Neurofibromin regulates corticostriatal inhibitory networks during working memory performance

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Neurofibromatosis type I (NF1) is one of the most common single-gene causes of learning disabilities. Here, we use behavioral working memory probes and electrophysiological studies in a mouse model of NF1 (Nf1 heterozygous null mutants; Nf1+/−) to demonstrate that (i) Neurofibromin regulates prefrontal and striatal inhibitory networks, specifically activity-dependent GABA release and (ii) is required for working memory performance, with inhibition-dependent working memory deficits seen in Nf1+/− mice. We find that increased inhibition in medial prefrontal cortex (mPFC) is sufficient to alter persistent activity in a biophysical model of an mPFC microcircuit, suggesting a possible mechanism for Nf1+/−-working memory deficits. Accordingly, working memory assays applied during functional MRI (fMRI) studies in human subjects with NF1 reveal hypoactivation of corticostriatal networks, which is associated with impaired working memory performance. Collectively, these integrative mouse and human studies reveal molecular and cellular mechanisms contributing to working memory deficits in NF1.

NF1 Regulates Activity-Dependent GABA Release in Prefrontal Cortex.

The Nf1+/− mutation was previously shown to alter GABAergic inhibition in the hippocampus in mice (6, 7). Because prefrontal dysfunction could account for the prominent executive function deficits associated with NF1, we studied inhibition in mPFC of Nf1+/− mice. We focused on this area because of its homology to human DLPFC and its key role in rodent working memory tasks (14, 15). We used whole-cell patch clamp to record spontaneous synaptic activity induced by local inhibitory and excitatory networks in single neurons within layer II/III in acute slices from adult mice. Our results show that in the mPFC of Nf1+/− mice, the frequency of spontaneous inhibitory postsynaptic currents (sIPSCs) is significantly higher (P = 0.02) than in WT littermate controls; this increase in frequency was observed across all sIPSC amplitudes (P < 0.001; Fig. 1A). Also, analysis of the cumulative distribution of the intervals between sIPSCs confirmed an increased frequency in Nf1+/− mice (P < 0.001; Fig. S1A). In contrast, the frequency of spontaneous excitatory postsynaptic currents (sEPSCs) recorded in the same neurons is not altered in Nf1+/− mice (P = 0.69; Fig. 1B). Importantly, the Nf1+/− mutation did not affect the kinetics of either sIPSCs or sEPSCs (Fig. S1B) in the mPFC. Because both sIPSCs and sEPSCs were recorded in the same neurons, these data also demonstrate that the ratio of excitatory to inhibitory drive onto those neurons is abnormally low in the Nf1+/− mice.

We next examined causes of increased inhibition in the Nf1+/− mice. Specifically, we asked whether the increased inhibition in the Nf1+/− mice is caused by a cell autonomous effect of NF1 deletion in interneurons. To assess this directly, we used the Cre/loxP system to create Dlx-Creox/+, Nf1+/− mice with a heterozygous deletion of Nfi restricted to inhibitory interneurons in the prefrontal cortex (7, 16). Dlx-Cre+/+, Nf1+/− mice show an increased frequency of sIPSCs (P = 0.006; Fig. 1C) compared with control NF11 revealed decreased neural activity in frontostriatal networks in NF1 patients that correlated with their degree of working memory impairment. These convergent cross-species findings support the hypothesis that working memory deficits associated with the NF1 mutation in both mice and humans are caused by loss of neurofibromin regulation of frontal corticostriatal inhibition.

Results

GABA | Ras | prefrontal cortex | learning disability | neurofibromatosis type I

Neurofibromatosis type 1 (NF1) is a valuable model for understanding mechanisms of learning disabilities (1). NF1 is a common genetic disorder (incidence 1:3,000) that results from mutations in a single gene (Nf1) that encodes the neurofibromin protein (2, 3). Specific deficits in the domains of visuospatial and executive functions are among the most common cognitive deficits associated with this syndrome (1, 4, 5). Previous mechanistic studies in a mouse model of NF1 (Nf1 heterozygous null mutants or Nf1+/−) demonstrated that neurofibromin modulates Ras-dependent GABA release in the hippocampus, which in turn modulates long-term potentiation (LTP) and hippocampal-dependent learning (6, 7). However, the mechanisms underlying frontal executive dysfunction in NF1, including prominent working memory deficits (5), are unknown. Therefore, to investigate mechanisms underlying working memory deficits associated with the NF1 mutation, we carried out parallel experiments in mice and humans.

Working memory is a cognitive construct involving the ability to hold and update information transiently in mind in the service of higher-order cognitive activities. Executive functions, including working memory, are thought to depend on common corticostriatal networks (6–11). Therefore, our experiments focused on frontal corticostriatal circuitry, with an emphasis on the dorsolateral prefrontal cortex (DLPFC) in humans, thought to be critical for working memory (12), and its functionally analogous structure in rodents, the medial prefrontal cortex (mPFC) (13, 14).

Here, we report Ras-dependent increases in GABA release in the mPFC and striatum of the Nf1+/− mouse model. Increased GABAergic inhibition is likely to be responsible for the working memory deficits that we found in the Nf1+/− mice because these deficits could be reversed with a drug that decreased inhibition. Further, functional MRI (fMRI) studies in human subjects with NF1 revealed decreased neural activity in frontostriatal networks in NF1 patients that correlated with their degree of working memory impairment. These convergent cross-species findings support the hypothesis that working memory deficits associated with the NF1 mutation in both mice and humans are caused by loss of neurofibromin regulation of frontal corticostriatal inhibition.
groups, demonstrating that Nf1 regulates inhibitory release from mPFC interneurons. This action of Nf1 within interneurons is confirmed pharmacologically: in the presence of CNQX (10 μM) and APV (50 μM), which blocks excitatory drive, Nf1+/− mice still showed a higher frequency of sIPSCs (P = 0.009; Fig. 1D) in mPFC. It is important to note that this difference is driven by the more frequent small amplitude sIPSCs (Fig. 1 C and D); large amplitude sIPSCs are comparatively rare, and therefore we could not determine whether their frequency is also different between mutants and controls.

We next examined how the Nf1+/− mutation affected interneuron function. It is possible that the increase in sIPSCs frequency is caused by activity-dependent changes, such as excitability. To address this possibility, we blocked action potentials in the slice with TTX (1 μM), and recorded miniature IPSCs (mIPSCs) in layer II/III pyramidal neurons. An analysis of the cumulative distribution of the intervals between mIPSCs revealed no differences between Nf1+/− mice and WT controls (P = 1; Fig. 1C). These data demonstrate that the increased frequency of inhibitory currents observed in Nf1+/− mice is activity-dependent. Cumulative amplitude distributions of mIPSCs in mPFC were also examined in TTX, and no differences between Nf1+/− mice and WT were found (P = 0.585; Fig. 1C). In combination, these experiments demonstrate that Nf1 affects the inhibitory system most prominently by regulating activity-dependent GABAergic release in mPFC.

We next investigated molecular mechanisms by which the Nf1 mutation leads to increased activity-dependent inhibition. Specifically, we focused on the Ras signaling pathway, as Nf1 encodes a negative regulator of Ras signaling (neurofibromin) (17, 18), and increased Ras activity has been previously reported in prefrontal cortex of Nf1−/− mice (19). To study whether increased inhibition in Nf1−/− mice is Ras dependent, we decreased Ras signaling in mPFC slices from Nf1−/− mice using bath application of U0126 (10 μM). U0126 functions as a noncompetitive inhibitor of MEK, a downstream effector of Ras. A significant effect of U0126 application on sIPSC frequency (P = 0.003; Fig. 2A) was seen. Importantly, U0126 eliminated the difference in sIPSC frequency between genotypes (P = 0.553) that was observed before U0126 application (P = 0.028). U0126 also decreased sEPSC frequency (P = 0.012; Fig. 2B, Right), but to the same degree across genotypes (P = 0.357; Fig. 2B, Left). These data suggest that increased GABA release from inhibitory networks in Nf1−/− mice is Ras/MEK dependent and that neurofibromin plays an important role in inhibitory, but not excitatory, neurons, to modulate Ras-dependent neurotransmitter release.

Levels of Inhibition Seen in Nf1−/− Mice Alter Persistent Activity in a Computational Model. As other mechanisms may be altered in Nf1−/− mice, we wished to further explore the significance of neurofibromin modulation of mPFC inhibitory networks. A computational approach allowed us to manipulate inhibition in isolation within a model prefrontal microcircuit in which we incorporated the background levels of sIPSCs and sEPSCs measured in Nf1−/− and WT mice. This allowed us to ask whether inhibitory changes seen in Nf1−/− mice could on their own be sufficient to significantly affect complex network properties thought to be important for working memory. We focused on the ability of prefrontal networks to sustain persistent activity following a transient stimulus, as this property is thought to be modulated by GABA-dependent inhibition (20–22), and is critical for working memory (23, 24). When using WT values for sIPSC and sEPSC frequency, persistent activity lasting more than 3 s was induced using synchronous, single pulse stimulation with a probability of 0.84 in our model microcircuit. When the frequency of background inhibitory activity was increased to levels seen in the Nf1−/− mPFC, the probability for induction of persistent activity was decreased by 35% (Fig. S2A). These results suggest that the increased inhibition seen in Nf1−/− prefrontal networks is sufficient to significantly alter the likelihood of induction of persistent activity.
tiation of persistent activity, a network property required for working memory in nonprimate and primate studies.

**Increased Inhibitory Tone in Striatum of Nf1+/− Mice.** In addition to prefrontal contributions to working memory, a distributed network of cortical and subcortical brain structures subserves the execution of behaviors involving working memory in nonprimate and primate studies. In particular, the striatum interacts closely with the prefrontal cortex in a circuit thought to be important for working memory (25). We therefore examined whether the Nf1+/− mice show increased inhibition in the striatum, similar to that seen in the mPFC. Whole cell recordings from medium-sized spiny neurons (MSNs) in the striatum were carried out and we analyzed frequency of siPSCs, sEPSCs, and mIPSCs (Fig. 3 and Fig. S3). As in mPFC, frequency of siPSCs onto MSNs is significantly higher than that seen in WT (P < 0.001; Fig. 3A), although frequency of mIPSCs is unchanged (Fig. S3A). In contrast, the frequency of sEPSCs recorded in the same neurons, is not different between WT and Nf1+/− mice (P = 0.56; Fig. 3B). These data indicate that the striatum of Nf1+/− mice shows an activity-dependent increase in frequency of inhibitory but not excitatory events.

**Behavioral Working Memory Deficits in Nf1+/− Mice.** Next, we examined the behavioral impact of increased inhibition in prefrontal cortex and striatum in Nf1+/− mice. We started by testing the mice in a behavioral task known to depend on the corticostriatal network, the delayed win-shift radial arm maze (15, 26, 27). This task is carried out in two phases such that the location of arms entered during the training phase that were previously visited during the testing phase. Testing phase errors on the testing phase identify a differential effect of picrotoxin on Nf1+/− and WT mice across days 1–6 (P = 0.009; Fig. 4E). Importantly, the dose of picrotoxin that improved the performance of Nf1+/− mice (within-phase errors on the testing phase; P = 0.047) did not affect WT (P = 0.77). Furthermore, picrotoxin did not affect across phase errors in either genotype (P = 0.184; Fig. S4). This result demonstrates the behavioral specificity of the picrotoxin manipulation and provides evidence that the increased inhibition in Nf1+/− mice disrupts behavior.

Increased within-phase errors in the delayed win-shift radial arm maze could potentially reflect failures of working memory, although failure of other higher level executive functions could also contribute. To specifically address working memory function in Nf1+/− mice, we tested performance on an operant spatial delayed nonmatch to sample task (29–31). In this task, mice are given serial
trials, each consisting of two phases (sample presentation and choice) separated by a variable delay of 3–10 s. Both Nf1+/− and WT mice showed comparable ability to accurately respond to the cue presented in the sample phase, indicating that sensorimotor function, motivation, and response selection were normal in Nf1+/− mice. However, in the choice phase of delayed testing, the Nf1+/− mice showed an impairment (P = 0.043; Fig. 4F): they showed poor accuracy across all delays tested, although WT mice showed higher performance at 3–6 s delays, but significantly decreased accuracy (P = 0.02) as delays increased up to 10 s. The delay dependent performance of the WT mice suggests that working memory can support performance on this task successfully across delays up to 6 s, with performance decreasing as the task becomes more challenging. However, in Nf1+/− mice, working memory cannot support accurate performance, even at intervals of 3–6 s.

This working memory impairment was also improved by a low dose (0.025 mg/kg) of picrotoxin (P = 0.037; Fig. 4G). Picrotoxin significantly improved the pattern of performance across delays in Nf1+/− mice (P = 0.025) but did not affect performance of WT mice (P = 0.389). In both tasks, the dose of picrotoxin used is similar to that previously seen to improve long-term memory in WT (7). Together, these studies suggest that Nf1+/− mice show behavioral deficits that are related to increased inhibition and specifically affect working memory.

**Working Memory-Associated Corticostriatal Dysfunction in Human NF1.** Using a combination of cognitive testing and fMRI, we next determined whether human patients with NF1 also demonstrate working memory deficits related to corticostriatal dysfunction. We used two visuospatial working memory tasks that parallel the operant delayed nonmatch to sample task used with Nf1+/− mice and are known to activate a network of cortical and striatal brain structures (32, 33). Within the prefrontal cortex, fMRI analysis focused on the DLPFC, an area in primates that is functionally homologous to mPFC in rodents (13, 14, 34). Additionally, regions of interest (ROIs) in frontal eye fields, parietal cortex, and caudate were examined, as these areas are consistently recruited during working memory tasks in human and nonhuman primates (35).

The first task assesses spatial working memory maintenance and manipulation (stMNN). NF1 individuals showed significantly lower accuracy across both the spatial working memory maintenance and manipulation conditions than controls (P = 0.019; Fig. 5A). During fMRI task performance, NF1 individuals showed significantly reduced neural activity relative to controls in both cortical (DLPFC, P = 0.036; frontal eye fields, P < 0.001; parietal cortex, P < 0.001; Fig. 5C) and striatal brain regions (caudate, P = 0.006; putamen, P = 0.015; Fig. 5C). Right DLPFC activation significantly predicted stMNN performance in NF1 individuals (F1,11 = 8.69, P = 0.013; r2 = 0.3119; Fig. 6A) in a robust regression analysis.

Consistent findings were observed in the second task, a parametric probe of spatial working memory capacity (SCAP) (32). NF1 individuals showed significantly lower accuracy relative to controls overall (P = 0.03; Fig. 5B), with a steeper decline in performance accuracy as memory load increased (P = 0.016). Using a robust regression analysis, task performance was again significantly predicted by right DLPFC activation in NF1 individuals (F1,11 = 50.86, P < 0.001; r2 = 0.5398; Fig. 6B). Additionally, NF1 individuals showed reduced neural activation compared with controls in two right-lateralized cortical areas critical for maintenance of spatial information in working memory, the right frontal eye fields (P = 0.002; Fig. 5C) and the right parietal cortex (P = 0.003; Fig. 5C). In controls, task-associated deactivation was observed in the medial PFC during task performance. However, this task-associated decrease was significantly attenuated in NF1 individuals (P = 0.020; Fig. 5C).

**Fig. 5.** Hypoactivation of corticostriatal structures and working memory deficits in NF1 individuals. (A) Individuals with NF1 (n = 14) show decreased testing phase accuracy in both working memory maintenance and manipulation conditions compared with age matched controls (n = 12; RM ANOVA group × % correct P = 0.019). (B) Individuals with NF1 show decreased testing phase accuracy in the spatial capacity task across loads (RM ANOVA group × % correct P = 0.03; *P < 0.05; **P < 0.01, Tukey’s post hoc). (C) Center: Regions of interest in which activation was measured during performance of both the SCAP and stMNN. Individual graphs: BOLD signal changes during each task (*P < 0.05; **P < 0.01, RM ANOVA group × region). Error bars represent SEM.

**Fig. 6.** Degree of DLPFC hypoactivation predicts degree of performance impairment in working memory maintenance tasks in individuals with NF1. Significant association between right DLPFC activation and task performance during (A) the maintenance condition of the MNN task (F1,11 = 8.69, P = 0.013; r2 = 0.32) and (B) the SCAP task (F1,11 = 50.86, P < 0.001; r2 = 0.54).
Overall, during SCAP performance, patients with NF1 evidenced abnormal regulation of prefrontal and related cortical structures.

As predicted by our mouse studies, our human studies indicate that blunted activation of corticostriatal networks, normally recruited during spatial working memory performance, is directly related to working memory impairments characteristic of NF1.

Discussion

Here, we report convergent evidence from parallel studies in transgenic mice and human patients regarding the mechanisms underlying working memory deficits associated with NF1 mutations. A key finding in the Nf1+/− mouse experiments is that neurofibromin is an important molecular regulator of interneuronal activity in the prefrontal cortex and striatum, brain regions critical for working memory performance. The convergence between our mouse behavioral, electrophysiological, and computational modeling results indicate that the increased inhibition observed in Nf1+/− mice disrupts corticostriatal activity and contributes to working memory deficits, a conclusion supported by an extensive literature that connects inhibition to working memory. These data highlight the importance of neurofibromin signaling to function of corticostriatal inhibitory networks during behavior, specifically during working memory in the Nf1+/− mice.

Our studies in human patients reveal striking convergence with findings in Nf1+/− mice. With working memory tasks highly homologous to those used in mice, we find that loss of neurofibromin is associated with corticostriatal dysfunction and that this dysfunction is directly correlated with working memory deficits. Performing parallel experiments across species offers the unique opportunity to gain deeper insight into the basis of neural changes seen using in vivo neuroimaging studies in humans. In particular, our convergent cross-species findings suggest that hypoaivation of corticostriatal structures, observed with fMRI in NF1 patients, may be caused by increased GABA(A) receptor signaling. Similar effects have been noted in other fMRI studies of healthy subjects given GABA(A) agonists (36). By directly examining parallel effects of Nf1 mutations across species in homologous neuronal circuits and behavioral tasks, we provide evidence that neurofibromin/Ras signaling in both mice and humans regulates working memory by modulating inhibition in prefrontal cortical and striatal networks.

Interestingly, altered inhibition relative to excitation has been seen in other animal models of neurodevelopmental disorders in which corticostriatal hypoactivation and working memory deficits are also found. For example, working memory deficits have been demonstrated in Down’s syndrome (37, 38), Fragile X (39), and Turner syndrome (40). These deficits were associated with corticostriatal hypoactivation in fMRI studies of individuals with Fragile X (39) and Turner’s syndrome (40). Animal models of these disorders have identified altered regulation of inhibition. Both Rett syndrome (Mecp2 mutant mice) (41) and Down’s syndrome (Ts65Dn mice) (42) animal models show increased inhibitory drive. The Fragile X animal model (Fmr1 knockout mice) shows reduced excitatory drive onto cortical GABAergic interneurons (43) as well as increased GABAergic inhibition in striatum (44). Taken together, these studies of other neurodevelopmental disorders are consistent with our findings in NF1. Across these models, abnormal inhibition alters multiple synaptic properties, from cortical UP states in the Fragile X mice to hippocampal LTP in the Ts65Dn mice (42). Within each disorder, inhibition-dependent changes likely interact with other molecular mechanisms affected by the disease (for example, enhanced responses to mGluR in Fragile X) (45) to generate cognitive symptoms. Our findings indicate that working memory has conserved and homologous underpinnings in both mouse and humans that could be leveraged to further study the complex molecular and cellular mechanisms disrupted by genetic mutations associated with developmental learning disabilities.

Methods

Animal Subjects. Nf1+/− and WT mice, as well as Dlx-Cre;Nf1lox/lox and their control littermates, were on a hybrid background of C57BL/6NTac and 129T2/SvEmsjl Dlx-Cre;Nf1lox/lox mice and controls were from a cross between a line expressing Cre under the Dlx promoter and a line expressing a floxed Nf1 allele (46). All experiments used littermates as controls and were done with the experimenter blinded to genotype. Behavioral experiments were conducted with animals food-deprived to 85% of their starting weight. Behavior and electrophysiology was carried out on mice of the same age range (3–8 mo old).

Animal Studies. Whole-cell patch-clamp recordings were made in acute slices containing mPFC or striatum as previously described (47). Event detection was carried out off-line using MiniAnalysis software. Threshold for event detection was set above the root mean square noise level (usually 10 pA for sIPSCs, 5 pA for sEPSCs). Analysis was carried out blind to genotype.

Delayed win-shift radial arm maze task was carried out on an eight arm apparatus. Access to each arm from a central platform was regulated by a guillotine door. Reward pellets (14 mg grain based pellet; Bio-Serv) were placed at the end of each arm. Mice were habituated to the apparatus for 2 d. The task consisted of one training phase, followed by a 2-min delay, followed by a testing phase. The training phase required mice to make single entries into four baited arms. The testing phase required mice to make single entries into new arms that were not visited during the training phase. This series was run once per mouse per day for 10 d. The delayed nonmatch to sample task was adapted from procedures previously described in mice (31). During training on the nonmatch to sample rule, two WT mice failed to meet the criterion level of 70% correct responding and were subsequently dropped from further testing. Picrotoxin and saline injections were carried out in the home cage 20 min before start of the training phase.

Biophysical Model. The compartmental model of a layer III/PFC pyramidal neuron was implemented in the NEURON simulation environment (48) and consisted of five compartments. The model was validated against experimental data recorded in this study (Fig. S2 C–E). Four pyramidal model neurons were connected in a recurrent circuit that also included one interneuron. Background inhibition and excitation in the model is based on experimental data described within this paper. Persistent activity was induced by providing a synchronous, single pulse stimulation of 40 synapses on the proximal dendrite of each model pyramidal neuron. The arrangement of these synapses varied randomly during 100 repetition trials.

Human Subjects. Fourteen individuals with NF1 and 12 controls with comparable demographics (Table S1) were recruited. A standard neuropsychological battery was performed (Table S2). NF1 and control individuals were recruited via advertisement and provided written consent for participation, as approved by the institutional review board of the University of California, Los Angeles, CA. NF1 individuals had been previously diagnosed by a physician with NF1; diagnoses were confirmed by clinical interview. Controls did not have Axis-I psychiatric or neurological disorders as assessed by the Structured Clinical Interview for DSM-IV. Exclusion criteria were significant and habitual substance use in the last 6 mo, history of head injury, mental retardation (IQ less than 70), and insufficient fluency in the English language.

Human Behavioral Task Paradigms. Study participants performed two spatial delayed response tasks, one contrasting maintenance and manipulation of spatial memaranda consisting of arrays of three dots [stMMN (32, 33)] and the other assessing parametrically increasing memory loads [SCAP (33)] consisting of arrays of increasing numbers of dots. Behavioral data were analyzed in SPSS, using repeated measures ANOVA.

fMRI Methods. All scans were acquired on a 3T Siemens Allegra scanner at University of California, Los Angeles, CA. Functional slices used an echo planar (EPI) sequence (TR/TE 3000/45 ms, 90° flip angle, 333-mm slices). A T2-weighted image for anatomical registration (TR/TE 5000/33 ms, 333-mm slices with 1-mm gap, 128 × 128 matrix, 200-mm FOV) was taken in the same AC-PC aligned plane as the functional scans. SCAP consisted of 180 scans with duration of 9 min each. All stMMN and SCAP data were preprocessed using FMRIB’s Software Library v 3.3 (FSL) (49). Subject individual analyses were carried out using FMRI Expert Analysis Tool (FEAT). EPI data were corrected for motion, spatially smoothed (5-mm FWHM Gaussian

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kernel), high pass filtered (72 s) and then registered, first to the subject's individual T2-image, then to the MNI-152 brain (50, 51). Time-series statistical analysis was performed using FMRIB's Improved Linear Model (FLM). Each condition was modeled for stMNN; each load was modeled separately for SCAP, with motion included as a covariate. Group analysis was carried out using FMRIB's Local Analysis of Mixed Effects (FLAME) (49, 52).

ROI Analysis. Functionally defined ROIs were created in the MNI-152 standard brain space. Using FuzzyAtomeq (http://www.fmrib.ox.ac.uk/fsl/links/fuzzyatomeq.html) the motion corrected, smoothed, and filtered data across each ROI was probed for percent signal change from baseline. To control for performance, using FMRIB SCAP, with motion included as a covariate. Group analysis was carried out using FMRIB's Local Analysis of Mixed Effects (FLAME) (49, 52).

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