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Status epilepticus enhances tonic GABA currents and depolarizes GABA reversal potential in dentate fast-spiking basket cells

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Status epilepticus enhances tonic GABA currents and depolarizes GABA reversal potential in dentate fast-spiking basket cells. J Neurophysiol 109: 1746–1763, 2013. —Temporal lobe epilepsy is associated with loss of interneurons and inhibitory dysfunction in the dentate gyrus. While status epilepticus (SE) leads to changes in granule cell inhibition, whether dentate basket cells critical for regulating granule cell feedforward and feedback inhibition express tonic GABA currents (I_{GABA}) and undergo changes in inhibition after SE is not known. We find that interneurons immunoreactive for parvalbumin in the hilar-subgranular region express GABA_A receptor (GABA_A,R) δ-subunits, which are known to underlie tonic I_{GABA}. Dentate fast-spiking basket cells (FS-BCs) demonstrate baseline tonic I_{GABA} blocked by GABA_A,R antagonists. In morphologically and physiologically identified FS-BCs, tonic I_{GABA} is enhanced 1 wk after pilocarpine-induced SE, despite simultaneous reduction in spontaneous inhibitory postsynaptic current (sIPSC) frequency. Amplitude of tonic I_{GABA} in control and post-SE FS-BCs is enhanced by 4,5,6,7-tetrahydroisoxazolo[5,4-c]pyridin-3-ol (THIP), demonstrating the contribution of GABA_A,R δ-subunits. Whereas FS-BC resting membrane potential is unchanged after SE, perforated-patch recordings from FS-BCs show that the reversal potential for GABA currents (E_{GABA}) is depolarized after SE. In model FS-BCs, increasing tonic GABA conductance decreased excitability when E_{GABA} was shunting and increased excitability when E_{GABA} was depolarizing. Although simulated focal afferent activation evoked seizure-like activity in model dentate networks with FS-BC tonic GABA conductance and shunting E_{GABA}, excitability of identical networks with depolarizing FS-BC E_{GABA} showed lower activity levels. Thus, together, post-SE changes in tonic I_{GABA} and E_{GABA} maintain homeostasis of FS-BC activity and limit increases in dentate excitability. These findings have implications for normal FS-BC function and can inform studies examining comorbidities and therapeutics following SE.

ACQUIRED TEMPORAL LOBE EPILEPSY occurring as a consequence of unprovoked seizures is marked by neuropathological changes in the dentate gyrus (Margerison and Corsellis 1966). Alterations in granule cell inhibition, resulting from loss or dysfunction of GABAergic interneurons, have been proposed to contribute to development of epilepsy (Cossart et al. 2005; Coulter 2001). The early period within a week after status epilepticus (SE) is characterized by enhanced entorhinal input to the dentate gyrus (Bragin et al. 2004; Kobayashi et al. 2003) and is associated with changes in granule cell GABA currents (I_{GABA}) (Kobayashi and Buckmaster 2003; Zhan and Nadler 2009) and GABA reversal potential (E_{GABA}) (Pathak et al. 2007). The cellular and synaptic changes that occur prior to development of spontaneous seizures have been the focus of several studies because of their potential to contribute to the epileptogenic process rather than being a side effect of epilepsy (Brooks-Kayal et al. 1998; Kobayashi et al. 2003; Pathak et al. 2007). SE leads to loss and structural reorganization of interneurons and alterations in their excitatory inputs (Zhang and Buckmaster 2009; Zhang et al. 2009). Although interneuronal inhibition underlies generation of brain rhythms and regulates network activity levels (Buzsaki 2006), whether interneuronal inhibition shows early changes after SE remains untested.

Fast-spiking basket cells (FS-BCs), a class of interneurons with perisomatic projections, are critical for maintaining the low excitability and sparse firing of dentate granule cells and contribute to feedforward and feedback dentate inhibition (Ewell and Jones 2010; Krausshaar and Jonas 2000). Dentate FS-BCs express the calcium-binding protein parvalbumin (PV), have a characteristic high-frequency nonadapting firing pattern (Harney and Jones 2002; Hefft and Jonas 2005), and are interconnected through high-fidelity GABAergic synapses (Bartos et al. 2001). Apart from synaptic GABA_A receptors (GABA_A,Rs), granule cells and certain interneurons express extra- and perisynaptic high-affinity GABA_A,Rs that contribute to “tonic” I_{GABA} (Farrant and Nusser 2005; Scimemi et al. 2005). GABA_A,Rs containing δ-subunits contribute to tonic I_{GABA} in dentate granule cells and molecular layer interneurons (Glykys et al. 2007; Mchedlishvili and Kapur 2006; Wei et al. 2003). Previous studies have shown that nonprincipal neurons in the dentate hilus express GABA_A,R δ-subunits (Peng et al. 2004); however, whether dentate FS-BCs express tonic I_{GABA} is not known. Since tonic I_{GABA} regulates neuronal excitability and can undergo activity-dependent changes during synaptic GABA spillover (Glykys and Mody 2007), the presence of tonic I_{GABA} in FS-BCs will impact their function during network activity. Moreover, FS-BC tonic I_{GABA} may be altered after SE, as has been observed in granule cells (Zhan and Nadler 2009; Zhang et al. 2007).

GABAergic inhibition is hyperpolarizing when E_{GABA} is negative to neuronal resting membrane potential (RMP) and shunting when E_{GABA} is close to RMP. Hippocampal and dentate interneuronal E_{GABA} has been shown to lie positive to RMP, contributing to shunting inhibition (Banke and McBain...
acquired with pCLAMP 10 at 10-kHz sampling frequency. Tonic and immersion objective. Recordings were obtained with Axon Instruments with a Nikon Eclipse FN-1 microscope, using a granule cell layer were performed with IR-DIC visualization technique. We examined whether dentate FS-BCs express tonic $I_{\text{GABA}}$ and whether FS-BC tonic $I_{\text{GABA}}$ and $E_{\text{GABA}}$ altered 1 wk after SE. Using single-cell models and large-scale network simulations, we characterized how the experimentally identified post-SE changes in FS-BC tonic inhibition influence FS-BC and dentate network activity.

MATERIALS AND METHODS

Pilocarpine status epilepticus. All procedures were performed under protocols approved by the University of Medicine and Dentistry of New Jersey Institutional Animal Care and Use Committee. Pilocarpine injection was performed as previously reported (Zhang et al. 2009). Young adult male Wistar rats between postnatal days 25 and 27 were injected with scopolamine methyl nitrate (1 mg/kg sc) 30 min before pilocarpine injection. SE was induced by injection of pilocarpine (300 mg/kg ip). After 1 h and 30 min of continuous stage 3 or greater seizures (Racine scale), diazepam (10 mg/kg ip) was administered and repeated as needed to terminate seizures. Control rats received scopolamine pretreatment followed by saline injection (ip) and diazepam after 2 h. Animals were video monitored (continuous video recording for 8 h on the day before experimentation with a PC333HR high-resolution camera coupled to a 4 Channel H.264 Pentaplex Digital Video Recorder) to rule out occurrence of spontaneous seizures. Unless otherwise stated, all anatomical and physiological studies were conducted on seizure-free rats 6–8 days after pilocarpine-SE and in age-matched, saline-injected control rats.

Slice preparation. One week (6–8 days) after saline injection or pilocarpine-induced SE, rats were anesthetized with isoflurane and decapitated. Horizontal brain slices (300 μm for patch clamp and 400 μm for field experiments) were prepared in ice-cold sucrose-artificial cerebrospinal fluid (sucrose-aCSF) containing (in mM) 85 NaCl, 75 sucrose, 24 NaHCO3, 25 glucose, 4 MgCl2, 2.5 KCl, 1.25 NaH2PO4, and 0.5 CaCl2 with a Leica VT1200S Vibratome (Wetzlar, Germany). The slices were sagittally bisected and incubated at 32±1°C for 30 min in a submerged holding chamber containing an equal volume of sucrose-aCSF and recording aCSF and subsequently held at room temperature (RT). The recording aCSF contained (in mM) 126 NaCl, 2.5 KCl, 2 CaCl2, 2 MgCl2, 1.25 NaH2PO4, 26 NaHCO3, and 10 d-glucose. All solutions were saturated with 95% O2-5% CO2 and temperature (RT). The recording aCSF contained (in mM) 126 NaCl, 2.5 KCl, 2 CaCl2, 2 MgCl2, 1.25 NaH2PO4, 26 NaHCO3, and 10 d-glucose. All solutions were saturated with 95% O2-5% CO2 and maintained at a pH of 7.4 for 1–6 h.

In vitro electrophysiology. For patch-clamp recordings, slices (300 μm) were transferred to a submerged recording chamber and perfused with oxygenated aCSF at 33 ± 1°C. Whole cell voltage- and current-clamp recordings from interneurons at the border of the hilus and granule cell layer were performed with IR-DIC visualization techniques with a Nikon Eclipse FN-1 microscope, using a X40 water-immersion objective. Recordings were obtained with Axon Instruments MultiClamp 700B (Molecular Devices, Sunnyvale, CA). Data were low-pass filtered at 3 kHz, digitized with DigiData 1440A, and acquired with pCLAMP 10 at 10-kHz sampling frequency. Tonic and synaptic $I_{\text{GABA}}$ were recorded in perfusing aCSF containing the glutamate receptor antagonist kynurenic acid (Kya, 3 mM; Tocris, Ellisville, MO). No additional GABA was included in the recording solution. Except in experiments presented in Fig. 5D, GABA transporter antagonists were not included in the recording solution. Recordings were obtained with microelectrodes (5–7 MΩ) containing (in mM) 125 KCl, 10 K-glucuronate, 10 HEPES, 2 MgCl2, 0.2 EGTA, 2 Na-ATP, 0.5 Na-GTP, and 10 phosphocreatine titrated to a pH of 7.25 with KOH. Biocytin (0.2%) was included in the internal solution for post hoc cell identification (Santhakumar et al. 2010). Recorded neurons were initially held at −70 mV, and the responses to 1.5-s positive and negative current injections were examined to determine active and passive characteristics. Cells with nonadapting, high-frequency firing for the entire duration of the current injection and low input resistance ($R_{\text{input}}$ <150 MΩ) were classified as FS-BCs (Heft and Jonas 2005). Neurons with adapting firing, high $R_{\text{input}}$ (>150 MΩ), and sag during negative current injection were considered non-fast-spiking interneurons (non-FS-Is) (Heft and Jonas 2005). Post hoc biocytin immunostaining and morphological analysis were used to definitively identify FS-BCs included in this study, on the basis of presence of axon terminals in the granule cell layer. After current-clamp recordings, cells were held in voltage clamp at −70 mV for analysis of GABA currents. Tonic $I_{\text{GABA}}$, steady-state currents blocked by the GABA$_{A}$R antagonist SR95531 (10 μM), was measured as described previously (Gupta et al. 2012) with custom macrons in IGOR Pro 7.0 software (WaveMetrics, Lake Oswego, OR). Briefly, the magnitude of tonic $I_{\text{GABA}}$ was calculated by plotting all-point histograms of relevant 30-s segments of data. These data were fit to Gaussian equations, constraining fits to values two bins more negative than the peak. This ensured that the tail of higher-amplitude values [representing spontaneous inhibitory postsynaptic currents (sIPSCs)] did not influence the fit (Santhakumar et al. 2006, 2010). Recordings were discontinued if series resistance increased by >20%. Cell capacitance was measured with the automated function in Multiclamp 700B. In some experiments, 4,5,6,7-tetrahydroisoxazolol[5,4-c]pyridin-3-ol (THIP, 1 μM), a selective GABA$_{A}$R agonist with a preference for δ-subunit-containing GABA$_{A}$Rs (Brown et al. 2002), or the GABA transporter-1 uptake inhibitor 1-[(diphenylmethylene)imino]oxo[ethyl]-1,2,5,6-tetrahydro-3-pyridinecarboxylic acid hydrochloride (NO-711, 10 μM) was included in the external solution. Individual sIPSCs were detected with custom software in IGOR Pro 7.0 (Gupta et al. 2012; Santhakumar et al. 2010). Events were visualized, and any “noise” that spuriously met trigger specifications was rejected. Cumulative probability plots of sIPSC parameters were constructed with IGOR Pro by pooling an equal number of sIPSCs from each cell.

Gramicidin-perforated-patch recordings were performed to prevent alteration of the intracellular chloride concentration by the pipette solution (Ebihara et al. 1995). Intracellular solution contained (mM) 135 KCl, 0.5 CaCl2, 5 Na$_2$EGTA, 10 HEPES, 2 MgCl2, and 2 Mg-ATP, with 0.2% biocytin, pH set to 7.2 with KOH and gramicidin D (100 μg/μl with 1% DMSO final concentration). Electrode tips were filled with gramicidin-free internal solution and back-filled with the solution containing gramicidin D. On formation of perforated patch, responses to positive and negative current injections were recorded in cell-attached mode for physiological identification. A cocktail containing tetrodotoxin (TTX, 1 μM), 6,7-dinitroquinoxaline-2,3-dione (DNQX, 20 μM), and d-(-)-2-amino-5-phosphonopentanoic acid (APV, 50 μM) (Tocris), to block Na$^+$ channels, AMPA receptors, and NMDA receptors, was used to isolate $I_{\text{GABA}}$. Perforated-patch recordings were obtained after series resistance had stabilized between 80 and 100 MΩ ~40 min after patch formation. Series resistance was monitored at 2 min intervals, and data were rejected when resistance suddenly decreased, indicating rupture of the perforated patch. In some initial experiments Alexa Fluor 488 (50–100 μM) was used in the pipette to confirm that the change in resistance was a reliable and adequate indicator of patch rupture. A Picospritzer (FMI-100, Dagan) was used to apply GABA (100 μM), containing blockers of synaptic transmission mentioned above) at 10 psi from a pipette residing 10–20 μm above the slice at the position of the recorded soma. Voltage ramps −130 to +10 mV over 200 ms applied from a holding potential of −60 mV in the absence and presence of GABA were used to determine $E_{\text{GABA}}$. The membrane voltage at which the current traces, obtained in the presence and absence of GABA, crossed was measured as the apparent $E_{\text{GABA}}$ (Billups and Attwell 2002). RMP was measured as the potential at which holding current = 0 pA. All measurements were corrected for a liquid junction potential and
voltage drop across series resistance. In a subset of cells, $E_{\text{GABA}}$ was also estimated by systematically varying the steady-state holding potential and estimating the reversal potential of GABA-evoked currents (Verheugen et al. 1999). After rupture of the patch, cells were filled and processed for biocytin immunostaining and morphological identification. Cells in which the access after patch rupture was inadequate were repatched with a gramicidin-free internal solution for biocytin fill. Tight-seal cell-attached recordings were obtained from FS-BCs with electrodes containing the gramicidin-free KCl-based internal solution in standard aCSF. Recordings were obtained in current-clamp mode with zero current injection. At the end of the recordings, the patch was ruptured to gain whole cell access for physiological identification and biocytin filling for post hoc morphological identification.

Field recordings were performed in an interface recording chamber (BSC2, AutoMate Scientific, Berkeley, CA) perfused with aCSF. Brain slices (400 μm) rested on filter paper and were stabilized with platinum wire weights. The tissue was continuously superfused with humidified 95% O$_2$-5% CO$_2$, and the temperature of the perfusing solution was maintained at 34°C with a proportional control heating unit (PTC03, AutoMate Scientific). Field recordings of evoked population spikes in the granule cell layer of the dentate gyrus were obtained with patch pipettes filled with recording aCSF. To evoke the field responses, constant-current stimuli (0.5–4 mA, 50 μs) were applied at 0.1 Hz through a bipolar 90-voltage drop across series resistance. The tissue was continuously superfused with 10% NGS and 0.3% Triton X-100 in PBS at RT for 1 h and incubated. In a subset of cells, $E_{\text{GABA}}$ was applied at 0.1 Hz through a bipolar 90-voltage drop across series resistance. In a subset of cells, 1748 STATUS EPILEPTICUS ALTERS DENTATE BASKET CELL TONIC INHIBITION

Williston, VT) on an Olympus BX51 microscope with a 100 oil objective with identical camera settings. Semiquantitative analysis of $G_{\text{GABA}}$ in the PV-labeled soma. Neurons were deemed colabeled if the staining for $G_{\text{GABA}}$ in the PV-labeled soma shared the outline of the PV-labeled soma and had a greater intensity than the hilar neuropil. The percentage of PV-labeled cells that were colabeled for $G_{\text{GABA}}$ was determined. Single-plane confocal images for illustration were obtained with a Nikon A1R laser confocal microscope with a ×60 water objective and identical camera settings. Semiquantitative analysis of $G_{\text{GABA}}$ and KCC2 fluorescence intensity in PV+ neurons was performed on images from an equal number of randomly selected PV+ neurons in the hilar-granule cell layer border from each section. Images were obtained with a Nikon A1R laser confocal microscope with a 1.2 NA ×60 water objective with identical camera settings and converted to R3G color mode. An ROI was traced around PV+ neurons (in the red channel), and the average grayscale intensity of $G_{\text{GABA}}$ was determined in the green channel (Δ-subunit). For estimation of KCC2 fluorescence intensity in PV+ neurons, the ROI was confined to the periphery of the PV-labeled profile in order to assess membrane expression of KCC2. Image analysis was performed with ImageJ v1.43u (National Institutes of Health) by an investigator blind to the treatment.

After physiological recordings, slices were fixed in 0.1 M phosphate buffer containing 4% paraformaldehyde at 4°C for 2 days. For post hoc immunohistochemistry, thick slices (300 μm) were incubated overnight at RT with anti-PV antibody (PV-28, 1:5,000, polyclonal rabbit, Swant) in 0.3% Triton X-100 and 2% NGS-containing PBS. Immunoreactions were revealed with Alexa Fluor 488-conjugated secondary goat antibodies against rabbit IgG (1:250), and biocytin staining was revealed with Alexa Fluor 594-conjugated streptavidin (1:1,000). Sections were visualized and imaged with a Nikon A1R laser confocal microscope with a 1.2 NA ×60 water objective. As a result of prolonged recordings and use of high-chloride internal solution, few cells showed somatic labeling for PV. When present, the expression of PV in the soma or dendrites was used as an added confirmation of cell identity. Cell reconstructions and morphological analyses were performed with Neurolucida V.10.02 (MBF Biosciences) and confocal image stacks.

Computational modeling. Single FS-BC models and dentate network simulations were implemented with the NEURON 7.0 simulation environment (Hines and Carnevale 1997). The biophysically realistic FS-BC model was adapted from earlier studies (Dyrhjeld-Johnsen et al. 2007; Santhakumar et al. 2005) and included a soma and two apical and basal dendrites each with four distinct compartments (a total of 17 compartments). Active and passive conductances were distributed as detailed previously (Santhakumar et al. 2005).
Sodium and fast delayed-rectifier potassium channels were restricted to the soma and proximal dendrites. Reversal potential of a nonspecific leak channel was set to ~75 mV to modify the basket cell RMP to match the data from perforated-patch recordings in the present study. Conductance of the nonspecific leak channel was not altered. Tonic \( g_{\text{GABA}} \) was modeled as a linear deterministic leak conductance with reversal \( (E_{\text{GABA}}) \) based on experimental data: ~74 mV or ~54 mV. Tonic \( g_{\text{GABA}} \) was distributed uniformly in all compartments and varied from 0 to 0.1 mS/cm². In some simulations, tonic \( g_{\text{GABA}} \) was restricted to the soma and proximal dendrite to determine whether the distribution of tonic \( g_{\text{GABA}} \) altered the magnitude of tonic \( I_E \) or \( R_m \) in model FS-BCs. In simulations performed to examine the biologically relevant range of tonic \( g_{\text{GABA}} \) (see Fig. 10B), FS-BCs were simulated with a somatic voltage clamp and \( E_{\text{GABA}} \) was set to 0 mV to model symmetrical chloride of our physiological recordings. Model FS-BCs were voltage clamped at ~70 mV, tonic \( g_{\text{GABA}} \) was varied between 0 and 5 mS/cm², and tonic \( I_E \) or \( R_m \) was measured as the difference in baseline current in the presence and absence of tonic \( g_{\text{GABA}} \). \( R_m \) was measured in response to ~100-pA current injection. To examine the effect of tonic \( I_E \) or \( R_m \) on excitability, model FS-BCs were activated by identical 200-Hz Poisson-distributed trains of excitatory synaptic inputs to the apical distal dendrite. Excitatory synaptic parameters were based on AMPA conductances in previous studies (Santhakumar et al. 2005), with the synaptic AMPA conductance \( (g_{\text{AMPA}}) \) set at 3 nS to simulate low activity levels and \( g_{\text{AMPA}} \) = 20 nS to simulate high activity levels. The effect of increasing tonic \( g_{\text{GABA}} \) on evoked firing of the model FS-BC was examined with \( E_{\text{GABA}} \) set at ~74 mV (control) and ~54 mV (after SE).

The large-scale, topologically and biophysically constrained model network used in this study was adapted from the 500-cell network described by Santhakumar et al. (2005) and expanded to include 1,000 granule cells, 30 mossy cells, 12 basket cells, and 12 hilar interneurons. The multicompartmental single-cell models, distribution and magnitude of active and passive properties, and synaptic conductances were based on Santhakumar et al. (2005) and were derived from anatomical and physiological data in the literature. The networks were topographically constrained, incorporating the axon distribution of the cell types, and simulated by distributing the neurons in a ring structure to avoid edge effects. Enhanced excitability in the early post-SE condition was modeled by including mossy fiber sprouting, simulated by adding synaptic connections from granule cells to the proximal dendrites of granule cells. Since the simulations were designed to test early stages of network excitability (1 wk after SE), the degree of mossy fiber sprouting in the model was set at 20% of the maximal sprouting observed in the pilocarpine model of epilepsy (Dyhrfjeld-Johnsen et al. 2007). In a second set of simulations, both sprouting and hilar neuronal loss were simulated with 20% sprouting as detailed above and a corresponding deletion of 20% of randomly selected hilar mossy cells and dendritically projecting interneurons as described in earlier studies (Santhakumar 2008; Santhakumar et al. 2005). Network models were simulated with 2.5-Hz spontaneous activity in all 1,000 granule cells during the entire duration of the simulation (3,500 ms). Spontaneous activity was simulated as independent Poisson-distributed spike trains to the perforant path input (Dyhrfjeld-Johnsen et al. 2007). A single, synchronous synaptic input to 100 granule cells and 2 local basket cells (at \( t = 2,001 \) ms) was used to simulate focally evoked network firing. The basket cell models (and only basket cell models) included tonic \( g_{\text{GABA}} \) (described above in the single-cell model) distributed uniformly in all compartments. In networks simulated with control GABA reversal, \( E_{\text{GABA}} \) of both tonic and synaptic GABA currents in all model FS-BCs was set to ~74 mV. Similarly, networks with the post-SE depolarized GABA reversal were simulated with \( E_{\text{GABA}} \) for tonic and synaptic GABA currents in basket cells set to ~54 mV. In a subset of simulations including sprouting and hilar cell loss (see Fig. 11E), \( E_{\text{GABA}} \) of synaptic and tonic \( g_{\text{GABA}} \) were set to different values. The \( E_{\text{GABA}} \) at GABA synapses on other neurons was not altered. Simulations included “synaptic spillover,” modeled to simulate increases in tonic \( g_{\text{GABA}} \) that accompany increases in extracellular GABA levels during neuronal activity (Glyksys and Mody 2007). Spillover was modeled by including three “spillover GABA conductances” with progressively decreasing amplitudes associated with each inhibitory synaptic connection to model FS-BCs. The spillover GABA conductances were modeled as synapses with slower rise (7 ms) and decay time (200 ms) constants (Rossi et al. 2003) and had 20%, 10%, and 5% of the peak conductance of the primary synapse. The effect of basket cell \( g_{\text{GABA}} \) and \( E_{\text{GABA}} \) on the average frequency of granule cell spontaneous activity and evoked activity were quantified during 1,000–2,000 ms and 2,001–3,500 ms, respectively. In initial simulations, instantiation of the network connections was randomized within preset topological constraints. Each set of simulations was run multiple times, and summary data are presented as means ± SE. In some simulations the random seed of the network connectivity was set to a specific value, and the effect of tonic \( g_{\text{GABA}} \) and \( E_{\text{GABA}} \) on dentate excitability was compared in structurally identical networks.

**Analysis and statistics.** Statistical analysis was performed by paired and unpaired Student’s t-test (Microsoft Excel 2007) or Kolmogorov-Smirnov (K-S) test (in IGOR Pro 7.0) for data that were not distributed normally or univariate and multivariate repeated-measures ANOVA (Systat) for experiments involving repeated measurements from the same sample. Significance was set to \( P < 0.05 \). Data are shown as mean ± SE or median and interquartile range (IQR) where appropriate.

**RESULTS**

**Early dentate cell loss and enhanced excitability in young adult rats after status epilepticus.** A single episode of SE following injection of pilocarpine leads to development of recurrent spontaneous seizures and has been used to model acquired epilepsy. In adult rats, pilocarpine-induced SE replicates the characteristic hippocampal cell loss and network reorganization observed in epileptic patients. The presence and extent of cell loss in younger rats may be variable (Raol et al. 2003). Therefore, we performed NeuN staining for neuronal nuclei in tissue from control rats and those subjected to SE to examine whether SE in the young adult rats (postnatal days 25–27) used in the present study resulted in dentate hilar cell loss. As illustrated in Fig. 1, comparison of sections prepared 1 wk after pilocarpine-induced SE (post-SE) and from age-matched saline-injected control rats revealed a significant decrease in NeuN-stained neurons in the dentate hilus after SE (Fig. 1, A and B; control: 377.5 ± 40.4 hilar neurons counted/section, a total of 4,251 cells counted in 15 sections from 3 rats; post-SE: 225.7 ± 18.8 hilar neurons counted/section, based on 2,789 cells counted in 15 sections from 3 rats; 40.2 ± 5.0% decrease, \( P < 0.05 \), Student’s t-test) consistent with observations in previous studies (Kobayashi et al. 2003; Mello et al. 1993; Zhang et al. 2012). Additionally, hippocampal sections from rats 1 wk after SE but not control rats had degenerating neurons stained by Fluoro-Jade C in CA1 and CA3 (data not shown), as observed in previous studies (Ekstrand et al. 2011). These data demonstrate that the young adult rats used in the present study show hilar cell loss within a week after SE. Previous studies have identified changes in dentate excitability and granule cell inhibition 1–2 wk after SE and prior to onset of epileptic seizures (Pathak et al. 2007). We examined the population spike (Fig. 1C) of dentate granule cells in response to perforant path stimulation to determine whether dentate network excitability was altered 1 wk after SE. Field recordings showed that the amplitude of the afferent-evoked granule cell population spike was enhanced 1 wk...
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Fig. 1. Early changes in dentate network function after pilocarpine-induced status epilepticus (SE). A and B: photomicrographs of NeuN-stained sections obtained from rats perfused 1 wk after saline injection or pilocarpine-induced SE demonstrate the presence of numerous NeuN-stained hilar neurons in the section from the control (A) and fewer hilar NeuN-stained neurons in a level-matched post-SE section (B). GCL, granule cell layer. Scale bars, 200 µm. C: representative traces of granule cell field responses evoked by perforant path stimulation in slices from control (top) and 1 wk after SE (bottom) illustrate the larger population spike amplitude in the post-SE dentate. Traces are an average of 4 trials in response to a 4-mA stimulus to the perforant path. Arrowheads indicate the location of the truncated stimulus artifact, and arrows point to the population spike. D: summary data demonstrate the post-SE increase in dentate-evoked excitability at various stimulation intensities. Error bars indicate SE. *P < 0.05 by repeated-measures ANOVA.

after SE compared with age-matched control rats (Fig. 1C). Summary data demonstrate the post-SE increase in dentate population spike amplitude at various stimulation intensities [Fig. 1D; control: n = 8 slices from 4 rats, post-SE: n = 13 slices from 6 rats; F(1,19) = 8.93, P < 0.05 by univariate repeated-measures ANOVA]. Even when the population spike amplitude was normalized by the simultaneously recorded field excitatory postsynaptic potential (fEPSP) slope, to account for differences in effective excitatory synaptic drive in response to stimulation, the fEPSP-normalized population spike amplitude was significantly greater in slices from post-SE rats [F(1,19) = 4.8, P < 0.05 by repeated-measures ANOVA]. Since identifying early SE-induced changes in dentate inhibition can provide mechanistic insights into dentate physiology in the latent period leading up to spontaneous seizures, we focused on inhibitory plasticity of perisomatically projecting PV-expressing basket cells 1 wk after SE.

GABA$_A$R δ-subunits are expressed in parvalbumin-positive dentate interneurons. In dentate granule cells, SE leads to alterations in synaptic and extrasynaptic GABA$_A$R expression and in synaptic and tonic I$_{GABA}$ (Mitchelldishvili and Kapur 2006; Peng et al. 2004; Zhan and Nadler 2009; Zhang et al. 2007). Hippocampal and dentate interneurons appear to express GABA$_A$Rs underlying tonic I$_{GABA}$ and demonstrate tonic I$_{GABA}$ (Glykys et al. 2007; Semyanov et al. 2003; Song et al. 2011). Studies in mice have revealed that GABA$_A$R δ-subunits, known to underlie tonic I$_{GABA}$, are expressed in presumed interneurons in the hilar-granule cell layer border (Peng et al. 2004). However, whether perisomatically projecting basket cells, which express PV and are critical for rapid and precise inhibition (Freund 2003; Hefft and Jonas 2005), express GABA$_A$R δ-subunits has not been determined. Confocal images of hippocampal sections, obtained from rats 1 wk after saline injection or pilocarpine-induced SE and immunostained for PV and GABA$_A$R δ-subunits, showed that PV-labeled neurons in the hilar-granule cell layer border (Fig. 2, A–D, right) consistently demonstrated somatic labeling for GABA$_A$R δ-subunits (Fig. 2, A and C, center). The intense GABA$_A$R δ-subunit labeling of granule cell dendrites made it difficult to isolate GABA$_A$R δ-subunit labeling of PV+ dendrites in the molecular layer. However, PV+ dendrites in the hilus of both the control (Fig. 2B) and post-SE (Fig. 2D) rats showed distinct colabeling for GABA$_A$R δ-subunit, consistent with somato-dendritic staining for GABA$_A$R δ-subunit observed in other neuronal types (Olah et al. 2009; Zhang et al. 2007). Absence of immunostaining in sections from Gabrd$^{	ext{−/−}}$ mice lacking the GABA$_A$R δ-subunit was used to confirm specificity of the GABA$_A$R δ-subunit primary antibody (data not shown). As illustrated by the summary plots, 99.1 ± 0.9% of the 115 PV+ neurons from control rats (18 slices from 3 rats) and 97.1 ± 2.9% of the 120 PV+ neurons from post-SE rats (18 slices from 3 rats) in the hilar-granule cell layer border (see MATERIALS AND METHODS) were colabeled for GABA$_A$R δ-subunit (Fig. 2E). The proportion of PV+ neurons labeled for the GABA$_A$R δ-subunit was not altered after SE (Fig. 2F; P > 0.05 by Student’s t-test). However, quantification of the fluorescence intensity for GABA$_A$R δ-subunit expression in PV-labeled neurons from control and post-SE rats revealed a significant increase in expression of GABA$_A$R δ-subunit after SE (Fig. 2F; grayscale intensity in arbitrary units: control 14.7 ± 5.7, n = 65 cells from 3 rats; post-SE 24.2 ± 15.8, n = 70 cells from 3 rats, P < 0.05 by Student’s t-test). These data demonstrate expression and post-SE enhancement of GABA$_A$R δ-subunits in PV+ interneurons in the hilar-granule cell layer border.

Expression of tonic GABA currents in dentate fast-spiking basket cells. In light of our immunostaining data demonstrating the expression of GABA$_A$R δ-subunits in PV+ interneurons (Fig. 2), we examined whether PV+ basket cells express functional tonic I$_{GABA}$. Dentate PV+ basket cells can be distinguished from the other interneurons in the hilar-granule cell
layer border on the basis of their characteristic morphological and physiological properties. Previous studies have established that dentate PV+ basket cells have axon collaterals largely localized in the granule cell layer, whereas axons of CCK- and somatostatin-expressing interneurons project to the molecular layer (Buckmaster et al. 2002; Hefft and Jonas 2005). Thus the distinctive axonal distribution in the granule cell layer can be used to morphologically identify PV+ interneurons. Apart from morphology, the typical high-frequency, nonadapting firing during depolarizing current injections has been used as a physiological marker of PV+ basket cells (Harney and Jones 2002; Hefft and Jonas 2005; Zhang and Buckmaster 2009).

Figure 3A illustrates a reconstructed FS-BC with the axon in the granule cell layer. The cell was filled during physiological recordings and processed for post hoc biocytin immunostaining. The FS-BC reconstructed in Fig. 3A, top, demonstrated characteristic high-frequency, nonadapting firing in response to a 500-pA depolarizing current injection, with low $R_{in}$ and numerous synaptic events during −100-pA hyperpolarizing current injection (Fig. 3A, bottom, and insets in Figs. 4B and 5A). Additionally, colabeling of the dendrites of the recorded neuron for parvalbumin (Fig. 3A, inset) confirm the identity of the cell as a PV+ FS-BC. Although our routine immunostaining identified PV labeling in dendrites of biocytin-filled FS-BCs, most FS-BCs did not show somatic labeling for PV. It is likely that the combination of high-chloride internal solution and long-duration recording contributed to the difficulty in detecting somatic PV labeling. Consequently, all FS-BCs in the present study were identified on the basis of a combination of fast-spiking, nonadapting physiology, the presence of axon collaterals in the granule cell layer, and dendritic PV labeling. In addition to FS-BCs, we recorded from non-FS-INs with somata in the hilar-granule cell layer border and axon collaterals distributed in the molecular layer (Fig. 3B, top). Physiological recordings from the same cell (Fig. 3B, bottom) illustrate the adapting firing pattern during positive current injection (+200 pA) and the presence of depolarizing sag and high $R_{in}$ in response to negative current injections (−100 pA).
Although the RMPs of FS-BCs and non-FS-INs were not different (RMP in mV: FS-BC \(H_11002^74.0\) \(H_11006^74.0\) in \(n_11005^10\) cells; non-FS-IN \(H_11002^72.3\) \(H_11006^2.4\) in \(n_11005^12\) cells, \(P_11022^0.05\) by Student’s \(t\)-test), the frequency of FS-BC firing in response to a \(800\)-pA current injection was significantly greater (frequency in Hz: FS-BC \(H_11006^112.1\) \(H_11005^7.9\) in \(n_11005^12\) cells; non-FS-IN \(H_11006^52.0\) \(H_11005^5.7\) in \(n_11005^8\) cells, \(P_11021^0.05\) by Student’s \(t\)-test) and the \(R_{in}\) lower (\(R_{in}\) in MΩ: FS-BC \(93.0\) \(10.6\) in \(n_11005^12\) cells; non-FS-IN \(233.3\) \(6.8\) in \(n_11005^8\) cells, \(P_11021^0.05\) by Student’s \(t\)-test) than in non-FS-IN.

Together, these morphological and physiological characteristics are consistent with earlier studies (Harney and Jones 2002) and were used to reliably distinguish FS-BCs from a potentially diverse set of non-FS-INs projecting to the molecular layer.

**Fig. 3.** Morphological and physiological characterization of dentate interneurons projecting to granule cell somata and dendrites. *A,* top: reconstruction of a fast-spiking basket cell (FS-BC) filled during recordings shows the typical morphology with soma and dendrites in blue and axon in GCL in black. ML, molecular layer. Bottom: membrane voltage traces from the same cell illustrate the fast-spiking, nonadapting firing pattern during a \(+500\) pA current injection and relatively low membrane hyperpolarization in response to a \(-100\) pA current injection. *Inset:* confocal image of biocytin-filled (BIO) soma and dendrites (arrowheads) of the cell in *A* (top) and labeling for PV in the dendrites (*middle*); *bottom:* merged image showing PV colabeling in the biocytin-filled dendrites (arrowheads). Scale bar, \(100\) μm. B, *top:* Neurolucida reconstruction of a non-fast-spiking interneuron (non-FS-IN) with axon in the ML. *Bottom:* membrane voltage traces from the same cell show the typical adapting firing during a \(+200\) pA current injection and membrane hyperpolarization and depolarizing sag (arrowhead) during a \(-100\)-pA current injection. Note the difference in membrane hyperpolarization in response to \(-100\)-pA current injections between the FS-BC (*A*) and non-FS-IN (*B*).

**Fig. 4.** Expression of tonic GABA currents (\(I_{GABA}\)) in dentate FS-BCs. *A:* representative voltage-clamp recordings (\(V_{hold} = -70\) mV) from a FS-BC in the hilar-GCL border illustrates the presence of tonic \(I_{GABA}\) blocked by SR95531 (gabazine, \(10\) μM). *B:* expanded 30-s traces of the boxed area in *A.* Gaussian fits to all-points histograms derived from the illustrated recording periods in control conditions, in the presence of \(3\) mM kynurenic acid (KyA), and after the addition of gabazine used to determine tonic current amplitude are shown on right. Dashed lines indicate Gaussian means, and the difference currents are noted. *Inset:* membrane voltage trace from the same cell shows fast-spiking firing. *C:* representative voltage-clamp recordings (\(V_{hold} = -70\) mV) from a non-FS-IN in the hilar-GCL border illustrates lack of tonic \(I_{GABA}\) on blocking GABAAR with SR95531 (\(10\) μM). *D:* expanded 30-s traces of the boxed area in *C.* Gaussian fits to all-points histograms used to determine tonic current amplitude are presented on right. *Inset:* membrane voltage trace from the cell in *C* and *D* shows adapting firing. *E:* summary data of tonic \(I_{GABA}\) amplitude in \(3\) mM kynurenic acid in FS-BCs and non-FS-IN. Individual data points are represented by gray dots. \(*P < 0.05\) by unpaired Student’s \(t\)-test.
We examined interneurons in the hilar-granule cell layer border for the presence of tonic $I_{\text{GABA}}$. Recordings were performed in the presence of the glutamate receptor antagonist KyA (3 mM), and tonic $I_{\text{GABA}}$ was measured as the baseline current blocked by the GABA$_A$R antagonist SR95531 (gabazine, 10 $\mu$M). Recordings were obtained at physiological temperature and in the absence of added GABA or GABA transporter inhibitors. As illustrated by representative recordings from an FS-BC in Fig. 4A and the magnified 30-s segments of the boxed areas in Fig. 4A (Fig. 4B), SR95531 caused a small but significant shift in the baseline current, indicating the presence of tonic $I_{\text{GABA}}$ in FS-BCs ($P < 0.05$, paired Student’s $t$-test). In contrast, non-FS-INS lacked tonic $I_{\text{GABA}}$ and showed little shift in baseline after addition of SR95531 (Fig. 4, C and D; $P > 0.05$, paired Student’s $t$-test). On average the magnitude of tonic $I_{\text{GABA}}$ in morphologically and physiologically identified FS-BCs was significantly greater than in non-FS-INS (Fig. 4E; tonic $I_{\text{GABA}}$ in pA: FS-BC $5.7 \pm 0.9$, $n = 17$; non-FS-INS: $0.5 \pm 0.9$, $n = 9$; $P < 0.05$ by Student’s $t$-test). Thus, consistent with the anatomical data showing labeling of GABA$_A$R $\delta$-subunits with PV+ interneurons (Fig. 2), our physiological data demonstrate that dentate FS-BCs express tonic $I_{\text{GABA}}$. Additionally, the results demonstrate that the magnitude of tonic $I_{\text{GABA}}$ in FS-BCs is greater than in non-FS-INS.

Status epilepticus enhances FS-BC tonic $I_{\text{GABA}}$. Since our immunostaining data (Fig. 2) indicate that the expression of GABA$_A$Rs containing $\delta$-subunits (Brown et al. 2002), enhanced tonic $I_{\text{GABA}}$, confirming the contribution of GABA$_A$R $\delta$-subunits (Fig. 5, A and C; tonic $I_{\text{GABA}}$ in pA: $5.3 \pm 2.0$ in KyA and $11.5 \pm 2.7$ in THIP, $n = 7$ cells, $P < 0.05$, paired Student’s $t$-test). Similarly, THIP enhanced tonic $I_{\text{GABA}}$ in FS-BCs from post-SE rats (Fig. 5, B and C; tonic $I_{\text{GABA}}$ in pA: $14.0 \pm 2.1$ in KyA and $21.9 \pm 2.1$ in THIP, $n = 7$ cells, $P < 0.05$, paired Student’s $t$-test). Importantly, the magnitude of FS-BC tonic $I_{\text{GABA}}$, both in KyA and in THIP, was significantly greater in post-SE rats compared with control rats (Fig. 5, A–C; $P < 0.05$, $t$-test). Application of THIP (1 $\mu$M) caused a significantly greater increase in baseline currents in FS-BCs from post-SE rats (baseline current increase in THIP in pA: control FS-BCs: $6.0 \pm 1.4$, $n = 8$ cells; post-SE FS-BCs $10.8 \pm 1.6$, $n = 10$ cells, $P < 0.05$ by Student’s $t$-test), confirming that increases in membrane expression of GABA$_A$R $\delta$-subunits contribute to post-SE increase in FS-BC tonic $I_{\text{GABA}}$. To eliminate the possibility that THIP, like GABA, may alter GABA$_A$R antagonist binding (Bianchi and Macdonald 2001) and confound estimation of tonic $I_{\text{GABA}}$, we also measured tonic $I_{\text{GABA}}$ in control aCSF in the presence of KyA and without perfusion of THIP. When normalized to cell membrane capacitance, to eliminate confounding effects due to differences in cell size, tonic $I_{\text{GABA}}$ in FS-BCs from post-SE rats was greater than those from control rats (Fig. 5D; tonic $I_{\text{GABA}}$ current density in pA/pF: control $0.10 \pm 0.02$, $n = 11$ cells; post-SE $0.23 \pm 0.05$, $n = 7$ cells, $P < 0.05$, Student’s $t$-test). Moreover, FS-BC tonic $I_{\text{GABA}}$ measured in the presence of the GABA transporter antagonist NO-711 (10 $\mu$M), to abolish potential post-SE differences in GABA transporter function, was also enhanced after SE (Fig. 5E; tonic $I_{\text{GABA}}$ in pA: control $14.7 \pm 2.6$, $n = 6$ cells; post-SE $24.6 \pm 3.1$, $n = 6$ cells, $P < 0.05$, Student’s $t$-test). Comparison of the intrinsic properties measured during whole cell recordings revealed no change in either $R_{\text{in}}$ or RMP between FS-BCs from control and post-SE rats ($R_{\text{in}}$ in MΩ: control $93.0 \pm 10.6$, $n = 12$ cells; post-SE $97.6 \pm 11.2$, $n = 9$ cells, $P > 0.05$, Student’s $t$-test; RMP in mV: control $-74.0 \pm 1.9$, $n = 10$ cells; post-SE $-75.0 \pm 2.4$, $n = 10$ cells, $P > 0.05$, Student’s $t$-test). $R_{\text{in}}$ was measured in response to a $-100$-pA current injection. Our measurements of FS-BC tonic $I_{\text{GABA}}$ demonstrate, unequivocally, that tonic $I_{\text{GABA}}$ is present in FS-BCs and is enhanced after SE. Our results indicate that SE-induced increases in the expression of GABA$_A$Rs with $\delta$-subunits likely contribute to increases in FS-BC tonic $I_{\text{GABA}}$ after SE.

Fig. 5. Tonic $I_{\text{GABA}}$ in dentate FS-BCs are enhanced after SE. A and B: segments (30 s) of representative voltage-clamp recordings ($I_{\text{GABA}} = -70$ mV) from control (A) and post-SE (B) FS-BCs illustrate the enhancement of tonic $I_{\text{GABA}}$, by addition of 4,5,6,7-tetrahydroisoaxolo[5,4-c]pyridin-3-ol (THIP, 1 $\mu$M). Tonic $I_{\text{GABA}}$ was measured as the current blocked by SR95531 (10 $\mu$M). Panels on right show Gaussian fits to all-points histograms of the 30-s recording periods in 3 mM kynurenic acid, after the addition of THIP (1 $\mu$M), and in SR95531. Dashed lines show Gaussian means, and the difference currents are noted. Insets: membrane voltage traces show fast-spiking firing of the respective cells. C: summary of the magnitude of FS-BC tonic $I_{\text{GABA}}$ in 3 mM kynurenic acid and after perfusion of THIP (1 $\mu$M) in controls and after SE. D: histogram presents baseline tonic $I_{\text{GABA}}$ recorded in 3 mM kynurenic acid normalized to the cell membrane capacitance. E: tonic $I_{\text{GABA}}$ in control and post-SE FS-BCs measured with the GABA transporter antagonist NO-711 (10 $\mu$M) in the presence of 3 mM kynurenic acid. *$P < 0.05$ by paired and unpaired Student’s $t$-test.
Spillover of synaptically released GABA into the extrasynaptic space is known to contribute to tonic $I_{GABA}$ (Glykys and Mody 2007). Therefore, one possibility is that the post-SE enhancement of tonic $I_{GABA}$ reflects post-SE increases in synaptic GABA release. Previous studies have demonstrated that in granule cells the frequency of sIPSCs is decreased after SE (Kobayashi and Buckmaster 2003), discounting the possible contribution of enhanced GABA spillover from inhibitory synapses to granule cells. To examine whether increases in synaptic GABA release to FS-BCs could contribute to the observed enhancement of tonic $I_{GABA}$ after SE, we analyzed the frequency of sIPSCs in cells examined for changes in tonic $I_{GABA}$ (in Fig. 5). As illustrated by representative traces (Fig. 6, A and B), there was a considerable decrease in the frequency of sIPSCs in FS-BCs from post-SE rats (Fig. 6B; sIPSC frequency in Hz: control median = 7.0, IQR = 3.7–13.7, mean ± SE = 10.9 ± 0.5, n = 10 cells; post-SE median = 5.0, IQR = 2.6–10.6, mean ± SE = 8.9 ± 0.6, n = 7 cells, $P < 0.05$, K-S test). Additionally, FS-BC sIPSC amplitude was reduced (in nP: control median = 100 ± 3.0, n = 10 cells; post-SE median = 50 ± 2.6, n = 7 cells, $P < 0.05$, K-S test). To avoid perturbations of the intracellular chloride concentration, we used gramicidin-perforated-patch recordings (Bil-lups and Atwell 2002; Pathak et al. 2007) and measured the RMP and reversal potential of GABA-evoked currents in parallel within the same FS-BC. Upon establishment of adequate access (see MATERIALS AND METHODS), we first determined the neuronal firing pattern in response to depolarizing current injections before perfusion of TTX (1 $\mu$M) to block action potential generation. Next, in the absence of added GABA, we recorded FS-BCs current responses to −130 to +10 mV voltage ramps from a holding potential of −60 mV (Fig. 7A). The potential at which the holding current was 0 pA was measured as the RMP (Fig. 7A). In interleaved recordings preformed at 40-s intervals, voltage ramps were repeated.

**Fig. 6.** Decrease in FS-BC spontaneous inhibitory post-synaptic current (sIPSC) frequency after SE: A: representative traces of voltage-clamp recordings from control (top) and post-SE (bottom) FS-BCs show the higher sIPSC frequency in the control FS-BC. Note the decrease in sIPSC frequency in the recording from the post-SE FS-BC. B and C: cumulative probability plot of the sIPSC instantaneous frequency (B) and charge transfer (C) in control and post-SE FS-BCs measured with symmetrical chloride from a holding potential of −70 mV in kynurenic acid (3 mM). The same number of individual events was selected from each cell to develop the cumulative probability distribution (control: n = 10 cells; post-SE: n = 7 cells).
during local pressure application of GABA (100 μM). The potential at which the currents generated during the voltage ramp in the absence and the presence of GABA crossed was measured as $E_{\text{GABA}}$. As shown in Fig. 7Aa, inset, RMP in control FS-BCs was not different from the potential at which the current traces obtained in the presence and absence of GABA crossed (in mV: $E_{\text{GABA}} = -74.2 ± 8.8$, RMP = $-77.6 ± 10.7$, in $n = 6$ cells, $P > 0.05$ by paired Student’s $t$-test). These data suggest that GABA$_A$R activation likely contributes to shunting inhibition in control FS-BCs. In control experiments, we found no difference between $E_{\text{GABA}}$ measured with voltage ramp and voltage step protocols in the same cell (in mV: $E_{\text{GABA}}$ with voltage ramp $= -77.28 ± 2.89$, $E_{\text{GABA}}$ with voltage step $= -80.0 ± 3.68$, in $n = 4$ cells, $P > 0.05$ by paired Student’s $t$-test), indicating that the voltage ramp protocol can be used to reliably estimate $E_{\text{GABA}}$. In contrast to control rats, $E_{\text{GABA}}$ measured in FS-BCs from post-SE rats was significantly more depolarized than RMP (Fig. 7Ab; in mV: $E_{\text{GABA}} = -53.3 ± 4.0$, RMP $= -73.7 ± 4.9$ in $n = 7$ cells, $P < 0.05$ by paired Student’s $t$-test). Overlapping interleaved traces from control experiments performed under identical conditions in the absence and during pressure application of vehicle without GABA confirmed the absence of pressure artifacts under our recording conditions (Fig. 7Ac).

We directly tested whether activation of GABA receptors leads to membrane depolarization in FS-BCs from post-SE rats. Although cell-attached recordings from FS-BCs showed spontaneous firing when external potassium was elevated to 3 mM and above (not shown), as observed by Fricker et al. (1999), few interneurons fired spontaneously under our recording conditions with 2.5 mM external potassium. To avoid confounding effects due to changes in RMP, cell-attached recordings from FS-BCs were obtained in standard aCSF containing 2.5 mM potassium. Cell-attached recordings showed that while pressure application of the GABA$_A$R agonist muscimol (50 μM) caused little change in membrane potential in control FS-BCs, it consistently depolarized FS-BCs from post-SE rats (Fig. 8; maximum membrane potential change in muscimol in mV: control $0.3 ± 1.3$, $n = 5$; post-SE $7.6 ± 2.1$, $n = 8$, $P < 0.05$ by Student’s $t$-test). Together, comparison of RMP and $E_{\text{GABA}}$ in FS-BCs from control and post-SE rats under identical experimental conditions suggests that activation of extrasynaptic GABA$_A$Rs may be shunting under control conditions and result in depolarizing currents after SE.

Previous studies have identified that FS-BC synaptic $E_{\text{GABA}}$ undergoes a developmental hyperpolarizing shift and have suggested that increases in the expression of the potassium-chloride cotransporter KCC2 (Sauer and Bartos 2010) may underlie this effect. Since changes in expression of KCC2 have been shown contribute to post-SE alterations in granule cell $E_{\text{GABA}}$ (Pathak et al. 2007), we examined FS-BCs for post-SE changes in KCC2 expression. Examination of sections immunostained for PV and KCC2 revealed the presence of KCC2 localized to the periphery of PV+ profiles in both control and post-SE rats (Fig. 9). Quantification of the fluorescence intensity for KCC2 subunit expression in PV+ neurons identified a significant post-SE decrease in KCC2 along the margins of PV+ profiles in the hilar-granule cell layer border (grayscale intensity in arbitrary units: control $15.7 ± 2.6$, $n = 29$ cells from 15 sections in 3 rats; post-SE $9.8 ± 1.1$, $n = 36$ cells from 15 sections in 3 rats, $P < 0.05$ by Student’s $t$-test). The observed decrease in KCC2 expression in PV+ neurons likely contributes to the depolarizing shift in FS-BC $E_{\text{GABA}}$ after SE.

**Impact of tonic $g_{\text{GABA}}$ and $E_{\text{GABA}}$ on FS-BC excitability: isolated basket cell simulations.** How do the presence and post-SE changes in tonic $g_{\text{GABA}}$, together with the depolarized $E_{\text{GABA}}$, modify FS-BC excitability? Given the concurrent changes in...
conductance and reversal of tonic $I_{\text{GABA}}$, and potential confounding SE-induced changes in the intrinsic physiology of FS-BCs (unpublished observations) that are as yet unknown, it is difficult to isolate how alterations in FS-BC tonic $I_{\text{GABA}}$, both singly and when coupled with depolarized $E_{\text{GABA}}$, modify network activity in biological experiments. Therefore, we adopted computational modeling to determine how systematic changes in FS-BC tonic $I_{\text{GABA}}$, in the presence and absence of the observed depolarizing shift in GABA reversal, influence FS-BC and dentate network excitability. A recent study conducted with generic single compartmental models showed that when $E_{\text{GABA}}$ is depolarizing, tonic $I_{\text{GABA}}$ increases neuronal excitability only in a narrow range of $I_{\text{GABA}}$. With additional increases in tonic $I_{\text{GABA}}$, the shunting effect of the conductance overwhelms the depolarizing currents, resulting in net inhibition (Song et al. 2011). To systematically examine how post-SE changes in tonic $g_{\text{GABA}}$ and $E_{\text{GABA}}$ influence FS-BC excitability, we implemented biophysically realistic multicompartmental simulations of FS-BCs (Santhakumar et al. 2005). As illustrated in Fig. 10A, the model FS-BC fired at a frequency of 100 Hz in response to a 800-pA current injection, simulating the experimentally determined characteristic high-frequency firing and low $R_{\text{in}}$ of biological dentate FS-BCs (for detailed comparison of the active and passive properties of the model FS-BC and biological basket cells see Table 3 in Santhakumar et al. 2005). To determine the magnitude of tonic $g_{\text{GABA}}$ that generates the biologically observed tonic $I_{\text{GABA}}$, we simulated voltage-clamp recordings in model FS-BCs and examined how increases in tonic $g_{\text{GABA}}$ influence the magnitude of tonic $I_{\text{GABA}}$ and $R_{\text{in}}$. Systematically increasing tonic $g_{\text{GABA}}$ from 5 nS/cm$^2$ resulted in negligible tonic $I_{\text{GABA}}$ and no change in $R_{\text{in}}$ in the model FS-BC up to a conductance of 0.5 μS/cm$^2$. Tonic $g_{\text{GABA}}$ of 1 μS/cm$^2$ resulted in 10-pA tonic $I_{\text{GABA}}$, comparable to experimentally observed tonic $I_{\text{GABA}}$ with $g_{\text{GABA}}$ contributing to <0.5% of the model FS-BC resting conductance. Increasing tonic $g_{\text{GABA}}$ to 10 μS/cm$^2$ resulted in 60-pA tonic $I_{\text{GABA}}$, a 3.3-ΜΩ decrease in $R_{\text{in}}$ (Fig. 10B), and a 2.7% decrease in FS-BC membrane conductance. When tonic $g_{\text{GABA}}$ was increased to 50 μS/cm$^2$ and above, tonic $I_{\text{GABA}}$ was >250 pA and $R_{\text{in}}$ decreased by >10 ΜΩ, which is well outside the range observed in our experimental data (Fig. 5). The effect of tonic $g_{\text{GABA}}$ on tonic $I_{\text{GABA}}$ and $R_{\text{in}}$ in model FS-BCs was unchanged even when the $g_{\text{GABA}}$ was restricted to the soma and proximal dendritic compartments (Fig. 10B). Since our experimental estimation of ~5-pA tonic $I_{\text{GABA}}$ in control FS-BCs is 1) without added GABA, 2) in continuously perfused slices, and 3) at physiological temperature, it is likely an underestimation. Note also that in the presence of the GABA transporter antagonist tonic $I_{\text{GABA}}$ in FS-BCs is >10 pA, despite the presumably low activity levels in slices perfused with glutamate receptor antagonists (Fig. 5E). Therefore, we expect that tonic $g_{\text{GABA}}$ between 1 and 10 μS/cm$^2$ (corresponding to 10- to 60-pA tonic $I_{\text{GABA}}$) in our simulations represents the range of biologically realistic tonic $I_{\text{GABA}}$ levels during neuronal activity.

Next, we examined the effect of tonic $I_{\text{GABA}}$ on the excitability of model FS-BCs during excitatory synaptic inputs. Model FS-BCs received 200-Hz Poisson-distributed excitatory inputs to their distal dendrites, to simulate perforant path AMPA-mediated synaptic inputs. The rise and decay of the AMPA synapse was constrained by experimental data as described previously (Santhakumar et al. 2005). $g_{\text{AMPA}}$ was set either at 3 nS to simulate low excitatory inputs that lead to minimal FS-BC firing (Fig. 10, C and D) or at 20 nS, which resulted in model FS-BC firing at >30 Hz in the absence of tonic $g_{\text{GABA}}$ (Fig. 10E). Model FS-BCs received identical input trains in each simulation. In the first set of simulations, the reversal potential for the tonic $g_{\text{GABA}}$ was set to −74 mV, the experimentally determined $E_{\text{GABA}}$ in control FS-BCs. Tonic $g_{\text{GABA}}$ was simulated as a uniformly distributed leak conductance in the range of 5 nS/cm$^2$ to 5 mS/cm$^2$. This wide range of tonic $g_{\text{GABA}}$ encompassed the biologically relevant range of 1–10 μS/cm$^2$, and extended to include conductance values that resulted in biphasic changes in neuronal excitability similar to those described in previous studies (Song et al. 2011). Tonic $g_{\text{GABA}}$ in the range of 1 μS/cm$^2$ to 1 μS/cm$^2$ decreased model FS-BC firing during both low (3 nS)- and high (20 nS)-conductance excitatory inputs and abolished firing at $g_{\text{GABA}}$ over 75 μS/cm$^2$ and 0.3 mS/cm$^2$, respectively (Fig. 10, C–E). Since RMP of the model FS-BC was −70 mV, the decrease in excitability when $E_{\text{GABA}}$ was −74 mV is consistent with shunting inhibition. When $E_{\text{GABA}}$ was set to −54 mV, the experimentally determined $E_{\text{GABA}}$ in post-SE FS-BCs, increasing tonic $g_{\text{GABA}}$ led to an initial increase in model FS-BC firing when tonic $g_{\text{GABA}}$ was over 1 μS/cm$^2$, which reached a maximum when tonic $g_{\text{GABA}}$ was 1 μS/cm$^2$. FS-BC firing declined with further increase in tonic $g_{\text{GABA}}$ and was eventually completely suppressed (Fig. 10, C–E). Model FS-BC firing during excitatory synaptic inputs delivered in the presence of depolarizing $E_{\text{GABA}}$ (−54 mV) and 1 μS/cm$^2$ to 1 μS/cm$^2$ tonic $g_{\text{GABA}}$ increased even though $E_{\text{GABA}}$ was more hyperpolarized than the model FS-BC action potential threshold (−40.5 mV). The simulation results are consistent with recent studies demonstrating that, in the presence of a depolarizing $E_{\text{GABA}}$, progressively increasing tonic $g_{\text{GABA}}$ leads to an initial neuronal depolarization followed by shunting inhibition with...
further increases (Song et al. 2011). However, our simulations demonstrate that in the biologically relevant range of 1–10 μS/cm² tonic $g_{GABA}$, the experimentally observed post-SE $E_{GABA} (-54$ mV) reverses the decrease in excitability mediated by tonic $I_{GABA}$.

In additional simulations, we restricted tonic $g_{GABA}$ to the somatic and proximal dendritic compartments to examine whether dendritic tonic $g_{GABA}$ influenced neuronal excitability by local modulation of synaptic inputs. In simulations performed without tonic $g_{GABA}$ in the distal dendritic compartments where the excitatory synapses were located, both the range of tonic $g_{GABA}$ values that modulated model FS-BC excitability and the biphasic nature of the changes were indistinguishable from simulations that included tonic $g_{GABA}$ in all compartments (not shown). These simulations results suggest that local dendritic modulation of synaptic inputs does not underlie the effect of tonic $g_{GABA}$ on model FS-BC excitability. Thus the isolated FS-BC simulations predict that post-SE increases in tonic $g_{GABA}$ in the biologically relevant range would, in the absence of concomitant changes in $E_{GABA}$ (at $E_{GABA} = -74$ mV), decrease FS-BC excitability during both low and moderate excitatory synaptic drive (Fig. 10C, left, and Fig. 10, D and E). Alternatively, a depolarizing shift in $E_{GABA}$, when not accompanied by changes in tonic $g_{GABA}$, would enhance FS-BC excitability (Fig. 10C, compare left and right at each tonic $g_{GABA}$; Fig. 10, D and E, compare firing at identical tonic $g_{GABA}$ levels). However, the combined effect of the concurrent increase in tonic $g_{GABA}$ and depolarized $E_{GABA}$ is to maintain the control levels of FS-BC excitability during low levels of excitatory synaptic drive (Fig. 10D).

Effect of seizure-induced FS-BC GABAergic plasticity on dentate excitability: dentate network simulations. Our experimental data have demonstrated two distinct SE-induced changes in the inhibitory tone of dentate FS-BCs, namely, an increase in tonic $I_{GABA}$ and a positive shift in the driving force for $I_{GABA}$ after SE. Since both tonic $I_{GABA}$ and $E_{GABA}$ are also altered in granule cells after SE (Pathak et al. 2007; Zhan and Nadler 2009; Zhang et al. 2007), we adopted computational simulations to determine how seizure-induced plasticity in FS-BCs tonic inhibition affects dentate network excitability independent of, and in the absence of, other confounding changes. First, we examined whether the post-SE changes in FS-BC $E_{GABA}$ altered dentate excitability in the absence of tonic $g_{GABA}$. The 1,000+ dentate neuronal network based on Santhakumar et al. (2005) was simulated as a ring and included 20% sprouting (see MATERIALS AND METHODS). The network included two major interneuronal populations, FS-BCs and hilar interneurons) from other FS-BCs and hilar dendritically projecting HIPP cells. Since simulating post-SE hilar neuronal loss by removing HIPP cells from the network would disproportionately enhance the role of FS-BCs, the first set of simulations were performed without introducing hilar neuronal loss. Networks received 2.5-Hz spontaneous activity for the entire duration of the simulation. A single synchronous input to 100 adjacent granule cells was used to simulate focal perforant path-evoked network activity. In the absence of tonic $g_{GABA}$, changing model FS-BC $E_{GABA}$ (of inhibitory synaptic inputs to model FS-BCs from other FS-BCs and hilar interneurons) from $-74$ mV to $-54$ mV did not change either the spontaneous background activity level or average granule cell firing in response to synchronous activation (Fig. 11, A1 and B1). This is consistent with the ability of the inhibitory synaptic conductance to provide shunting inhibition even when $E_{GABA}$ is more depolarized than RMP (Song et al. 2011; Vida et al. 2006).

Next, we introduced tonic $g_{GABA}$ with activity-dependent synaptic spillover conductance (see MATERIALS AND METHODS) in model FS-BCs. In one group of simulations, $E_{GABA}$ for both tonic and synaptic $g_{GABA}$ was set to $-74$ mV as observed in controls. Tonic $I_{GABA}$ was increased between 3 μS/cm², contributing to 18-pA tonic $I_{GABA}$ similar to levels observed in

Fig. 9. SE decreases expression of KCC2 in PV interneurons. Confocal images from sections labeled for KCC2 (left) and PV (center) are shown. Merged images (right) show colabeling of KCC2 particularly in the periphery of the PV-positive neurons from a control (top) and a post-SE (bottom) rat. Boxed areas are shown at higher magnification in insets. Images were obtained with identical camera settings. Scale bars, 10 μm.
post-SE FS-BC, and 10 μS/cm². Tonic $g_{\text{GABA}}$ in model FS-BCs did not alter the background spontaneous activity (Fig. 11, A and C). Strikingly, while increasing FS-BC tonic $g_{\text{GABA}}$ did not alter overall network excitability in response to focal afferent activation in some network instantiation (summarized in Fig. 11D as “low activity”), it significantly increased average granule cell firing and resulted in self-sustained, recurrent, seizure-like activity in a subset of simulated networks [Fig. 11, B and C]. Notably, unlike simulations with shunting $E_{\text{GABA}}$, networks simulated with depolarizing $E_{\text{GABA}}$ showed no increase in evoked granule cell activity at all tonic $g_{\text{GABA}}$ values tested [Fig. 11, B and D; between-subject analysis for effect of $E_{\text{GABA}}$: $F_{(1,11)} = 345.3, P < 0.05$ by univariate repeated-measures ANOVA; interaction between network structure and $E_{\text{GABA}}$: $F_{(1,11)} = 355.2, P < 0.05$ by multivariate repeated-measures ANOVA]. To isolate the impact of FS-BC tonic $g_{\text{GABA}}$ and $E_{\text{GABA}}$ from network structural features, we set the randomization seed to a constant value to simulate structurally identical networks. Spike raster plots from simulations of structurally identical networks (illustrated in Fig. 11A) showed that when FS-BC $E_{\text{GABA}}$ was at $−74$ and $−54$ mV, $g_{\text{GABA}}$ had higher granule cell firing and recurrent network activity compared with networks with depolarizing ($−54$ mV) FS-BC $E_{\text{GABA}}$ at all tonic $g_{\text{GABA}}$ values tested.

Fig. 10. Effect of GABA reversal and tonic GABA conductance ($g_{\text{GABA}}$) on model FS-BC excitability. A: responses of biophysically realistic multicompartmental model FS-BC to depolarizing and hyperpolarizing current injections illustrate nonadapting firing and low input resistance ($R_{\text{in}}$). B: summary plot shows tonic $I_{\text{GABA}}$ and $R_{\text{in}}$ in model FS-BCs as a function of tonic $g_{\text{GABA}}$. In simulations with perisomatic $g_{\text{GABA}}$, tonic GABA channels were distributed only in the soma and proximal dendrite. Simulations incorporated voltage-clamp recording conditions with symmetrical chloride and $V_{\text{hold}} = −70$ mV. Shaded region represents biologically relevant tonic $I_{\text{GABA}}$ and $g_{\text{GABA}}$ range. C: membrane voltage traces illustrate firing in FS-BC simulations during 200 Hz during identical Poisson-distributed excitatory inputs when tonic $g_{\text{GABA}}$ is systematically increased from 0 to 1 mS/cm². Peak amplitude of input synaptic conductance ($g_{\text{AMPA}}$) was 3 nS. Simulations were performed with control (−74 mV, left) and post-SE (−54 mV, right) $E_{\text{GABA}}$ values ($E_{\text{GABA}}$). D: summary plot of FS-BC firing evoked by 200-Hz excitatory synaptic inputs (3 nS peak conductance) in the presence of increasing tonic $g_{\text{GABA}}$ in FS-BC with $E_{\text{GABA}}$ set at $−74$ and $−54$ mV. E: summary data show effect of $E_{\text{GABA}}$ on FS-BC firing during excitatory synaptic activation at 200 Hz at increasing FS-BC tonic $g_{\text{GABA}}$. Peak conductance of synaptic inputs was 20 nS.
seed values that resulted in high-activity networks) including loss of dentate hilar neurons (mossy cells and HIPP cells) and mossy fiber sprouting. As in networks without cell loss, average granule cell firing was increased when FS-BC $E\text{_{GABA}}$ was $-74$ mV and remained low when $E\text{_{GABA}}$ was set to $-54$ mV (Fig. 11E). While our measurements of $E\text{_{GABA}}$ during exogenous GABA application identified a post-SE depolarizing shift, given findings that the $E\text{_{GABA}}$ within a neuron can differ between compartments (Baldi et al. 2010; Romo-Parra et al. 2008), it remains possible that synaptic GABA reversal may differ from the reversal of tonic $I\text{_{GABA}}$. In an additional simulation in which synaptic $E\text{_{GABA}}$ was held either at $-54$ mV or at $-74$ mV independent of the $E\text{_{GABA}}$ of the tonic $g\text{_{GABA}}$, networks with a depolarizing shift in the $E\text{_{GABA}}$ of tonic $g\text{_{GABA}}$ still consistently demonstrated lower excitability than identical networks with $-74$ mV tonic $E\text{_{GABA}}$ [between-subject analysis for effect of tonic $E\text{_{GABA}}$: $F(1,11) = 32.68, P < 0.05$; synaptic $E\text{_{GABA}}$: $F(1,11) = 0.29, P > 0.05$; interaction
between tonic and synaptic $E_{\text{GABA}}$: $F_{1,11} = 0.46, P > 0.05$ by 2-way repeated-measures ANOVA). Taken together, the simulations show that the depolarization of FS-BC $E_{\text{GABA}}$ that occurs after SE can maintain FS-BC excitability despite increases in tonic $g_{\text{GABA}}$ and could limit dentate network excitability after SE.

**DISCUSSION**

The ability of tonic GABAergic inhibition to regulate excitability of projection neurons suggests that tonic $I_{\text{GABA}}$ may be modulated to prevent epileptogenesis (Meldrum and Rogawski 2007). Expression and seizure-induced changes in tonic $I_{\text{GABA}}$ among dentate perisomatic interneurons critical for feedback inhibition can influence how modulating tonic inhibition will affect dentate function. Our study demonstrates that (1) GABA$_A$R $\delta$-subunits, known to underlie tonic $I_{\text{GABA}}$, are expressed in PV+ interneurons in the hilar-granule cell layer border; 2) FS-BCs express tonic $I_{\text{GABA}}$ that is enhanced 1 wk after SE; 3) increase in GABA$_A$R $\delta$-subunit expression contributes to post-SE enhancement of FS-BC tonic $I_{\text{GABA}}$; 4) in contrast to shunting GABAergic inhibition in controls, there is a positive shift in FS-BC $E_{\text{GABA}}$ 1 wk after SE, resulting in a depolarizing driving force for $I_{\text{GABA}}$; 5) corresponding to the post-SE depolarizing shift in FS-BC $E_{\text{GABA}}$, there is a reduction in the expression of the potassium-chloride cotransporter KCC2 in PV+ interneurons 1 wk after SE; and 6) in computational simulations incorporating tonic $g_{\text{GABA}}$ in model FS-BCs, whereas shunting $E_{\text{GABA}}$ reduces FS-BC firing and enhances dentate network excitability, depolarizing $E_{\text{GABA}}$ reverses decreases in model FS-BC activity and prevents increases in network excitability.

**Expression of tonic $I_{\text{GABA}}$ in dentate FS-BCs.** Perisomatically projecting interneurons expressing PV are critical for precision and timing of network activity (Freund 2003). Since tonic inhibition influences neuronal excitability, gain, and fidelity of information transmission (Duguid et al. 2012; Mitchell and Silver 2003; Song et al. 2011) and is additionally augmented by synaptic spillover during neuronal activity (Glykys and Mody 2007), FS-BC tonic $I_{\text{GABA}}$ is ideally suited to regulate feedback inhibition and dentate throughput during behaviorally relevant neuronal activity. The ability of interneuronal tonic $I_{\text{GABA}}$ to regulate gamma oscillations (Mann and Mody 2010) and the central role for FS-BCs in dentate gamma oscillations (Bartos et al. 2002) suggest that modulation of FS-BC tonic $I_{\text{GABA}}$ by alcohol and neurosteroids, which act on GABA$_A$R$_S$ containing $\delta$-subunits (Stell et al. 2003), may contribute to the impact of these drugs on memory function. Moreover, activity-dependent enhancement of FS-BC tonic $I_{\text{GABA}}$ could provide the transient suppression of feedback inhibition needed to induce long-term potentiation in the dentate gyrus (Arima-Yoshida et al. 2011).

There are regional differences in the GABA$_A$R $\delta$-subunits mediating interneuronal tonic $I_{\text{GABA}}$, with $\alpha_2$-subunits responsible for tonic $I_{\text{GABA}}$ in CA1 interneurons and $\delta$-subunits in CA3 and molecular layer interneurons (Glykys et al. 2007; Mann and Mody 2010; Semyanov et al. 2003). Unlike PV interneurons in the neocortex (Olah et al. 2009), and like molecular layer interneurons in the dentate (Glykys et al. 2007), we find that dentate PV interneurons express GABA$_A$R $\delta$-subunits. Similar to fast-spiking interneurons in the murine neocortex (Krook-Magnuson et al. 2008), we show that tonic $I_{\text{GABA}}$ in dentate FS-BCs is modulated by a GABA$_A$R $\delta$-subunit-specific agonist. Since dentate interneuron-like profiles show robust expression of GABA$_A$R $\alpha_1$-subunits (Fritschi et al. 1999), and GABA$_A$R$_S$ containing $\alpha_1$- and $\delta$-subunits can mediate tonic $I_{\text{GABA}}$ (Glykys et al. 2007), it is possible that GABA$_A$R$_S$ with $\alpha_1$- and $\delta$-subunits underlie FS-BC tonic $I_{\text{GABA}}$. Thus, although both granule cell and FS-BC tonic $I_{\text{GABA}}$ are mediated by GABA$_A$R $\delta$-subunits, differential contribution of $\alpha_1$-subunits to FS-BC and $\alpha_2$-subunits to granule cell tonic $I_{\text{GABA}}$ may provide targets for cell-specific modulation of tonic inhibition.

**SE-induced plasticity of FS-BC tonic inhibition.** Perisomatic inhibition is pivotal in regulating granule cell excitability in both normal and epileptic dentate gyrus (Coulter and Carlson 2007; Kraushaar and Jonas 2000). Granule cell tonic $I_{\text{GABA}}$ is known to be altered after SE (Rajasekaran et al. 2010; Zhan and Nadler 2009; Zhang et al. 2007). Since FS-BC tonic $I_{\text{GABA}}$ is increased before the occurrence of behavioral seizures, and not as a compensatory response to changes in the epileptic network, FS-BC tonic $I_{\text{GABA}}$ could play a role in development of epilepsy. Indeed, earlier studies have suggested that cellular and synaptic plasticity prior to the onset of spontaneous seizures contribute to epileptogenesis (Brooks-Kayal et al. 1998; Kobayashi et al. 2003; Pathak et al. 2007). Both increases in extracellular GABA due to synaptic spillover or compromised GABA transporter function and changes in GABA$_A$R expression could enhance tonic $I_{\text{GABA}}$ (Farrant and Nusser 2005). However, sIPSC frequency in both FS-BCs (Fig. 6) and granule cells (Kobayashi and Buckmaster 2003) is reduced after SE, indicating that synaptic GABA spillover does not underlie the post-SE increases in FS-BC tonic $I_{\text{GABA}}$. Additionally, even when GABA transporters were blocked, post-SE FS-BCs had larger tonic $I_{\text{GABA}}$, indicating that factors other than compromises in GABA transporter function underlie the increase in tonic $I_{\text{GABA}}$. Our semiquantitative analyses demonstrate increase in the expression of GABA$_A$R $\delta$-subunit in PV+ interneurons after SE. Although extensive efforts were taken to pair sections from control and experimental animals and to use identical methods during staining and imaging, admittedly quantification of immunofluorescence intensity is associated with limitations including nonlinear antigen-antibody reaction and changes in fluorescence arising from tissue properties and may not accurately reflect protein levels. Crucially, THIP, a GABA$_A$R $\delta$-subunit-selective agonist under our experimental conditions (Brown et al. 2002; Gupta et al. 2012), causes greater baseline current shift in FS-BCs from post-SE rats, validating the semiquantitative immunofluorescence analysis and confirming that enhanced GABA$_A$R $\delta$-subunit expression contributes to increases in FS-BC tonic $I_{\text{GABA}}$ after SE. Thus the presence and SE-induced plasticity of FS-BC tonic $I_{\text{GABA}}$ must be considered when developing seizure therapies targeting tonic inhibition.

Whether tonic $I_{\text{GABA}}$ enhances or decreases neuronal excitability depends on complex interactions between GABA driving force and conductance (Song et al. 2011). Several studies have demonstrated that interneuronal $E_{\text{GABA}}$ can provide shunting inhibition when $E_{\text{GABA}}$ lies between RMP and action potential threshold (Banke and McBain 2006; Song et al. 2011; Vida et al. 2006). Consistent with earlier observations of shunting synaptic $E_{\text{GABA}}$ in FS-BCs (Sauer and Bartos 2010; Vida et al. 2006), the difference between FS-BC RMP and
$E_{GABA}$ determined during GABA application in controls was not significant. Methodological differences including use of whole cell recording conditions, younger animals, and CNQX, which unlike structurally related quinoxalinediones depolarizes hippocampal interneurons by nonspecific mechanisms (McBain et al. 1992), may have resulted in more depolarized RMP values in earlier studies (Geiger et al. 1997; Vida et al. 2006). It is important to note that the RMP values in our study are in the same range as the RMP of hippocampal interneurons measured with less invasive cell-attached recordings (Fricker et al. 1999; Verheugen et al. 1999). Moreover, RMP and $E_{GABA}$ values in control FS-BCs in our study are similar to the RMP and IPSC reversal potential reported in hippocampal interneurons from animals over postnatal day 30 (Banke and McBain 2006) and likely result from developmental hyperpolarizing shift in interneuronal RMP and $E_{GABA}$ (Banke and McBain 2006; Sauer and Bartos 2010). Importantly, we show that FS-BC $E_{GABA}$ is depolarized relative to the RMP 1 wk after SE, resulting in a depolarizing GABA driving force, as previously reported in granule cells (Pathak et al. 2007). Curiously, despite post-SE changes in FS-BC $I_{GABA}$ and $E_{GABA}$, FS-BC RMP remains unchanged. It is possible that changes in non-GABAergic ionic conductances compensate for changes in membrane conductance when tonic $g_{GABA}$ is altered (Brickley et al. 2001). While the trauma of slicing can impact neuronal chloride homeostasis (Dzhala et al. 2012), our use of young adult animals and high-sucrose slicing solution minimizes these effects. Additionally, our inclusion of only morphologically intact neurons, the hyperpolarized FS-BC RMP, and the lack of difference between RMP in control and post-SE FS-BCs suggest that slicing-induced trauma is unlikely to underlie the post-SE depolarization of FS-BC $E_{GABA}$. FS-BC $E_{GABA}$ undergoes a developmental hyperpolarizing shift believed to result from increases in the expression of the potassium-chloride cotransporter KCC2 (Sauer and Bartos 2010). While cognizant of the limitations of semiquantitative analysis of fluorescence intensity, we find consistent post-SE changes in FS-BC $I_{GABA}$ and reversal potential reflect homeostatic plasticity to maintain FS-BC activity levels in the network (Howard et al. 2007; Turrigiano et al. 1994). In dentate network models with recurrent mossy fiber collaterals, including FS-BCs with simulated tonic $I_{GABA}$ consistently resulted in low evoked granule cell activity when FS-BC $E_{GABA}$ was depolarizing. Curiously, evoked granule cell activity in some but not all network implementations was greatly enhanced and developed into seizure-like recurrent firing when $E_{GABA}$ was $\sim$74 mV. Since the average number of cell type-specific connections to each neuron in a given class of cells was not different between the networks, it is likely that the presence of a few highly connected hub cells and formation of feedback excitatory loops (Morgan and Soltesz 2008) may have contributed to the differences in activity patterns between the networks with otherwise identical connectivity statistics. While our simulations predict that post-SE changes in FS-BC inhibition can limit early SE-induced increases in dentate excitability when sprouting is low, structural reorganization and cell loss associated with epileptogenesis could undermine this potential compensatory mechanism. Additionally, maintenance of normal neuronal and network excitability despite altered $g_{GABA}$ and reversal is consistent with recent studies suggesting that multiple combinations of intrinsic parameters can result in similar circuit performance (Marder 2011). However, all parameter combinations may not be identically robust in maintaining normal activity during different network perturbations. Thus, while the observed changes in tonic $I_{GABA}$ and $E_{GABA}$ maintain normal FS-BC and dentate activity during focal afferent activation, these changes may compromise FS-BC responses during certain patterns of network perturbations and may impact generation of network rhythms contributing to memory impairments associated with seizure disorders.

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**DISCLOSURES**

No conflicts of interest, financial or otherwise, are declared by the author(s).
AUTHOR CONTRIBUTIONS

Author contributions: J.Y., A.P., and T.I. performed experiments; J.Y., A.P., and F.S.E. analyzed data; J.Y., A.P., and V.S. interpreted results of experiments; J.Y., A.P., and F.S.E. prepared figures; J.Y., A.P., F.S.E., T.I., and V.S. prepared final version of manuscript; F.S.E. and V.S. edited and revised manuscript; V.S. conceived and designed of research; V.S. drafted manuscript.

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