Repeated Stress Alters Dendritic Spine Morphology in the Rat Medial Prefrontal Cortex

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ABSTRACT

Anatomical alterations in the medial prefrontal cortex (mPFC) are associated with hypothalamic-pituitary-adrenal (HPA) axis dysregulation, altered stress hormone levels, and psychiatric symptoms of stress-related mental illnesses. Functional imaging studies reveal impairment and shrinkage of the mPFC in such conditions, and these findings are paralleled by experimental studies showing dendritic retraction and spine loss following repeated stress in rodents. Here we extend this characterization to how repeated stress affects dendritic spine morphology in mPFC through the utilization of an automated approach that rapidly digitizes, reconstructs three dimensionally, and calculates geometric features of neurons. Rats were perfused after being subjected to 3 weeks of daily restraint stress (6 hours/day), and intracellular injections of Lucifer Yellow were made in layer II/III pyramidal neurons in the dorsal mPFC. To reveal spines in all angles of orientation, deconvolved high-resolution confocal laser scanning microscopy image stacks of dendritic segments were reconstructed and analyzed for spine volume, surface area, and length using a Rayburst-based automated approach (8,091 and 8,987 spines for control and stress, respectively). We found that repeated stress results in an overall decrease in mean dendritic spine volume and surface area, which was most pronounced in the distal portion of apical dendritic fields. Moreover, we observed an overall shift in the population of spines, manifested by a reduction in large spines and an increase in small spines. These results suggest a failure of spines to mature and stabilize following repeated stress and are likely to have major repercussions on function, receptor expression, and synaptic efficacy. J. Comp. Neurol. 507: 1141–1150, 2008.

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Indexing terms: dendritic spine, morphometry, plasticity, prefrontal cortex, stress

This article includes Supplementary Material available via the Internet at http://www.interscience.wiley.com/jpages/0021-9967/suppmat. Grant sponsor: National Institutes of Health; Grant number: MH58911 (to B.S.M., J.H.M., P.R.H.); Grant number: MH60734 (to S.L.W.); Grant number: DC05669 (to S.L.W.); Grant number: RR16754 (to S.L.W., P.R.H.). Grant sponsor: Anxiety Disorders Association of America Young Investigator Award (to J.J.R.); NARSAD Young Investigator Award (to J.J.R.).

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Received 6 July 2007; Revised 4 September 2007; Accepted 30 October 2007

DOI 10.1002/cne.21588

Published online in Wiley InterScience (www.interscience.wiley.com).
Life stress has been demonstrated to play a critical role in the etiology of numerous psychiatric illnesses, such as depression, posttraumatic stress disorder, and anxiety (Sapolsky, 1996; Heim et al., 1997; McEwen, 1998; Newport and Nemeroff, 2000; Caspi et al., 2003), and in other instances it may precipitate the onset of psychotic disorders (Corcoran et al., 2001; Phillips et al., 2007). Functional imaging studies in humans suggest that medial prefrontal cortex (mPFC) dysfunction is correlated with abnormalities in neuroendocrine regulation as well as with cognitive and affective changes that are symptomatic in many stress-related mental illnesses (Newport and Nemeroff, 2000; Shin et al., 2001, 2004). Rodent models have provided much information about prefrontal cortical function and its role in emotional stress regulation. One overarching principle that has emerged from this research is that the mPFC has the capacity to modulate the stress-responsive hypothalamic-pituitary-adrenal (HPA) axis (Diorio et al., 1993; Figueiredo et al., 2003; Radley et al., 2006a) yet is also a target of the effects of repeated stress (Cook and Wellman, 2004; Radley et al., 2004, 2006b; Murmu et al., 2006; Cerqueira et al., 2007a). Detailed investigation of how repeated stress affects neuronal morphology in the mPFC may provide a cellular and synaptic basis for the neuroendocrine and behavioral manifestations of stress, particularly with respect to long-lasting pathology. Several reports have shown that repeated stress induces the retraction and debranching of apical, but not basal, dendrites in mPFC pyramidal neurons (Cook and Wellman, 2004; Radley et al., 2004, 2005; Brown et al., 2005; Liston et al., 2006). In addition, repeated stress also reduces apical dendritic spine density in the mPFC (Radley et al., 2006b).

In recent years, much progress has been made in understanding the relationship between dendritic spine structure and function. Dendritic spines are highly motile structures that have the capacity to undergo rapid changes in response to physiological and environmental stimuli (Woolley and McEwen, 1994; Kirov et al., 2004; Yuste and Bonhoeffer, 2004; Zhang et al., 2005). In this regard, the shape of dendritic spines has been shown to provide for biochemical compartmentalization that is critical for regulating synaptic plasticity (for review see Nimchinsky et al., 2002). Repeated-stress-induced dendritic retraction in the mPFC has recently been shown to correlate with functional deficits in attentional control (Liston et al., 2006). Thus, an analysis of how chronic stress modifies dendritic spine morphology may help in elucidating the cellular mechanisms that give rise to such mPFC-dependent behavioral alterations.

Recently, we developed a novel algorithm, Rayburst, for automated three-dimensional shape analysis from laser scanning microscopy images (Rodriguez et al., 2006a). Rayburst defines and casts a multidirectional core of rays from an interior point to the surface of a solid, allowing precise quantification of anisotropic and irregularly shaped 3D structures. The development of this Rayburst-based approach has allowed for the implementation of very-high-resolution three-dimensional (3D) morphometric analyses of dendritic spines on mPFC pyramidal neurons, which permits rapid and automated digitization, 3D reconstruction, and geometric analyses on deconvolved optical stacks of dendritic segments (Wearne et al., 2005; Rodriguez et al., 2006). In this study, we utilized the Rayburst-based automated approach to examine the effect of repeated restraint stress on dendritic spine morphology in Lucifer yellow-filled pyramidal neurons in layers II/III of the rat mPFC. Part of this work has been previously reported in abstract form (Radley et al., 2006c).

MATERIALS AND METHODS

Animals and treatments

Adult male Sprague-Dawley albino rats (300–350 g by the first day of the experimental treatments) were group housed in a temperature-controlled room on a 12-hour light/dark cycle with the light period beginning at 7:00 AM. Food and water were available ad libitum. Rats were adapted to handling for 7 days before any manipulations were performed. All procedures were conducted in accordance with the National Institutes of Health Guide for the care and use of laboratory animals and were approved by the Rockefeller University and Mount Sinai School of Medicine Institutional Animal Care and Use Committees.

Rats were placed in wire mesh restrainers for 6 hours starting at 10:00 AM. Stress exposure was timed to occur in the middle of the lights-on phase of the diurnal cycle. The restraint stress procedure was performed daily for 21 days. After the last day of the repeated stress procedure, rats were anesthetized with chloral hydrate (350 mg/kg, i.p.) and perfused via the ascending aorta at a rate of 100 ml/minute with 100 ml ice-cold 1% paraformaldehyde in 0.1 M phosphate-buffered saline (PBS; pH 7.4), followed by fixation with 4% paraformaldehyde with 0.125% glutaraldehyde in 0.1 M PBS. The brains were removed and postfixed for 2 hours in the same fixative.

Intracellular dye injections

Serial 250-μm-thick coronal sections were collected from the frontal pole using a Vibratome and stored in 0.1 M PBS. Sections were briefly immersed in a fluorescent nucleic acid stain (4,6-diamidino-2-phenylindole; Sigma, St. Louis, MO) to display the cytoarchitectonic features of the prefrontal cortical subfields of interest for intracellular filling. Sections were mounted on nitrocellulose filter paper, submerged in a tissue culture dish containing 0.1 M PBS, and viewed under fluorescence using a fixed-stage microscope. Intracellular fluorescent dye filling was carried out by microinjecting layer II/III pyramidal neurons in the dorsal mPFC with 5% Lucifer yellow (Molecular Probes, Eugene, OR). Injections were made by iontophoresis through micropipettes (1–2 μm inner diameter) under a DC current of 1–6 nA for 10 minutes. The sections were then mounted onto glass slides and coverslipped in PermaFluor.

Neuronal and dendritic reconstructions

To ensure that the neuronal reconstructions and data analyses were performed by an experimenter unaware of the treatment condition, each animal was coded by an independent observer, and the code was not broken until all analyses were completed. The selection method for dendritic segmental spine morphologic analysis was designed to minimize any possible bias. Initially, pyramidal neuron dendritic arbors were reconstructed in 3D using a computer-assisted morphometry system consisting of a Zeiss (Thornwood, NY) Axioplan 2 photomicroscope equipped with an Applied Scientific Instrumentation (Eu-
gene, OR) MS-2000 XYZ computer-controlled motorized stage, a DAGE-MTI DC-330 video camera, a Gateway computer, and morphometry software (MBF Biosciences, Williston, VT). Neurons were visualized, and the dendritic tree was reconstructed with a Zeiss Apochromat ×40 objective with a numerical aperture (N.A.) of 1.4 and Neurolucida software (MBF Biosciences). Neurons had to exhibit complete filling of the dendritic tree, as evidenced by well-defined endings, and display intact primary, secondary, and tertiary branches in order to be considered for further analysis.

Two-dimensional dendritic tracing diagrams that were rendered for each neuron from Neurolucida (MBF Biosciences) were printed, and concentric circles were drawn at radial increments of 75 μm relative to the soma (Fig. 1). Dendritic branches falling within the domains bounded by these concentric circles (i.e., <75 μm, 75–150 μm, >150 μm) were randomly selected for visualization on the confocal laser scanning microscope (CLSM; Zeiss LSM 410, Oberkochen, Germany). Lucifer yellow was excited by an Ar/Kr laser at 488 nm (attenuation set at 10). Settings for pinhole size, aperture, gain, and offset were optimized initially and then held constant throughout the study to ensure that all images were digitized under the same illumination conditions at a resolution of 256 × 256 pixels. Strict criteria for dendritic segment selection were used prior to reconstruction on the CLSM: 1) each had to reside within a depth of 50 μm from the top surface of the section, because of the limited working distance of the optical system; 2) each had to be either parallel to or at acute angles relative to the coronal surface of the section; and 3) there was no overlap with other branches that would obscure visualization of spines. Each designated segment (25–50 μm in length) that met these criteria was located in the microscope field, and confocal stacks of 100–250 digital images separated by a z-step of 0.1 μm were captured using a Zeiss Plan-Achromat ×100 (1.4 N.A.) oil-immersion objective on the CLSM. All confocal stacks included at least 1 μm above and below the identified dendritic segment. On average four high-resolution z-stacks were captured per neuron (a total of 50 neurons were used for spine analysis: 25 from control and 25 from repeatedly restrained animals, N = 5 animals per group). The confocal stacks were deconvolved with AutoDeblur (version 8.0.2.; Autoquant, Troy, NY). For data presentation, composites from deconvolved optical stacks were exported first to Adobe Photoshop (version 7; Adobe Systems, San Jose, CA) for adjustments to optimize/balance contrast and brightness and then to Canvas (version 8; Deneba Systems, Miami, FL) for assembly and labeling.

The Rayburst-based automated system digitizes, reconstructs in three dimensions, and performs geometric analyses on deconvolved confocal stacks of dendritic segments in two principal steps (Fig. 2; Wearne et al., 2005; Rodriguez et al., 2006). First, the medial axis of the dendritic segment is extracted using a dynamically adjusting local segmentation technique. Next, voxels in the vicinity of the medial axis are clustered into spines by building layers from local high points toward the dendrite surface. A 3D Rayburst Sampling algorithm run from the center of mass of each spine cluster extracts spine geometric features, returning volume and surface area for spines oriented at any angle with respect to the dendrite (Rodriguez et al., 2006). The open-source code for implementing the algorithm is freely available for download from the Computational Neurobiology and Imaging Center, Mount Sinai School of Medicine, website http://www.mssm.edu/cnic/tools.html (Wearne and Hof). It is also available from the Journal of Comparative Neurology website as an on-line supplement. The program is implemented in ANSI C and as such is platform-independent. It will run on any computer once it is compiled with any compliant C compiler.
Additional details and performance tests on specific machines were previously reported (Rodriguez et al., 2006).

**Statistical analysis**

Values from each morphometric parameter (spine volume, spine surface area, spine length) on each dendritic segment were averaged, and then averaged for each animal, and values per animal were used for the comparison of means. Each treatment (control and 21 days stress) consisted of five animals, with five Lucifer yellow-filled neurons analyzed from each animal and four dendritic segments analyzed for a given neuron. The normality of the population of dendritic segments was examined with the Kolmogorov-Smirnov test. Given that the distributions for each morphometric parameter analyzed were Gaussian, the group means were compared with a mixed-design ANOVA with one within (apical: 75–150 μm, 75–150 μm, 75–150 μm; basal: 75 μm, 75–150 μm)- and one between (control, stress)-group variable, followed by post hoc pairwise comparisons with Bonferroni’s correction. In addition, to determine whether changes in spine dimensions occurred equally across the entire spectrum of size measurements or are more pronounced in specific subpopulations of spines (e.g., the smallest and largest), all spine parameters from both groups were pooled and the 25th and the 75th percentiles for each parameter were determined. Then, for a given size parameter, the differences in proportions of spines below the 25th percentile, as well as above the 75th percentile, were examined between groups. The χ² test was used to compare the proportions below the first quartile cutoffs between the groups as well as the proportions above the third quartile cutoffs (Knafo et al., 2005). Grouped data were expressed as the mean ± SEM based on one aggregate, or average, per animal.

**RESULTS**

Previous reports have shown that the repeated restraint procedure (6 hours/day, 21 days) used in the present study activates the stress-responsive HPA axis and produces modest, but significant, increases in adrenal weights compared with unstressed rats (Watanabe et al., 1992; Magarinos and McEwen, 1995). Although we did not assess either of these indices of stress activation, we did verify that 21 days of repeated daily restraint stress resulted in a 15% reduction in body weight compared with unstressed controls (P < 0.005; see Watanabe et al., 1992).

The precise location of Lucifer yellow-filled pyramidal neurons in layers II/III of the dorsal mPFC was evaluated with reference to standard cytoarchitectonic parcellations of the region (Krettek and Price, 1977; Vogt and Peters, 1981). The two fields of primary interest are the dorsal subdivision of the anterior cingulate area (ACd) and the prelimbic area (PL). With the fluorescent nucleic acid counterstain (4,6-diamidino-2-phenylindole), the ACd is characterized by a sparse layer II and a broad layer V, distinguishing it from PL, which contains a more homogeneous layer V comprising larger nuclei. Layer I is nearly devoid of neurons, in contrast to the cell-dense layers II/III, permitting clear establishment of laminar borders. Figure 3 provides an example of the locations for each of the Lucifer yellow-filled pyramidal neurons in the dorsal mPFC that were subject to the dendritic tracing and spine morphometric analysis.

In a previous report, repeated stress was shown to induce spine loss in apical dendrites of dorsal mPFC pyramidal neurons (Radley et al., 2006b). Although this earlier study reported spine density estimates from confocal stacks of three-dimensionally reconstructed dendritic segments (25–50 μm in length), spine densities were manually determined by counting the number of spines for each.
length of dendritic segment and expressing values in terms of spines per micrometer. Changes in mean spine density as a function of treatment status were readily determined from the Rayburst-based system, because an initial step in the algorithm involves the extraction of the medial axis of the dendritic segment and recognition of spines using a dynamically adjusting local segmentation technique. Fifty branches from the data set were randomly subjected to counting using both manual and Rayburst-based approaches. The Rayburst-based spine analysis system yielded spine densities that were 87% similar to the manual counting approach.

Repeated restraint stress produced an overall 11% decrease in spine density throughout apical and basal dendrites of dorsal mPFC pyramidal neurons in layers II/III ($F_{1,8} = 8.3; P < 0.05$; Fig. 4, upper left) and an overall 12% decrease in apical dendritic spine density ($P < 0.05$; Fig. 4, middle left). There was also a main effect for dendritic spine density relative to distance from the soma ($F_{4,8} = 10.1; P < 0.005$) and no interaction between experimental treatment and distance from the soma ($F_{4,8} = 1.6; P = 0.2$). By location on the apical dendritic arbor, the stress-induced reduction in spine density was most pronounced at >150 µm relative to the soma (15% decrease; $P < 0.05$; Fig. 4, middle left). Although stress did not have any significant overall effect on basal dendritic spine density, a 13% decrease in spine density was present at radial distances of <75 µm ($P < 0.05$; Fig. 4, lower left).

The dendritic spine morphometric parameters of volume, surface area, and length were assessed for each dendritic spine using the Rayburst-based automated approach. In total, 187 dendritic segments were analyzed (92 control, 97 repeated stress) and 17,078 dendritic spines (8,091 control, 8,987 repeated stress). Based on five animals/group and five neurons/animal, means of 3.7 and 3.9 dendritic segments/neuron were analyzed for the control and repeated stress groups, respectively. In un-stressed controls, on average 12 apical and 6.4 basal dendritic segments were analyzed per animal (totaling 5,983 and 2,708 spines, respectively), and, in the repeatedly stressed group, on average 12.4 apical and 7 basal dendritic segments were analyzed per animal (totaling 5,587 and 3,400 spines, respectively).

Repeated restraint stress resulted in an overall 10% decrease in mean dendritic spine volume ($F_{1,8} = 7.3; P < 0.05$; Fig. 5) and surface area ($F_{1,8} = 15.3; P < 0.005$; Fig. 6) and no effect on spine length ($F_{4,8} = 2.2; P = 0.2$; Fig. 7) in pyramidal neurons in layers II/III of the dorsal mPFC. With regard to mean spine volume, there was also a main effect for distance from the soma ($F_{4,8} = 3.9; P < 0.05$) and no interaction ($F_{4,8} = 0.6; P = 0.6$), whereas no effects for distance ($F_{4,8} = 1.3; P = 0.3$) or interaction ($F_{4,8} = 0.6; P = 0.7$) were evident for mean spine surface area.

Fig. 3. Anatomical localization of Lucifer yellow-filled layer II/III pyramidal neurons in mPFC for subregional analysis. Atlas plates (modified from Swanson, 1992) of coronal sections through a similar level of the mPFC from animals that received intracellular Lucifer yellow injections. Neurons from layers II/III in the dorsal mPFC were identified by using a fluorescent nucleic acid stain, followed by iontophoretic injections with Lucifer yellow. The location for each filled neuron is designated for each treatment with a triangle (control, blue; stress, red). ACd, anterior cingulate cortex, dorsal subdivision; ACv, anterior cingulate cortex, ventral subdivision; PL, prelimbic area; IL, infralimbic area.
Similar to the effects of stress on dendritic spine density, the reduction in mean spine volume and surface area is most prominent in the more distal aspect (>150 μm) of apical dendrites. Although there was no overall effect of stress on basal dendrites, a significant reduction of spine density was present at <75 μm from the soma (lower left). The spine density values for each dendritic segment were quantified using the Rayburst-based automated approach and were similar to previous estimates that involved manually counting spines from optical stacks by controlling the plane of focus for z-step increments and marking spines as they appear on the dendritic segment (Radley et al., 2006b). Examples of dendritic segments are shown in the middle (control) and right (21 days stress) columns. *Differs significantly from unstressed controls at \( P < 0.05 \). Scale bar = 10 μm.

Fig. 4. Repeated restraint stress reduces dendritic spine density in layer II/III pyramidal neurons. Repeated restraint stress (21 days, 6 hours/day) induces decreases in overall (upper left) and apical dendritic spine density (middle left) dendritic spine density. This effect is most prominent in more distal portions (>150 μm) of apical dendrites. Although there was no overall effect of stress on basal dendrites, a significant reduction of spine density was present at <75 μm from the soma (lower left). The spine density values for each dendritic segment were quantified using the Rayburst-based automated approach and were similar to previous estimates that involved manually counting spines from optical stacks by controlling the plane of focus for z-step increments and marking spines as they appear on the dendritic segment (Radley et al., 2006b). Examples of dendritic segments are shown in the middle (control) and right (21 days stress) columns. *Differs significantly from unstressed controls at \( P < 0.05 \). Scale bar = 10 μm.

To determine whether there was an overall shift in the spine size distribution following repeated stress or a disproportionate change in the incidence of spines of smaller volume, surface area, and length, the frequencies of spines in the top and bottom quartile for the subject population as a whole (i.e., control and repeated stress groups) were analyzed. For volume and surface area, the proportions of spines on apical and basal dendrites following repeated stress were higher in the lowest quartile, and lower in the highest quartile, relative to unstressed controls (\( P < 0.001 \); Table 1). A similar shift in spine length distribution
was also present, but the effect was limited to apical dendrites \( (P < 0.001; \text{Table 1}) \). Spines on apical dendritic segments at radial distances >150 \( \mu \)m from soma exhibited the most pronounced shift in frequency distribution in all three dimensions examined (Table 1, Fig. 8).

**DISCUSSION**

These results show that dendritic spine morphology in mPFC pyramidal neurons is altered in response to repeated restraint stress. This analysis was made possible by using a Rayburst-based spine analysis system (Wearne et al., 2005; Rodriguez et al., 2006), which affords the rapid and automated calculation of spine morphometric parameters in 3D. Mean values for spine volume and surface area underwent an 11\% reduction on apical and basal dendritic arbors in pyramidal neurons following repeated stress. Such decreases in spine size are likely to have major repercussions on function, receptor expression, and efficacy. The more distal aspect of apical dendritic arbors exhibited the most pronounced decreases in all three dendritic spine morphometric parameters (volume, surface area, length). This region also revealed significant shifts in the frequency distribution of spines, such that there were significantly fewer spines represented in the upper quartile, which suggests a selective failure of spine synapses to either mature or stabilize.

One important aspect of this study is that it underscores the advantage of employing automated, 3D spine analysis methods over more traditional approaches for the analysis of spine morphometric parameters. Previously, we and others have calculated spine morphometric features manually, by measuring spine head/neck length and diameters, and calculating spine head volume based on the assumption that spine heads are ellipsoidal (Knafo et al., 2005; Hao et al., 2006). Although this method may be employed in fluorescent dye-labeled or Golgi-impregnated neurons, measurements are limited to spines that are parallel to the plane of section, excluding spines that are oriented at oblique angles or obscured by residual glow immediately above or below the dendritic segment. Fur-
thermore, this approach limits the ability to characterize spines of different morphologic classes, such as long filopodial-type or short stubby spines. The Rayburst-based approach performs high-resolution analyses on deconvolved optical stacks of dendritic segments obtained from the confocal laser scanning microscope, allowing for each spine to be completely reconstructed in 3D, regardless of the plane of orientation, morphologic classification, or position relative to the dendritic segment. Finally, automated morphometric analyses can be performed very rapidly (i.e., 2–3 seconds) for a given dendritic segment (i.e., 80–100 spines), yielding spine volume, surface area, and length for each spine almost instantaneously, whereas manual approaches of measurement and calculation of spine head volume for the same segment may take several hours (Radley, Hof, Wearne, unpublished observations).

One limitation of characterizing spine morphometric features at the light and confocal microscopic level is that it is not possible to visualize synapses made onto dendritic shafts or cell bodies. Whereas electron microscopic (EM) serial section reconstruction of synapses remains the highest standard for the morphologic characterization for spines and synapses, reconstruction of sufficiently large data sets for robust statistical analysis at the EM level remains infeasible with the current technology.

Another interesting difference that we observed in the present study was that spine densities in proximal basal dendrites (<75 μm from the soma) were reduced following repeated stress, whereas, in a previous report, significant reductions in spine densities were limited to apical dendrites (Radley et al., 2006b). One possibility is that refinements in the experimental design may reveal this difference, e.g., sampling of the dendritic arbor at 75 μm vs. 50 μm radial distances from the soma in the present and previous report, respectively. Another possibility is that the manual counting methods used for determining spine densities by Radley et al. (2006b) were less consistent than that resulting from the Rayburst-based analysis. We also found that the Rayburst-based automated approach yielded spine density values that were an 87% match to the manual counting approach. Although it is tempting to state that the automated approach is 87% "accurate" compared with the manual counting approach, another explanation is that the manual counting is more subjective.

![Fig. 8. Probability distributions for spine volume and surface area for distal apical dendritic segments. The distributions were derived from high-resolution imaging of spines on apical dendritic segments at radial distances greater than 150 μm relative to the soma and analyzed by the Rayburst-based approach (control = 1,629 spines; 21 days stress = 1,438 spines). The dashed vertical lines in each histogram show the 25th and 75th percentiles of the entire spine population (i.e., 3,067 spines). These frequency distributions illustrate how repeated stress results in an overall downward shift in spine morphometric parameters.](image-url)
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Whereas an individual can reproduce spine density values from a given dendritic segment with virtually 100% accuracy, considerable variation (up to 35% in some instances) in spine density reporting has been noted among observers (Hof and Wearne, unpublished observations).

Previous work from our group has shown that repeated stress induces apical dendritic retraction and spine loss in layer II/III mPFC pyramidal neurons (Radley et al., 2004, 2005, 2006b; Laison et al., 2006). Other groups that have examined the effects of repeated stress (Cook and Wellman, 2004; Brown et al., 2005) and corticosteroids (Wellman, 2001; Cerqueira et al., 2007b) on dendritic morphology in layer II/III pyramidal neurons in mPFC have also observed similar alterations in dendritic length and branch number, particularly in distal apical dendrites. These effects are not likely to be limited to layer II/III pyramidal neurons but are apt to be generalized to the entire pyramidal neuron population in mPFC. Liu and Aghajanian (2006) recently showed that repeated stress decreased apical dendritic branching and spine density in layer V pyramidal neurons in mPFC, particularly in the more distal aspect of apical dendrites. Repeated stress and prolonged corticosteroid administration both result in a decrease in the volume of layer I in the dorsal mPFC (Cerqueira et al., 2005, 2007a), the region containing apical dendrites from both pyramidal cell populations in layers II/III and V. In support of the present findings, in nearly all of the aforementioned studies, the effects of repeated stress on structural plasticity were most prominent in distal apical dendrites. Collectively, these studies suggest that distal apical dendrites are most sensitive to the effects of repeated stress, resulting in a significant decrease in the population of excitatory synapses in distal apical dendrites residing in superficial layer I near the pial surface. It is important to note that chronic stress-induced plasticity in mPFC is not considered to be part of a pathological sequel leading to neurotoxicity, because repeated stress has not been found to result in a decrease in cell number or volume in pyramidal layers II/III or V in mPFC (Cerqueira et al., 2007a), and that these morphologic changes are reversible following a stress-free recovery period (Radley et al., 2005).

The reductions in spine volume and surface area, and a shift in spine size such that large spines became less prevalent following repeated stress, allow us to consider these changes in the context of spine stability and plasticity. Larger spines have been shown to be more stable and less motile (Trachtenberg et al., 2002; Kasai et al., 2003; Holtmaat et al., 2005) and, as a result, less plastic than smaller spines (Gruitzendler et al., 2002). Spine size is also correlated with synapse size (Harris and Stevens, 1989), and larger postsynaptic densities have more α-amino-3-hydroxy-5-methyl-4-isoxazole propionate (AMPA) receptors (Nusser et al., 1998; Kharazia and Weinberg, 1999; Takumi et al., 1999), consistent with the relative stability attributed to larger spines (Kasai et al., 2003). In contrast, N-methyl-D-aspartate (NMDA) receptors are more prevalent in small spines (Kasai et al., 2003) and are more sensitive to Ca2+-mediated signaling than large spines (Nimchinsky et al., 2004). Spine size is also related to the distribution of endocytic proteins, with such proteins closer to the synaptic edge in small spines (Racz et al., 2004), which might facilitate receptor insertion or promote changes in synaptic membrane structure in activated synapses. Although it is difficult to ascertain the functional consequences of the observed decrease in spine size and smaller proportion of large spines following repeated stress, when considered with spine loss, these morphologic changes are suggestive of a failure of spines to mature and stabilize.

In the context of plasticity, Kasai et al. (2003) have proposed that small spines represent “learning spines” that can either retract or become stabilized in the context of learning. In this view, large spines are “memory spines” that are stabilized and retain information over the long term. Based on this information alone, it would appear that our analysis reveals that stress increases the proportion of small spines, a subset of which could be poised for an increase in size and stability and consolidation in the context of learning (Matsuzaki et al., 2004). However, because we found that stress also reduces the number of large spines and overall spine number, it seems more plausible that repeated stress may limit spine liability that would, in turn, result in the stabilization and retention of more mature morphologies, such as might be important for memory formation (Matsuzaki et al., 2004). We and others have shown that repeated stress produces mPFC-dependent learning impairments (Liston et al., 2006; Cerqueira et al., 2007). This suggests that repeated stress may impair mPFC-dependent learning through an inability of spines to undergo plasticity-induced shifts from the smaller “learning” to larger “memory” phenotypes. Holmes and colleagues have shown that shorter intervals of stress are also capable of impairing the extinction of fear conditioning, a form of learning that is also dependent on the mPFC (Izquierdo et al., 2006). Although none of these studies has correlated learning with spine morphology in mPFC following stress, both acute and repeated stress-induced mPFC-dependent learning impairments corresponded to decreased apical dendritic arborization in pyramidal neurons in this region (Izquierdo et al., 2006; Liston et al., 2006). Indeed, one interesting possibility is that short-term and repeated stress affects mPFC-dependent learning through differential effects on spine number and morphology.

ACKNOWLEDGMENTS

We thank W.G.M. Janssen and K. Kelliher for expert technical assistance, Drs. J. Hao and W. Lou for help with the statistical analysis, and Dr. D.L. Dickstein for help with software development.

LITERATURE CITED


