An Abrupt Developmental Shift in Callosal Modulation of Sleep-Related Spindle Bursts Coincides With the Emergence of Excitatory-Inhibitory Balance and a Reduction of Somatosensory Cortical Plasticity

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Transecting the corpus callosum of postnatal day (P)1–6 rats disinhibits the production of spindle bursts (SBs) within primary somatosensory cortex (S1), most notably during periods of sleep-related myoclonic twitching. Here we investigated developmental changes in this callosally mediated disinhibition and its association with cortical plasticity. Recordings in P2-15 subjects revealed that callosotomy-induced disinhibition is a transient feature of early development that disappears abruptly after P6. This abrupt switch was accompanied by sharp decreases in myoclonic twitching and equally sharp increases in spontaneous SBs and in the number of GABAergic and glutamatergic presynaptic terminals in S1. Expression of the K⁺/Cl⁻ cotransporter 2 (KCC2) also increased across these ages. To determine whether these developmental changes are associated with alterations in cortical plasticity, pups were callosotomized at P1, P6, or P8, and tested over the subsequent week. Regardless of age, callosotomy immediately disrupted SBs evoked by forepaw stimulation. Over the next week, the P1 and P6 callosotomy groups exhibited full recovery of function; in contrast, the P8 group did not exhibit recovery of function, thus indicating an abrupt decrease in cortical plasticity between P6 and P8. Together, our data demonstrate that callosotomy-induced disinhibition is a transient phenomenon whose disappearance coincides with the onset of increased intrinsic connectivity, establishment of excitatory-inhibitory balance, and diminished plasticity in S1. Accordingly, our findings indicate that callosotomy-induced disinhibition of twitch-related SBs is a bioassay of somatosensory cortical plasticity and, in addition, support the hypothesis that myoclonic twitches, like retinal waves, actively contribute to cortical development and plasticity.

Keywords: plasticity, myoclonic twitching, somatosensory cortex, REM sleep, glutamate, GABA, KCC2

Activity-dependent processes guide the development of interhemispheric connectivity by the corpus callosum during the prenatal and early postnatal period in rats (Mizuno, Hirano, & Tagawa, 2007; Wang et al., 2007). In rodents, callosal fibers cross the midline soon after birth, after which they make contact with the cortical subplate and invade the contralateral cortex during the first postnatal week (Innocenti & Price, 2005; Ivy & Killackey, 1981; Wise & Jones, 1976). We recently found that transecting the corpus callosum (i.e., callosotomy, CCx) of postnatal day (P)1–6 rats disinhibits spontaneous spindle bursts (SBs) in the forelimb region of primary somatosensory cortex (S1; Marcano-Reik & Blumberg, 2008); CCx also disrupted evoked S1 responses to contralateral forepaw stimulation. These findings suggested that the corpus callosum—soon after birth and at least through P6—exerts a net inhibitory influence over homotopic cortical circuits and plays a modulatory role in sensory processing.

SBs are brief 5- to 25-Hz oscillatory bursts and are among the most prevalent early cortical activity patterns (Khazipov et al., 2004). SBs occur in S1 during waking but most prominently during periods of sleep-related myoclonic twitching, and they can be specifically evoked by proprioceptive stimulation (Marcano-Reik & Blumberg, 2008). In addition to S1, SBs are produced in visual (Hanganu, Staiger, Ben-Ari, & Khazipov, 2007) and barrel (Minlebaev, Ben-Ari, & Khazipov, 2007) cortex, but they have not been detected in non-sensory cortical regions (Seelke & Blumberg, 2010). Models of cortical injury and recovery typically involve direct damage to specific areas within cortical lobes (Kolb, Halliwell, & Gibb, 2010). Such studies have revealed that the long-term consequences of cortical damage depend in part on the stage of brain development at the time of injury. For example, in rats, functional recovery after cortical injury is better when the injury is produced during the second postnatal week than during the first postnatal week, that is, after neurons have completed their migration to superficial cortical layers. However, if SBs contribute to the development of somatosensory representations in S1, as has been suggested (Khazipov et al., 2004), then the dramatic CCx-induced disinhibition of SB activity that we observed before P6 might indicate a state of heightened plasticity.

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Here we investigated developmental changes in CCx-induced disinhibition and its association with somatosensory cortical plasticity. We document the sudden disappearance after P6 of CCx-induced disinhibition, accompanied by behavioral, cellular, and molecular changes that are associated with a reduction of one form of cortical plasticity. We propose that CCx in early development offers a new and unique model system for examining the effects of early perturbations of cortical circuitry on somatosensory processing and recovery of function. All together, these findings support the hypothesis that myoclonic twitches serve as a primary source of sensory input driving activity-dependent processes in somatosensory cortex during early postnatal development, similar to the role played by retinal waves in the development of visual cortex (Katz & Shatz, 1996; Wong, 1999).

Materials and Methods

All experiments were performed in accordance with National Institutes of Health guidelines for the care and use of animals in research and were approved by the Institutional Animal Care and Use Committee of the University of Iowa. All efforts were made to minimize the number of animals used.

For all experiments, mothers and their litters were housed in standard laboratory cages (48 × 20 × 26 cm) in the animal colony at the University of Iowa. Food and water were available ad libitum and all animals were maintained on a 12-hr light–dark cycle with lights on at 0700 h. All experiments took place during the lights-on period. Litters were culled to 8 pups within 3 days of birth (day of birth = P0).

Developmental Changes in Cortical Activity in Sham and Callosotomized Pups

Subjects. A total of 64 pups from 32 litters were used. Subjects were tested on P2, P4, P6, P7, P8, P9, P12, or P15 (n = 4 per age). Males and females were equally represented among the subjects.

Surgery. On the day of testing, 2 littermates underwent CCx or sham surgery (with counterbalancing of the order of surgery), as described previously (Marcano-Reik & Blumberg, 2008). Briefly, under isoflurane anesthesia, a 2 to 3 mm opening was created in the skull, halfway between bregma and lambda and parallel and lateral to midline. In CCx subjects, a thin surgical knife was inserted to a depth of approximately 5 mm and was swept in an anterior-to-posterior direction to transect the corpus callosum. Sham surgeries were identical except that the surgical knife was not inserted. In all subjects, 2 recording sites were prepared for placement of cortical surface electrodes over the left and right S1s. Each site consisted of 2 holes separated by approximately 2 mm and centered over the forelimb region of S1, approximately 1 mm rostral to bregma and 2 to 3 mm lateral to midline.

As described previously (Karlsson, Gall, Mohns, Seelke, & Blumberg, 2005; Marcano-Reik & Blumberg, 2008), a custom-built stainless steel apparatus, designed to attach to the earbar holders of a stereotaxic apparatus (David Kopf Instruments, Tujunga, CA), was secured to the skull with cyanoacrylate adhesive. Bipolar stainless steel electrodes (50-µm diameter, California Fine Wire, Grover Beach, CA) were inserted into the left and right nuchal muscles. A ground wire was implanted anterior to the nuchal electromyo-
followed by a 3% formalin solution. Brains were postfixed for at least 48 h in a formalin–sucrose solution before being sliced in the coronal plane (50-μm sections). Light microscopy was then used to assess the extent of damage to the corpus callosum. Similar to what was reported previously using the same method (Marcano-Reik & Blumberg, 2008), approximately 90% of the corpus callosum was typically transected, beginning at the anterior portion and extending posteriorly.

Data analysis. For each subject, 15-min periods of synchronized data comprising digital records of behavior, nuchal EMG activity, and EEG activity were created and analyzed using Spike2 software, as described previously (Marcano-Reik & Blumberg, 2008). For all analyses of sham and CCx littermate subjects, paired (within-subjects) t tests were performed using JMP 5.0 software (SAS, Cary, North Carolina). For all tests, alpha was set at 0.05 and Bonferroni corrections were used to correct alpha for multiple comparisons.

Myoclonic twitches were counted by referring to both the behavioral record (i.e., limb and tail twitching) and the left and right nuchal EMG activity. Thresholds were set in Spike2 to analyze nuchal EMG activity (Mohs & Blumberg, 2008). Cursors were used to identify individual twitches in the EMG records and relate them to behaviorally scored twitches. A memory buffer was created with all the marked twitches, and an analysis was run to count all the events during the 15-min recording period.

Spontaneous and evoked SBs were identified by referring to both the raw EEG records of the left and right S1s as well as filtered records (bandpass: 5–40 Hz). Using criteria similar to those described previously (Khazipov et al., 2004; Marcano-Reik & Blumberg, 2008), SBs were defined as comprising at least 3 complete oscillations, approximately 100 ms in duration, and containing at least 1 oscillation that exceeded 50 μV in amplitude (from baseline to peak). In contrast with spontaneous SBs, evoked SBs were embedded within large, slow potentials with amplitudes ≥100 μV. Finally, SB oscillation frequency and duration were measured in sham and CCx subjects as described previously; consistent with our previous study (Marcano-Reik & Blumberg, 2008), SB oscillation frequency, but not duration, increased after CCx.

SB latencies were measured and compared for all subjects, as described previously (Marcano-Reik & Blumberg, 2008). For this analysis, 20 “anchor” SBs in the left S1 recording were selected at random and its duration measured. Then, the latencies between the anchor SB and the prior (L−) and subsequent (L+) SBs in the right S1 recording were determined. Mean values of L− and L+ were always statistically indistinguishable.

Effect of GABAergic Agonists and Antagonists on Cortical Activity

Subjects and surgical preparation. A total of 32 P4 rats from 8 litters were used. On the day of testing and under isoflurane anesthesia, 2 littermates experienced sham surgery and 2 additional littermates experienced CCx surgery. All subjects were prepared for testing as described above (n = 8 per group). Surgeries were staggered to minimize the latency between surgery and testing.

Procedure. After recovery from surgery, the pup was transferred to the recording apparatus. After acclimation, the recording session began with a 1-μl infusion of the vehicle, artificial cerebrospinal fluid (ACSF), into the left or right S1 and behavior was scored for 15 min immediately after the infusion. After the behavioral recording period, the subject received either a 1-μl infusion of the GABA_A receptor agonist, muscimol hydrobromide (0.01M mixed with ACSF; Sigma-Aldrich, St. Louis, MO), or a 0.5-μl infusion of the GABA_A antagonist, bicuculline methiodide (10 μmol/l; Sigma-Aldrich). The volumes and concentrations for muscimol (Campolattaro & Freeman, 2008) and bicuculline (Jones & Barth, 2002) are based on previously published work. After the infusion, another 15-min period of behavioral scoring was conducted. The remaining 3 littermates were tested in succession and the order of testing (i.e., sham vs. CCx; muscimol vs. bicuculline; left vs. right S1) was randomized.

Infusions were delivered using a Hamilton microsyringe with a 25-gauge needle (Model 7001; Hamilton, Reno, NV) secured to the stereotaxic apparatus and mounted directly above the infusion site. The syringe was lowered just below the cortical surface into a predrilled hole located halfway between the two Ag/AgCl electrode sites. The infusion rate for all infusions was 0.1 μl/s. These methods are identical to those used in a previous report (Marcano-Reik & Blumberg, 2008), and it has been shown previously that surface and intracortical infusions do not differ significantly in their effects on SB activity (Hanganu et al., 2007).

Three of the muscimol subjects received an infusion of the same concentration of fluorescently labeled muscimol (Allen et al., 2008; BODIPY TMR-X conjugate; Invitrogen, Eugene, OR). At the end of the recording session, the pups were overdosed with an intraperitoneal injection of sodium pentobarbital and perfused transcardially with phosphate-buffered saline, followed by a 3% formalin solution. Brains were postfixed for at least 48 h in a formalin–sucrose solution before being sliced in the coronal plane with a sliding microtome (50-μm sections). A fluorescent microscope was used to visualize the distribution of muscimol throughout the brain.

Data analysis. For all subjects, paired (within-subjects) t tests were performed using JMP 5.0 software (SAS, Cary, NC). For all tests, alpha was set at 0.05, and Bonferroni corrections were used to correct alpha for multiple comparisons.

Western Blot Analysis of K⁺Cl⁻ Cotransporter 2 (KCC2) Expression Across Age

Procedure. Cortical tissue including the S1 region of P5, P6, P7, P8, and P9 rats (n = 2 per age) was homogenized in RIPA buffer (150 mM NaCl, 1% Triton X-100, 0.5% deoxycholic acid, 0.1% SDS, 50 mM Tris-HCl, pH 8.0) with Complete, EDTA-free protease inhibitor cocktail (Roche Applied Science, Indianapolis, IN). Concentration was determined by BCA analysis. Samples were incubated in sample buffer (80 mM Tris-HCl, 2% SDS, 10% glycerol, 5.3% β-mercaptoethanol, 0.025% bromophenol blue) at 75 °C for 10 min. Samples were run, 10 μg each, on a 7.5% polyacrylamide gel and then transferred to a nitrocellulose membrane. The membrane was blocked in TBST (Tris Buffered Saline, 0.1% Tween-20) and 5% milk, then incubated overnight at 4 °C with antibodies against KCC2 (Rabbit anti-KCC2, Upstate Cell Signaling Solutions, Lake Placid, NY; 1:1000 dilution in TBST) and β-tubulin (Mouse anti-β-tubulin, Sigma-Aldrich; 1:1000 dilution in TBST). The membrane was washed in TBST,
and probed with corresponding HRP-conjugated secondary antibodies (Goat anti-Rabbit HRP IgG HL, Jackson Immunoresearch, West Grove, PA; 1:10,000 dilution in TBST with 5% milk; Goat anti-Mouse HRP IgG HL, Pierce Biotechnology, Rockford, IL; 1:5000 dilution in TBST with 5% milk) and imaged using SuperSignal West Pico Chemiluminescent Substrate (Pierce Biotechnology, Rockford, IL).

Scanned images of blots were quantified using the “Analyze Gels” functions in ImageJ (National Institutes of Health). The intensity of KCC2 bands was quantified for each lane and normalized to the β-tubulin signal. Data were then expressed as arbitrary units after normalization to the P5 signal.

Disruption and Recovery of Evoked Responses to Forepaw Stimulation After Callosotomy

Subjects and surgical preparation. A total of 132 P1-15 infant rats from 18 litters were used. In one set of 6 litters, 60 littermates (10 per litter) received sham or CCx surgery at P1 and were tested in pairs at P1, P2, P4, P6, or P8. In a second set of 6 litters, 36 littermates (6 per litter) received sham or CCx surgery at P6 and were tested in pairs at P6, P9, or P13. In a final set of 6 litters, 36 littermates (6 per litter) received sham or CCx surgery at P8 and were tested in pairs at P9, P12, or P15 (this dataset was supplemented with previously collected data from 12 subjects and 6 litters in which pups received sham or CCx surgery on P8 and were tested in pairs the same day). Thus, at each age, 6 sham and 6 CCx littermates from 6 litters were tested. Males and females were equally represented among the subjects.

Surgeries were identical to those described above. However, for those subjects that received sham or CCx surgeries at P1, P6, or P8 and were to be tested at a later age, the incision was closed with Vetbond and pups, along with their other littermates, were transferred to a humidified incubator maintained at thermoneutrality (35 °C) to recover for 1 h. At the end of the recovery period, all pups, with the exception of the 2 subjects to be tested that same day, were placed back in the home cage with their mother. At the appropriate time, 2 littermates were transferred to an electrically shielded chamber for acclimation and neurophysiological and behavioral testing. All procedures for recording evoked and spontaneous activity were the same as described above.

When pups were returned to their mother and homecage, daily observations were conducted to ensure that they were healthy and exhibited visible milk bands. Body weights were not different between sham and CCx subjects 1 week after surgery (at P8 after surgery at P1: shams = 20.2–24.1 g; CCx = 19.1–24.5 g; at P13 after surgery at P6: shams = 30.6–36.9 g; CCx = 30.5–40.2 g; at P15 after surgery at P8: shams = 37.4–42.7 g; CCx = 38.9–43.2 g).

Data analysis. For all subjects, paired t tests were performed at each age using JMP 5.0 software (SAS). For all tests, alpha was set at 0.05, and Bonferroni corrections were used to correct alpha for multiple comparisons.

Immunofluorescence

Procedure. To determine normal changes in presynaptic terminals and receptor populations, additional subjects were anesthetized with isoflurane and received sham surgery [i.e., the same surgical procedures as those described above] on P6, P7, P8, or P9 (n = 3 at each age]. Subjects were perfused transcardially with cold 0.1M phosphate buffered saline (PBS) followed by 2% paraformaldehyde in PBS (pH = 7.4). Following perfusion the brains were extracted and postfixed in cold 2% paraformaldehyde in PBS overnight. The brains were then cryoprotected in 30% sucrose in PBS at 4 °C and frozen in OCT compound (Tissue-Tek; Sakura-Finetek) using dry ice/ethanol-cooled isopentane.

Coronal cryostat sections (12 μm) at the level of the corpus callosum were cut on a Leica CM1850 cryostat and collected with the aid of a Leica CryoJane tape transfer system. Tissue sections were air-dried and then blocked in 2.5% bovine serum albumin, 0.1% Trition X-100 in PBS for 1 h, followed by overnight incubation at 4 °C with primary antibodies: guinea pig antivesicular glutamate transporter-1 (VGLUT-1; 1:500, Chemicon) and mouse antiglutamic acid decarboxylase (GAD; mAb GAD-6; 1:10 Developmental Studies Hybridoma Bank) diluted in the same blocking solution. Sections were washed in PBS and incubated for 1 h at room temperature with the appropriate secondary antibodies conjugated to Alexa 488 and 568 (Invitrogen). Sections were washed in PBS containing the nuclear counterstain DAPI (4′, 6′ diamidino-2 phenylindole) and mounted in Gel/Mount (Biomeda) aqueous mounting media.

Image acquisition and analysis. Quantification of presynaptic (VGLUT-1-positive or GAD-positive) puncta was performed on 6 sections per animal from 3 different animals at each time point (i.e., 18 fields for each marker). Images were taken of fields within S1 cortex using 63X Plan Apo objectives on a Leica DM5000B digital epifluorescence microscope. Digital images were captured in Adobe Photoshop and similarly adjusted for brightness and contrast. Images were thresholded in NIH ImageJ and synaptic puncta were counted by using the Analyze Particle module; the accuracy of these automated quantifications was confirmed in a portion of each field by manual counts.

Data analysis. To control for field-by-field variations in cell density, we normalized the number of presynaptic terminals to the number of cell nuclei in each field. The proportion of cells that labeled with antibodies to the neuronal marker NeuN was also determined and did not change between P6 and P9 (data not shown), confirming that total cell density is proportional to neuronal density. Analyses of variance (ANOVs) were separately conducted for each marker (i.e., VGLUT-1 and GAD) and cortical depth (i.e., layers 2–3 and 4–6) using JMP 5.0 software (SAS). Student’s t test was used for post hoc comparisons. Alpha was set at 0.05 and the Bonferroni correction procedure was used.

Results

Abrupt Disappearance of CCx-Induced Disinhibition of Spontaneous SBs After P6

As found previously (Marciano-Reik & Blumberg, 2008), CCx doubled the rate of spontaneous SBs in P1-6 subjects (Figure 1, Top; ps < .001, n = 4/age). This disinhibition was reflected in shorter latencies separating SBs produced by the two hemispheres (Figure 1, Bottom; p < .0001). (Given that rates of twitching were unaffected by CCx (see below) and that SBs largely cluster during periods of myoclonic twitching, it appears that CCx increases the probability that twitches trigger SBs in the contralateral S1.) Then,
abruptly at P7, CCx no longer disinhibited spontaneous SBs in relation to shams. Coincident with this abrupt disappearance of CCx-induced disinhibition, the rate of SBs in both sham and CCx subjects increased sharply and peaked at P8 before steadily decreasing over the next several days.

We next confirmed that the age-related effects of CCx described above were not attributable to alterations of sleep or rates of twitching. Given that disconnection of the cerebral cortex alone does not alter sleep-wake cycling in P8 rats (Karlsson, Kreider, & Blumberg, 2004), we expected CCx to have no effect on the percentage of time asleep in relation to shams; indeed, no effect was found (Figure 2, Top). Also, because twitches are produced within the brainstem and are not dependent on forebrain structures (Kreider & Blumberg, 2000), we anticipated that CCx would not alter or influence their occurrence, which is what was observed (Figure 2, bottom). Importantly we found that over several days beginning at P6, the rate of twitching rapidly declined in both sham and CCx subjects.

**GABAergic Modulation of Spontaneous Spindle Bursts in Intact and Callosotomized Subjects**

We hypothesized that CCx-induced disinhibition of SBs through P6 results from a reduction of excitatory drive onto GABAergic interneurons. To test this hypothesis, we unilaterally infused artificial cerebrospinal fluid (ACSF) just below the S1 cortical surface in P4 rats, followed by an infusion of the GABA_A agonist muscimol or antagonist bicuculline (n = 8 subjects/group). In shams, muscimol did not significantly influence spontaneous SB production whereas bicuculline significantly increased spontaneous SBs (p < .001, n = 8) across the 15-min test period in relation to ACSF (Figure 3A). On a percentage basis, muscimol decreased SBs by only 11.3% whereas bicuculline increased SBs by 60% in relation to ACSF. This finding of a floor effect with respect to GABAergic inhibition suggests that the corpus callosum exerts a powerful inhibitory influence on S1 networks at this age.

If the corpus callosum exerts its effects at P4 by preferentially exciting GABAergic interneurons, then it should be possible to compensate for the reduction of GABAergic modulation after CCx.
by infusing muscimol. This is exactly what was found. Specifically, in relation to ACSF, muscimol and bicuculline significantly reversed and exacerbated the effect of CCx-induced disinhibition on spontaneous SBs. On a percentage basis, muscimol decreased SBs by 71% and bicuculline increased SBs by 64.5% in relation to ACSF (Figure 3A).

In a subset of the muscimol subjects (n = 3), fluorescently labeled drug was infused to confirm the desired distribution of the drug (Figure 3B). Muscimol was clearly localized near the site of infusion, although some diffusion to other cortical and subcortical areas may have occurred. Diffusion to the contralateral hemisphere was not detected; indeed, consistent with this observation, neither muscimol nor bicuculline infusions affected spontaneous SB production in the nondrug hemisphere (data not shown).

To assess developmental changes in GABAergic modulation of SBs, we examined in P10 subjects the effects of muscimol and bicuculline infusion. However, in contrast with its effects at P4, infusion of bicuculline at P10 produced a hyperexcitable state that included epileptiform activity. That such activity would occur after bicuculline infusion at P10 is consistent with a previous report examining age-related differences in epileptiform activity in neocortical slices (Wong & Yamada, 2001).

KCC2 Upregulation at the Beginning of the Second Postnatal Week

The inhibitory and excitatory effects of muscimol and bicuculline, respectively, on SB activity at P4 are consistent with GABA exerting its postsynaptic effects through either shunting inhibition or hyperpolarization. Upregulation of the K⁺/Cl⁻ cotransporter KCC2, through its control of the Cl⁻ reversal potential, is thought to be a critical factor in the postnatal emergence of GABA’s hyperpolarizing effects (Sipilä, Blaesse, & Kaila, 2010). Although KCC2 is known to be upregulated sometime during the first two postnatal weeks in the cortex (Blaesse et al., 2006; Wang et al., 2002), protein levels have not been examined on a day-by-day basis. Therefore, we assessed KCC2 levels in cortex from P5, P6, P7, P8, and P9 rats using Western blot analysis (Figure 4A). As shown in Figure 4B, levels of KCC2 increased abruptly after P7.

Rapid Developmental Changes in GABAergic and Glutamatergic Presynaptic Terminals

We examined whether the rapid changes in SB production between P6 and P9 are accompanied by anatomical changes in GABAergic and glutamatergic presynaptic terminals in S1. Using an additional set of sham subjects (n = 3 pups/age; 6 fields/pup), we stained cryosections with antibodies directed against glutamic acid decarboxylase (GAD) and vesicular glutamate transporter-1 (VGLUT1) in layers 2–3 and 4–6 in S1. We controlled for any variation in cell density by normalizing the number of presynaptic terminals to the number of cell nuclei in each field (see Materials and Methods). We found that GABAergic and glutamatergic presynaptic terminals increased rapidly and significantly at P7 in relation to P6, especially in the deep layers (Figure 5A; ps < .0001).

The marked increase in GAD and VGLUT1 terminals between P6 and P7 can be seen clearly in representative immunofluorescent fields from layers 2–3 and 4–6 (Figure 5B). Along with this sharp increase in the number of presynaptic terminals, cortical cells appeared more widely spaced at P7 and later, resulting in a reduced number of cells per microscope field. This increased spacing is almost certainly because of the rapid expansion of the neuropil and continued growth of the cortex, as no apoptosis was observed between P6 and P7 (based on immunostaining for cleaved caspase-3; data not shown). As noted above, we normalized our synaptic terminal counts to the cell density in each microscope field; thus, these changes in cell density cannot account for the observed increase in the number of presynaptic terminals. Indeed, the absolute (i.e., not normalized) number of presynaptic terminals also exhibited a sharp increase after P6 despite the decrease in cell density. Thus, our histological findings indicate a rapid and steep increase in cortical innervation after P6 that coincides with the loss of the disinhibitory effects of CCx, the upregulation of KCC2, and the developmental decrease in myoclonic twitching.
Immediate Disruptive Effects of Callosotomy on Evoked Spindle Bursts Are Consistent Across Age

We measured the immediate effect of CCx on SB responses in the S1 forelimb region to peripheral stimulation to the plantar surface of the contralateral forepaw. In sham subjects, forepaw stimulation reliably evoked an SB response at all ages (Figure 6); in contrast, CCx significantly reduced the probability of an evoked response by 30% to 50% ($p < .0001$).

Developmental Changes in Recovery of Function After Callosotomy

To assess recovery of function across the early postnatal period, we performed sham or CCx surgeries at P1, P6, or P8 and tested littermates on the same day or up to one week later. As shown in Figure 7A and 7B, CCx at P1 and P6 immediately disrupted evoked responding to forepaw stimulation; over the ensuing week, evoked responding in the CCx pups gradually recovered to sham levels. Also, as expected, at P1 and P6 CCx doubled the rate of spontaneous SBs in relation to shams; over the ensuing week, spontaneous SBs in CCx subjects gradually declined to levels at or below those of shams (data not shown).

As shown in Figure 7C, CCx at P8 also produced an immediate disruption of evoked responding. However, in contrast to pups experiencing CCx at P1 and P6, recovery of function was not observed; instead, steady deterioration of evoked responding occurred over the ensuing week. As expected, disinhibition of spontaneous SBs by CCx was not observed in these subjects.

Figure 5. Changes in GABAergic and glutamatergic presynaptic terminals in postnatal day (P)6–9 rats. (A) Number of primary somatosensory cortex (S1) glutamatergic (VGLUT-1) and GABAergic (GAD) presynaptic terminals in layers 2–3 (left) and 4–6 (right), normalized to the number of cell nuclei, increase significantly at P7 in relation to P6. * Significant difference from value at P6. Means ± SEM. (B) Sections through S1 cortex in P6, P7, P8, or P9 sham-operated animals were stained with antibodies against inhibitory (GAD, green) and excitatory (VGlut-1, red) presynaptic terminals; nuclei were counterstained with DAPI (blue). The density of both GABAergic and glutamatergic terminals in layers 2–3 (top row) and layers 4–6 (bottom row) increases sharply between P6 and P7, with little subsequent change over the next 2 days.

Figure 6. Callosotomy (CCx) disrupts the spindle burst response to peripheral stimulation of the contralateral forepaw in postnatal day (P)2–15 rats. In sham subjects, the response to stimulation of the forepaw reliably evokes an SB in contralateral primary somatosensory cortex (S1) in P2–P15 infant rats; however, CCx immediately decreases the response to forepaw stimulation at all ages. * Significant difference between groups. Means ± SEM.
Discussion

We have documented contemporaneous behavioral, neurophysiological, anatomical, and molecular changes that suggest a role for sleep-related motor activity and its associated sensory feedback in modifying the structure and function of cortical circuits during early development. Specifically, we saw a rapid decrease in myoclonic twitching after P6, at which time CCx-induced disinhibition of spontaneous SBs also disappeared abruptly. In addition, both GABAergic and glutamatergic innervation of superficial and deep cortical layers increased sharply at P7 as the rate of spontaneous SBs in both sham and CCx subjects increased dramatically, peaked at P8, and then decreased over the next week. KCC2 expression also increased across these ages, suggesting that developmental changes in GABAergic functioning are associated with changes in the intrinsic excitability of local cortical circuits involved in the production of SBs. These behavioral, neurophysiological, neuroanatomical, and molecular changes after P6 correlated with an abrupt reduction in somatosensory cortical plasticity. Of note, the recovery of reliable evoked SBs after CCx was predicted by CCx-induced increases in the production of spontaneous SBs, thus suggesting that sensory feedback from myoclonic twitching is a participating factor in cortical plasticity and recovery of function.

Myoclonic twitch movements of the limbs, tail, and head are produced by phasic, rapid activation of skeletal muscles (Gramsbergen et al., 1970; Karlsson & Blumberg, 2002; Seelke, Karlsson, Gall, & Blumberg, 2005; for review, see Blumberg & Seelke, 2010). Neurons within the mesopontine region participate in the production of twitches during the early postnatal period (Karlsson et al., 2005; Kreider & Blumberg, 2000). Recently, it was discovered that sensory feedback from twitching produces SBs in somatosensory cortex (Khazipov et al., 2004); moreover, it appears that it is the proprioceptive feedback from limb twitching that triggers somatosensory SBs (Marcano-Reik & Blumberg, 2008). In addition, SBs are produced in visual (Hanganu et al., 2007) and barrel (Minlebaev et al., 2007) cortex, and hippocampal activity is also modulated by sensory feedback from twitching (Mohns & Blumberg, 2008, 2010). Thus, it is now clear that sensory feedback from twitching modulates neural activity throughout the neuraxis—from spinal cord (Petersson, Waldenström, Fåhraeus, & Schouenborg, 2003) to forebrain—which supports the hypothesis that sleep-related endogenous activity shapes and refines neural circuits in early development and across the lifespan (Blumberg & Lucas, 1996; Blumberg & Seelke, 2010; Corner et al., 1980; Roffwarg, Muzio, & Dement, 1966).

From the data in Figure 2—and given that circadian effects on sleep-wake activity at these ages are relatively small (Gall, Todd, Ray, Coleman, & Blumberg, 2008)—we estimate that the nuchal muscle twitches over 38,000 times per 24 h at P6; this number decreases to 9,500 at P9, a fourfold decrease over 3 days. Within the forelimb region of somatosensory cortex, we estimate that nearly 4,000 SBs occur per 24 h at P6, a value that jumps to over 15,000 at P8 before declining to levels below 4,000 by P12. These values highlight the considerable quantity of endogenous activity produced and experienced by infant rats over the first two postnatal weeks. Thus, in early development, twitching may serve a similar role for the somatosensory cortex as retinal waves serve for the visual cortex (Katz & Shatz, 1996; Wong, 1999). However, further work is needed to establish a role for twitching in the

Figure 7. Effect of callosotomy (CCx) on evoked spindle burst responses to contralateral forepaw stimulation and recovery of function across different ages. Subjects received sham (squares) or CCx (triangles) surgery at (A) postnatal day (P)1, (B) P6, or (C) P8, with subsequent testing over the ensuing week. In subjects that received CCx at P1, P6, and P8, evoked responding was immediately disrupted. Recovery of function was observed after 7 days in pups that received CCx at P1 or P6 (shaded boxes) but not in pups that received CCx at P8. * Significant difference between groups. Means ± SEM.
development and refinement of cortical and subcortical circuits and to determine whether its role involves permissive or instructive influences (Craig, 1999).

Our findings suggest that callosal fibers inhibit spontaneous SBs during the first postnatal week via excitatory effects on GABAergic interneurons (see Khazipov, 2009, for further evidence of a role for GABA in SB modulation). Although activation of GABA_A receptors typically depolarizes postsynaptic cells in immature cortex before the developmental upregulation of KCC2, GABA-mediated inhibition can still occur as a result of shunting inhibition of glutamatergic excitatory postsynaptic currents (Akerman & Cline, 2007; Ben-Ari, 2002; Lamza, Palva, Rausuvuori, Kaila, & Taira, 2000; Owens & Kriegstein, 2002; Sipilä et al., 2010). For example, similar to what was found here, in the newborn hippocampus muscimol and bicuculline decrease and increase network activity, respectively (Lamsa et al., 2000). However, the pharmacological manipulations used here are not sufficient to discern the mechanism by which GABA inhibits postsynaptic activity.

The fact that KCC2 is upregulated in the cortex at the beginning of the second postnatal week suggests that GABA does indeed exert depolarizing effects at earlier ages. This upregulation is also consistent with previous findings in cortical tissue across a broader range of ages. Specifically, KCC2 mRNA in cortex increases between P1 and P15 (Wang et al., 2002) and KCC2 immunoreactivity is very low at P4 and uniformly high at P12 (Blaesse et al., 2006). In rat neocortical brain slices, the GABA_A receptor reversal potential approaches the resting membrane potential sometime after P4 (Owens, Boyce, Davis, & Kriegstein, 1996). Similar to what we found here in cortex, KCC2 expression in hippocampus also increases sharply between P5 and P9 (Rivera, Voipio, & Kaila, 2005).

The callosal inhibition observed here through P6 might be attributable in part to a transient GABAergic population of callosal neurons comprising approximately 21% of the callosal bundle in infant rats (Cobas, Alvarez-Bolado, & Fairen, 1988; Kimura & Baughman, 1997). Significantly, these GABAergic neurons largely disappear at or before P6 and few such fibers are detected in adults (Fabri & Manzoni, 2004). The connectivity of these transient neurons in the contralateral hemisphere is not yet known.

Figure 8 presents a model depicting hypothesized changes in intrinsic cortical circuitry and callosal influences on that circuitry between P6 and P10. At P6, the local circuit producing SBs comprises glutamatergic pyramidal cells and GABAergic interneurons; at this age, GABAergic inhibition occurs through shunting. The fact that CCx disinhibits SB activity at this age suggests net excitation in the intrinsic cortical circuit that is counteracted by a net inhibitory influence provided by callosal excitation of GABAergic interneurons and/or direct inhibition of pyramidal cells by the transient population of GABAergic callosal neurons. By P10, after KCC2 upregulation, the emergence of hyperpolarizing GABA, and the other changes in intrinsic connectivity described here, we hypothesize that excitatory-inhibitory balance is now expressed within the intrinsic cortical circuit. In addition, we hypothesize that callosal fibers, which are thought to exert both excitatory and inhibitory effects in adults (Bloom & Hynd, 2005; Makarov, Schmidt, Castellanos, Lopez-Aguado, & Innocenti, 2008), exert a more balanced influence on the intrinsic circuit by P10. Thus, we view excitatory-inhibitory balance as arising from complex interactions among intrinsic cortical circuits and extrinsic influences that include the corpus callosum.

In addition to their neurophysiological effects, glutamate and GABA may act as trophic signals in early development that, among other things, stimulate the release of brain-derived neurotrophic factor (BDNF; Akerman & Cline, 2007; Ben-Ari, 2002; Marmigere, Rage, & Tapia-Arancibia, 2003; Marty, Berzaghi Mda, & Berninger, 1997; Owens & Kriegstein, 2002; Xiong et al., 2002). BDNF is produced by cortical pyramidal cells and, in an activity-dependent manner, regulates synaptic interactions among glutamatergic pyramidal cells and GABAergic interneurons (Turrigiano, 1999). As a consequence, twitch-triggered SBs in the developing neocortex may, through the release of glutamate, GABA, and BDNF, contribute to the somatosensory cortical plasticity observed here during the first postnatal week. Moreover, the endogenously generated SBs may provide the activity upon which KCC2 upregulation and the emergence of hyperpolarizing GABA depend. This activity may be mediated by GABA itself (Ganguly, Schinder, Wong, & Poo, 2001), by glutamate (Kanold, 2009), or by other factors (Ludwig, Li, Saarma, Kaila, & Rivera, 2003).

After P6, as rates of twitching sharply decline and GABA no longer exerts depolarizing effects, we hypothesize that associated declines in the activity-dependent release of BDNF will lead to relative increases in excitatory drive onto pyramidal cells, as has been hypothesized for cultured cortical networks (Turrigiano, 1999). These adjustments in network organization, which may also
include the observed relative increases in glutamatergic presynaptic terminals (Figure 5), may contribute to the peak in SB activity at P8 and the emergence of excitatory-inhibitory balance (Akerman & Cline, 2007; Hensch, 2005; Owens & Kriegstein, 2002).

In light of all of these foregoing issues, we predicted that CCx-induced disinhibition during the first postnatal week would serve as a bioassay of somatosensory cortical plasticity. To test this prediction, we performed sham or CCx surgeries at P1, P6, or P8 and tested littermates immediately or over the ensuing week. Although the evoked response to contralateral forepaw stimulation is typically very reliable, CCx consistently disrupted evoked responses to forepaw stimulation. Only pups receiving CCx at P1 or P6—ages at which CCx disinhibits spontaneous SB activity—exhibited recovery of function. In contrast, the evoked SB responses of pups that received CCx at P8 deteriorated further over the week. Although we do not yet know whether CCx-induced disinhibition of spontaneous SBs is causally related to recovery of function in S1, it does appear that disinhibition can be used as a bioassay of plasticity-promoting conditions conducive to recovery of function. Moreover, the present findings suggest a new model for exploring somatosensory cortical plasticity—involving disruption of the local circuit by CCx—that complements other models of cortical plasticity (Aton et al., 2009; Feldman, 2009; Feller & Scanzi, 2005).

It is interesting that CCx before P6 doubles the expression of spontaneous SBs but decreases the likelihood of evoked SBs to contralateral forepaw stimulation (compare the developmental profiles in Figure 1, Top, and Figure 6). We have previously reported other pronounced differences between spontaneous and evoked SBs (Marcano-Reik & Blumberg, 2008). Such differences could reflect the contributions of nonoverlapping neural circuits to the production of spontaneous and evoked SBs, including perhaps the production of an efference copy in association with spontaneous SBs. Efference copy provides a mechanism by which animals distinguish sensations arising from self-generated or passive movements (Blakemore, Wolpert, & Frith, 2000; Cullen, 2004). Thus, we hypothesize that efference copy associated with self-generated twitches, perhaps communicated through the cholinergic basal forebrain (Juliano, Ma, & Eslin, 1991; Kilgard & Merzenich, 1998; Ma, Hohmann, Coyle, & Juliano, 1989; Rasmusson & Dykes, 1988; Tremblay, Warren, & Dykes, 1990), prepares the neocortex for twitch-related sensory feedback, thereby contributing to sensorimotor development and somatotopic organization. Of note, cholinergic afferents from the basal forebrain facilitate spindle burst activity in primary visual cortex in P5-6 rats (Hanganu et al., 2007).

All together, the changes in twitch behavior and associated SB activity observed here suggest that the first postnatal week in rats ends with an increase in intrinsic cortical synaptic connectivity, a transformation in GABA’s functional properties, the establishment of excitatory-inhibitory balance, and a diminution of one form of somatosensory cortical plasticity. These findings also suggest that the corpus callosum participates in the activity-dependent development and refinement of cortical networks and point the way to an understanding of how malformations of the corpus callosum may contribute to a variety of neurological and psychiatric disorders (Paul et al., 2007).

References


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