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ORIGINAL ARTICLE

A critical role for NMDA receptors in parvalbumin interneurons for gamma rhythm induction and behavior

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Synchronous recruitment of fast-spiking (FS) parvalbumin (PV) interneurons generates gamma oscillations, rhythms that emerge during performance of cognitive tasks. Administration of *N*-methyl-p-aspartate (NMDA) receptor antagonists alters gamma rhythms, and can induce cognitive as well as psychosis-like symptoms in humans. The disruption of NMDA receptor (NMDAR) signaling specifically in FS PV interneurons is therefore hypothesized to give rise to neural network dysfunction that could underlie these symptoms. To address the connection between NMDAR activity, FS PV interneurons, gamma oscillations and behavior, we generated mice lacking NMDAR neurotransmission only in PV cells (PV-Cre/NR1f/f mice). Here, we show that mutant mice exhibit enhanced baseline cortical gamma rhythms, impaired gamma rhythm induction after optogenetic drive of PV interneurons and reduced sensitivity to the effects of NMDAR antagonists on gamma oscillations and stereotypies. Mutant mice show largely normal behaviors except for selective cognitive impairments, including deficits in habituation, working memory and associative learning. Our results provide evidence for the critical role of NMDAR in PV interneurons for expression of normal gamma rhythms and specific cognitive behaviors.

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Introduction

Fast-spiking (FS) parvalbumin (PV) interneurons are gamma-aminobutyric acid (GABA)-ergic cells that express the Ca²⁺-binding protein PV and receive N-methyl-D-aspartate (NMDA)-dependent excitatory input from pyramidal cells.¹ FS–PV interneurons regulate the activity of neural networks through GABA-ergic inhibition of local excitatory neurons, and synchronous activity of FS interneurons generates gamma oscillations (30-80 Hz).²⁻⁶ Normal gamma oscillations are correlated with performance of a

variety of cognitive tasks, including the allocation of attention and working memory.⁷ Consistent with the importance of gamma oscillations, and with the more general requirement of appropriate inhibition in neural circuits, FS–PV interneurons have been proposed to underlie the cognitive disturbances associated with psychiatric disorders.⁸

The NMDA receptor (NMDAR) has also been directly implicated in the emergence of gamma rhythms, as NMDAR antagonists can disrupt⁹ or potentiate,¹⁰ gamma rhythms in slices and *in vivo*.^{11–15} This aspect of the glutamatergic system is also implicated in cognitive disturbances by the psychotomimetic effects of NMDAR antagonists,^{16,17} which recapitulate core features of schizophrenia, most notably cognitive deficits in planning, attention, learning and memory.^{18,19} Further, schizophrenic patients display aberrant induction of gamma oscillations during cognitive tasks.^{7,20,21}

These intersecting lines of evidence strongly implicate PV interneurons and NMDAR in the expression of gamma rhythms in normal cognitive

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function and in disease. To directly test the impact of NMDAR function in PV interneurons on gamma oscillation expression and behavior, we generated mice lacking NMDAR only in PV neurons. We then probed the resting-state expression of gamma oscillations, and their induction through optogenetic and pharmacological means. We also measured a variety of behaviors ranging from sensorimotor gating to cognitive dimensions such as working memory. We found that selective deletion of NMDAR from PV cells leads to an enhanced resting-state expression of gamma oscillations and a deficit in gamma induction. Further, we found a selective pattern of behavioral disturbance that spares several metrics but impairs expression of habituation, working memory and other learning indices. These data strongly support the role of NMDAR on PV cells in normal gamma functionality in the *in vivo* mammalian neocortex, and indicate a specific role for it in selective aspects of learning and memory.

Subjects and methods

Animals

All procedures were conducted in accordance with the National Institutes of Health guidelines and with the approval of the Committee on Animal Care at MIT, Cambridge, MA, USA. We generated PV-Cre/NR1f/f mice by crossing of PV-Cre mice²² with mice carrying 'floxed' NR1 alleles.²³ In awake electrophysiology and behavior tests, male PV-Cre/NR1f/f mice and littermate male controls (NR1f/f) were used. For slice electrophysiology and optogenetic experiments in anesthetized animals, age-matched male PV-Cre mice were used as controls.

Immunohistochemistry

Free-floating sections (30 µm) were prepared and immunostained.⁴ The following primary antibodies were used: PV PVG-214 (Swant, Bellinzona, Switzerland; 1:2000), enhanced yellow fluorescent protein (EYFP) (GFP-1020 Aves, Tigard, OR, USA; 1:500). Antibody staining was revealed using species-specific fluorophore-conjugated secondary antibodies (Cy5 from Jackson, West Grove, PA, USA, Alexa 488 from Molecular Probes, San Diego, CA, USA).

Quantification

Cre recombination was quantified in PV-Cre mice crossed to the R26R-EYFP Cre reporter mouse line. Free-floating sections stained with antibodies against PV and EYFP were used. For quantification of recombination in S1, every PV cell was counted and scored for co-labeling with EYFP in 1190.30 \times 1190.30 \times 30 μm images including all six cortical layers. For quantification of recombination in hippocampus every PV cell in dentate gyrus, CA1, CA2 and CA3 was counted and scored for co-labeling with EYFP.

For quantification of the number and distribution of PV cells in S1 in NR1f/f and PV-Cre/NR1f/f mice and every PV cell was counted as describe above and assigned to layers 2/3 or 4–6 based on its position in relation to the layer 4 barrels.

Slice electrophysiology

AAV DIO channelrhodopsin-2 (ChR2)-mCherry⁴ was injected into hippocampus of 5- to 7-week-old PV-Cre and PV-Cre/NR1f/f mice. At 7-10 days after viral transduction, transverse hippocampal slices (400 μ m thickness) were prepared as described.^{25,26} The tungsten bipolar electrode (FHC, Bowdoin, ME, USA) was placed in the stratum radiatum or oriens and the Schaffer collateral/commissural fibers were stimulated at 0.1 Hz. Picrotoxin (0.15 mM, Sigma, St Louis, MO, USA) was dissolved in artificial cerebrospinal fluid (aCSF) to block GABA_A receptor-mediated synaptic transmission for whole-cell patch clamp recordings. α-Amino-3-hydroxy-5-methyl-4-isoxazolepropionic acid receptor-mediated excitatory postsynaptic current (EPSCs) were recorded at -70 mV and NMDARmediated EPSCs were recorded at +40 mV with the same stimulus strength in the presence of NBQX (0.005 mm, Tocris, Ellisville, MO, USA).

Anesthetized electrophysiology

AAV DIO ChR2-mCherry was injected into barrel cortex of adult (8–12 weeks old) PV-Cre or PV-Cre/NR1f/f mice as described earlier. Electrophysiological *in vivo* recordings were performed 1–3 weeks after viral injections. Extracellular single-unit and local field potential (LFP) recordings were made with tetrodes or stereotrodes. Stimulus control and data acquisition was performed using software custom written in LabView (National Instruments, Austin, TX, USA) and Matlab (The Mathworks, Natick, MA, USA) by Ulf Knoblich.

Light stimulation was generated by a 473 nm laser and light pulses were given via a 200 μ m diameter, unjacketed optical fiber at the cortical surface 75–200 μ m from the recording electrodes.

Unit and local field potential analysis used software custom written in Igor Pro (Wavemetrics, Portland, OR, USA) by JAC. Spontaneous anesthetized LFP measurements were made during periods with no light stimulation. For each light stimulation frequency, we measured relative power²⁷⁻²⁹ in an 8 Hz band centered on that frequency. Relative power was used to account for differences in LFP amplitude between recording sites and between electrodes. Relative power was calculated by measuring the ratio of power within the band of interest to total power (1-100 Hz) in the power spectrum of the unfiltered LFP (for details see Supplementary Information; extended Materials and Methods). For each recording site, we measured power from 10 to 30 LFP traces under each condition. Example power spectra are population averages. We also measured the power ratio:

$P_{\text{light}}/P_{\text{baseline}}$

where P_{light} is the relative power in a frequency band in the presence of light stimulation and P_{baseline} is the power in that band in the absence of light stimulation.

Spike waveforms of regular spiking (RS) and FS cells were characterized. In each case, FS measurements were made from cells expressing ChR2 and driven by light pulses and RS measurements were made from non-driven, spontaneously active cells. Statistical significance was assessed with the Mann-Whitney test. All numbers are given as mean ± s.e.m., except where otherwise noted.

Awake electrophysiology

Eight NR1f/f control and eight PV-Cre/NR1f/f mice (8–10 weeks) were used for awake electrophysiology. Teflon-coated tungsten electrodes (impedance of $100 \,\mathrm{k}\Omega$) were implanted: two electrodes were placed bilaterally in primary somatosensory (barrel) cortices 1.5 mm posterior to bregma and 3.5 mm from the midline. In each hemisphere, a signal electrode was implanted 0.5 mm below the cortical surface and a reference electrode was implanted 1.75 mm below the cortical surface. A stainless steel screw over right posterior parietal cortex served as ground. Recording sessions took place in an empty box to which animals had not been previously exposed. After a 5-10-min habituation period, 20 min of baseline data were recorded for each animal. After the baseline period, four NR1f/f control and four PV-Cre/NR1f/f mice were injected intraperitoneally with 0.5 mg kg⁻¹ of MK-801 and four NR1f/f control and four PV-Cre/NR1f/f mice were injected intraperitoneally with saline and another 40 min of data were recorded. The behavioral state of all animals was scored by an observer blind to the genotype, and the sessions for ten of the animals were monitored with a video camera. All analysis was performed offline in Matlab. Normalized relative power indicates power normalized by both total power and by relative power during the baseline period, to highlight pre/post changes. Power change is the logarithm of the normalized relative power, which facilitates comparisons across frequency bands. Statistical significance was assessed with the Mann-Whitney test (between-group comparisons) and the Wilcoxon signed-rank test (within-group comparisons).

Mouse behavior

All behavioral tests were performed blind for the genotype of the mice. Male PV-Cre/NR1f/f mice and littermate male controls (NR1f/f) were used for all behavioral tests. Behavioral parameters were analyzed by t-tests for the two genotypes, except for otherwise noted, and corrected for multiple comparisons. P < 0.05 was considered significant. All results are presented as mean ± s.e.m.

Open field analysis

Activity in a novel open field was measured in monitors with sets of 16 light beam arrays. During 60 min the hardware detected beams broken by the animal, with the software determining the location and activity of the animal. For the pharmacological treatment, one set of mice was first monitored in the open field for 30 min. Directly thereafter MK-801

(M107; Sigma) (0.2 or $0.3 \,\mathrm{mg\,kg^{-1}}$) was injected intraperitoneally and the mice were monitored for additionally 60 min, in the same boxes as before.

Acoustic startle and prepulse inhibition

For testing of sensorimotor gating, startle response and prepulse inhibition (PPI) were determined using the Startle Monitor System (SM100; Hamilton Kinder, San Diego, CA, USA). The animals were habituated to the experimental equipment for two days. Day 3, the PPI testing day, each animal was exposed to 65 dB ambient noise for 5 min followed by the testing session. The PPI paradigm consisted of trials with presentation of a startle stimulus alone and trials where a prepulse of varying intensity preceded the startle stimulus by 100 ms. Trials were presented in blocks, with each block consisting of one startle stimulus alone trial, each of the prepulse-pulse stimulus trials, and a no stimulus trial, in a fixed pseudo randomized order. A total of six blocks were presented in a session and corresponding responses were averaged for each mouse and trial separately.

Habituation

Habituation of the startle reflex was assessed by comparing the mean startle amplitude of four pulses of a 120 dB white noise stimulus presented at the beginning and the end of the startle session using the equation (100–(mean startle block 2/mean startle block 1) \times 100)).

T-maze test

For testing of working memory, a modified T-maze, the discrete paired-trial variable-delay alternation task, was used.30

Contextual and cued fear conditioning

Context-dependent fear conditioning: the animals were placed in a novel conditioning chamber for 3 min where they were exposed to a foot shock. After 24 h, animals were returned to the same chamber and contextual memory was assessed as freezing during a 3-min test period. Tone-dependent (cued) fear conditioning: the animals were placed in the conditioning chamber for 3 min after which a 20-s tone (auditory cue) followed which co-terminated with a foot shock. After 24 h, animals were returned and cued fear learning was assessed. Before placement of the animals in the conditioning chamber, the environment in the chamber had been changed (visual, tactile and olfactory cues) to present the animals with a new context for the test.

Water maze

The spatial reference task was a water maze task performed in a circular tank (diameter 1.8 m) filled with opaque water. A fixed platform (10 cm diameter) was hidden below the water's surface in the center of the target quadrant. After training for several days, the platform was removed (probe trial). Reversal training started the day after the probe trial, followed by a probe trial for the new target quadrant.



Results

Genetic ablation of NMDAR signaling in PV interneurons

Previous mouse models of NMDAR hypofunction have targeted all neuronal classes, resulting in severe behavioral abnormalities. To directly address the function of NMDAR specifically in PV interneurons in network function and cognitive behavior, we created mice lacking the NMDAR subunit NR1 only in PV-expressing cells. Mice with Cre recombinase expression in PV cells (PV-Cre mice)²² were crossed with mice carrying floxed alleles of the NR1 subunit (NR1f/f mice).²³

Characterization of PV-Cre/NR1f/f mice

PV-Cre/NR1f/f mice were viable, developed normally and did not exhibit growth abnormalities or other gross anatomical changes (data not shown). Credependent recombination from the PV locus was specific to PV-expressing cells and followed the postnatal onset of PV expression³³ (Figure 1a). PV-Cre-driven recombination in somatosensory cortex and hippocampus was detected at postnatal day 13 (P13) with increased recombination at 29 days (P29) and almost complete recombination at 8 weeks (Figure 1a, Supplementary Figure 1). Whole-cell recordings in hippocampal slices *in vitro* confirmed the functional loss of NMDAR currents in PV cells in PV-Cre/NR1f/f mice (five cells in four PV-Cre/NR1f/f

mice, seven cells in five control mice, P=0.03, unpaired t-test; Figures 1b and c). We found the cortical architecture (layers and barrels in somatosensory cortex), migration and differentiation of PV interneurons to be normal in PV-Cre/NR1f/f mice (Figures 1d and e; P=0.65 for layers 2/3 and P=0.53 for layers 4–6, two-tailed unpaired t-test; n=3 per genotype).

Spontaneous and induced gamma oscillations are altered in PV-Cre/NR1f/f mice

We have previously described that activation of FS-PV interneurons enhances gamma oscillations in neocortex in vivo,4 in agreement with substantial correlative and theoretical previous evidence.34,35 Somatosensory cortex represents a local circuit with well-described anatomy and function, arguably being the most widely employed model of neocortical function in the rodent. FS-PV interneurons are local circuit neurons, and recent studies in somatosensory cortex have provided substantial progress for our understanding of FS behavior in gamma rhythm generation in general cortical network functions in vivo.4,6 There is also a substantial previous work measuring receptive fields of FS interneurons in this specific cortical area. 36–38 Recently, it was shown that NMDAR antagonists increase the power of basal gamma oscillations in rodents.13 Abnormal gamma was found to be ongoing simultaneously all over the

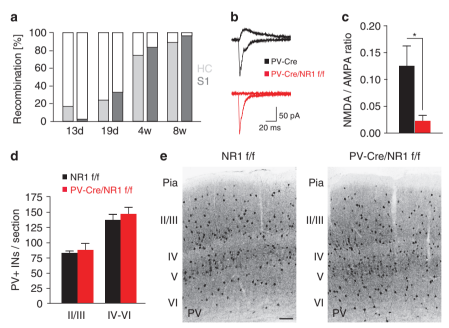


Figure 1 Genetic ablation of NMDA receptor (NMDAR) specifically in parvalbumin (PV) interneurons. (a) Quantification of recombination from the PV locus in somatosensory cortex and hippocampus at different time points. (b, c) NMDAR-mediated synaptic transmission in PV interneurons is abolished in PV-Cre/NR1f/f mice. (b) Sample EPSC traces mediated by the AMPAR (downward) and NMDAR (upward) from a control PV-Cre mouse and a PV-Cre/NR1f/f mouse. (c) NMDAR EPSC/AMPAR EPSC ratio in control PV-Cre and PV-Cre/NR1f/f mice. (d) Distribution of PV interneurons in NR1f/f and PV-Cre/NR1f/f mice, respectively, at 11 weeks. (e) Immunohistochemistry for PV interneurons in somatosensory cortex of an adult NR1f/f and PV-Cre/NR1f/f mouse, respectively. *P < 0.05; error bars, mean \pm s.e.m. Scale bar: (e) 200 µm. See also Supplementary Figure 1. AMPAR, α -amino-3-hydroxy-5-methyl-4-isoxazolepropionic acid receptor; EPSC, excitatory postsynaptic current.

cerebral cortex (prefrontal, frontal, parietal and occipital areas) in free-moving, sedated and anesthetized animals.¹³ The dysfunction in information processing found in schizophrenic patients is not limited to high-order association cortex, such as prefrontal cortex, but also influences sensory cortex.³⁹

To identify potential changes in baseline cortical rhythms, we recorded spontaneous LFPs in layers 2/3 and 4 of primary somatosensory (barrel) cortex of anesthetized PV-Cre/NR1f/f mice (n=8 sites in five control mice and 10 sites in six PV-Cre/NR1f/f mice; Figures 2a and b). We found a significant decrease in

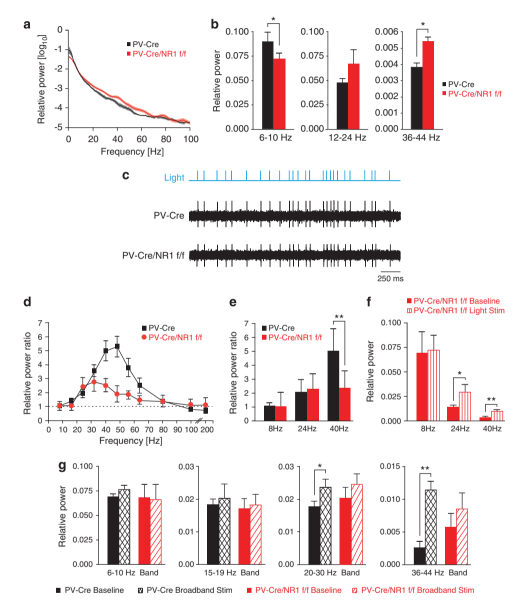


Figure 2 Spontaneous and induced cortical gamma oscillations require NMDA receptor (NMDAR) in parvalbumin (PV) interneurons. (a, b, d-g) Local field potential (LFP) activity in anesthetized control (black) and PV-Cre/NR1f/f (red) mice. (a, b) Spontaneous LFP activity. (a) Relative power from 1-100 Hz in control and PV-Cre/NR1f/f mice. (b) Relative LFP power in the 6-10, 12-24 and 36-44 Hz frequency bands. (c) Single-unit recordings during optogenetic activation of FS-PV+ interneurons. Middle trace: control mice, lower trace: PV-Cre/NR1f/f mice. In each case, a series of 1 ms light pulses (blue trace) was given in a random pattern drawn from a broadband distribution (5-200 Hz). In both cases, each light pulse evoked a single spike from the cell. Both cells followed the light stimulus with a high degree of reliability. (d-g) Optogenetic activation of FS-PV interneurons in somatosensory cortex in control and PV-Cre/NR1f/f mice. (d) Mean power ratio in each LFP frequency band in response to light activation of channelrhodopsin-2 (ChR2)-expressing FS-PV interneurons at varying frequencies. PV-Cre/NR1f/f mice generate significantly less 30-60 Hz oscillations (gamma) than control mice. (e) Comparison of the effect of activating FS-PV interneurons in the control and PV-Cre/NR1f/f mice at 8, 24, and 40 Hz on relative LFP power in those frequency bands. (f) Relative LFP power in the 8, 24 and 40 Hz frequency bands in the PV-Cre/NR1f/f mice during spontaneous (solid bars) and light-evoked (striped bars) activity. (g) Relative power in the 6-10, 15-19, 20-30 and 36-44 Hz frequency bands in response to broadband light stimulation in control and PV-Cre/NR1f/f mice. *P<0.05, **P<0.01; error bars, mean \pm s.e.m.



relative power^{27–29} in the 6–10 Hz theta frequency band (P < 0.05; Figure 2b) and a significant increase in relative power in the 36–44 Hz gamma frequency band (P < 0.05; Figure 2b) in PV-Cre/NR1f/f mice compared with control mice, whereas LFP activity at slightly lower frequencies in the 12–24 Hz beta-frequency band was not significantly altered in baseline conditions in PV-Cre/NR1f/f mice (Figure 2b). RS, putative excitatory neurons in PV-Cre/NR1f/f mice showed a significantly higher spontaneous firing rate (n = 16 cells; 0.49 ± 0.09 Hz) than did RS cells in control mice (n = 18 cells; 0.25 ± 0.06 Hz; Mann–Whitney test; P < 0.05; data not shown).

We next tested the role of NMDAR in FS-PV interneurons for the induction of gamma oscillations in cortical networks by direct recruitment of FS-PV interneurons through the specific expression of the light-activated channel ChR2, as described previously.4 We drove ChR2-expressing FS-PV cells at a range of frequencies (8-200 Hz) in bouts of 3 s of 1 ms light pulses (n=7 sites in five control mice and 12 sites in six PV-Cre/NR1f/f mice; Figures 2c-f). Across our sample, FS-PV interneurons in control and PV-Cre/NR1f/f mice were driven with high reliability by light pulses (Figure 2c). Light pulses in the gamma range resulted in significant amplification of relative LFP power at those frequencies in control mice (P < 0.01; Figure 2d). Driving FS-PV interneurons lacking NMDAR revealed a specific disruption in the ability of the local network to induce gamma oscillations, resulting in a significantly reduced gamma activity enhancement compared with control mice (Figure 2d). The specificity of this disruption to the gamma band in PV-Cre/NR1f/f mice was highlighted when FS-PV cells were driven at 8, 24 and 40 Hz, respectively (Figure 2e). The oscillatory disruption was not complete as optogenetic stimulation at 24 or 40 Hz each induced significantly increased activity compared with baseline conditions in PV-Cre/NR1f/f mice (P < 0.05; Figure 2f).

We used randomly patterned light stimulation (broadband stimulation⁴) of FS–PV interneurons to probe the emergent properties of the local cortical circuit. This paradigm, in agreement with periodic light stimulation, evoked a significant increase in 20–30 Hz and 36–44 Hz activity in control (P<0.05) but not PV-Cre/NR1/f/f mice (P>0.05; n=7 sites in five control and 10 sites in six PV-Cre/NR1f/f mice; Figure 2g).

To probe the impact of NMDAR deletion on recruitment of FS-PV interneurons, we analyzed spike latency and variance in spike timing following optical stimulation in PV-Cre/NR1f/f mice. We found a diminished spike synchronization of FS units in response to light activation (Supplementary Figures 2b, c and g) compared with control mice (Supplementary Figures 2e-g) with significantly increased spike latency and variance in spike timing. As shown in this example, we observed reduced inhibition of RS neurons early in the gamma cycle (4–10 ms after light pulse; Supplementary Figures 2a and c) as well

as diminished entrainment of the RS neurons at the end of the cycle (around 20 ms after light pulse; Supplementary Figures 2a and c) compared with control mice (Supplementary Figures 2d and f).

Computational modeling, described below, suggests that reduced synchronization of the FS-PV population results in higher gamma band activity during baseline conditions as well as a reduced ability to recruit and synchronize the local network during higher levels of excitatory activity (Supplementary Figure 3a).

To test whether deficits in cortical network activity were also present in awake behaving animals, we performed LFP recordings in layers 2/3 and 4 of somatosensory cortex in freely moving PV-Cre/NR1f/f (n=7) or control mice (n=7) during behavior in a novel environment. We observed prominent gammaband oscillations in both genotypes (Figure 3a). During a baseline recording period, PV-Cre/NR1f/f mice exhibited a nonsignificant trend toward elevated spontaneous LFP in the 30-80 Hz gamma range independent of specific behaviors such as grooming or active exploration, compared with control mice (Figure 3b). PV-Cre/NR1f/f mice showed significantly longer mean event durations for spontaneous gamma oscillations compared with control mice (P < 0.005)without changes in the number of gamma events (Figure 3c).

To explore the role of NMDAR in PV interneurons on evoked gamma activity in the awake state, we challenged PV-Cre/NR1f/f mice with an acute administration of the non-competitive NMDAR antagonist MK-801 (0.5 mg kg⁻¹; intraperitoneal; Figures 3d–g) or saline (data not shown). In agreement with previous findings,14 control mice acutely displayed a significant increase in LFP in the 30-50 Hz gamma range (post1, 5-15 min after MK-801 administration; \bar{P} <0.05, Wilcoxon signed-rank test; Figure 3e). PV-Cre/NR1f/f mice, in contrast, displayed a significant reduction in gamma-band activity after NMDAR antagonist treatment (Figure 3e). Control mice and PV-Cre/NR1f/f mice displayed a significant induction in the 6-10 Hz frequency band relative power at later time points (post2, 25-35 min after MK-801 administration; Supplementary Figures 4a and b). In the 12-24 Hz band, the relative power remained unchanged in control animals during the same time frame, but was significantly reduced in PV-Cre/NR1f/f animals (Supplementary Figures 4c and d). Saline treatment did not induce any significant changes in locomotor behavior or relative power in any frequency band between genotypes throughout the duration of the recording sessions (data not shown).

To interpret the findings relative to established theories of gamma emergence, we conducted simulations using a modified form of an established neural circuit model. We simulated NMDAR dysfunction in FS–PV interneurons and found emergence of higher frequency oscillations in baseline activity, analogous to the experimental data (Supplementary Figure 3a and Supplementary Text; Modeling results).

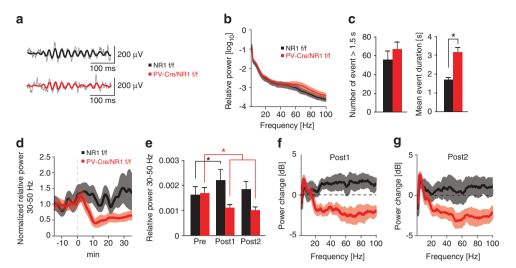


Figure 3 Baseline cortical oscillations and NMDA receptor (NMDAR) antagonist induced gamma rhythms in awake behaving animals. (a–g) Local field potential (LFP) activity in somatosensory cortex in awake control and parvalbumin (PV)-Cre/NR1f/f mice. (a) Examples of single-trial gamma activity from a PV-Cre/NR1f/f and control mouse. Thin lines: LFP filtered between 5–300 Hz; thick lines: LFP filtered in the 30–50 Hz gamma frequency range. (b, c) Awake baseline activity. (b) Mean power spectra 1–100 Hz. Lighter regions indicate s.e.m. (c) Characteristics of 30–50 Hz gamma events. Mean event duration is significantly increased in freely moving PV-Cre/NR1f/f mice, whereas number of events are not. (d–g) NMDAR antagonist (MK-801) challenge. (d, e) Average relative power in the 30–50 Hz gamma frequency band 15 min before administration of MK-801 (dashed line), to 35 min after. Lighter regions indicate s.e.m. Administration of MK-801 gives a significant increase in relative gamma power in control mice and a significant reduction in relative gamma power in PV-Cre/NR1f/f mice. (f) Average power changes (dB) between pre and post1. Lighter regions indicate s.e.m. (g) Average power changes (dB) between pre and post2. Pre, 5–15 min before MK-801; post1, 5–15 min after MK-801; post2, 25–35 min after MK-801. *P<0.05; error bars, mean ± s.e.m. See also Supplementary Figures 3 and 4.

We also observed a significant reduction in gamma-band activity following simulation of pharmacological NMDAR inhibition in a PV-Cre/NR1f/f network (Supplementary Figure 3b and Supplementary Text; Modeling results). In addition, we found an increase in the latency and variation of the FS-PV response to simulated light drive that may be responsible for the smaller enhancement of gamma power in this condition (Supplementary Figure 3c and Supplementary Text; Modeling results).

Behavioral effects of NMDAR deletion in PV cells We characterized the PV-Cre/NR1f/f mice in a series of paradigms to assess general locomotor and exploratory behaviors as well as cognitive tasks evaluating learning and memory. The PV-Cre/NR1f/f mice and their littermate controls (NR1f/f) were introduced into an automated novel open field environment at an age of 7 weeks (n=11 per genotype). We did not observe any consistent behavioral differences between PV-Cre/NR1f/f and control mice over the entire 60 min trial or in any 5 min block after analysis of 18 different parameters, including total distance (Figure 4a), locomotion, stereotypy and center/margin time (data not shown).

We questioned whether PV-Cre/NR1f/f mice at later stages develop behavioral abnormalities in the open field and therefore assessed mice of 11-12 weeks of age (n=9 per genotype). We found a nonsignificant trend toward a decrease in total distance traveled over

the 60-min period for PV-Cre/NR1f/f mice compared with control mice (P>0.05; Mann–Whitney test; Figure 4b). Analysis of time spent in the center (Figure 4c) or margin of the open field, vertical movements (data not shown) or stereotypy counts (Figure 4d) did not reveal differences between the genotypes over the 60-min period.

Our electrophysiological findings indicate that the PV-Cre/NR1f/f mice have a selective deficit in gamma emergence following optogenetic drive of FS-PV interneurons or pharmacological NMDAR inhibition. To investigate potential behavioral differences in response to acute NMDAR inhibition, we exposed naïve PV-Cre/NR1f/f and control mice (n=11) per genotype) to the open field for 30 min and then challenged them with administration of the NMDAR antagonist MK-801 at a low dose (0.3 mg kg⁻¹) followed by continued exposure to the open field for 60 min. Control mice responded over time to MK-801 administration with increased horizontal activity and induction of stereotypies as previously reported³¹ (Figures 4e and f). In contrast, PV-Cre/NR1f/f mice showed a markedly reduced sensitivity to MK801 (Figures 4e and f), consistent with our electrophysiological findings that PV interneurons are an important target for the non-competitive NMDAR antagonists.

PPI is associated with deficiencies in sensory information filtering and has been well characterized in schizophrenia.^{41,42} PPI deficits have repetitively been found in schizophrenia patients and their

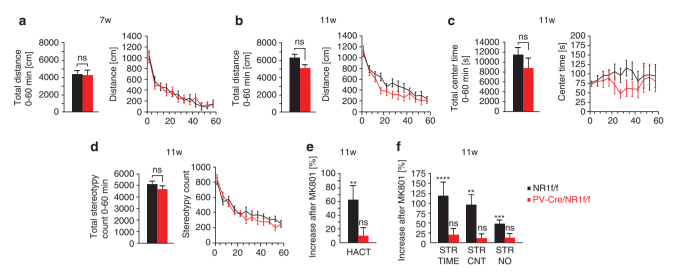


Figure 4 Parvalbumin (PV)-Cre/NR1f/f mice display no behavioral changes in the open field but reduced sensitivity to pharmacological NMDA receptor (NMDAR) treatment. (a) Total distance traveled over 60 min in the open field of PV-Cre/NR1f/f and control mice at 7 weeks. (b-d) Open field behavior over 60 min at 11 weeks. (b) Total distance traveled over 60 min in the open field of PV-Cre/NR1f/f and control mice at 11 weeks. (c) No significant difference in the time spent in the center of the field between the genotypes. (d) PV-Cre/NR1f/f mice do not display stereotypy behavior. (e) MK-801 treatment induces significant increase in horizontal activity only in control mice. (f) MK-801 treatment results in marked increase in stereotypy behavior only in control mice. HACT, horizontal activity; STR TIME, stereotypy time; STR CNT, stereotypy count; STR NO, stereotypy numbers; ns P > 0.05, **P < 0.01, ***P < 0.001, ***P < 0.0001; error bars, mean \pm s.e.m.

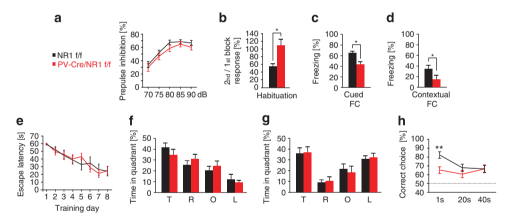


Figure 5 Loss of NMDA receptor (NMDAR) in parvalbumin (PV) interneurons results in selective cognitive disruptions. (a) PV-Cre/NR1f/f mice show no deficiency in sensorimotor gating as measured by prepulse inhibition (PPI). (b) PV-Cre/NR1f/f display deficiencies in habituation. (c, d), PV-Cre/NR1f/f mice exhibit impaired freezing behavior both to a tone-dependent (cued) and a contextual version of fear conditioning. (e) PV-Cre/NR1f/f mice perform similarly to control mice during training in a hidden platform version of water maze. (f) Time spent (%) in each quadrant during the water maze probe trial. There is no significant difference between the genotypes. (g) Time spent (%) in each quadrant during the water maze probe trial after reversal training. There is no significant difference between the genotypes. (h) PV-Cre/NR1f/f mice perform at a similar accuracy levels independent of working memory load in the discrete paired-trial variable-delay T-maze task. PPI, prepulse inhibition; FC, fear conditioning; T, target quadrant; R, right quadrant; O, opposite quadrant; L, left quadrant. *P<0.05, **P<0.01; error bars, mean \pm s.e.m. See also Supplementary Figure 5.

unaffected first-degree relatives. Interneurons are considered to be important for phase-dependent inhibition and circuit oscillations that in turn regulate information processing. Our evaluation of PPI in PV-Cre/NR1f/f mice compared with control mice did not reveal any significant changes in PPI over several prepulse intensities (70–90 dB) (Figure 5a, n=17 control mice, 15 PV-Cre/NR1f/f mice). Importantly, the startle response was not different between control

mice and PV-Cre/NR1f/f mice (Supplementary Figures 5a and b), thereby excluding sensorimotor defects.

 $Loss\ of\ NMDAR\ in\ PV\ cells\ results\ in\ selective\ cognitive\ disruptions$

Several studies have analyzed the role of NMDAR in pyramidal neurons during learning and memory by pharmacological NMDAR inhibition⁴³ or with genetic approaches.^{23,44} The contribution of interneurons to learning and memory processes has been proposed⁴⁵ but not genetically evaluated, and interneurons display similar NMDAR-dependent electrophysiological properties as pyramidal neurons.⁴⁶ Habituation is considered to be one of the simplest forms of learning and is disrupted by NMDAR antagonists. 41,47 Accordingly, we observed a significant reduction in acoustic startle habituation in PV-Cre/NR1f/f mice, supporting the NMDAR-dependent contribution of PV interneurons to learning and memory (Figure 5b and Supplementary Figure 5c).

To investigate the role of NMDAR in PV cells for associative learning processes, we exposed mice to cued and contextual fear conditioning paradigms (n=5 per genotype). Mice were exposed to a neutral conditioned stimulus (sound) in a novel environment paired with an aversive foot shock as the unconditioned stimulus, PV-Cre/NR1f/f mice exhibited impaired associative learning both in a tone-dependent trial 24 h later (P < 0.05, Mann–Whitney test; Figure 5c) and a contextual version of the test (P < 0.05, Mann-Whitney test; Figure 5d).

To evaluate spatial reference memory, we used a water maze task with a fixed hidden platform consisting of two trials per day with a 5-min intratrial interval. Infusion of NMDAR antagonists has revealed a critical role for NMDAR during learning and memory for spatial reference memory,43 although the specific contribution of NMDAR on interneurons has not been addressed. The rate of learning in the water maze task over 8 days was not impaired in PV-Cre/ NR1f/f mice compared with controls (Figure 5e; escape latency, P = 0.8794, two-way analysis of variance; n = 8 control and 11 PV-Cre/NR1f/f mice). The probe trial on day 9 did not reveal significant differences in target quadrant preference compared with control mice (P=0.33 for target quadrant, unpaired t-test; Figure 5f). To address potential impairments in reversal learning, we introduced the hidden platform in a different quadrant and continued the learning paradigm for 4 days. Learning rates for the new platform location over 4 days were not significantly different between genotypes (P = 0.513, two-way analysis of variance) and a probe trial on the fifth day did not reveal target quadrant preference differences between genotypes (P = 0.9 for target quadrant, unpaired t-test; Figure 5g).

Working memory deficits in schizophrenic patients,48-50 and computational models implicate NMDAR as a critical component of working memory,51 with inhibition being important in shaping the time-dependent firing of pyramidal cells in neocortical circuits during cognitive tasks.⁵² Further, NMDAR antagonists disrupt working memory in humans⁵³ and rodents.⁵⁴ We assessed working memory in a T-maze (the discrete paired-trial variabledelay T-maze task;³⁰ n = 10 control and nine PV-Cre/ NR1f/f mice). Both genotypes behaved similarly during the 10 days of training (Supplementary Figure 5d). After training, we performed 10 trials per mouse

each day with varying intratrial intervals, to evaluate effects of working memory load on performance. The PV-Cre/NR1f/f mice performed similarly to control mice at 20- and 40-s intratrial intervals (Figure 5h). In contrast, the 1-s intratrial interval revealed a marked difference between genotypes, with PV-Cre/NR1f/f mice remaining at 65% accuracy whereas control mice were significantly better with 82% accuracy (P<0.01; Figure 5h). PV-Cre/NR1f/f mice therefore performed at similar accuracy levels independent of working memory load.

Discussion

We have investigated the in vivo function of NMDAR specifically in FS-PV interneurons in regulating cortical brain rhythms and cognitive functions (Supplementary Table 1). This work is based on a longstanding hypothesis connecting PV interneuron dysfunction, NMDAR hypofunction and disturbances in brain rhythms associated with cognitive tasks/functions. We find that NMDAR signaling in FS-PV interneurons is critical for the regulation of gamma oscillations during baseline conditions as well as for gamma rhythm induction. The data we present on optogenetic drive in the superficial cortical layers are specific to FS-PV interneurons, as PV-expressing cells in these laminae are only FS interneurons. That said, PV-expressing neurons are present throughout the brain. One alternative cell type that could impact our findings is PV-expressing thalamic neurons, which typically project to the granular layers in cortex. There is correlative (neurophysiological), causal (optogenetic) and computational (modeling) evidence that neocortical gamma oscillations depend crucially on local FS interneurons, but these studies also suggest that the tonic level of excitation to the neocortical circuit is a key. As such, alternations in these thalamic neurons could have impacted, for example, our baseline data.

The inability of the cortical network to induce additional gamma oscillations by direct activation of FS-PV interneurons might indicate an impairment of network flexibility. The results suggest that PV-Cre/ NR1f/f mice exhibit spontaneous and evoked network abnormalities similar to those observed after low does administration of NMDAR antagonists.¹³ This is similar to findings in psychiatric patients, who display aberrant recruitment of cortical circuits and diminished evoked gamma rhythm in response to cognitive and sensory tasks.55 The reduced gammaband activity after NMDAR antagonist treatment in PV-Cre/NR1f/f mice supports the hypothesis that FS-PV interneurons are an important target for pharmacological NMDAR blockade associated with altered gamma rhythms, 13,56 consistent with our computational model of the PV-Cre/NR1f/f cortical circuit.

We have further found a dissociation between the requirement for NMDAR in FS-PV interneurons during baseline behavior versus demanding cognitive tasks. Although the small age-dependent effects in the open field may be of interest in light of behavioral changes associated with transitions from adolescence to adulthood, our results suggest a subtle behavioral effect at most of NMDAR deficiency in PV interneuron in the unchallenged state. This finding is in contrast to the phenotypes of hyperlocomotion and stereotypical behaviors in mice with general NMDAR hypofunction.^{31,32}

Working memory includes executive components such as goal maintenance and interference control.⁵⁷ It is difficult to establish the explicit role of executive components in rodent working memory tasks, and in addition, several cognitive processes and memory systems may be used in conjunction in the tasks.⁵⁷ Although the selective working memory deficit in performance of the PV-Cre/NR1f/f mice at short delays in the discrete paired-trial variable-delay T-maze task most likely represent a complex network deficit, these data suggest that activation of NMDAR on PV interneurons is required for rapid initiation of the working memory encoding phase. In contrast, PV-Cre/NR1f/f mice displayed intact spatial reference memory, as measured in the Morris water maze task. In line with this, mice with deletion of the α -amino-3hydroxy-5-methyl-4-isoxazolepropionic acid (AMPA) receptor GluR-A specifically in PV-expressing interneurons⁵⁸ do not show impaired spatial reference memory, but have reduced kainate-induced gamma oscillations in hippocampal slices. Our results point to a specific role for NMDAR in PV interneurons in memory tasks involving the neocortex, amygdala and hippocampus. A potential unifying feature of these forms of memory is their dependence on intact gamma rhythms, which are enhanced during exposure to novel environments,⁵⁹ but might have a lesser role in long-term learning in the water-maze task or the longer intratrial intervals in the T-maze task.

Evidence supporting the important role of NMDAR in interneurons has recently been provided through genetic deletion of NMDAR in a mixed population of GABAergic interneurons, including PV interneurons. For This deletion results in a wide range of behavioral phenotypes, including novelty-induced hyperlocomotion, PPI deficits and anxiety-like effects, which were not observed in PV-Cre/NR1f/f mice. Our model of a targeted adolescent disruption of NMDAR transmission specifically in PV interneurons provides a specific framework for understanding the role of NMDA transmission in PV interneurons for oscillatory activities in neuronal ensembles.

An important question is to what extent the observed circuit deficits depend on specific NMDAR-related activities, or represent a general hypoexcitability of PV interneurons conferred by diminished glutamatergic drive. It has been shown that reduced excitatory recruitment of PV interneurons through loss of α -amino-3-hydroxy-5-methyl-4-isoxazolepropionic acid (AMPA) receptors affects ongoing and induced gamma oscillations in hippocampal slices, 58 with corresponding deficits in work-

ing memory but not spatial reference memory. Our *in vivo* physiological, behavioral studies and computational modeling suggest that such deficits, found in our mouse model as well, could reflect a general decrease in excitability of PV interneurons.

There is direct evidence for NMDAR hypofunction in psychiatric patients. ⁶¹ Additionally, human genetic data for schizophrenia-associated genes have implicated the NMDAR signaling pathway and disruption of the neuregulin-1/ErbB4 pathway. ^{62–64} PV interneurons express ErbB4 protein ⁶⁵ and the neuregulin-1/ErbB4 pathway has a role in gamma oscillations. ⁶⁶ However, we emphasize that the PV-Cre/NR1f/f mouse does not represent a model of schizophrenia, but rather contributes to the investigation of possibly one dimension of schizophrenia that includes cognitive defects. Further, under even the best circumstances, a mouse model is limited in its ability to recapitulate all complex cognitive dimensions that have evolved in humans.

Conflict of interest

The authors declare no conflict of interest.

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