Chronic Desipramine Prevents Acute Stress-Induced Reorganization of Medial Prefrontal Cortex Architecture by Blocking Glutamate Vesicle Accumulation and Excitatory Synapse Increase

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Abstract

Background: Although a clear negative influence of chronic exposure to stressful experiences has been repeatedly demonstrated, the outcome of acute stress on key brain regions has only just started to be elucidated. Although it has been proposed that acute stress may produce enhancement of brain plasticity and that antidepressants may prevent such changes, we still lack ultrastructural evidence that acute stress-induced changes in neurotransmitter physiology are coupled with structural synaptic modifications.

Methods: Rats were pretreated chronically (14 days) with desipramine (10 mg/kg) and then subjected to acute foot-shock stress. By means of serial section electron microscopy, the structural remodeling of medial prefrontal cortex glutamate synapses was assessed soon after acute stressor cessation and stress hormone levels were measured.

Results: Foot-shock stress induced a remarkable increase in the number of docked vesicles and small excitatory synapses, partially and strongly prevented by desipramine pretreatment, respectively. Acute stress-induced corticosterone elevation was not affected by drug treatment.

Conclusions: Since desipramine pretreatment prevented the stress-induced structural plasticity but not the hormone level increase, we hypothesize that the preventing action of desipramine is located on pathways downstream of this process and/or other pathways. Moreover, because enhancement of glutamate system remodeling may contribute to overexcitation dysfunctions, this aspect could represent a crucial component in the pathophysiology of stress-related disorders.

Keywords: stress, glutamate, synapse, desipramine
Introduction
Emotional arousing and stressful experiences represent constitutive parts of our daily life; tight and efficient regulation of the systems in charge of mediating the stress response is therefore required to ensure adaptation (Sterling and Eyer, 1988). However, when dysregulation of the stress response network or too much stress output occur, an overload of the system and its physical and psychological manifestations such as depressed mood and inflammation may be promoted (McEwen and Stellar, 1993). Among the key targets for stress hormones is the prefrontal cortex (PFC) (Meaney and Aitken, 1985; McEwen, 2007), a limbic and psychological manifestations such as depressed mood and psychological manifestations such as depressed mood and inflammation may be promoted (McEwen and Stellar, 1993). Among the key targets for stress hormones is the prefrontal cortex (PFC) (Meaney and Aitken, 1985; McEwen, 2007), a limbic region mediating highly evolved executive functions, including working memory and cognitive flexibility (McEwen and Morrison, 2013). Depending on the severity and duration of the stimulus, stressful episodes have been reported to exert different effects on the PFC (Popoli et al., 2012). Preclinical studies in animal models of stress have shown that chronic exposure to repeated stressors produces a number of physiological, cognitive, and structural deficits within medial PFC (mPFC); these changes cover suppression of glutamate receptor expression and function, reduced glial metabolism, and decreased expression of synaptic proteins as well as impaired decision-making and working memory (Dias-Ferreira et al., 2009; Banasr et al., 2010; Li et al., 2011; Yuen et al., 2012). Chronic stress has also been shown to induce profound structural remodeling of mPFC layer II-III pyramidal neurons, including dendritic shortening, spine loss, and neuronal atrophy (Cook and Wellman, 2004; Radley et al., 2004). In contrast, short-term activation of the stress response systems, as produced by acute stressors, has been reported to cause plasticity-enhancing effects, including potentiation of glutamate transmission in conjunction with enhanced working memory and increase of excitatory amino acid levels and of glutamate release in parallel with increased release probability in layer III pyramidal neurons of mPFC (Moghaddam, 1993; Yuen et al., 2009; Musazzi et al., 2010). Less, however, is known about the effects of acute stress on mPFC structural remodeling. It is well established that excitatory synapses play a major role in mediating synaptic transmission, plasticity, and memory; on the other hand, it has recently been demonstrated that stress and glucocorticoids, the major stress hormones, can actively and dynamically regulate these synapses (Kruegers et al., 2012; Timmermans et al., 2013). Thus, in the present study we aimed at investigating the immediate effects of acute stress on presynaptic and synaptic remodeling of layer II-III pyramidal neurons within the mPFC. It has recently been demonstrated that a single foot-shock (FS) stress exposure may produce hippocampal synapse loss 24 hours after stress (Hajszan et al., 2009). Moreover, we have recently shown that acute FS stress may strongly and rapidly increase the number of glutamate vesicles available for release within mPFC layer II-III (Treccani et al., 2014); however, whether the previously observed rearrangement of synaptic vesicles towards the presynaptic membrane may be coupled with changes in the number of mPFC synapses has never been explicitly demonstrated soon after acute stress. In this study, we also aimed to determine whether pretreatment with the traditional antidepressant desipramine (DMI) may affect any such change. It has recently been demonstrated that pretreatment with the norepinephrine reuptake inhibitor DMI dampens acute FS stress-induced increase of depolarization-evoked glutamate release from synaptic terminals of frontal and PFC cortex, and enhancement of the release probability in layer III pyramidal neurons within mPFC (Musazzi et al., 2010). However, it has never been explicitly demonstrated whether the preventive action of DMI on FS stress-induced enhancement of glutamate transmission may be coupled with any structural changes of mPFC after acute FS stress.

Materials and Methods
Animals and Drug Treatment
Male Sprague-Dawley rats (175–200 g, Charles River, Calco, Italy) were housed 2 per cage in a light-controlled room (under a 12-h-light/12-dark cycle; lights on at 7:00 AM) at room temperature (22°C) with food and water ad libitum. After 5 days of housing, one-half of the animals were subjected to chronic treatment (14 days) with DMI (10 mg/kg) delivered in drinking water. Drug solutions were changed every 2 days according to the animals’ weight and water intake. All experimental procedures involving animals were performed in accordance with the European Community Council Directive 86/609/EEC and approved by Italian legislation on animal experimentation (Decreto Ministeriale 116/1992).

Stress Paradigms and Corticosterone Levels
Twenty-four hours after the last drug or vehicle administration, animals were subjected to 2 different acute stressors. For the FS stress, rats were placed in a plastic chamber for 40 minutes, and 20 minutes of total actual shock (0.8 mA) was delivered with a random inter-shock length between 2 and 8 seconds (Vollmayr and Henn, 2001). Control animals were kept in the stress apparatus for the same interval of time (40 minutes) without delivering the shocks. For the acute restraint (RS) stress, rats were immobilized in air-accessible cylinders for 1 hour (Deolindo et al., 2013). Stress exposure was conducted between 8:00 AM and 12:00 AM in a room different from that used for housing.

Trunk blood corticosterone (CORT) levels were estimated using the IDS ELISA kit (Immunodiagnostic System, Herlev, Denmark). Inter- and intra-assay coefficient of variation was <10% (supplementary Material).

Confocal Microscopy of Synapses
Four male Sprague-Dawley rats (175–200 g) were treated for 14 days with either vehicle or DMI, as described above. Following a 24-hour wash-out, rats were deeply anesthetized with pentobarbital sodium (Unikem A/S, Copenhagen, Denmark; intraperitoneal, 75 mg/kg) and transcardially perfused with 100 mL heparinized (10 mg/mL saline) NaCl 0.9%, pH = 7.4) for 5 minutes followed by 2% fresh paraformaldehyde in 0.1 M phosphate buffered saline (pH = 7.4, 4°C). Brains were paraffin-embedded and processed for immunofluorescence staining against Piccolo and glutamate vesicular transport 1 protein (VGLUT1); for quantitative analysis, sections were examined in a 2-photon Zeiss Confocal LSM 510 META microscope (supplementary Material).

Total Number of Synapses and Synaptic Vesicles Using Electron Microscopy
Directly after the end of acute stress, animals were anesthetized and transcardially perfused with heparinized (10 mg/mL) saline (NaCl 0.9%, pH = 7.4) for 5 minutes followed by fixative containing 2.5% glutaraldehyde and 2% fresh paraformaldehyde in 0.1 M cacodylate buffer (pH 7.4). Series of 100-µm-thick sections were cut coronally on a vibratome 3000 (Vibratome, St. Louis, MO); based on a systematic, uniform random sampling
principle and a sampling fraction of one-sixth, 2 series were chosen. Given their exquisite sensitivity to chronic and acute stressors (Radley et al., 2004; Musazzi et al., 2010), mPFC layer II-III containing the apical dendrites of the vast majority of cortical pyramidal neurons were unambiguously identified and processed for electron microscopy (supplementary Material) (Nava et al., 2014).

Synaptic vesicle quantification was performed on a series of 45-nm-thick sections sampled from 2 regional areas per animal. Briefly, neurotransmitter-containing vesicles were classified into docked vesicles, when overlapping with the presynaptic membrane; all nondocked vesicles were defined as reserve pools. Synaptic vesicles were quantified in perforated (PSs) (Figure 1a) and nonperforated (NPSs) (Figure 1b) synapses. Quantification was performed as previously described (supplementary Material) (Nava et al., 2014).

For synapse number estimation, four epon blocks were sliced into series of 65-μm-thick serial sections with a MC MT-7 ultramicrotome (Tucson, AZ), and for each block two regions were sampled and imaged in a FEI Morgagni transmission electron microscope at final magnification of 18000x. Synapses were classified into axo-spinous, including PSs (Figure 1a) and NPSs (Figure 1b) synapses, based on the presence of a perforation on their postsynaptic density (black arrow in 1a), and axo-dendritic, namely axo-shafts (AXSs) (Figure 1c-d), synapsing directly onto the dendritic shaft (asterisk in Figure 1c-d) (Geinisman et al., 2001). For quantification, a modified physical dissector was adopted (Sterio, 1984) (supplementary material). All animals were assigned code numbers and decoded only after completion of the quantification.

mPFC Volume Estimation
mPFC total volume was obtained by applying the Cavalieri estimator in combination with the 2D-nucleator (6 half-lines; newCAST software) on thionin-stained sections (supplementary Material) (Gundersen and Jensen, 1987).

Preparation of Figures
Electron micrographs, as shown in the figures, were adjusted for contrast and brightness using Adobe Photoshop (San Jose, CA). Three-dimensional reconstructions were performed by applying of the RECONSTRUCT software (Fiala, 2005).

Statistical Analysis
All results were expressed as mean and relative coefficient of variation and calculated as the standard deviation of the estimates, divided by the mean. N represented the number of animals per group, and n represented the number of sections per animal analyzed for immunofluorescence study.

Coefficients of error (CEs) for synapse and vesicle number estimation were estimated as previously described (Nava et al., 2014) (supplementary Tables S1 and S2). Data were analyzed using 2-way (stress x treatment) and 1-way (stress subtypes) analysis of variance. When analysis of variance revealed significant group differences, the Bonferroni posthoc test was employed. For the immunofluorescence study, an unpaired t test was used. Statistical analysis and graphical representation was carried out using Systat 12.0 and GraphPad Prism 5.0. The degrees of significance were considered *P<.05, **P<.01, and ***P<.001.

Results
Glutamatergic Synaptic Terminal Distribution within mPFC
It is believed that excitatory synapses represent the majority of synapses within neocortex (Sanacora et al., 2012). However, to verify this, we assessed the relative distribution of mPFC excitatory terminals in rats subjected to chronic treatment with vehicle or DMI. A presynaptic active zone was visualized with antibody to Piccolo protein (Figure 2a) (Dondzillo et al., 2010); glutamate-specific synaptic terminals were identified using antibody to VGLUT1 (Figure 2b), regarded as expressed primarily in glutamatergic neurons and also in glial cells within cortical areas (Sanacora et al., 2012). In line with this, transporter VGLUT1 and Piccolo showed a high level of colocalization (Figure 2c) in both

Figure 1. Electron micrographs identifying subpopulations of synapses analyzed. Axo-spinous synapses were classified into perforated (a) and nonperforated (b) based on the presence of a discontinuity (black arrow in a) in the postsynaptic density. Axo-dendritic synapses (c-d) were identified by the presence of a mitochondrion on the postsynaptic compartment (asterisk in c and d). Scale bar = 500 nm.
vehicle- and DMI-treated groups, with an average of 92.3% and 92.1%. Unpaired t test revealed no significant difference in the degree of colocalization of the 2 proteins between the 2 groups (Figure 2d). Taken together, these results confirm that the large majority of mPFC synaptic contacts are excitatory.

Acute Stress-Induced Increase of Docked Vesicles Is Partially Prevented by Chronic Pretreatment with DMI

At the synaptic level, presynaptic terminals (blue region in Figure 3a) are enriched with neurotransmitter-containing vesicles of varying release competence (Dobrunz, 2002), so that vesicles docked to the active zone (cyan in Figure 3a) are thought as ready to be released (white spheres in Figure 3a) (Schikorski and Stevens, 2001); instead, nondocked vesicles (purple spheres in Figure 3a) comprise the reserve pool.

Acute FS stress has been reported to induce a strong and rapid increase of glutamate release from frontal and PFC synaptic terminals. This effect was paralleled by increased release probability and changes of excitatory postsynaptic current (EPSC) kinetics in layer III pyramidal neurons, two electrophysiological measurements of pre- and postsynaptic activity, respectively (Musazzi et al., 2010). Moreover, we have recently shown that an FS stress-induced increase of glutamate release occurs in parallel with an increased number of glutamate vesicles available for release within mPFC (Treccani et al., 2014). Chronic treatment with DMI dampened the stress-induced increase of glutamate release and release probability, but not the kinetics of EPSC, suggesting a mechanism of action limited at the presynaptic level (Musazzi et al., 2010). Thus, to evaluate the effect of chronic DMI on stress-induced glutamate vesicle redistribution within the presynaptic compartment, we quantified the number of docked and reserve pool vesicles in stressed animals pretreated with chronic DMI. At the end of acute FS stress exposure, the average number of docked vesicles was increased almost 2-fold in PSs (+90.3%; N = 6; P < .001), and this increase was partially prevented by chronic DMI (−6.9%; N = 6; P < .05) (Figure 3b, left panel).

No effect of either stress or DMI was observed in NPSs (N = 6; P, ns) (Figure 3b, right panel). In contrast, the average number of reserve pool vesicles significantly decreased after stress in NPSs (−16.0%; N = 6; P < .05) (Figure 3c, right panel) with no effect on PSs (N = 6; P, ns) (Figure 3c, left panel).

These results confirm that acute FS stress induces a dramatic and selective increase in the number of docked vesicles available for release in PSs, as previously reported (Treccani et al.,...
Moreover, the results show that, in PSs, the stress-induced major redistribution of synaptic vesicles towards the presynaptic membrane is attenuated by DMI pretreatment. Overall, these results suggest that, by reducing the stress-induced increase of docked vesicles, chronic DMI acts at the presynaptic level by preventing the stress-induced potentiation of a morphological correlate of excitatory synaptic strength.

Acute Stress Induces a Rapid Increase of mPFC Excitatory Synapse Number, Which Is Prevented by Chronic DMI

An altered number of docked vesicles has been observed during long-term potentiation in conjunction with fast synaptic remodeling, as shown by an increased number of synapses in hippocampal Cornu Ammonis 1 as soon as 30 minutes after the stimulus induction (Bourne and Harris, 2011; Bourne et al., 2013). Moreover, high vulnerability of synapses to acute stressful events has previously been reported in studies showing hippocampal synaptic marker increase and synapse loss, respectively, shortly and 24 hours after stress termination (Hajszan et al., 2009; Sebastian et al., 2013).

Hence, to investigate the effects of acute FS stress and DMI treatment on mPFC excitatory terminal remodeling, we quantitated the number of mPFC asymmetric synapses regarded as excitatory by means of physical disector and electron microscopy. Interestingly, the total number of excitatory synapses was found to be rapidly and dramatically increased by 42.6% (N = 6; P < .001) after 40 minutes of FS stress compared with the sham group, whereas chronic administration of DMI partially prevented the stress-induced increase (+13.3% compared with control group; N = 6; P < .01) (Figure 4a).

To further verify whether the observed synaptogenic effect was specific for FS stress or common to other acute stress models, we included rats previously subjected to 1 hour of RS stress in our analysis. When comparing the total number of asymmetric synapses across different stress subtypes, acute RS stress also increased asymmetric synapses by 22% compared with the sham group (N = 6; P < .05) (Figure 4a).

We have recently shown that the majority of mPFC asymmetric spine synapses, namely PSs and NPSs, are endowed with heterogeneous structural features that may reflect differences in synaptic efficacy. For example, NPSs have shown smaller bouton volume and lower numbers of neurotransmitter-containing vesicles compared with PSs (Nava et al., 2014). Moreover, asymmetric synapses may undergo differential activity-dependent remodeling, depending on the subtype. For example, long-term potentiation, a synaptic model of memory, has been shown to selectively increase the number of PSs only (Geinisman et al., 1991, 1993).

Therefore, to clarify whether the synaptogenic effect of acute stress is selective for a specific synaptic population, we further classified asymmetric synapses between axospinous, including NPSs and PSs, and axo-dendritic contacts, namely AXSs (Figure 4c), in our serial section electron microscopy analysis (Figure 4b). Small NPSs were increased after FS stress compared with the sham group (+39.6%; N = 6; P < .001) regardless of treatment with DMI. Comparison across stressor subtypes showed...
Figure 4. Total number of medial prefrontal cortex (mPFC) excitatory synapses is rapidly and selectively increased after 40 minutes of acute foot shock (FS) stress and after 1 hour of restraint (RS) stress, whereas treatment with chronic desipramine (DMI) partially prevents FS stress-induced synaptogenesis. (a) Total number of mPFC synapses was found to be rapidly increased at the end of 40 minutes of FS stress exposure ($P < .001$); however, FS stress-induced synaptogenesis was partially prevented by treatment with chronic DMI ($P < .01$). RS stress also increased the total number of synapses compared with sham vehicle-treated (Veh) group ($P < .05$). (b) For synapse quantification, a modified physical disector was applied (Tang et al., 2001); a grid was superimposed over 15 serial sections; section 1 was look-up section, sections from 2 to 10 were used as the reference section of the disectors, whereas the last 5 sections were used as look-up sections to ensure that all counted postsynaptic densities were included in the section series. Synaptic profiles touching the inclusion (green) lines were included in the analysis, whereas those touching the exclusion (red) lines were excluded (scale bar = 1 μm). Synapses were classified between axo-spinous, synapsing onto spines, including nonperforated (blue dots) and perforated (orange dots). Contacts synapsing directly onto the dendritic shaft were classified as axo-shaft (AXSs) (yellow dots). (c) Application of disector technique allowed unbiased estimates of the number of neocortical synapses, reconstructed here as spheres, within neuropil volume. Blue spheres represent nonperforated synapses (NPSs), orange spheres perforated synapses (PSs), and yellow spheres AXSs. Representative estimates of the number of synapses are reported as the number of 3D spheres for each experimental group: Sham+Veh (n = 21); FS+Veh (n = 37); Sham+DMI (n = 18); FS+DMI (n = 26); RS+Veh (n = 35). (d) Acute FS stress led to a strong significant increase in small NPSs ($P < .001$) regardless of DMI treatment. The number of NPSs was also increased in RS animals compared with the sham group ($P < .01$). (e) The total number of great PSs was shown to be stable among experimental groups. (f) AXSs were found strongly increased after acute FS stress ($P < .001$), whereas treatment with chronic DMI prevented the stress-induced increase ($P < .05$). (g) mPFC volume estimated by the Cavalieri estimator did not show any significant difference among experimental groups. (h) Corticosterone concentration was found significantly increased after both FS and RS stress, regardless of previous DMI treatment ($P < .001$). Compared with RS-stressed animals, rats subjected to FS stress showed higher levels of CORT ($P < .05$).
that RS stress also significantly increased the number of NPSs (+41.2%; N = 6; P < .01) (Figure 4d). Interestingly, big PSs were stable across groups (N = 6; P, ns) (Figure 4e). Moreover, only acute FS stress, but not RS stress, induced a strong increase in AXSs (+51.2%; N = 6; P < .001), and this effect was partially prevented by treatment with chronic DMI (+6.01%; N = 6; P < .05) (Figure 4f). When estimating overall mPFC volume, no significant effect of stress-induced new synapse formation was observed (Figure 4g).

To determine whether the increase of mPFC asymmetric synapses in acutely stressed animals is correlated with the elevated levels of adrenal corticosteroid hormones, we performed enzyme-linked immunosorbent assay to measure CORT levels. As shown in Figure 4h, compared with unstressed animals, rats exposed to acute stressors had significantly higher blood concentration of CORT (FS stress: +532%, P < .001; RS stress: +390%, P < .001) regardless of previous treatment with DMI (Musazzi et al., 2010). When comparing among stressor subtypes, CORT levels were found to be significantly higher after FS compared with RS stress animals (+29.1%, P < .05) (Figure 4h).

Thus, an acute severe stressor rapidly induces strong sprouting of asymmetric synaptic contacts, mainly by increasing the number of small and less efficacious NPSs; on the other hand, the number of glutamate synapses is only modestly increased after exposure to a less severe stressor. Moreover, DMI pretreatment, ineffective in the absence of stress, partially prevents the stress-induced increase. These results suggest that an acute stressor can rapidly boost the mPFC glutamate system by enhancing the number of glutamate release sites and apposed postsynaptic specializations. On the other hand, the preventive effect of DMI on stress-induced synapse sprouting, but not on CORT increase, suggests that the site of action of DMI in the previously observed modulation of stress-induced glutamate release (Musazzi et al., 2010) is downstream of CORT release, as previously found (Conti et al., 2004; Musazzi et al., 2010), and may directly involve the blockade of new pre- and postsynaptic element formation.

**Acute FS Stress Induces a Generalized Increase of Glutamate Vesicles Available for Release within mPFC, Which Is Prevented by DMI Pretreatment**

The observed effects of FS stress and DMI pretreatment on synaptogenesis and presynaptic ultrastructural plasticity showed a high degree of selectivity towards different synaptic subtypes. Thus, acute stress increased the number of docked vesicles in PSs only and increased the number of NPSs and AXSs. Therefore, we examined how the stress-induced increase in number of NPSs could affect the total number of docked vesicles and, possibly, excitatory synaptic strength within mPFC by combining the data obtained on docked vesicles and synapse number estimation.

The total number of docked vesicles, calculated by multiplying the total number of synapses within mPFC with the average number of vesicles, was significantly increased after FS stress in both PSs (+78.1%; N = 6; P < .01) and NPSs (+145.4%; N = 6; P < .001) compared with the control group, with a more remarkable effect on the latter subtype. Treatment with DMI, ineffective in the absence of stress, fully and partially normalized the stress-induced increase in PSs (−11.9%; N = 6; P < .001) and NPSs (+31.4%; N = 6; P < .01), respectively (Figure 5a, left and right panels).

![Figure 5.](image-url) Acute foot shock (FS) stress dramatically increases morphological correlates of synaptic strength by increasing the number of vesicles available for release in medial prefrontal cortex (mPFC) perforated (PSs) and nonperforated (NPSs) synaptic population. (a) The total number of docked vesicles in PSs within mPFC was significantly increased by almost 2-fold after acute FS stress (P < .01); treatment with chronic desipramine (DMI) completely prevented the stress-induced increase in the number of docked vesicles within mPFC PSs (left panel) (P < .001). (b) Similarly, in mPFC NPSs, FS stress significantly increased the total number of docked vesicles (P < .001), whereas treatment with DMI partially prevented the stress-induced effects (right panel) (P < .01). In FSs, the total number of mPFC reserve pool vesicles was not affected by stress or DMI treatment in either PSs or NPSs. (c) Three-dimensional representation of synaptic vesicles (white spheres) docked to the active zone (cyan) for each experimental group. Cube side = 50 nm.
panels). No effect of either stress or treatment was observed on reserve-pool vesicles in PSs and NPSs (Figure 5b, left and right panels). These results suggest that, following the stress-induced increase of synapses, acute FS stress produces a strong and generalized increase in the number of vesicles available for release in both synapse subtypes, which is strongly normalized by DMI pretreatment (Figure 5c).

Discussion

The results from the present study suggest that a single brief exposure to stress potentiates excitatory neurotransmission by inducing presynaptic structural plasticity, as shown by an increased number of docked vesicles and large sprouting of excitatory synapses. Moreover, these changes were partially blocked by DMI pretreatment. Our ultrastructural findings were localized in layers II-III of mPFC, which contains predominantly somata of pyramidal neurons, and is known to be particularly sensitive to chronic stress insults (McEwen and Morrison, 2013). With regards to the effects of acute stress on mPFC structural plasticity, the first result of the present study showed that acute FS stress strongly increases the number of docked vesicles in PSs only, confirming our previous observation on the selective effect of acute FS stress (Treccani et al., 2014). These results suggest that following acute FS stress, more vesicles from the reserve pool become competent to fuse with the presynaptic membrane and empty their content by being released from the cytoskeletal matrix (Schikorski and Stevens, 2001; Sudhof, 2004). In line with this hypothesis, acute FS stress has shown to increase presynaptic release probability (Musazzi et al., 2010). We also observed for the first time that a single severe stressor can rapidly induce a strong increase in the number of asymmetric synapses, whereas synapse increase is only modest after a less severe stressor. This finding deserves particular attention considering that the general synapse increase was differentially provided for the 2 stressors by the counting of AXSs and NPSs. Interestingly, AXSs were found increased only after FS (+51.2%) but not RS stress. Most likely due to their relatively higher strength through their strategic location directly on dendritic shafts rather than on spines (Figure 1c-d), we postulate that potentiation of AXSs only after FS stress may be linked to significantly higher levels of CORT found after FS compared with RS stress, as well as to a different degree of complexity among the 2 stressors (Maras et al., 2014). Small NPSs were increased after both FS (+39.6%) and RS stress (+41.2%), with greater stability being observed among PSs. Hippocampal PSs mediate greater synaptic efficacy compared with small NPSs by expressing a larger number of α-amino-3-hydroxy-5-methyl-4-isoxazole propionate receptors (AMPARs) (Ganeshina et al., 2004b). Because NPSs express fewer AMPARs and are regarded as less efficacious (Ganeshina et al., 2004a, 2004b), we hypothesize that the observed stress-induced increase of NPSs is consistent with an increase in the so-called silent synapses, containing mainly N-methyl-D-aspartate receptors (NMDARs) and few or no AMPARs (Isaac et al., 1995). Despite being functionally silent at resting potentials, these synapses represent a new substrate for the formation, by insertion of AMPARs, of new functional synapses more likely to be potentiated (Suvrathan et al., 2014). In line with this hypothesis, a number of recent studies strongly suggest postsynaptic changes in glutamate receptors, as shown by an increased level of surface NMDARs and AMPARs in PFC slices examined 1 to 4 hours poststress and altered EPSC kinetics likely occurring at the postsynaptic level (Yuen et al., 2009; Musazzi et al., 2010). According to a well-accepted model of synaptic remodeling following long-term potentiation, an increased number of synapses is provided by splitting small NPSs into large PSs (Geinisman, 2000). However, in the present study, the stress-induced increase in small NPSs and AXSs was not accompanied by a corresponding decrease in PSs. Therefore, we postulate that additional AXSs and NPSs observed after acute stress are formed mostly ex novo rather than being converted from another synapse subtype. Although a number of in vitro studies have shown that activity-dependent synaptogenesis can occur within 2 hours from the application of the stimulus (Bourne et al., 2013), only few studies investigated the formation of new synaptic contacts ex vivo, reporting 24 hours as the shortest interval required for the formation of large efficacious boutons (Medvedev et al., 2012). Interestingly, our findings provide the first ex vivo evidence that activity-dependent synaptogenesis of small and less efficacious NPSs can occur locally ex novo as early as 40 minutes after the application of a severe stressful stimulus. This shows remarkable dynamics of synapse structure observed in a short time-frame following application of the stressor. Given that the effects of acute stress on synapse number were observed immediately after the end of the stress protocol, it is conceivable that these changes may be accounted for by changes in local protein translation in dendrites. One of the crucial mediators of activity-dependent synaptic plasticity is the neurotrophin brain derived neurotrophic factor involved in the translation of a subset of dendritic mRNAs at synapses, including cytoskeletal proteins involved in synaptic rearrangement (Leal et al., 2014). This new knowledge can be of great importance in clarifying the dynamics leading to the long-term behavioral and structural impairments that exposure to CORT may produce (Gourley et al., 2013).

A number of previous studies have shown that tricyclic antidepressants, including DMI, can reduce glutamate release and transmission under both basal conditions in hippocampus (Bonanno et al., 2005; Tokarski et al., 2008) and in the presence of an acute stressful challenge in prefrontal and frontal cortex (Musazzi et al., 2010). In the last study, electrophysiological recordings showed that DMI exerts its effect mainly by acting at the presynaptic level, with little or no effect at the postsynaptic side (Musazzi et al., 2010). In the present study, we examined whether DMI may also dampen the enhanced structural plasticity induced by acute FS stress; this was done by investigating the size of synaptic vesicle pools, as presynaptic markers, and the number of glutamate synapses as an overall measure of synaptic changes. Here we showed that DMI treatment prevents the effects of acute FS stress on structural plasticity. Although subchronic treatment (5–7 days) with DMI has been shown to reverse the detrimental effects of inescapable FS on hippocampal spine synapse number (Hajszan et al., 2009), the present results suggest that DMI may also have a preventive action on the effects of stress on structural plasticity, as shown by normalized vesicle docking and small synapse sprouting.

On the other hand, the dampening action of DMI on the stress-induced structural potentiation was not accompanied by a reduction of CORT levels. Thus, we suggest that the potential mechanism underlying the blockade of stress effects by chronic DMI must be located on pathways downstream of CORT release or on alternative pathways. Several mechanisms may account for the DMI-induced prevention of the stress-induced elevation of structural plasticity. As previously discussed (Musazzi et al., 2010), by increasing noradrenaline availability, DMI can induce downregulation of chronically activated autoreceptors inhibiting noradrenaline release and leading in turn to disinhibition of
noradrenaline exocytosis (Raiteri et al., 1986; Starke et al., 1989). In parallel, activation of α1-adrenergic heteroreceptors located on glutamatergic neurons does not seem to lead to their down-regulation, thereby inhibiting glutamate release (Raiteri et al., 1983; Kamisaki et al., 1992; Pittaluga et al., 2007). In particular, a major role in these processes may be played by synapsin I, which is known to control the trafficking of synaptic vesicles between the reserve and the readily releasable pool, depending on phosphorylation state (Greengard et al., 1993; Ceccaldi et al., 1995). Because we have recently shown that acute FS stress increases levels of Phospho-Ser9 synapsin I (et al., 1995). Because we have recently shown that acute FS stress increases levels of Phospho-Ser9 synapsin I in synaptic membranes (Treccani et al., 2014), it will be interesting to analyze the effects of DMI pretreatment on the acute stress-induced increase of Phospho-Ser9 synapsin I. Overall, this effect of DMI on stress-induced vesicle accumulation may account for the previously described blockade of acute stress-induced enhancement of glutamate release and transmission by antidepressants (Musazzi et al., 2010).

Moreover, the effect of DMI on stress-induced structural plasticity was found double-sided. Besides limiting the accumulation of presynaptic–located docked vesicles, DMI was found to act at the postsynaptic level by limiting the number of highly electron-dense PSDs, regarded as containing postsynaptic receptors (Gray, 1959). Antidepressants have long been known to regulate glutamate receptors by limiting NMDAR function through decreased protein expression level of its subunits (Pittaluga et al., 2007) and, conversely, potentiating AMPAR-mediated transmission (Barbon et al., 2006). Therefore, it will be interesting to assess the effect of DMI pretreatment on the postsynaptic surface level of glutamate receptors.

By limiting the number of glutamate vesicles available for release and the number of excitatory synapses as potential glutamate release sites, DMI may help prevent FS stress-induced boosting of the mPFC glutamate system and subsequent hyper-glutamate–induced noxious effects (Sanacora et al., 2008; Popoli et al., 2012).

In conclusion, by increasing vesicle docking and forming new small synapses, acute stress promotes an immediate form of ultrastructural plasticity of mPFC pyramidal neurons. This heterogeneous population of neurons tends to segregate inputs with extracortical afferents (from the mediadorsal nucleus of the thalamus and hippocampal CA3) clustering on distal dendrites (layer I) (Swanson and Cowan, 1977; Groenewegen, 1988) and synapses of local cortical circuits clustering on proximal apical and basal dendrites (Scheibel and Scheibel, 1970). The acute FS stress-induced enhancement of structural plasticity as observed here was circumscribed to mPFC layer II-III containing primarily the proximal portions of the apical and basilar arbor: in turn, this may suggest a shift in emphasis from subcortical to intracortical information, with important implications for the functioning of mPFC and cognitive behaviors mediated by it.

Although further research is needed to assess how long the stress-induced synaptogenesis and presynaptic potentiation can be sustained and which pathways mediate the preventive effects of DMI, these results support our initial hypothesis that acute stress-induced presynaptic plasticity occurs in conjunction with increased synaptic remodeling, suggesting in turn an overall FS stress-induced enhancement of mPFC structural plasticity. In addition to the previously observed preventive effect on stress-induced increase of glutamate release, pretreatment with classic antidepressant DMI can strongly counteract FS stress-induced mPFC structural plasticity. This preventing action of DMI may be directly linked with its therapeutic effect in the treatment of mood and anxiety disorders.

While our observations on stress-induced structural plasticity confirms the high vulnerability of mPFC to even a single stress exposure, our data on the preventive effects of DMI suggest synapse remodeling as a neuronalanatomical marker predicting the effectiveness of antidepressant compounds on preventing long-term noxious consequences.

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Statement of Interest

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