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Enhanced Excitatory Synaptic Connectivity in Layer V Pyramidal Neurons of Chronically Injured Epileptogenic Neocortex in Rats

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Formation of new recurrent excitatory circuits after brain injuries has been hypothesized as a major factor contributing to epileptogenesis. Increases in total axonal length and the density of synaptic boutons are present in layer V pyramidal neurons of chronic partial isolations of rat neocortex, a model of posttraumatic epileptogenesis. To explore the functional consequences of these changes, we used laser-scanning photostimulation combined with whole-cell patch-clamp recording from neurons in layer V of somatosensory cortex to map changes in excitatory synaptic connectivity after injury. Coronal slices were submerged in artificial CSF (23°C) containing 100 μM caged glutamate, APV (2-amino-5-phosphonovaleric acid), and high divalent cation concentration to block polysynaptic responses. Focal uncaging of glutamate, accomplished by switching a pulsed UV laser to give a 200–400 μs light stimulus, evoked single- or multiple-component composite EPSCs. In neurons of the partially isolated cortex, there were significant increases in the fraction of uncaging sites from which EPSCs could be evoked ("hot spots") and a decrease in the mean amplitude of individual elements in the composite EPSC. When plotted along the cortical depth, the changes in EPSCs took place mainly between 150 and 200 μm above and below the somata, suggesting a specific enhancement of recurrent excitatory connectivity among layer V pyramidal neurons of the undercut neocortex. These changes may shift the balance within cortical circuits toward increased synaptic excitation and contribute to epileptogenesis.

Key words: epilepsy; neocortex; synaptic transmission; caged compound; electrophysiology; excitability

Introduction

Neuronal injuries after cortical trauma, stroke, and prolonged seizures result in axonal sprouting in the hippocampus and neocortex (Tauck and Nadler, 1985; Salin et al., 1995; Perez et al., 1996; Sutula et al., 1998; Esclapez et al., 1999; Golari et al., 2001; Santhakumar et al., 2001; Buckmaster et al., 2002; Marchenko et al., 2004). This process may underlie the capacity of neuronal circuits to recover from damage (Stroemer et al., 1998; Carmichael et al., 2001; Carmichael, 2003; Lee et al., 2004); however, formation of new excitatory circuitry may also contribute to pathophysiological processes such as epileptogenesis (Purpura and Housepian, 1961; Sutula et al., 1992; Dudek and Spitz, 1997; McKinney et al., 1997; Prince et al., 1997; Prince, 1999).

Recurrent excitatory synapses formed by sprouting axons of pyramidal neurons and dentate gyrus granule cells (Perez et al., 1996; Sutula et al., 1998; Zhang et al., 1999; Wenzel et al., 2000; Buckmaster et al., 2002; Smith and Dudek, 2002) are associated with enhanced functional excitation in hippocampal slices from epileptic humans and rat models of temporal lobe epilepsy (Wuarin and Dudek, 1996; Patrylo and Dudek, 1998; Molnar and Nadler, 1999; Lynch and Sutula, 2000; Nadler, 2003; Scharfman et al., 2003). These injury-induced circuit reorganizations may contribute to epileptogenesis (Wuarin and Dudek, 1996, 2001; Okazaki et al., 1999) (but see Kotti et al., 1997; Longo and Mello, 1997; Sloviter, 1992).

Axonal sprouting also is present in partially isolated neocortex, a chronic model of posttraumatic epileptogenesis (Salin et al., 1995) (for review, see Graber and Prince, 2006). Increases in the total length, number of collateral branches, and the density of boutons occur in axons of layer V pyramidal neurons and are most prominent within layer V, where interictal epileptiform activity is initiated (Hoffman et al., 1994). Additional experiments are required to determine whether such anatomical changes underlie functional alterations such as increases in the amplitude or frequency of spontaneous EPSCs or in the spatial extent and synchronization of excitation within cortical neuronal networks of partially isolated epileptogenic neocortex (Li and Prince, 2002; Marchenko et al., 2004).

The low "hit rate" for recording synaptically coupled layer V pyramidal cells in paired recordings, even from closely spaced neurons (Markram et al., 1997), makes such an approach unfeasible for quantitatively comparing functional connectivity across significant distances in normal versus chronically injured cortex.
As an alternative, we used laser-scanning photostimulation of cortical slices exposed to caged glutamate, combined with whole-cell patch-clamp recording (Katz and Dalva, 1994; Molnar and Nadler, 1999; Dantzker and Callaway, 2000; Shepherd et al., 2003) to map and quantify changes in monosynaptic excitatory connectivity. The strength and distribution of synaptic inputs from cortical layers II–VI onto layer V pyramidal neurons in slices from control and partially isolated cortex were compared. Data indicate that there is an increase in the efficacy and extent of recurrent excitatory synaptic connectivity among layer V pyramidal neurons of the undercut neocortex. Portions of these results have been published in abstract form (Jin et al., 2005b).

Materials and Methods

Surgical procedures. All experiments were performed according to protocols approved by the Stanford Institutional Animal Care and Use Committee. Neocortical slices from 28 Sprague-Dawley rats aged postnatal day 35 (P35) to P41 (P0, date of birth) were used for in vitro recordings. Partially isolated islands of neocortex (“undercuts”) were produced in 11 anesthetized rats at P21, using previously described techniques (Hoffman et al., 1994; Li and Prince, 2002). Rats were deeply anesthetized with ketamine (80 mg/kg, i.p.) and xylazine (Rompun, 8 mg/kg, i.p.), mounted in a stereotaxic frame, the scalp incised and retracted, and a portion of frontoparietal cortex of the left hemisphere exposed by moving a 3 × 5 mm bone window centered on the coronal sulcure, leaving the dura intact. A partial isolation of an island of sensorimotor cortex was made as described previously. A 30 gauge needle, bent at approximately a right angle 2.5–3 mm from the tip, was inserted tangentially through the dura, just beneath the pial vessels, parasagittally 1–2 mm from the interhemispheric sulcus, and lowered to a depth of 2 mm. The needle then was rotated 120–135° to produce a contiguous white matter lesion, elevated to a position just beneath the pia, and removed. The skull opening was then covered with sterile plastic wrap (Saran Wrap), and the skin was sutured. Rats were allowed to recover for at least 2 weeks, a latency at which most in vitro slices containing a portion of the injured area generated evoked epileptiform activity in previous experiments (Hoffman et al., 1994; Graber and Prince, 1999).

Slice preparation and recording. Animals were anesthetized with pentobarbital (55 mg/kg, i.p.) and decapitated, and the brain was rapidly removed and placed in ice-cold (4°C) oxygenated slicing solution containing the following (in mM): 230 sucrose, 2.5 KCl, 1.25 NaH2PO4, 10 MgSO4 · 7H2O, 10 glucose, 0.5 CaCl2 · 2H2O, and 26 NaHCO3. The area of the cortical lesion was easily identified in the left hemisphere. Coronal slices (300 μm) were cut with a vibratome (Lancer Series 1000; Vibratome Company, St. Louis, MO) through the lesioned sensorimotor cortex and from the same region in control animals, and maintained using standard techniques (Li and Prince, 2002). After ~1 h incubation at 36°C in standard artificial CSF (ACSF), slices were held at room temperature. The ACSF contained the following (in mM): 126 NaCl, 2.5 KCl, 1.25 NaH2PO4, 2 CaCl2 · 2H2O, 2 MgSO4 · 7H2O, 26 NaHCO3, and 10 glucose; pH 7.4 when saturated with 95% O2–5% CO2.

Patch electrodes were pulled from borosilicate glass tubing (outer diameter, 1.5 mm) and had an impedance of ~4–7 MΩ when filled with intracellular voltage-clamp solution containing the following (in mM): 120 Cs-glucuronate, 10 KCl, 11 EGTA, 1 CaCl2 · 2H2O, 2 MgCl2 · 6H2O, 10 HEPES, 2 NaATP, 0.5 NaGTP, and 0.5% biocytin. In current-clamp experiments, we used a K-glutamate-based intracellular solution containing the following (in mM): 95 K-glucuronate, 40 KCl, 5 EGTA, 0.2 CaCl2 · 2H2O, 10 HEPES, and 0.5% biocytin. The osmolarity of the pipette solutions was adjusted to 285–295 mOsm and pH to 7.3 with 1 M HEPES.

Single slices were transferred to a recording chamber where they were minimally submerged in high divalent ACSF that contained the following (in mM): 121 NaCl, 2.5 KCl, 1.25 NaH2PO4, 4 CaCl2, 4 MgSO4 · 7H2O, 26 NaHCO3, and 10 glucose. Patch-clamp recordings were made from visually identified layer V pyramidal cells in undercut cortex or the same region in control slices, using infrared video microscopy (Zeiss Axioskop; Carl Zeiss, Oberkochen, Germany) and a 63× water-immersion lens (Achromplan 63×, 0.9 W; Carl Zeiss) with differential interference contrast optics and an Axopatch 200A amplifier (Molecular Devices, Foster City, CA). Access resistance was measured in voltage-clamp mode from responses to 5 mV depolarizing voltage pulses. Recordings with access resistance <30 MΩ and without significant changes (~25%) during the recording were used for data analysis. The responses were low-pass filtered at 2 kHz and recorded on hard disk for later analysis. EPSCs were measured at a holding potential of ~70 mV, close to reversal for fast inhibition.

Polysynaptic recurrent excitation (Prince and Tseng, 1993; Lynch and Sutula, 2000) is common in different models of epileptogenesis, which raises the concern that polysynaptic activity could be evoked by glutamate uncaging and result in incorrect maps of excitatory synaptic input. To address this issue, we used a high-divalent ACSF containing 4 mM Ca2+, 4 mM Mg2+ (Shepherd et al., 2003), and 10 μM 2-amino-5-phosphonovaleric acid (APV) to eliminate potential polysynaptic activation via NMDA receptors. We tested the effectiveness of this solution in blocking polysynaptic activity under conditions of reduced inhibition. When the control slices were perfused in normal ACSF containing 10 μM GABAA receptor antagonist bicuculline, polysynaptic events occurred spontaneously. These polysynaptic events were completely abolished when the perfusate was switched to one containing 10 μM bicuculline, 10 μM APV, and the increased divalent cation concentrations (see above) (data not shown), suggesting that this perfusion solution was effective in reducing polysynaptic transmission. Furthermore, examination of action potential (AP) firing evoked by photostimulation revealed that, whereas direct uncaging stimuli clearly could evoke one or more APs, no such firing was induced by evoked or spontaneous EPSPs in either control and undercut slices. These control data ruled out the possibility of polysynaptic activation from photostimulated neurons under our recording conditions. Addition of 10 μM APV also eliminated NMDA-mediated long-lasting plateau potentials (Wei et al., 2001), which could otherwise interfere with the detection of individual AMPA receptor-mediated synaptic events.

After the electrophysiological recordings, slices containing biocytin-filled neurons were fixed and processed with the standard avidin–biotin peroxidase method (Horikawa and Armstrong, 1988; Tseng et al., 1991) or immunofluorescent staining. Labeled neurons were examined under light or confocal microscopy to verify their morphology and location.

Photolysis of caged glutamate and stimulus patterns. An uncaging setup was built according to the principles described by Tsai et al. (2002) (see also Shepherd and Svoboda, 2005). An upright microscope (Axioskop) with a fixed stage and submersion slice chamber was mounted on a locally made mobile stage that allowed coincident longitudinal displacement of the microscope and the laser unit relative to the slice. The slice was imaged with a monochrome charge-coupled device (CCD) video camera (AxioCam, San Dieter GmbH, Braunschweig, Germany) and a frequency-tripled Nd:YVO4 laser (Spectra 3500 pulsed laser; 100 kHz repetition rate; DPSS Lasers, San Jose, CA) was used to provide 50–100 mW of 355 nm light. The UV laser was coupled to the epifluorescence port of the microscope through several mirrors and a lens assembly that focused the light with 1:2 amplification onto a dichroic mirror, reflecting the light beam to the back aperture of a 5× objective (Fluar 5×; Carl Zeiss). The movement of the laser beam was precisely controlled with mirror galvanometers (model 6210; Cambridge Technology, Cambridge, MA), trigged by scanning and data acquisition software developed by I. R. Huguenard.

A concentration of 100 μM caged glutamate (methyl 1-[5-(4-amino-4-carboxybutanoyl)]-7-nitroindoline-5-acetate) and 10 μM APV (both from Sigma, St. Louis, MO) were added to 20 ml of recirculating high divalent ACSF at the beginning of experiments. Once a whole-cell recording was established, the lens was switched to the 5× objective. Focal photolysis of caged glutamate was accomplished by switching the UV laser to give a 200–400 μs light stimulus. The mapping stimulus patterns consisted of 264 positions on an 11 × 25 grid. Spacing between adjacent rows and columns was set to 50 μm, resulting in a 500 × 1200 μm mapping area in the slices. The grid was carefully aligned to cover cortical layers II–VI. When a single map did not cover all cortical layers, two shorter grids of 500 μm wide were used to extend over the lower and upper half of the cortex separately. We used a random stimulus sequence pattern at a 1 s interstimulus interval. Each recorded trace consisted of a
100 ms prestimulus baseline and a 500 ms poststimulus period. We repeatedly recorded evoked responses from a few individual uncaging spots with 5–20 s interstimulus interval, and found that direct and synaptic responses from a single uncaging spot were quite consistent and reproducible (n = 4). In most mapping experiments, repeated photostimulation of the same cortical regions resulted in comparable maps of synaptic input with small variation.

Data acquisition and analysis. Responses evoked by glutamate uncaging were analyzed within a time window between 12 ms and 200 ms after photostimulation. This time window was chosen because all excitatory responses attributable to direct effects of uncaged glutamate on the recorded neuron peaked within 12 ms after photostimulation (see Fig. 1). This time window was chosen because all excitatory responses attributable to direct effects of uncaged glutamate on the recorded neuron peaked within 12 ms after photostimulation (see Fig. 1) and thus could be separated from the later occurring synaptic responses. All directly evoked APs occurred within a window between 12 and 200 ms after the onset of photostimulation, and most of them fell within the first 100 ms. These differences in latency allowed us to include all evoked synaptic responses in the data analysis. EPSCs evoked by uncaging were analyzed using PSC detection software, Wdetecta (Ulrich and Huguenard, 1996) (http://huguenardlab.stanford.edu/apps/wdetecta/). The detector parameters, including differentiation window size and event amplitude, were carefully adjusted and tested for recorded traces of each map to ensure correct detection of synaptic events. Once proper settings were determined, the program automatically and accurately detected synaptic events and generated a number of quantitative parameters of the evoked EPSCs.

We used three measures of detected EPSCs to quantify the characteristics of synaptic connectivity. For spots that evoked EPSCs, the “composite EPSC amplitude” was defined as the sum of the amplitudes of all detected synaptic events during the 200 ms time window in each recording trace (see Fig. 1G, amplitude of b + c). The “EPSC number” refers to the total number of events in a response (see two events in Fig. 1G). The “mean individual EPSC amplitude” was obtained by dividing composite EPSC amplitude by EPSC number (see Fig. 1G, b + c)/2. As an indicator of the extent of synaptic connectivity, composite amplitude for each spot of photostimulation was used to construct maps of synaptic inputs for individual neurons (see Fig. 4B, C).

We used a modified Sholl analysis to compare the intensity of synaptic inputs between the control and undercut groups. This involved calculating the EPSC amplitude for each cell evoked by uncaging stimuli delivered within concentric 100 μm rings centered on its soma (see Fig. 4D). The “region-normalized EPSC amplitude” was obtained by dividing the sum of all composite EPSC amplitudes evoked within a given circular Sholl region by the total number of stimulus sites within the region (see Fig. 4E, F). This holistic measure reflects strength of functional synaptic coupling within a region and incorporates both number and amplitude of evoked events. “Hot spots” were defined as sites on the stimulation grid from which a synaptic event could be evoked. To compare the size of the synaptic input maps, the number of hot spots within a given Sholl circular area was divided by the total number of stimulated sites within the same area to give “fractional hot spots” (see Fig. 4G, H). Analysis of region-normalized EPSC amplitude and fractional hot spots along vertical and horizontal axes revealed possible changes in interlaminar and intralaminar connectivity, respectively. The mean EPSC number, composite EPSC amplitude, and individual EPSC amplitude were also analyzed using the modified Sholl method by separately dividing the sum of EPSC number, composite EPSC amplitude, and individual EPSC amplitude in a given circular Sholl area by the number of hot spots in the same Sholl region.

Statistical significance for group means was determined with two-tailed Student’s t test with p < 0.05. Results from modified Sholl analyses were analyzed with two-way ANOVAs. Additional comparisons between control and undercut groups at each individual distance from the soma were made with Student’s t test (Kittlerberger and Mooney, 1999). Data are presented as mean ± SEM. Origin and Microsoft Excel software were used to perform all statistical analyses.

Results
Coronal brain slices were prepared from control and undercut rats aged P35–P41, 14–20 d after partial isolations in lesioned animals. Transcortical cuts were easily seen in slices under a low-power objective. Undercutting lesions were present in white matter closely adjacent to layer VI, and a small area of cavitation was often associated with the undercut. Cortical pyramidal neurons were visually identified based on their location, pyramidal-shaped somata, and a single emerging apical dendrite extending toward the pial surface. Their pyramidal cell identities were later confirmed by morphological images from biocytin staining (see Fig. 3). Maps of uncaging-evoked synaptic events were obtained from pyramidal neurons of layers Va and Vb within the partially isolated cortex and from homologous areas of control cortex from age-matched untreated rats, whereas recordings of evoked AP firing in current clamp were from layers II–VI of the control and undercut cortex. Based on their dendritic morphology and our previous results (Jin et al., 2005a), layer V pyramidal neurons included both regular spiking neurons and intrinsically burst firing neurons. It is known that excitatory synaptic input onto these two type of pyramidal neurons are different in layers II/III, IV, and VI (Schubert et al., 2001). Because axonal sprouting in undercut rats occurs mainly in layer V (Salin et al., 1995), we did not differentiate data from these two groups of cells for analysis.

Spatial resolution of photostimulation
To test the spatial resolution and determine the timing of direct neuronal activation, we applied uncaging stimuli to 350 × 350 μm regions at 50 μm intervals under voltage clamp in control brain slices, perfused with ACSF containing 1 μM TTX and 10 μM APV to block synaptic activity and NMDA receptor-mediated responses. Under these conditions, direct activation of non-NMDA glutamate receptors was mapped for individual neurons (Fig. 1A, B). All direct responses initiated with little or no delay from the onset of the uncaging stimulus and peaked within 8–12 ms (Fig. 1C). The maximal direct activation resulted from glutamate uncaging was in the perisomatic area, with one or two spots eliciting the largest response (Fig. 1B). This result indicates that spatial resolution of direct activation was <50 μm and that response onset after direct activation (Fig. 1C) was similar to values described in previous reports (Schubert et al., 2001; Shepherd et al., 2003). These findings allowed us to separate synaptic responses from those attributable to direct activation by detecting events with latencies 10–12 ms after uncaging in mapping experiments (Fig. 1D–G).

Our previous results showed that layer V pyramidal neurons in the partial cortical isolation have increased input resistance and membrane time constant (Jin et al., 2005a; Prince and Tseng, 1993), as well as a steeper relationship between AP frequency and depolarizing current (Prince and Tseng, 1993) (see also Tseng and Prince, 1996). Such changes in intrinsic properties might result in enhanced excitability and increased AP firing when neurons are depolarized by uncaged glutamate. To rule out this possibility and further determine the spatial resolution of photostimulation in undercut and control cortical slices, we obtained whole-cell and loose-seal recordings from pyramidal neurons in cortical layers II–VI in current-clamp mode to generate maps of evoked AP firing. Neurons were photostimulated in an area of 500 × 500 μm with 40 μm spacing and a laser flash duration of 200–400 μs. All neurons recorded fired APs when stimuli were located close to the soma or apical or basal dendrites (Fig. 2A, B). When the duration of laser uncaging was substantially increased to 400 ~ 800 μs, evoked APs appeared with stimuli at greater distances from somata (data not shown). There were no significant differences between the control (n = 19) and undercut (n = 16) groups of cells in total spike number per neuron (5.5 ± 0.9 in
control; 5.8 ± 0.8 in undercut), mean number of AP-generating spots per neuron (2.8 ± 0.3 in control; 3.6 ± 0.3 in undercut), and number of APs per spot (2.0 ± 0.2 and 1.7 ± 0.3 in control and undercut groups, respectively) (Fig. 2C). Therefore, under our experimental conditions, there was no significant increase in excitability of pyramidal neurons in the undercut cortex that would affect responses of presynaptic cells to glutamate uncaging, and maps of synaptic input generated by photostimulation. In all cortical layers, all APs occurred with stimuli within an effective radius of <100 μm from somata (mean, 80.3 ± 6.8 μm in control and 87.5 ± 5.9 μm in undercut; p > 0.05). There were also no significant differences in mean effective radius between the control and undercut groups in layer V pyramidal neurons and in pyramidal neurons of other cortical layers (Fig. 2D). The above result suggests that most pyramidal neurons in a slice can be depolarized to threshold for firing APs in response to glutamate uncaging with a spatial resolution close to 100 μm, confirming that we were able to generate synaptic input maps with sublaminar resolution.

**Increased excitatory connectivity in partially isolated cortex**

We previously found significant increases in axonal length, collaterals, and synaptic bouton density in layer V pyramidal neurons of the undercut cortex (Salin et al., 1995) as well as increases in the frequency and amplitude of spontaneous EPSCs (sEPSCs) and frequency of miniature EPSCs (Li and Prince, 2002), supporting the hypothesis that formation of new recurrent synaptic connections contributes to the enhancement of excitatory synaptic connectivity. To further test this hypothesis, we used the laser-scanning photostimulation technique to detect changes in monosynaptic excitatory connectivity in layer V pyramidal neurons.

Whole-cell voltage-clamp recordings were obtained from 18 control and 17 undercut neurons that were filled with biocytin and retroactively identified as layer V pyramidal cells (Fig. 3). To evaluate the frequency and amplitude of spontaneous EPSCs in our experiment condition, we recorded sEPSCs for 2–3 min in voltage-clamp mode at −70 mV at the beginning of each experiment. The mean sEPSC frequency averaged from cells of the undercut group (3.2 ± 0.7 Hz; n = 17) was slightly higher than that in the control group (2.7 ± 0.3 Hz; n = 18), but the difference was not statistically significant (p > 0.05, Student’s t test), ruling out the possibility of sEPSC making significant contribution to the change in synaptic input maps in undercut neurons. In contrast, the amplitude of sEPSC in the undercut group was smaller than the control (−11.0 ± 0.9 pA in undercut vs −14.3 ± 1.1 pA in control; p < 0.05, Student’s t test). This latter difference would

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**Figure 1.** Spatial resolution of photostimulation and generation of excitatory synaptic input maps. A–C, Direct activation profile under voltage-clamp recording of a layer V pyramidal neuron in ACSF containing TTX and APV (see Materials and Methods). A, Responses to photostimulation of a layer V pyramidal neuron generated by photostimulation in a 400 × 400 μm square area with 50 μm spacing. Superimposed responses are shown in A, B, Map generated from peak amplitudes of direct responses in A. Response intensity, in picocuries, is color coded according to the scale on the right. B, D–F, Triangle, Site of somata. C, Direct responses evoked by glutamate in cell of A and B. Responses peaked within the first 10 ms after the laser stimulus (red rectangular area). D–G, Responses from another pyramidal neuron to laser uncaging of glutamate. D, Color-coded map of composite amplitude of both direct and synaptic responses evoked by glutamate uncaging (see Materials and Methods) at a holding potential of −70 mV in a bathing solution containing 4 mM Ca²⁺/4 mM Mg²⁺. E, Map of direct activation responses constructed by detecting peak events within 10 ms of glutamate uncaging. The direct activation profile coincided with the presumed position of the portion of the dendritic tree of this pyramidal neuron. F, A synaptic map of the neuron of D–G, generated by excluding direct activation responses of E (i.e., those before 10 ms) from the total responses of D. Grid spacing is 50 μm for D–F. Scale on right, Composite amplitude of triggered events (in picocuries). G, A representative trace containing both direct and synaptic responses. Direct response (a) to photostimulation occurred within the first 10 ms (red rectangle), synaptic responses (b, c) were detected within the first 200 ms (between the end of the red rectangle and the dashed red line) after photostimulation.
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Figure 2. Direct excitation profiles of pyramidal neurons in control and undercut cortex. A, B, The 500 × 500 μm grids were photostimulated by laser spots with a 40 μm spacing, and the average number of evoked action potentials evoked by each spot was plotted in 19 control (A) and 16 undercut neurons (B). Most of the APs were activated by stimuli close to the soma (triangles) of the recorded neurons. Scale to the right, Average number of APs evoked by spots. C, D, Analysis of direct excitation profiles. C, There were no significant differences between the control (open bars; n = 19) and undercut (black bars; n = 16) in the average total number of APs per neuron (left graph), average active spots per neuron (middle), or number of spikes per spot (right). D, There were also no significant differences in the mean distance from the soma of all of the spots that could evoke action potentials (effective radius) between the control and undercut groups in layer V pyramidal neurons and in pyramidal neurons in other cortical layers (layers II–IV and VI; p > 0.05, one-way ANOVA). The mean effective radius was larger in layer V pyramidal neurons than in pyramidal neurons of other layers, but no significant differences were observed between control and undercut groups in either layer (p > 0.05, one-way ANOVA). Blue open circles, Control; red squares, undercut. Error bars indicate SEM.

Figure 3. Confocal images of biocytin-filled layer V pyramidal neurons in control (A) and undercut (B) 300-μm-thick coronal neocortical slices. FITC was conjugated to the biocytin reaction product to produce green fluorescence. Both control and undercut neurons had large somata and a thick primary apical dendrite, characteristic of deep-lying layer V pyramidal cells. Scale bar: (in A for both neurons) 100 μm.
amplitude within each Sholl area (data not shown). The mean individual EPSC amplitude was significantly smaller in the undercut group (−16.5 ± 0.3 vs −12.5 ± 0.6 pA in the control and undercut groups, respectively; p < 0.001, Student’s t test), consistent with result from sEPSC recording (see above). The cumulative probability plot of individual EPSC amplitudes from control and undercut neurons (Fig. 5C) showed a significant left shift for undercut neurons [Kolmogorov–Smirnov (K-S) test, p < 0.0001], confirming the mean individual amplitude result. Mean individual EPSC amplitudes were generally smaller in all Sholl regions, but significant differences between the two groups were only observed at distances of 400–500 μm (−17.1 ± 1.2 pA in control vs −12.3 ± 0.9 pA in undercut; p < 0.005) and 800–900 μm (−16.5 ± 2.3 pA control and −11.0 ± 0.6 pA undercut; p < 0.05) (data not shown).

To determine alterations in the laminar source of the excitatory synaptic input, we further analyzed the changes in region-normalized EPSC amplitude and fraction of hot spots relative to the cortical depth of the uncaging stimulus. There were significant differences between the control and undercut group in region-normalized EPSC amplitude and mean fraction of hot spots (F = 40.1, p < 0.0001; and F = 61.1, p < 0.0001, respectively; two-way ANOVA) (Fig. 6A,B). Consistent with the above modified Sholl analysis (Fig. 5) and results of previous experiments on the distribution of axonal sprouting in the partially isolated cortex (Salin et al., 1995), the most significant increases in region-normalized EPSC amplitude and mean fraction of hot spots were centered in an area between 175 μm below and 125 μm above the somata of pyramidal neurons in layer V of the undercut cortex (p < 0.05 ~ p < 0.005, Student’s t test) (Fig. 6A,B). There were also significant increases in region-normalized EPSC amplitude in areas 375 μm (p < 0.05) and 725 μm (p < 0.01) superficial to the somata in layer V, and significant increases in mean fraction of hot spots at four depths between −375 and 875 μm (p < 0.05 ~ 0.005) (Fig. 6B). The data suggest that, compared with control, layer V pyramidal neurons in the undercut cortex receive more widespread excitatory synaptic input, most prominently from adjacent pyramidal neurons in layer V, but also from excitatory cells in upper cortical layers. To test the horizontal extent of the increase in the excitatory synaptic connectivity, we plotted the data along the horizontal direction, medial and lateral to the recorded cells. A two-way ANOVA analysis indicated that neurons in the undercut cortex had a higher region-normalized EPSC amplitude and a greater fraction of hot spots (F = 22.6, p < 0.0001; and F = 42.0, p < 0.0001, respectively; two-way ANOVA) (Fig. 6C,D). Compared with controls, neurons in the undercut cortex had a higher
region-normalized EPSC amplitude from laser uncaging at distances of 100 and 200 μm medial and 250 μm lateral to the somata (p < 0.05, Student’s t test) (Fig. 6C), as well as a greater fraction of hot spots at distances of 50–150 and 250 μm medial, and 50–150 and 250 μm lateral to the somata (p < 0.05 ~ p < 0.01, Student’s t test) (Fig. 6D). Two-way ANOVA analysis further indicated that there were no significant differences in either region-normalized EPSC amplitude or fraction of hot spots relative to the proximity of uncaging sites to the transcortical lesion, which was clearly visible in the slice.

Discussion
Reorganization of cortical connectivity consequent to injury is regarded as a key mechanism in epileptogenesis (see references above). Resulting small changes in the degree of recurrent exci-
tation or inhibition in cortical circuits can lead to the emergence of epileptiform activity (Traub and Wong, 1982; Chagnac-Amitai and Connors, 1989). We used scanning laser photostimulation and whole-cell recording to determine whether the axonal sprouting of layer V pyramidal neurons in the partial neocortical isolation model of posttraumatic epileptogenesis (Salin et al., 1995) is associated with enhanced functional excitatory connectivity. Results indicate that layer V pyramidal cells in the injured cortex receive an excitatory synaptic input that is increased in spatial extent and intensity and is derived principally from presynaptic neurons in layer V. The increased number of EPSCs evoked by each stimulus in the undercut cortex is balanced by the decreased amplitude of individual events, so that the composite EPSC is similar in amplitude in control and injured cortex. However, it is likely that repetitive EPSCs would translate into a larger AP output by layer V pyramidal neurons in vivo, where resting potential is significantly more depolarized (Destexhe and Pare, 1999).

Laser-scanning photostimulation for analysis of circuit reorganization

The combination of UV photostimulation and whole-cell recordings has been used effectively in studies of hippocampal (Molnar and Nadler, 1999; Shao and Dudek, 2004) and neocortical connectivity (Callaway and Katz, 1993; Katz and Dalva, 1994; Kotter et al., 1998; Staiger et al., 2000; Schubert et al., 2001; Callaway, 2002; Dodt et al., 2003; and others). The technique is highly efficient for mapping details of interlaminar and intercolumnar connections in neocortex (Callaway, 2002; Dodt et al., 2003; Shepherd et al., 2003; Shepherd and Svoboda, 2005) and has been applied to demonstrate enhanced connectivity among both CA1 pyramidal neurons and dentate granule cells in models of temporal lobe epilepsy (Molnar and Nadler, 1999; Shao and Dudek, 2004). The advantages and potential limitations have been discussed previously (Shepherd and Svoboda, 2005, and references therein); however, additional issues arise when lesioned and control cortex are to be compared. The density of surviving neurons, their intrinsic excitability, presynaptic terminal and receptor functions, and other factors, such as the efficiency of clearance of released glutamate, may each be altered in the injured cortex, potentially affecting the numbers of cells responding to the uncaging stimulus, the size of the evoked EPSCs, and the resulting maps. Furthermore, responses to injury can vary over time (Fishman and Mattu, 1993; Marchenko et al., 2004), and the current study focused on only one time point. The experiments that demonstrated sprouting in the partially isolated cortex were also done at ~2–3 weeks after injury, but sprouting may be a dynamic process (Marchenko et al., 2004) so that different results might be obtained if mapping were applied earlier or later after injury. Some of these variables have been considered (Fig. 2), but others that may have contributed to the differences between control and partially isolated cortex are yet to be examined.

Enhanced excitation

The hypothesis that there is increased functional excitatory connectivity within the partially isolated cortex is supported by the increased density of hot spots (Figs. 5B, 6B, D), and an increase in the region-normalized EPSC amplitude that results from both the increase in hot spots and an increased incidence of multipeaked flash-evoked EPSCs in these neurons (Figs. 5A, 6A, C). The lack of significant increase in sEPSC frequency and decreased amplitude of sEPSC and single evoked EPSCs in the undercut cortex were unexpected, as previous results indicated that the frequency and amplitude of sEPSCs, and the input–output slope of evoked EPSCs were increased (Li and Prince, 2002). The reasons for this discrepancy may lie in the different experimental conditions in the two series of experiments, including the low temperature and elevated divalent cation concentration in the current study that could have had differential effects on excitatory synaptic transmission in the injured cortex. The EPSC frequencies recorded in the current study at room temperature (~23°C; 2.7 ± 0.3 Hz in control and 3.2 ± 0.7 Hz in undercut) were lower than those reported in an previous study in which recordings were made at 32°C (7.4 ± 0.6 in control and 9.3 ± 0.7 in undercut) (Li and Prince, 2002). Thus, in both studies, there is a trend toward increased frequency of sEPSCs in undercut cortex, but the previous study did demonstrate an increase in sEPSC amplitude that was not reproduced here. A potential explanation for smaller EPSC amplitude in our current study would be that the newly sprouted connections are made selectively onto distal dendritic regions from which the EPSCs would be subjected to significant electrotonic filtering and thus be of smaller apparent magnitude. This result would be consistent with a slowed rise time, as expected for electrotonic filtering; however, it was not possible to reliably resolve rise times in the multipeaked composite responses evoked by the uncaging stimuli. Resolution of this issue will require additional study.

Because the mean composite EPSC amplitude in our mapping experiments is a summation of all EPSCs evoked from a given presynaptic spot, the postsynaptic response would be affected by the number of neurons being activated, the number of APs fired per stimulated neuron, the total number of synaptic contacts between presynaptic and postsynaptic neurons, the site(s) of synaptic contacts, the release probability at presynaptic terminals, and the density and structure of postsynaptic receptors (Shepherd and Svoboda, 2005). Under our experimental conditions, contamination of results by unrecognized polysynaptic responses evoked by stimulation of epileptogenic cortex (Prince and Tseng, 1993) would be unlikely (see Materials and Methods). Also, increased intrinsic excitability known to occur in pyramidal neurons of partially isolated cortex (Prince and Tseng, 1993; Jin et al., 2005a), did not affect the number of APs induced by direct glutamate-induced depolarization (Fig. 2), making enhanced presynaptic excitability an unlikely cause for the increase in multipeaked EPSCs. The multipeaked EPSCs likely reflect events evoked by release of glutamate from the synaptic terminals of several neurons in the area of the hot spot. The increased incidence of multipeaked EPSCs would then be attributable to either a greater number of activated neurons, or increased connectivity from activated cells onto the recorded neuron. In light of the known decrease in density of pyramidal cells in the partially isolated cortex (Prince and Tseng, 1993; Jin et al., 2005a), the latter possibility appears most likely. The distribution of hot spots in our experiments (Fig. 6B) is consistent with the anatomical data showing that sprouting is most prominent within layer V, the lamina in which interictal epileptiform events originate (Prince and Tseng, 1993; Hoffman et al., 1994). The increase in the fraction of sites from which EPSCs are evoked strongly suggests the formation of new synapses that make a major contribution to the stronger and spatially more extensive functional excitatory connections between layer V pyramidal neurons. Because dendritic trees of layer V pyramidal neurons do not undergo outgrowth after undercut lesion (Salin et al., 1995) and there is no increase in the number of evoked APs of layer V cells when stim-
ulated in the upper layer (Fig. 2D), it is likely that axonal sprouting and formation of new synapses in excitatory neurons of more superficial layers also contribute to the significant increases in region-normalized EPSC amplitude and fraction of hot spots in upper cortical layers (Fig. 6A, B).

Functional implications

Our results demonstrate significant increases in functional excitatory connectivity onto layer V pyramidal neurons of undercut cortex, thus supporting the hypothesis that the axonal sprouting found in anatomical experiments (Salin et al., 1995; Marchenko et al., 2004) enhances cortical excitability in this model of post-traumatic epileptogenesis. The spatial distribution of anatomical sprouting and altered uncaging responses of presynaptic cells in layer V is consistent with the origin of interictal epileptiform activity in that lamina (Hoffman et al., 1994). The new recurrent excitatory connections between layer V pyramidal neurons likely result in enhanced synchronization within the cortical network and contribute to the generation of epileptiform activity in the undercut animals both in vitro and in vivo (for review, see Graber and Prince, 2006). Extensive synaptic reorganization and resulting enhanced excitatory connectivity is also a key mechanism underlying epileptogenesis in temporal lobe epilepsy (Wong et al., 1984; Perez et al., 1996; Wuarin and Dudek, 1996; Sutula et al., 1998; Zhang and Houser, 1999; Wenzel et al., 2000; Lehmann et al., 2001; Buckmaster et al., 2002; Smith and Dudek, 2002). Other factors, such as less effective GABAergic postsynaptic inhibition attributable to a variety of mechanisms (Ribak et al., 1982; Rice et al., 1996; Buckmaster and Jongen-Reilo, 1999; Kobayashi et al., 2003), although not evident here, would work hand-in-hand with enhanced recurrent excitation to promote seizures (Traub and Wong, 1982).

Axonal sprouting is likely a ubiquitous response to cortical injury (Maxwell et al., 1990; Fishman and Mattu, 1993), and yet posttraumatic epilepsy occurs infrequently (Annegers et al., 1980), emphasizing the multifatorial nature of epileptogenesis. Our results show that this regenerative process results in functionally stronger and spatially expanded cortical connections that may constitute a substrate for functional recovery after neuronal injuries (Carmichael et al., 2001; Ramirez, 2001; Chockworee et al., 2004) as well as a mechanism underlying epileptogenesis.

References


