293 cell line reduced FL protein by 30%, compared to control cells expressing GFP. The expression of FL mRNA was unchanged (Fig. 4B). These results confirmed that the presence of FL protein inversely correlates with the abundance of FMRP and provided the rationale to examine the effect of FMRP loss on BMPR2 in *FMR1*-null mice. Mouse embryonic

Table 1. List of potential FMRP binding sites identified in the BMPR2 transcripts by RIP sequence. On the basis of a previous research (*16*), potential FMRP binding sites were searched in the BMPR2 transcripts. Its core sequence (WGGA or ACUK) is underlined. Location of the RIP primers and the enrichment of RNA fragments are shown. NA, not applicable.

Domain	Start	End	FBM sequence	FBM type	RIP primer	Enrichment	T-to-C conversion/ read count	Enrichment
ECD	203329583	203329597	NA	NA			0.83	
ECD	203329616	203329667	NA	NA			0.71	
ECD	203329669	203329687	NA	NA			0.22	
ECD	203332327	203332341	NA	NA			0.83	
ТМ	203378479	203378507	NA	NA			0.69	
KD	203383687	203383725	ACAACAUUGCCCGCUUUAUAG UUGGAGAUGAGAGAGUCA	WGGA	1820	No	0.95	No
KD	203383744	203383768	AAUAUUUGCUUGUGAUGGAGUACUA	WGGA			0.76	
KD	203384831	203384856	AGUCUCCACACAAGUGACUGGGUAAG	ACUK			0.36	
KD	203384869	203384893	UGCUCAUUCUGUUACUAGAGGACUG	Mixed	2041	No	0.33	No
KD	203395532	203395544	NA	NA			0.25	
KD	203395592	203395623	UUAUUAGUGACUUUGGACUGU CCAUGAGGCUGA	Mixed			0.62	
KD	203397352	203397369	UGUGAACUUGAGGGACUG	ACUK			0.5	
KD	203397390	203397420	UAGACAU <u>GUAU</u> GCUCU <u>UGGA</u> CUA AUCUAUUG	WGGA			0.88	
KD	203417463	203417487	AUCGAAGACUGUUGGGACCAGGAUG	Mixed	2513	No	0.83	No
KD	203417515	203417538	CUGAGGAAAGGAUGGCUGAACUUA	Mixed	2513	No	0.71	No
KD	203417570	203417588	NA	NA			0.6	
CTD	203420039	203420070	UACAUUGAAGACUCUAUCCAU CAUACUGACAG	ACUK	2736	Yes	0.65	Yes
CTD	203420082	203420103	NA	NA			0.71	
CTD	203420438	203420465	AAUCUCAUGGAGCACUCUCUUAAACAGU	WGGA	3116	Yes	0.8	Yes
CTD	203420507	203420530	CUUUACCCACUCAUAAAACUUGCA	ACUK	3240	Yes	1	Yes
CTD	203420532	203420570	UAGAAGCAACUGGACAGCAGG ACUUCACACAGACUGCAA	Mixed	3240	Yes	0.56	Yes
CTD	203420588	203420610	NA	NA			0.78	
CTD	203420887	203420913	NA	NA			0.38	
CTD	203420939	203420961	UUUAUUGGUGAGGACACCCGGCU	WGGA	3630	Yes	0.63	Yes
CTD	203420995	203421016	UUUACUGAGACGAGAGCAACAA	ACUK	3630	Yes	0.67	Yes
CTD	203421033	203421052	GU <u>GUUC</u> UGGAUCGUCUUGUG	WGGA			0.82	
CTD	203421200	203421222	CCUAAUUCUCUGGAUCUUUCAGC	WGGA			0.73	
CTD	203424647	203424668	AUAUAGGAAUGAACUGUCUGUG	Mixed	4125	Yes	0.63	Yes



Fig. 4. *FMR1* is a negative regulator of FL BMPR2 protein expression. (A) Immunoblotting and qRT-PCR for BMPR2 in COS-7 cells transfected with siRNAs against *FMR1* (si-*FMR1*) or a control sequence (si-Ctr). Data are means \pm SD of three experiments. **P* < 0.01 by Student's *t* test. (B) Immunoblotting and qRT-PCR for BMPR2 in stable HEK293 cell lines with doxycycline (Dox)–inducible expression of FLAG-tagged FMRP or GFP (control). Protein and mRNA abundance was normalized to GAPDH. Data are means \pm SEM of three experiments. **P* < 0.05 by ANOVA with post hoc Tukey's test. (C) Immunoblotting and qRT-PCR for BMPR2 in brain lysates from P7 *FMR1*^{-/-} or WT mice. Data are means \pm SEM of seven individual mice. **P* < 0.05 by Student's *t* test. Asterisk (*) indicates a nonspecific band in the FMRP immunoblot panel. In (A) to (C), protein and mRNA abundance was normalized to GAPDH.

fibroblasts (MEFs) derived from FMR1-null mice exhibited ~2-fold increase in BMPR2 (FL) protein compared to MEFs derived from wild-type littermates (fig. S3A). Upon BMP treatment, the phosphorylation of SMAD1/5, a hallmark of the "canonical" pathway downstream of BMPR2, was also increased (fig. S3A), whereas TGFB signaling was not affected, suggesting that the effect of the loss of FMRP is BMP-specific (fig. S3B). Next, we collected brain samples from FMR1-null or control (wild-type) mice at postnatal day 7 (P7) and subjected them to immunoblot and gRT-PCR analysis of BMPR2 abundance and expression, respectively. At protein but not at mRNA level, a greater amount of FL isoform was detected in the brain of FMR1-null mice in comparison with wild-type mice (Fig. 4C). Unlike FL, the amount of Δ CTD isoform was similar in both FMR1-null and wildtype mice (Fig. 4C). These results confirm an increase of FL isoform in the FMRP-depleted mouse brain and, thus, suggest that the increase in the CTD-mediated signal might contribute to the development of the neuronal phenotypes associated with FXS.

Abnormalities at the NMJ in a loss-of-expression *FMR1 Drosophila* mutant are rescued by reducing Wit

The Drosophila NMJ is a powerful model system for uncovering and characterizing genetic and molecular mechanisms that regulate synaptic formation (36, 42). A loss-of-expression mutation of dFMR1, the Drosophila ortholog of FMR1, causes overgrowth of synaptic boutons in the larval NMJ (43). In contrast, loss-of-expression mutants of the Drosophila ortholog of BMPR2 Wit exhibit reduced numbers of synaptic boutons (31, 37, 38). Because both Wit and dFMR1 are expressed in presynaptic motor neurons, we speculated that translational regulation of *Wit* by dFMR1 might be important for synaptic growth and stability at the NMJ. More precisely, if the overgrowth of boutons in dFMR1 mutants was due to increases of Wit and BMP signaling in presynaptic motor neurons, this phenotype could be rescued by reducing Wit abundance. To test this hypothesis, we crossed heterozygous ($wit^{412/+}$) or homozygous (wit^{412/B11}) wit mutants with heterozygous dFMR1 mutants (dFMR1^{D113/+}) and analyzed the number of mature synaptic boutons and branching synapses by immunofluorescence staining of the postsynaptic marker discs large (DLG; Fig. 5A, green) and phalloidin (Fig. 5A, red) on muscle 6/7 in the abdominal segment 3 (A3) of third instar larvae. We also costained samples with DLG (Fig. 5B, red) and the presynaptic marker horseradish peroxidase (HRP) (Fig. 5B, green) to distinguish mature synaptic boutons, which are costained with both HRP and DLG, from "synaptic footprints," which are stained only with DLG (32). As previously reported (43), loss of expression of one copy of dFMRI ($dFMRI^{D113/+}$) resulted in ~75% increase in the number of mature boutons and ~60% increase in branching compared to wild-type animals (Fig. 5C). When one allele of wit was mutated in the heterozygous dFMR1 mutant background (dFMR1^{D113/+}; wit $^{A12/+}$), the bouton number decreased by ~20% (Fig. 5C). Mutation of both wit alleles in the dFMR1 mutant background (dFMR1^{D113/+}; wit^{A12/B11}) reduced both boutons and branching numbers to indistinguishable quantity from those observed in *wit* mutants ($wit^{A12/H}$ or $wit^{A12/B11}$) with wild-type background (Fig. 5C). Thus, the augmented synaptic morphology in the dFMR1 mutant is, at least in part, due to increased Wit abundance and signaling in presynaptic neurons, and both can be decreased by reducing wit gene dosage. Considering that they are orthologs, the epistatic relationship between dFMR1 and wit in the regulation of synaptic development at the NMJ is consistent with the translational regulation of BMPR2 by FMRP.

CTD-LIMK1–dependent signal promotes augmented dendritogenesis in *FMR1*-null neurons

In the Drosophila NMJ, presynaptic expression of a Wit mutant lacking the CTD fails to rescue synaptic stability in wit-null mutants (32). This failed rescue has been attributed to the failure of the Wit Δ CTD mutant to activate dLIMK1 (the Drosophila ortholog of LIMK1), which is essential for promoting the stability of neuromuscular synapses (32). To compare the effect of FL and Δ CTD in BMP-mediated actin remodeling and filopodia formation, we expressed each isoform in the neural crest-derived neuroblastoma cell line N1E-115 (N1E) and subjected them to BMP7 (also known as Osteogenic protein-1) stimulation followed by phalloidin staining (Fig. 6A). BMP7 was used in this experiment because it is expressed in the developing nervous system and is a potent inducer of dendritic morphogenesis in N1E cells (29) and rat sympathetic neurons (44). Phalloidin staining in control cells showed that BMP7 induced the formation of filopodia, which are fundamental for neurite outgrowth (Fig. 6A). In comparison, FLexpressing cells showed an increased intensity of both basal and BMP7induced phalloidin staining and filopodia formation (Fig. 6A, top). We observed proper plasma membrane localization (45) and similar amounts of ΔCTD and FL proteins (Fig. 6A, bottom left), which correlated to similar induction of the ID1 gene (Fig. 6A, bottom right), a readout of the



Fig. 5. Epistasis between *dFMR1* and wit during the formation of NMJ in *Drosophila*. (A) Confocal images of muscle 6/7 in segment A3 from larvae of the indicated genotype. Boutons and muscle were stained with DLG antibody (green) and Alexa Fluor 568–conjugated phalloidin (red), respectively. Scale bars, 50 µm. (B) As in (A), except presynaptic and postsynaptic boutons and muscle were stained with HRP (green) and DLG (red) antibodies and Alexa Fluor 647–conjugated phalloidin (blue), respectively. Scale bars, 50 µm. (C) Bouton (left) and branch (right) numbers in muscle 6/7 in segment A3. Data are means ± SD of the following numbers of larvae: n = 11 for WT, n = 27 for $dFMR1^{D113M/+}$, n = 41 for $wit^{A12/+}$; $dFMR1^{D113M/+}$, n = 10 for $wit^{A12/B11}$; $dFMR1^{D113M/+}$, and n = 10 for $wit^{A12/B11}$. *P < 0.05; **P < 0.01 by ANOVA with post hoc Tukey's test.

SMAD-dependent signaling. However, Δ CTD did not mimic the FL effect but, rather, inhibited BMP7-mediated filopodia formation (Fig. 6A), indicating that Δ CTD may act as a dominant negative against endogenous BMP7induced noncanonical signaling activation. These results suggest that FL, but not Δ CTD, promotes actin remodeling presumably through CTD-mediated activation of LIMK1 and phosphorylation of cofilin, which is fundamental for dendritogenesis and synaptogenesis.

To test whether filopodia formation is augmented upon down-regulation of FMRI, we transfected N1E cells with si-FMRI and stained with phal-

loidin after BMP7 stimulation. Filopodia formation was increased in si-FMR1 cells with and without BMP7 stimulation (Fig. 6B) but reverted to the morphology of control cells when LIMK1 was simultaneously silenced by siRNA (Fig. 6B). These results suggest that the increase of the FL-LIMK1 signal leading to altered actin dynamics may be an underlying cause of dendritic spine abnormalities in cortical and hippocampal neurons in animal model of FXS and patients (46-49). To test whether perturbation of LIMK1 activity could modulate actin dynamics in FMRPdepleted cells, we exposed si-FMR1-treated N1E cells to LIMKi-3, a noncytotoxic small-molecule inhibitor of LIMK1 (50), and then treated them with BMP7 and scored for filopodia formation (Fig. 6C). As a control, cells were treated with LDN-193189 (LDN), a potent inhibitor of the BMP type I receptor-mediated, SMAD-dependent signaling pathway (51, 52). LIMK-i inhibited BMP7-mediated filopodia formation (Fig. 6C), similarly to si-LIMK1 (Fig. 6B). Phosphorylation of cofilin, but not SMAD1/5, was reduced by LIMK-i, indicating an effective and specific inhibition of a BMPR2 noncanonical pathway mediated by LIMK1 activity by LIMK-i treatment (fig. S4A, top). On the contrary, LDN effectively inhibited both SMAD1/5 phosphorylation and transcriptional activation of the SMAD target gene Id3 (fig. S4, A and B) but did not inhibit filopodia formation (Fig. 6C). Thus, the LIMK1-cofilin pathway, but not the SMAD pathway, plays an essential role in promoting BMP7-BMPR2-mediated filopodia formation.

Reduction of the BMPR2-LIMK1 pathway rescues abnormalities of neuronal development in *FMR1*-null mice

To examine whether the aberrant dendritogenesis in FMR1-null neurons (53, 54) can be rescued by inhibition of LIMK1, we isolated cortical neurons from FMR1-null or wild-type mice at P0, followed by BMP7 and LIMK-i or LDN treatment for 24 hours (Fig. 7A). BMP7 treatment significantly increased the dendrite number in FMR1-null neurons, whereas it had only a modest effect on wild-type neurons (Fig. 7A). Treatment with LIMK-i, but not LDN, abolished the BMP7-dependent increase of dendritogenesis in FMR1-null neurons (Fig. 7A). The density of spines on the dendrites of cortical and hippocampal neurons is abnormally high in FMR1-null mice (46-49) and FXS patients (55). Staining with phalloidin revealed a ~40% increase in spine density in FMR1-null cortical neurons compared to that in wild-type neurons after 16 days in culture (Fig. 7B), resembling the morphology described in vivo (46-49). LIMK-i treatment reversed the abnormal dendritic spine morphology of FMR1-null neurons to the wild-type phenotype (Fig. 7B), suggesting that LIMK-i treatment could ameliorate the abnormally high turnover of dendritic protrusions observed in FMR1-null neuron (54). These results suggest that FMRP-deficient neurons develop irregular dendrites and spines at least in part as a result of increased FL BMPR2-LIMK1 signaling activity.

Thus, we postulated that reducing the abundance of FL BMPR2 or the activation of LIMK1 in vivo might prevent some aspects of the FMR1-null phenotype. We first tested whether a reduction of the BMPR2 gene dosage could rescue the neuronal morphology in FMR1-null mice. We crossed FMR1 homozygous-null and BMPR2 heterozygous mice to generate FMR1^{-/-}; BMPR2^{+/-} mice. BMPR2 homozygous-null mice die during early embryogenesis and could not be tested. As previously reported (46-49), granule neurons in the dentate gyrus (DG) of FMR1-null mice exhibit an increased dendritic spine density (Fig. 7C, left) with a higher percentage of long thin (>2 μ m) immature spines compared to control (wild-type) mice (Fig. 7C, right). The fraction of long immature spines of DG neurons observed in the $FMR1^{-/-}$; $BMPR2^{+/-}$ mice was similar to that seen in wild-type littermates or BMPR2 heterozygous mice (Fig. 7C). These results demonstrate that genetic perturbations of the BMPR2 signaling pathway are sufficient to modify the abnormal dendrite morphology in an FXS mouse model and in the fly, implying that the pathway is evolutionally conserved.



Fig. 6. BMPR2 modulates actin remodeling through activation of the LIMK1-cofilin pathway. (A) Alexa Fluor 568–conjugated phalloidin staining in N1E cells infected with adenovirus (Ad) expressing GFP (control), FL, or Δ CTD, and treated with BMP7 (10 ng/ml) for 24 hours. Scale bars, 40 µm. Insets are magnified ×2. Data are means ± SEM of ≥33 cells per condition. **P* < 0.05, ***P* < 0.01 by ANOVA with post hoc Tukey's test. A fraction of N1E cells was subjected to immunoblotting for BMPR2 (bottom left) and qRT-PCR for *ID1* expression (bottom right), each was normalized to *GAPDH*. Data are means ± SD of three experiments. **P* < 0.001 by ANOVA with post hoc Tukey's test. (**B** and **C**) Phalloidin staining in N1E cells transfected with targeted siRNA(s) or control (si-Ctr) and treated 48 hours later with BMP7 (10 ng/ml) (B) for 24 hours or LIMKi-3 (LIMK-i; 3 µM) (C) or BMP type I receptor kinase inhibitor LDN (100 nM) for 12 hours and then BMP7 (10 ng/ml) for 24 hours. Scale bars, 50 µm. Blue arrows indicate filopodia. Phalloidin stain was blindly assessed and quantitated by ImageJ. Data are means ± SEM of ≥26 (B) or 29 (C) cells per condition. ***P* < 0.01 by ANOVA with post hoc Tukey's test. ns, not significant.

Next, we examined whether postnatal administration of the LIMK1 inhibitor could rescue the abnormal morphology of hippocampal neurons in the *FMR1*-null mouse. LIMK-i or vehicle (dimethyl sulfoxide) was injected into the cranium of P1 and P4 mice, followed at P7 by Golgi's staining of the central nervous system (CNS) and biochemical analysis. Brain samples from *FMR1*-null mice treated with LIMK-i showed markedly reduced abundance of phosphorylated cofilin (fig. S5), indicating an effective inhibition of the catalytic activity of LIMK1. As previously re-

ported (46–49), compared to those in wild-type mice, granule neurons in the DG of *FMR1*-null mice exhibited increased dendritic spine density (Fig. 7D) and an increased proportion of long, thin (>2 μ m), and immature spines (Fig. 7E). In *FMR1*-null mice treated with LIMK-i, both spine density (Fig. 7D) and the proportion of immature spines (Fig. 7E) were reversed to those observed in mock-treated wild-type mice (46–49). These results demonstrate that genetic and pharmacological perturbations of the BMPR2-LIMK1-cofilin pathway are sufficient to modify the abnormal dendrite morphology of an FXS mouse model.

Induction of the BMPR2-LIMK1-cofilin pathway is seen in brain tissue from FXS patients

Finally, to test the hypothesis that the activation of the BMPR2-LIMK1-cofilin pathway found in the mouse FXS model also occurs in human FXS patients, we performed immunohistochemical (IHC) analysis to compare the amount of BMPR2 in the prefrontal cortexes of FXS patients and of gender- and age-matched control individuals. We observed ~2-fold more FL BMPR2 in FXS compared to control cortexes in the upper to middle layers (Fig. 8A). Consistent with the result of IHC, an immunoblot analysis confirmed that loss of FMRP in FXS patients' cortexes correlated with a ~3-fold increase of FL BMPR2 protein (Fig. 8B) and 16-fold increase of phosphorylated cofilin in FXS cortexes compared to control cortexes (Fig. 8B). These results demonstrate that loss of FMRP leads to an increase of FL BMPR2, which in turn augments the phosphorylation of cofilin, presumably through activation of LIMK1, in the brain tissue of FXS patients. Thus, they support our model of a critical contribution of the BMPR2-LIMK1 signaling axis to the pathology of human FXS and offer a novel opportunity to a therapy for FXS and possibly other ASD conditions.

DISCUSSION

The BMP signaling pathway controls morphogenesis of nearly every tissue and organ by coordinating basic properties of the cell, such as differentiation, proliferation, motility, morphology, and death, either during development and in the adult (27, 28). Here, we demonstrate that the *BMPR2* mRNA is a target of translational regulation by FMRP and provide evidence supporting a link between augmented BMP signaling and neurological disorder in humans. The epistatic relationship between *FMR1* and *BMPR2* and the physiological significance of the FMRP-mediated down-regulation of

BMPR2 during neuronal development have been conserved during evolution from *Drosophila* to mammals. In particular, the noncanonical signaling pathway downstream of BMPR2, which includes LIMK1, appears to play an essential role in the development of the neuropathology of patients with FXS and in the mouse model of FXS. Tempering this pathway, either by reducing the *BMPR2* gene dosage or applying a small-molecule inhibitor of LIMK1, rescues the neuromorphological abnormality in the FXS mouse model, indicating that the BMPR2-LIMK1 axis is an intervention site for



Fig. 7. Inhibition of BMPR2-LIMK1 pathway rescues abnormal neuronal development in the mouse model of FXS. (A) Confocal microscopy detecting the number of dendrites (arrowheads) in neurons isolated from the cortex of FMR1-/- or littermate WT P0 mice, cultured for 3 days, and then transfected with a GFP expression plasmid and treated 24 hours later with LIMK-i (3µM) or LDN (100 nM) for 12 hours, followed by BMP7 (10 ng/ml) for 24 hours. Cells were stained and imaged at day 6 in vitro. Data are means + SEM from ≥30 GFPpositive neurons. *P < 0.01 by ANOVA with post hoc Tukey's test. Scale bar, 30 μm. (B) Alexa Fluor 568conjugated phalloidin staining (left) and confocal microscopy detecting spine density (right) in neurons isolated from the cortex of FMR1-/- or littermate WT P0 mice, cultured for 14 days, and then treated with LIMK-i (3µM) for 12 hours, followed by BMP7 (10 ng/ml) for 24 hours. Insets are magnified ×2. The minimum was counted. Data are means + SEM from ≥25 neurons. *P < 0.05 by ANOVA with post hoc Tukey's test. Scale bar, 30 μm. (C) Golgi staining (left) in brain tissue isolated at P7 from FMR1^{-/-}, FMR1^{-/-}; BMPR2^{+/-}, BMPR2^{+/-}, or WT littermate mice. Dendritic spines in DG neuron were classified as either short $(<2 \mu m)$ or long immature (>2 μm , red arrowheads), and the percentage of long immature spines was calculated (right). Data are means + SEM from more than seven DG neurons from three mice. *P < 0.01by ANOVA with post hoc Tukey's test. Scale bar, 5 µm. KO, FMR1 homozygous-null mice; Het, BMPR2 heterozygous mice. (D and E) Golgi staining (left) in brain tissue isolated at P7 from FMR1^{-/-} or littermate WT mice treated with LIMK-i by intracerebroventricular injection at P1 and P4. Spine density was calculated as the total number of spines in a 30-µm stretch (D). The percentage of long immature spines (>2 µm) indicated by red arrowheads was calculated (E). Data are means and SEM from more than seven DG neurons from three mice. *P < 0.01 by ANOVA with post hoc Tukey's test. Scale bars, 10 μ m (D) and 5 μ m (E).

development of an FXS and ASD therapy. A previous study, which finds that *BMPR2* deletion in the mouse hippocampus and forebrain produce reduced anxiety-related behaviors and increased exploration, supports a role of BMPR2 signaling in modulating cognitive and behavioral phenotypes (*56*).

CTD_{seq} contains nine FBMs (three ACUK, four WGGA, and two mixed; Table 1). Although the mRNA sequence encoding the KD (KDsea) also contains FBMs (two ACUK, three WGGA, and four mixed; Table 1), association of FMRP with KDseq was not detectable (Fig. 3A), strongly suggesting the potential involvement of other RNA sequence or structural elements adjacent to FBMs for recruitment and stable association of FMRP with target mRNAs. Deletion analysis of individual FBMs suggests that FMRP binds multiple sites and represses translation without affecting mRNA stability. Furthermore, deletion of one FBM ($\Delta 4125$) or two FBMs (Δ 3240) produces a smaller reduction of BMPR2 protein compared to the deletion of all nine FBMs (Δ CTD), suggesting that the recruitment of FMRP to multiple FBMs in CTDseq results in more potent repression of BMPR2 translation. Several modes of translational inhibition by FMRP have been reported (57). In some cases, FMRP binds the coding sequence, stalls polysomes, and represses active translation, whereas on some other targets, FMRP binds the 5' or 3' untranslated region and inhibits translation initiation (12-15). More recently, FMRP has been shown to bind directly to the L5 protein on the 80S ribosome and inhibit translation (58). We speculate that FMRP stalls polysomes or directly inhibits the activity of the 80S ribosome associating with the BMPR2 mRNA, as previously proposed (26). Ribosome profiling analysis in the presence or absence of FMRP will give a snapshot of the translation status of the BMPR2 mRNA and help understand the mode of translational inhibition by FMRP.

It has been reported that higher amounts of *FMR1* mRNA and FMRP correlate with a more aggressive form of breast cancer (59). This correlation was attributed to FMRP binding to E-cadherin and vimentin mRNAs and regulating epithelial-to-mesenchymal transition, a hallmark of cancer invasion and metastasis (57, 59–62). Conversely, loss of BMP signaling in mammary carcinomas can accelerate metastasis, indicating that BMP plays a tumor suppressor role (63). Therefore, it is conceivable that increased *FMR1* transcripts might permit more invasive breast cancer by reducing the tumor suppressive



Fig. 8. Induction of the BMPR2-LIMK1-cofilin pathway is seen in brain tissue from FXS patients. (A) Representative images of BMPR2 staining on the middle layer prefrontal cortex from FXS patients and control subjects (CTR) (top). Insets are magnified ×2. Average intensity of BMPR2 signals of lower, middle, or upper layer of prefrontal cortex of three FXS and three controls was quantitated (bottom). Data are means ± SD of three samples. *P < 0.01 by Student's *t* test. (B) Immunoblotting in lysates from frontal cortexes from three FXS patients and three control subjects. Data are means ± SEM of three samples. *P < 0.01 by Student's *t* test. P-cofilin, phosphorylated cofilin; T-cofilin, total cofilin.

activity of the BMP signal. Another disease potentially associated with the FMRP-BMPR2 axis is pulmonary artery hypertension (PAH). More than 80% of patients with heritable PAH and ~20% with sporadic PAH carry a loss-of-expression or loss-of-function mutation in the *BMPR2* gene (64), but the penetrance of PAH among *BMPR2* carriers is less than 20% (64). Therefore, we speculate that expression of genetic variants of *FMR1* or dysregulation of FMRP might determine the development of PAH among a small subset of *BMPR2* carriers.

The kinase PAK1, which phosphorylates and activates LIMK1, also appears to play a role in FXS (65). It has been reported that PAK1 binds the KH2 domain of FMRP and phosphorylates it, allowing it to be active (66). Furthermore, expression of a dominant-negative PAK1 mutant in the forebrain (67) or pharmacological inhibition of PAK1 rescues irregular dendritic spine morphology and behavioral abnormalities in FMR1-null mice (68), supporting the inhibition of PAK1 as a potential therapy for FXS and ASDs. However, there is yet no evidence of abnormal activity and/or amount of PAK1 in human patients with FXS or other neurological diseases. PAK1 is ubiquitously expressed, activated by various extracellular stimuli, and upstream of various essential biological processes, including cell proliferation, motility, and apoptosis, in addition to the control of the actin cytoskeleton (69). Thus, compared to LIMK1, PAK1 might be a less ideal therapeutic target for FXS because of the expected secondary effects of its inhibition. In contrast, LIMK1, which is expressed mainly in the CNS (70), appears to phosphorylate only cofilin, which directly controls actin polymerization (71). LIMKi-3, which we used in this study, was originally identified as an anticancer drug that inhibits LIMK1 and LIMK2 activity (median inhibitory concentration equals 7 and 8 nM for LIMK1 and LIMK2, respectively) and blocks tumor cell invasion (72). Under the condition used in mouse primary neuron culture, we detected no cytotoxicity; however, development of LIMK1-specific inhibitor might be more desirable for FXS therapy because of the ubiquitous presence of LIMK2.

MATERIALS AND METHODS

Animals

B6.129P2-*FMR1*^{tm1Cgr}/J, FVB.129P2-*Pde6b*⁺ *Tyr*^{c-ch} *FMR1*^{tm1Cgr}/J, FVB.129P2-*Pde6b*⁺ *Tyr*^{c-ch}/AntJ (The Jackson Laboratory), B6.129S4(Cg)-*BMPR2*^{tm1.1En1}/Mmnc (MMRC), and Tg(ACTB-cre)2Mrt mice were maintained according to the University of California, San Francisco (UCSF) Laboratory Animal Resource Center guidelines. To obtain *FMR1* knockout littermates, we used wild-type and knockout (*FMR1*^{-/-}) male mice derived from *FMR1* heterozygous female and hemizygous male mating. To obtain *FMR1*^{-/-}; *BMPR2*^{+/-} mutant mice and littermates, we first mated B6.129S4(Cg)-*Bmpr2*^{tm1.1En1}/Mmnc mice with Tg(ACTB-cre)2Mrt mice, followed by mating with *FMR1* knockout mice, and finally *FMR1*^{+/-}; *BMPR2*^{+/-} female and *FMR1*^{-/-}; *BMPR2*^{+/-} male mice were mated. Genotyping was performed following the The Jackson Laboratory protocol.

Drosophila lines and genetic crosses

Flies were raised at 25°C. Control wild-type flies used in this study were the w¹¹¹⁸ strain. Mutant alleles wit⁴¹², wit^{B11}, and dFMRI^{D113M} were obtained from the Bloomington Stock Center. These lines were crossed to generate wit^{412/+} [wit⁴¹²/+], dFMRI^{D113M/+} [dFMRI^{D113M/+}], wit^{412/+}; dFMRI^{D113M/+} [wit⁴¹²; dFMRI^{D113M/+}], wit^{412/B11} [wit⁴¹²/wit^{B11}], and wit^{412/B11}; dFMRI^{D113M/+} [wit⁴¹²; dFMRI^{D113M/+}].

Cell culture and transfection

COS-7, HEK293, N1E, and PAC1 cells were purchased from American Type Culture Correction and maintained in Dulbecco's modified Eagle's medium (DMEM) containing 10% fetal bovine serum (FBS). HEK293 cell lines stably expressing FLAG-HA-*FMR1* isoform 1 and eGFP were reported previously (22). Primary MEFs were prepared from embryonic day 14.5 *FMR1^{-/-}* and control wild-type mouse embryos and cultured in DMEM containing 10% FBS. PASMCs from heterozygous Δ ex12 and control wild-type mice were obtained from K. D. Bloch and cultured as reported previously (39). Postnatal (P0) cortical neurons were prepared from littermate *FMR1^{-/-}* and wild-type mice and cultured in Neurobasal Medium (Life Technologies). Cells were cultured at 37°C in the presence of 5% CO₂. Plasmid and siRNA transfections were performed using FuGENE 6 (Promega) or Lipofectamine 2000 (Life Technologies) and Lipofectamine RNAiMAX (Life Technologies) reagents, respectively.

TALEN-mediated targeted gene mutagenesis in PAC1 cell line

TALEN arms were designed using TAL Effector Nucleotide Targeter 2.0 (Cornell University). PAC1 cells were transfected with two targeting TALEN effectors [TAL_17Q (NN HD NI HD NI NN NG NN NG NN HD NG NN NI NN NN NI NN NI) and TAL_18R (NG NG HD NG HD NG HD HD HD NI NG NI NG HD NI] and with pcDNA3.1-eGFP in a ratio of 5:5:1. Forty-eight hours after transfection, eGFP-positive cells were sorted by flow cytometry, and single cells were reseeded into 96-well plates. After single-cell cloning, the mutations present in each clone were examined by Sanger sequencing.

Plasmid constructs

The N-terminal FLAG-tagged FL BMPR2, BMPR2 mutants, and negative control were subcloned into pcDEF3 for overexpression experiments. The BMPR2 mutants were constructed by a PCR-based approach. The BMPR2 530X was constructed by inserting a FLAG coding sequence and stop codon between amino acids 529 and 530. The MBP-BMPR2-FLAG and MBP-ΔCTD-FLAG were constructed by replacing the extracellular domain of BMPR2 with MBP. pcDNA3.1-eGFP was constructed by inserting the eGFP sequence into the pcDNA3.1 vector.

siRNAs and adenovirus

The siRNAs targeting human, monkey, mouse, and rat *FMR1* (5'-GGAUGAU-AAAGGGUGAGUUdTdF3' and 5'-GCUAGAAGCUUUCUCGAAUdTdF3') and mouse *LIMK1* (5'-UAGUACUGGUGUGAAAGGGAGACdTdT-3') were previously reported (73–75) and purchased from Dharmacon. The control siRNA was purchased from QIAGEN. Adenovirus vectors expressing FLAG-HA-FMRP isoform 1 and eGFP were prepared using AdEasy Adenoviral Vector System (Agilent Technologies). The adenovirus vectors that express FL and Δ CTD BMPR2 were previously reported (45).

qRT-PCR analysis

Total RNAs were isolated using TRIzol (Life Technologies), and real-time PCR was performed as described previously (76) using the oligodeoxynucleotide primers listed in table S1. The primers for human or mouse *ID1* and *GAPDH* were reported previously (77).

Immunoblot analysis

Cells and mouse tissues were lysed in lysis buffer [tris-buffered saline (TBS) containing 1% NP-40, 1 μ M NaF, and 1 mM EDTA). The frontal cortexes of FXS patients and controls were lysed in RIPA (radioimmunoprecipitation assay) buffer. Lysed samples were separated by SDS–polyacrylamide gel electrophoresis, transferred to nitrocellulose membrane (Millipore), immunoblotted with antibodies, and visualized using a LI-COR Imaging System. The antibodies used were directed against FLAG epitope tag (M2, Sigma), FMRP (ab17722, Abcam), BMPR2 KD (19087-1-AP, Proteintech Group), total SMAD1/5/8 (BMR 00479, Bio Matrix Research), phospho-SMAD1/5/8 (#9511, Cell Signaling), cofilin (#5175, Cell Signaling), phospho-cofilin (SC-12912-R, Santa Cruz), β -actin (A5441, Sigma), and GAPDH (MAB374, Millipore).

Phalloidin staining

Eight-well chamber slides (Nunc Lab-Tek) were coated with poly-D-lysine (0.1 mg/ml) overnight, followed by laminin (20 µg/ml) for 1 hour at 37°C. N1E cells (50,000) were seeded onto each chamber, transfected or infected with siRNAs or adenovirus vectors, and treated with BMP7 with or without inhibitors [LIMKi-3 (Millipore) and LDN] as indicated in the figure legends. The cells were fixed, permeabilized, stained with Alexa Fluor 568–conjugated phalloidin (Life Technologies), as described in the manufacturer's protocols, and observed under a point scanning confocal microscope (Leica). The phalloidin signal on filopodia was traced and quantified using the ImageJ program by a blinded investigator.

Immunofluorescence microscopy

Primary cortical neurons were isolated from littermate $FMRI^{-/-}$ and wildtype P0 mice, and 80,000 cells were plated on 12-mm poly-D-lysine-coated coverslips (Corning). The cells were transfected with pcDNA3.1-eGFP plasmid using Lipofectamine 2000 (Life Technologies) after 3 days in vitro. Twenty-four hours after transfection, the cells were treated with LIMK-i for 12 hours, followed by BMP7 (10 ng/ml) treatments for 24 hours. The cells were fixed with 4% PFA/4% sucrose in phosphatebuffered saline (PBS) for 10 min, permeabilized with 0.1% Triton X-100 in PBS for 10 min, blocked with 3% bovine serum albumin/0.02% Triton X-100 in PBS, and stained with GFP antibody (AB13970, Abcam) overnight at 4°C, followed by incubation with an Alexa Fluor 488–conjugated secondary antibody. Confocal images were acquired using a point scanning confocal microscope (Leica). Images were from a projection of z sections.

RIP assay

RIP assay was performed as reported previously (11). Briefly, HEK293 cells were transfected with vectors expressing wild-type or mutant BMPR2. Forty-eight hours after transfection, the cells were washed with PBS containing cycloheximide (100 µg/ml), and the cell dishes were placed on an ice pack and ultraviolet (UV)-cross-linked in ice-cold PBS containing cycloheximide (100 μ g/ml) for three times at 400 mJ/cm² (254-nm UV light) using an UVC 500 cross-linker (Hoefer). The cells were collected, lysed in SDS-immunoprecipitation buffer, sonicated, and centrifuged at 15,000 rpm for 15 min at 4°C. The supernatant was heated to 90°C for 10 min with shaking and diluted nine times by volume with CSK buffer. The lysate was treated with ribonuclease T1 (5 U/ml) (Thermo Scientific) and deoxyribonuclease I (DNase I; 0.8 U/ml) (Roche) for 5 min at 37°C, and SUPERAse In (100 U/ml) (Life Technologies) was added to stop the reaction. After preclear with Protein A Dynabeads (Life Technologies), FMRP and RNA complexes were immunoprecipitated using Protein A Dynabeads bound to either FMRP antibody (ab17722, Abcam) or control Ig overnight at 4°C. After immunoprecipitation, the beads were washed with high-stringency buffer, high-salt buffer, low-salt buffer (twice), and NT2 buffer, followed by two washes with PK buffer. The beads were resuspended in PK buffer containing proteinase K (0.8 mg/ml) (Roche), incubated for 20 min at 37°C with shaking, mixed with an equal volume of PK-7 M urea buffer, and incubated for 20 min at 37°C with shaking. The RNAs were purified by acid phenol/chloroform extraction, followed by ethanol precipitation. The RNA solution was treated with DNase I (1U/ml) for 30 min at 37°C, purified by acid phenol/chloroform extraction and ethanol precipitation, and used for reverse transcription using SuperScript III Reverse Transcriptase (Life Technologies), followed by real-time PCR. The oligodeoxynucleotide primers used for qRT-PCR are listed in table S2.

Larva immunofluorescence and image quantitation

Wandering late third instar larvae were dissected for analysis of NMJ phenotype. After dissection, the tissues were fixed with 8% formaldehyde in PBS for 10 min, washed with PBT (PBS containing 0.1% Triton X-100), and blocked with 5% normal donkey serum in PBT for 1 hour at room temperate. The larvae were incubated with DLG (4F3, Developmental Studies Hybridoma Bank) antibody overnight at 4°C, followed by incubation with Alexa Fluor 488– or Alexa Fluor 555–conjugated secondary antibodies against mouse Ig, Alexa Fluor 488–conjugated HRP antibody (123-545-021, Jackson ImmunoResearch), and Alexa Fluor 568–conjugated (A12380, Life Technologies) or Alexa Fluor 647–conjugated (#8940, Cell Signaling) phalloidin. Confocal images were acquired using a point scanning confocal microscope (Leica). Images used for quantification of NMJ bouton number and branch number were derived from a projection of z sections. For quantification of the branch number, branches with at least three boutons were counted blindly.

Golgi staining

The mice received LIMKi-3 (20 μ g per pup) by intracerebroventricular injection at P1 and P4. At P7, the mice were anesthetized, and the brains were quickly removed and rinsed in distilled water to be used for Golgi's staining. Golgi's staining was performed as described in the manual (FD Rapid GolgiStain Kit, FD NeuroTechnologies). Shortly, the brains were incubated in solution A + B for 3 weeks, followed by 3-day incubation in solution C. We

stained 100- μ m brain sections and performed spine density and morphological analyses under a microscope at ×100 magnification. Quantitation of dendritic spine number in 30 μ m (>30 μ m from cell body) was performed blindly.

FXS patient brain immunohistochemistry

FXS patient and control frontal cortexes were fixed by formalin, and the samples in paraffin were sectioned at 7 µm. Antigen retrieval was performed with 1× diva solution (heat-induced epitope retrieval buffer, Biocare Medical) at 110°C for 8 min, followed by 3% H₂O₂ treatment for 7 min. The samples were incubated with blocking buffer (TBS buffer containing 10% donkey serum and 0.025% Triton X-100) for 1 hour, primary antibodies (in TBS buffer containing 10% donkey serum and 0.0025% Tween 20) for 15 hours at 4°C, biotin-conjugated secondary antibodies (Jackson ImmunoResearch; in TBS buffer containing 10% donkey serum and 0.0025% Tween 20) for 1 hour at room temperature, and DAB Substrate (Vector Laboratories Inc.) for 5 min, followed by counterstaining with Harris hematoxylin (Thermo Scientific), dehydration with ethanol, clearing with xylene, and mounting with Permount (Fisher). An antibody against BMPR2 (Ab78422, Abcam) was used. The brain layers were categorized as upper (I and III), middle (III to V), and lower (V and VI) layers. To quantify the amount of protein in each tissue sample, the percentage of the stained area inside the cell was measured using ImageJ.

Statistical analysis

Statistical analysis was performed using the Prism 5.01 GraphPad package and reviewed by M. Nojima at the University of Tokyo. Statistical test and significance are denoted in the figure legends.

SUPPLEMENTARY MATERIALS

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- Fig. S1. Translational regulation of BMPR2 through the mRNA sequence encoding the CTD. Fig. S2. FMRP binds BMPR2- CTD_{seq} and suppresses translation.
- Fig. S3. BMP4-SMAD1/5 signaling is increased, but TGFβ-SMAD2/3 signaling is not altered in *FMR1*-null cells.
- Fig. S4. LIMK-i and LDN effectively inhibit LIMK1 and BMPR1 kinase activity in N1E cells. Fig. S5. In vivo administration of LIMK-i inhibits phosphorylation of cofilin in mouse brain. Table S1. qRT-PCR primers.

Table S2. RIP PCR primers.

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Acknowledgments: We would like to dedicate this article to K. D. Bloch who passed in 2014. We thank K. Kawamura, A. Mundkur, and S. Manghise for technical supports. We also thank L. Jan for sharing *FMR1*-null mice and M. Nojima for reviewing the statistical analysis. Finally, we thank D. Hart, G. Davis, R. Edwards, and all members of the Hata Laboratory for the critical discussion and reading of the manuscript. **Funding:** R.K. is a recipient of a fellow-ship from Japan Society for the Promotion of Science. This work was supported by grants from the NIH (HL093154 and HL108317) and Foundation LeDucq to A.H., and from the NIH (HD040661) to P.J.H. Author contributions: R.K., S.R., S.K., J.L., Y.L., P.L., V.M.-C., and J.A.-T. designed and performed the experiments and interpreted the data. R.K., K.D.B., T.B.K., M.A., P.J.H., R.H., and G.L. designed the experiments, interpreted the data, and edited the manuscript. A.H. conceived the project, designed the experiments, interpreted the data, and wrote the manuscript. Competing interests: The authors declare that they have no competing interests. Data and materials availability: The plasmids require a material transfer agreement from UCSF, USA.

Submitted 2 March 2016 Accepted 10 May 2016 Final Publication 7 June 2016 10.1126/scisignal.aaf6060

Citation: R. Kashima, S. Roy, M. Ascano, V. Martinez-Cerdeno, J. Ariza-Torres, S. Kim, J. Louie, Y. Lu, P. Leyton, K. D. Bloch, T. B. Kornberg, P. J. Hagerman, R. Hagerman, G. Lagna, A. Hata, Augmented noncanonical BMP type II receptor signaling mediates the synaptic abnormality of fragile X syndrome. *Sci. Signal.* **9**, ra58 (2016).

Augmented noncanonical BMP type II receptor signaling mediates the synaptic abnormality of fragile X syndrome Risa Kashima, Sougata Roy, Manuel Ascano, Veronica Martinez-Cerdeno, Jeanelle Ariza-Torres, Sunghwan Kim, Justin Louie, Yao Lu, Patricio Leyton, Kenneth D. Bloch, Thomas B. Kornberg, Paul J. Hagerman, Randi Hagerman, Giorgio Lagna and Akiko Hata (June 7, 2016) *Science Signaling* 9 (431), ra58. [doi: 10.1126/scisignal.aaf6060]

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