Title
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Permalink
https://escholarship.org/uc/item/8p65282b

Journal
PloS one, 10(2)

ISSN
1932-6203

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Publication Date
2015

DOI
10.1371/journal.pone.0117266

Peer reviewed
Deletion of Fmr1 Alters Function and Synaptic Inputs in the Auditory Brainstem

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Abstract

Fragile X Syndrome (FXS), a neurodevelopmental disorder, is the most prevalent single-gene cause of autism spectrum disorder. Autism has been associated with impaired auditory processing, abnormalities in the auditory brainstem response (ABR), and reduced cell number and size in the auditory brainstem nuclei. FXS is characterized by elevated cortical responses to sound stimuli, with some evidence for aberrant ABRs. Here, we assessed ABRs and auditory brainstem anatomy in Fmr1-/- mice, an animal model of FXS. We found that Fmr1-/- mice showed elevated response thresholds to both click and tone stimuli. Amplitudes of ABR responses were reduced in Fmr1-/- mice for early peaks of the ABR. The growth of the peak I response with sound intensity was less steep in mutants that in wild type mice. In contrast, amplitudes and response growth in peaks IV and V did not differ between these groups. We did not observe differences in peak latencies or in interpeak latencies. Cell size was reduced in Fmr1-/- mice in the ventral cochlear nucleus (VCN) and in the medial nucleus of the trapezoid body (MNTB). We quantified levels of inhibitory and excitatory synaptic inputs in these nuclei using markers for presynaptic proteins. We measured VGAT and VGLUT immunolabeling in VCN, MNTB, and the lateral superior olive (LSO). VGAT expression in MNTB was significantly greater in the Fmr1-/- mouse than in wild type mice. Together, these observations demonstrate that FXS affects peripheral and central aspects of hearing and alters the balance of excitation and inhibition in the auditory brainstem.

INTRODUCTION

Fragile X syndrome (FXS) is the most common single-gene inherited form of autism. FXS results from an expansion of the CGG repeats in the promoter region of the FMR1 gene, which reduces the amount of fragile X mental retardation protein (FMRP) produced. FMRP acts as a modulator of mRNA translation and has numerous target genes [1, 2]; its loss results in unregulated production of synaptic proteins [3]. Individuals with FXS display cognitive impairments, hyperactivity, seizures, aberrant dendritic spine morphology and several autism-related symptoms [4–8]. FXS subjects also display a general enhancement of response to sensory stimuli. When presented with an array of stimuli that spanned several sensory modalities, individuals with FXS...
show elevated electrodermal responses across all types of stimuli [9, 10]. Notably, there is a heightened response to auditory stimuli. When subjects with FXS were presented with sound stimuli as part of an oddball discrimination task, cortical responses were consistently elevated [11–15]. Whether heightened cortical response in FXS is a phenomenon unique to the cortex or results from dysfunction of other auditory brain structures is unknown. However, electrophysiological and anatomical evidence suggest impaired auditory brainstem function. Auditory brainstem responses (ABRs) performed on individuals with FXS have reported longer absolute peak latencies and inter-peak intervals [16–19]. Of particular interest, prolonged I-III interpeak interval (Ferri et al., 1986) and III-V interpeak interval (Arinami et al., 1988) have been reported in FXS. Peak III results from the activation of the superior olivary nuclei in response to sound; therefore altered peak III latency may suggest some dysfunction of the superior olivary complex in FXS patients. Other studies attribute ABR anomalies in FXS to impaired peripheral auditory processing [20], or sedatives which may have been used during the testing process [21].

In addition to these physiological observations, molecular and anatomical changes have been associated with the auditory brainstem in FXS. Aberrations in brainstem anatomy have been found in the post mortem tissue of subjects with FXS and autism [22, 23]. A survey of autistic men and boys revealed reduced cell number in medial superior olive (MSO) of several subjects. Autistic MSO cell bodies were smaller and more variable in their orientation than in controls. Of note, an individual included in this study who was diagnosed with FXS and autism also displayed the MSO anomalies found in his autistic counterparts [22]. Other work examining the auditory brainstem of autistic individuals found a 77% decrease in the number of MSO neurons, a 67% decrease in the number of lateral superior olive (LSO) neurons, and a 45% decrease in the number of medial nucleus of the trapezoid body (MNTB) neurons as well as altered composition of neuronal populations within nuclei [24]. Within the MNTB, the Kv3.1b potassium channel is typically expressed in a tonotopic gradient. In FXS, the Kv3.1b gradient is lost, leading to improper coding of sound stimuli and impaired sensory modulation in the MNTB [25]. Additionally, FMRP is present in 83% of MSO cells and has been found to concentrate at dendritic branch points of MSO cells across species [26, 27]. Excessive dendritic branching and impaired dendritic pruning have been reported in animal models of FXS [28, 29].

In this study, we examined ABRs and quantified excitatory and inhibitory inputs to auditory brainstem nuclei in the Fmr1−/− (KO) mouse, an animal model of FXS. Similar to individuals with FXS, Fmr1−/− mice have aberrations in dendritic spine number and morphology, hyperactivity, audiogenic seizures, repetitive behavior, communication deficits, and difficulties with social interactions, making them a useful model for studying FXS-related dysfunction [30–33]. Our results suggest that loss of Fmr1 causes reduction in the ABR response, with both central and peripheral components. Fmr1−/− mice displayed elevated ABR thresholds and showed reduced growth of responses with stimulus intensity compared to WT animals. We found that VCN and MNTB cell size was decreased in Fmr1−/− mice compared to wild type animals, and MNTB showed increased expression of vesicular GABA transporter protein (VGAT) relative to vesicular glutamate transporter protein (VGLUT) expression within the MNTB.

**MATERIALS AND METHODS**

**Animals**

Adult male Fmr1−/− mice and wild type littermates on the FVB strain were used for ABR measurements and anatomical studies. All procedures were approved by the University of California, Irvine Institutional Animal Care and Use Committee.
ABR Recordings

ABRs were performed inside a 102cm x 98cm x 81cm sound-attenuating chamber (Industrial Acoustics Company, NY). Before recording ABR responses, the ABR system was calibrated under computer control from 5kHz to 80kHz in 5Hz steps using a 0.5-inch condenser microphone model no. 4134 (Bruel & Kjaer). Mice were anesthetized with a solution of ketamine (75–85mg/kg) and xylazine (0.1–0.5 mg/kg) and placed on a Kopf Model 900 Small Animal stereotaxic instrument (Tujunga, CA). Body temperature was monitored and maintained with a custom built heating pad. Sound was presented using a Tucker-Davis Technologies (TDT) MF1 Multi-Function Speaker (Alachua, FL) through an ear tube placed in animals’ left ear. Pin electrodes were placed subdermally to probe brainstem responses. The positive (active) electrode was placed at the vertex, the negative (reference) electrode was placed in the right cheek, and a ground electrode was placed at the base of the tail. The pin electrodes were connected to a TDT RA4PA 4-channel Medusa preamplifier, which connected to a TDT RA16 Medusa Base Station.

Sound stimuli were generated using Tucker-Davis Technologies SigGen software and presented under the control of the TDT BioSig platform. Presentation of sound stimuli was driven by a TDT RP2.1 enhanced real time processor and signal level was controlled using a TDT PA5 programmable attenuator. Sound was amplified by a TDT SA1 stereo power amp. All sounds were presented 500 times at a rate of 29 stimuli per second. Mice were first presented with 100 μs click stimuli between 80 and 10 dB SPL, decreasing in 5 dB SPL steps. Next, mice were presented with 8, 12, 16, 24, and 32 kHz tone stimuli. Each tone was 3ms long and sound intensity was reduced in 5 dB SPL intervals, beginning at 80 dB SPL and ending at 10 dB SPL.

ABR Analysis

ABR traces were analyzed for threshold, peak latency, peak amplitude, and inter-peak interval in response to click and tone stimuli. Threshold was defined as the lowest intensity at which any peaks could be discerned. Peak latency was the time from the onset of a stimulus to the time at which each peak reached its apex. Peak amplitude was measured as the voltage at the apex of each peak relative to the noise floor. The inter-peak interval was the time between the apex of one peak and the apex of another peak.

To test for genotype differences in peak latency and peak amplitude, latency and amplitude data were collected for peaks I, II, III, and IV. We analyzed input-output functions to compare WT and KO animals in their response amplitudes (μV) and latencies (msec) as sound intensity (dB SPL) increased. We performed two-way repeated measure ANOVA (JMP 9.0, SAS Institute, Cary, NC) on each peak to determine the effect of genotype and sound intensity and to identify interactions between these variables.

Nissl Staining and Analysis

Deeply anesthetized animals were perfused with 0.9% saline followed by 4% paraformaldehyde in phosphate buffered saline (PBS) before brainstems were harvested. Dissected brainstem specimens were post-fixed with 4% paraformaldehyde for at least 2 hours, incubated overnight in a 30% sucrose solution in PBS, and sectioned in the coronal plane at 16 μm on a cryostat. Sections were collected on chrome-alum coated glass slides in a 1-in-10 series and dried on a 37°C slide warmer. Mounted sections from one set of alternate sections were stained with 1% thionin and coverslipped. The ventral cochlear nucleus (VCN), MNTB, and LSO were digitally imaged on a Zeiss Axioskop2 microscope using Axiovision acquisition software.

To determine cell density in the VCN and LSO, a 9530 μm² outline was placed over a portion of a VCN or LSO image in a minimum of three sections per brain. ImageJ cell counter was
used to count all cells within the outline. The number of cells was divided by the area of the outline to find cell density. Cell density in MNTB was found by placing a 14405 μm² outline over the medial MNTB and lateral MNTB. The number of cells in both the medial and lateral outlines were counted using ImageJ cell counter and divided by the area of the outline to find cell density. Cell densities were found in both the left and right VCN and MNTB. Left and right values were averaged to produce final VCN, medial MNTB, and lateral MNTB cell densities.

To determine cell area, outlines of Nissl labeled cells in each nucleus were analyzed in ImageJ. At least 10 cells were measured in each section, using a minimum of three sections distributed through the rostrocaudal extent of the nucleus. Cells were selected for measurement if the nucleus was discernable within the section. Cross sectional area was recorded and cell areas were averaged to obtain a single value for each nucleus for each brain.

**VGLUT and VGAT Immunostaining**

Cryosectioned brainstems were used for immunofluorescence. Mounted sections were surrounded with a hydrophobic Pap pen barrier and rinsed. For antigen retrieval we placed slides in 0.1% sodium dodecyl sulfate in PBS for 5 minutes. Slides were then rinsed with PBS and incubated at room temperature in a humid chamber with blocking solution (4% bovine serum albumin, 0.4% Triton X in PBS) for 1 hour. Slides were then incubated with primary antibodies diluted in blocking solution. We used a rabbit polyclonal antibody that recognizes VGAT (Phosphosolutions, Aurora, CO) at 1:200 and a guinea pig polyclonal antibody that recognizes VGLUT2 (Millipore, Temecula, CA) at 1:2500. Slides were washed in PBS then incubated at room temperature for 2 hours in AlexaFluor (Invitrogen, Carlsbad, CA) secondary antibodies (goat anti-guinea pig Alexa 594 and goat anti-rabbit Alexa 488) diluted 1:500 in blocking solution. Slides were rinsed in PBS, and coverslipped using ProLong Gold antifade reagent with DAPI (LifeTechnologies, Eugene, OR) as a mounting medium.

**Imaging and Analysis**

Slides were viewed on a Zeiss AxioSkop-2 epifluorescence microscope. Images of regions containing the VCN, LSO and MNTB were processed using ImageJ software. For this analysis we included at least three sections spanning the rostrocaudal extent of each nucleus. We outlined the nuclei and found the total area immunopositive for VGLUT and VGAT. Using ImageJ, images were separated by red (VGLUT), green (VGAT), and blue (DAPI) color-channels. The thresholds on the red and green channels were adjusted to highlight the stained portions of the image on each channel. The ImageJ Analyze Particles function was used to summate all of the VGLUT- or VGAT-stained objects of an image. We defined an index (ISP) to characterize relative levels of each synaptic protein for each nucleus: $ISP = \frac{(VGLUT-VGAT)}{(VGLUT+VGAT)}$

Values of $ISP$ range between -1 and 1. Positive values of $ISP$ indicate more excitatory puncta than inhibitory puncta, and negative values would indicate more inhibition. Absolute values of $ISP$ reflect differences in the relative levels of these two proteins. To determine statistical significance for anatomical data, we performed one-way ANOVA with post-hoc two-tailed t-tests using Bonferroni correction to adjust for multiple comparisons.

**RESULTS**

**ABR measurements**

A total of 25 Fmr1 -/- mice and 20 wild type mice were used for ABR measurements. Click or pure tone stimuli were presented at intensities beginning at 80 dB SPL decreasing in 5 dB SPL...
steps to 10 dB SPL (Fig. 1A). Traces generated in response to both types of stimuli were analyzed for threshold, peak amplitude, absolute peak latency, and inter-peak latency. Peak V could be identified after click stimulation but not in most traces generated using tone stimuli.

**ABR thresholds are elevated in KO mice**

The mean threshold for the ABR in KO mice was 27.5 ± 1.88 dB SPL (± SEM, n = 24), which was significantly higher than those of WT mice when tested with clicks (18.5 ± 2.06 dB SPL (n = 20; two-tailed t-test, p < 0.0024; Fig. 1B). KO mice also had elevated thresholds compared to controls at each of the five pure tone stimuli tested (Fig. 1C; Table 1), with significance ascertained after Bonferroni correction.
ABR amplitude

To assess how peak amplitude may change with stimulus intensity, we generated input-output functions for each peak, where the mean peak amplitude was plotted against stimulus intensity (Fig. 2). Peak latency and amplitude were analyzed for each peak for intensities at 40, 50, 60, 70, and 80 dB SPL following click stimuli. For peak I, two-way repeated measure ANOVA revealed a significant effect of genotype on peak amplitude ($F_{1,40} = 5.94, p = 0.0193$) and also showed a significant interaction between genotype and sound intensity ($F_{4,37} = 3.98, p = 0.0087$). The peak I responses in Fmr1-/- mice were smaller than those of wild type mice and increased more slowly with sound intensity. We next tested whether this effect was evident using pure tone stimuli. Genotype significantly reduced peak I responses to stimuli at 8 kHz ($F_{1,39} = 6.76, p < 0.0131$), 12 kHz ($F_{1,34} = 11.363, p < 0.0019$), and 16kHz ($F_{1,25} = 10.88, p < 0.0029$). The interaction between genotype and intensity was also seen for 12 kHz stimuli ($F_{4,31} = 2.79, p < 0.0433$); $p$ values for all comparisons are shown in S1 Table. The effects for tone stimuli further support effect of Fmr1 deletion on decreased peak I amplitude and gain.

Analysis of peak II following click stimuli did not show an effect of genotype on peak response ($F_{1,40} = 4.0428, p = 0.0511$) and no interaction between genotype and sound intensity ($F_{4,37} = 0.8590, p = 0.498$). Consistent with this observation, effect of genotype did not reach significance for any of the tone frequencies presented (S1 Table). Peak III amplitude was significantly reduced in KO mice when click tones were presented ($F_{1,33} = 9.6590, p = 0.0039$) and when 8 kHz tone stimuli were used ($F_{1,17} = 8.0925, p = 0.0112$), but not at other frequencies. The interaction between genotype and sound intensity did not reach significance for peak III amplitude. We did not see effects of genotype on peak amplitude for click stimuli for peak IV or peak V, and no effect of interaction between genotype and sound intensity was seen (S1 Table).

ABR latency

We examined how peak latency varied with stimulus intensity in wild type and Fmr1-/- mice. Two-way repeated measure ANOVAs revealed no main effect of genotype on peak latency for any peak using click stimuli. Responses to tone stimuli were largely similar. Significant effects were seen only in two cases: 16 kHz stimulation at peak II and 12 kHz stimulation at peak III (S1 Table), and interpeak measurements of latency did not show main effects of genotype for any stimulus.

Histological analysis of auditory brainstem nuclei

To probe anatomical anomalies potentially related to central components of the ABR changes seen in Fmr1-/- mice, we performed histological analysis on Nissl stained brainstem sections. The VCN, MNTB, and LSO of 7 wild type and 7 Fmr1-/- mice were analyzed for cell density
and cell area (Fig. 3I). We found no significant differences in cell density within the VCN (WT = 0.004 ± 0.00016 cells/μm², KO = 0.004 ± 0.00029, cells/μm²; p = 0.704; Fig. 3B), MNTB (WT = 0.0063 ± 0.00033 cells/μm², KO = 0.0069 ± 0.001 cells/μm²; p = 0.2494; Fig. 3E), or LSO (WT = 0.019 ± 0.001 cells/μm², KO = 0.019 ± 0.001 cells/μm²; p = 0.8872; Fig. 3H).
Cell area measurements showed significant reduction in cell size in VCN and MNTB, but not in LSO. In VCN (Fig. 3C) the mean cell area for wild type mice was $222 \pm 8.7 \, \mu m^2$ ($n = 8$) while the area for $Fmr1^{-/-}$ mice was $183 \pm 5.8 \, \mu m^2$ ($n = 9$; two-tailed t-test with Bonferroni correction, $p = 0.0017$). In MNTB (Fig. 3F) the mean cell area for wild type mice ($223 \pm 13 \, \mu m^2$, $n = 9$) was greater than that for $Fmr1^{-/-}$ mice ($172 \pm 6.6 \, \mu m^2$, $n = 7$; $p = 0.002$). In LSO (Fig. 3I), wild type cell area ($214 \pm 15 \, \mu m^2$, $n = 8$) was similar to that seen in $Fmr1^{-/-}$ mice ($182 \pm 12 \, \mu m^2$, $n = 8$; $p = 0.102$).

Inhibitory and excitatory synaptic proteins in the auditory brainstem nuclei

To estimate relative levels of synaptic input in the auditory brainstem, we performed double immunofluorescence for VGLUT and VGAT. We quantified the area of VGLUT-positive and VGAT-positive inputs to cells in the VCN, MNTB, and LSO. Regions of VCN (Fig. 4A–B), MNTB (Fig. 4C–D), and LSO (Fig. 4E–F) were selected, and thresholds adjusted such that the total area of VGLUT or VGAT within a region was highlighted, and the label density, or fractional areal coverage, was computed by dividing the area with label by the area of the sampled region.
In VCN the fractional coverage of VGLUT did not differ between wild type mice (0.22 ± 0.00074, n = 8) and Fmr1-/- mice (0.025 ± 0.00116, n = 10; p = 0.0913). Similarly, VGAT did not differ between wild type mice (0.023 ± 0.0007, n = 8) and Fmr1-/- mice (0.024 ± 0.001, n = 10; p = 0.3919) in VCN (Fig. 5A). In MNTB VGLUT density was similar in wild type mice (0.217 ± 0.01, n = 9) and Fmr1-/- mice (0.19 ± 0.019, n = 11; p = 0.3559). However, MNTB showed a significant increase of VGAT in Fmr1-/- mice (0.126 ± 0.013, n = 11) compared to wild type controls (0.072 ± 0.008, n = 9; p = 0.0034) (Fig. 5B). In LSO, the increase in VGLUT and VGAT density in Fmr1-/- mice was not significant after Bonferroni correction (VGLUT: WT, 0.023 ± 0.0036, n = 7; KO 0.036 ± 0.0034, n = 9; p = 0.0176; VGAT: WT, 0.024 ± 0.0040, n = 7; KO, 0.041 ± 0.005, n = 9; p = 0.0272; Fig. 5C).

Thresholded VGLUT and VGAT regions were then summed and an index ISP comparing VGLUT to VGAT within each region tested was calculated (Fig. 5D). No significant differences were seen in VCN (WT = -0.022 ± 0.023, n = 8; KO = 0.009 ± 0.012, n = 10, p = 0.2575) or LSO (WT = -0.024 ± 0.036, n = 7, KO = -0.047 ± 0.033, p = 0.6444). In contrast, MNTB in Fmr1-/- mice revealed a reduced ISP relative to control (WT = 0.51 ± 0.031, n = 9; KO = 0.19 ± 0.06, n = 11; p = 0.0003), suggesting greater VGAT expression relative to VGLUT expression than in wild type animals.

DISCUSSION

Individuals with FXS demonstrate a variety of difficulties in auditory processing [11–13, 15, 16, 18, 34]. The ABR is a non-invasive test that reliably describes auditory brainstem function. Given their ease of use, ABRs may also serve as a metric for gauging the efficacy of possible treatments or therapies for auditory processing disorders, such as those seen in FXS. Fmr1-/- mice replicate many symptoms of FXS, including auditory processing dysfunction within the cortex [35–37], and might therefore serve as model for FXS-related anomalies in auditory
processing. In this study, we performed ABR tests on Fmr1−/− mice, examined cell distribution in the auditory brainstem nuclei, and quantified excitatory and inhibitory inputs in these nuclei.

In the ABR measurements we found heightened response thresholds in Fmr1−/− mice compared to wild type mice in response to both click and tone stimuli. Peak I amplitudes were reduced in Fmr1−/− mice, and gain with respect to stimulus intensity was reduced compared to wild types. These observations suggest a loss of auditory function originating in the periphery. In addition, we found that peak III amplitude was also reduced in Fmr1−/− mice, possibly reflecting the observed decrease in peripheral input. This effect may also signify impairment in brainstem function, as peak III is thought to reflect activity in the superior olivary complex [38–40]. In contrast to ABR studies performed on human participants with FXS, we did not find evidence for differences in peak latency between wild type and in Fmr1−/− mice. A lack of latency effect may be attributable to the reduced size of the mouse pathway and/or to differences between the species in their auditory processing or in their response to mutations in Fmr1.

Consistent with a central auditory defect, we found that Fmr1−/− mice showed reduced cell area in VCN and MNTB. Reduced cell size in auditory nuclei has been reported in human anatomical studies [22–24] and FMRP protein expression has been demonstrated in the human auditory brainstem nuclei [27]. In addition, we showed that VGAT expression was increased in MNTB in Fmr1−/− mice, suggesting that cells in this nucleus receive more inhibitory inputs.
than in wild type animals. An increase in VGAT relative to VGLUT expression was seen in MNTB but not in VCN or LSO. Enhancement of inhibition in MNTB is potentially related to the observed reduction in ABR amplitude for peak III. Because MNTB is a sign-reversing relay nucleus, an interesting possibility is that MNTB cells receive less synaptic drive and thus provide less inhibition at subsequent stages of processing.

Individuals with FXS demonstrate generally exaggerated responses to sensory stimuli [9] and unusual cortical activity in response to sound [11–13, 15, 34]. In addition to increased responses to sensory stimuli, findings from pre-pulse inhibition testing show both individuals with FXS and Fmr1+/– mice experience deficits in sensorimotor gating [41, 42]. Children with FXS experience corticotemporal seizures [43]. These observations, together with reports that Fmr1+/– mice experience audiogenic seizures and produce a greater cortical response to sound [35], suggests greater excitability in the auditory pathways.

While increased excitability might arise from early auditory regions, cochlear hearing loss has been shown to elicit central increases in gain, which may lead to hyperacusis [44, 45]. In Fmr1+/– mice, hyperacusis might thus originate at the level of the superior olivary complex or at later processing areas, and might develop as a consequence of gain increases that compensate for the reduced input. Similarly, audiogenic seizures likely originate at higher processing levels. Previous studies of audiogenic seizures in Fmr1+/– mice have demonstrated the involvement of dorsal nucleus of the lateral lemniscus and posterior intralaminar nucleus of the thalamus [30]. The lateral lemniscus and inferior colliculus comprise the fourth and fifth ABR peaks, respectively [38, 40], which no longer showed amplitude decreases in Fmr1+/– mice in the present study.

Increased excitability is thought to be a major factor in the symptoms presented in FXS [46–49], and modulation of the GABAergic system remains a target for therapeutic treatment [50]. Nevertheless, excessive excitation is not ubiquitous in all brain regions. Notably, several studies report enhanced long term depression in the hippocampus of Fmr1+/– mice [3, 51–53]. Overactive mGluR1/5 coupled with a loss of FMRP regulation initiates a biochemical cascade that results in accelerated removal of AMPA receptors from the postsynaptic membrane.

Evidence of alterations in GABAergic input in Fmr1+/– mice have also been reported in the striatum, cortex, brainstem, and diencephalon [54]. Normal glutamatergic synaptic transmission was found within the striatum of Fmr1+/– mice, but neurotransmitter release was elevated at GABAergic terminals, resulting in an increase in sIPSCs and mIPSCs. Increased expression of GAD65 and GAD67 was found in the cortex, hippocampus, brainstem, and diencephalon, suggesting excessive GABA production in those brain regions. However, areas demonstrating increased GAD65 and GAD67 also show a decrease in the GABAAR subunit β, which is required to form functional GABAARs.

The MNTB is a major source of inhibitory input to the superior olivary complex. The ventral cochlear nucleus sends glutamatergic input to the MNTB through the calyces of Held, and the MNTB in turn delivers glycinergic input to the LSO [55, 56]. The MNTB supplies essential inhibitory input for processing differences in sound intensity and timing. Notably, MNTB function is known to be impaired in Fmr1+/– mice. Kv3.1 voltage-dependent potassium channels allow extremely rapid repolarization of neurons following action potentials, and therefore enable neurons to produce many action potentials very quickly. Kv3.1 is typically distributed in a gradient within MNTB that follows the tonotopic gradient. In Fmr1+/– mice, Kv3.1 is not expressed along a gradient; rather expression is uniform throughout MNTB. The loss of Kv3.1 expression gradient results in an inability to follow high frequency stimulus [25]. The sodium-activated potassium channel Slack is also highly expressed in MNTB [57], where it improves temporal fidelity. FMRP binds the cytoplasmic C-terminal of Slack, and can stimulate Slack channel activity. Loss of FMRP results in greatly reduced potassium current and may thus
impair the temporal precision of action potentials [57]. Coupled with our discovery of increased VGAT expression within MNTB, these findings may explain a decrease in synchrony of population activity, which may manifest as an elevation in ABR threshold.

In addition to imbalances in inhibitory and excitatory input, individuals with FXS also display changes in white and gray matter that may affect the degree and timing of population responses within different nuclei. FXS patients consistently exhibit increases in caudate nucleus, thalamus, and parietal lobe volume [58–61]. Temporal lobe white matter and frontostriatal tracts are also more extensive in FXS [61, 62]. Of particular interest, both total brainstem volume and brainstem white matter volume are enhanced in FXS individuals [59]. FXS is associated with widespread effects on cell size, cell number, fiber tract volume, and fiber density. In the auditory brainstem, alterations in cell and dendritic morphology (Kulesza et al., 2010; Beebe et al., 2014) suggest substrates for the effects of FMRP reduction on auditory function in individuals with FXS.

Our results show elevated ABR thresholds, decreased peak I and III amplitudes, and enhanced inhibition to MNTB in a mouse model of FXS. These observations provide evidence that auditory brainstem function is altered by inactivation of Fmr1. While central deficits may arise as a consequence of peripheral deficits, FXS may produce several independent effects. Given the diversity of FMRP targets, including those known to occur in the brainstem, our observation of both peripheral and central auditory effects in Fmr1−/− mice suggests that the mutation acts at numerous points along the auditory pathway.

Supporting Information
S1 Table. P values for effects on ABR input-output functions.

Acknowledgments
The authors are grateful to Dr. Tom Lu for technical assistance and to Dr. Arnold Starr for valuable advice on ABR data acquisition and analysis.

Author Contributions
Conceived and designed the experiments: SER KSC. Performed the experiments: SER SM KSC. Analyzed the data: SER SM KSC. Contributed reagents/materials/analysis tools: SER SM KSC. Wrote the paper: SER SM KSC.

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