Toward a limbic cortical inhibitory network: implications for hypothalamic-pituitary-adrenal responses following chronic stress

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A network of interconnected cell groups in the limbic forebrain regulates hypothalamic-pituitary-adrenal (HPA) axis activation during emotionally stressful experiences, and disruption of these systems is broadly implicated in the onset of psychiatric illnesses. A significant challenge has been to unravel the circuitry and mechanisms providing for regulation of HPA output, as these limbic forebrain regions do not provide any direct innervation of HPA effector cell groups in the paraventricular hypothalamus (PVH). Recent evidence will be highlighted that endorses a discrete region within the bed nuclei of the stria terminalis serving as a neural hub for integrating and relaying HPA-inhibitory influences to the PVH during emotional stress, whereas the prevailing view has involved a more complex organization of multiple cell groups arranged in parallel between the forebrain and PVH. A hypothesis will be advanced that accounts for the capacity of this network to constrain the magnitude and/or duration of HPA axis output in response to emotionally stressful experiences, and for how chronic stress-induced synaptic reorganization in key cell groups may lead to an attrition of these influences, resulting in HPA axis hyperactivity.

Keywords: bed nuclei of the stria terminalis, prefrontal cortex, hippocampus, ventral subiculum, HPA axis, paraventricular nucleus of the hypothalamus, plasticity, dendritic spine

INTRODUCTION

Stress may be broadly defined as the constellation of physiological and behavioral responses to any challenge that overwhemls, or is perceived to overwhelm, selective homeostatic systems of the individual (Selye, 1980; Day, 2005). A hallmark feature of stress entails activation of the hypothalamic-pituitary-adrenal (HPA) axis. This neuroendocrine cascade is initiated when visceromotor neurons in the paraventricular nucleus of the hypothalamus (PVH) stimulate the release of pituitary adrenocorticotropic hormone (ACTH) into the bloodstream, which, in turn, activates glucocorticoids (GC; cortisol in humans, corticosterone in rodents) secretion from the adrenal gland (Antoni, 1986). GCs are the end-products of HPA axis activation, and facilitate catabolic processes throughout the body during stress by increasing energy metabolism and utilization. GCs also have activating effects on cardiovascular output, and inhibit non-essential processes, such as immune and reproductive functions. Finally, HPA axis activation during stress alters cognitive and emotional processes relevant for behavioral adaptation (e.g., Shors et al., 1992; McIntyre et al., 2003).

Despite the critical role that stress plays for adaptive coping and survival of the individual, it is widely implicated in the onset of psychiatric disease, most notably depression and post-traumatic stress disorder (Kessler, 1997; Yehuda, 2002). Initial studies revealed that patients hospitalized for major depressive illness commonly manifested hypercortisolemia and HPA axis insensitivity to GC receptor agonist treatment (i.e., dexamethasone suppression test; Carroll et al., 1976). A wealth of research implicates elevated GCs in compromised brain function, disruptions in the neural circuits imparting negative feedback control over the HPA axis, and further endangerment of brain regions targeted by GCs (for reviews, see Sapolsky et al., 1986; Conrad, 2008). Since the neural substrates providing restraining influences over the stress axis are also regions that play important roles in cognition and emotion, elevated GC levels and HPA axis dysregulation may be key steps in producing the disordered thought and affect that characterize stress-related mental illnesses.

Animal models of repeated stress (e.g., chronic variable stress, chronic intermittent stress, chronic social defeat stress) have proven useful for modeling HPA axis hyperactivity and depression-like behaviors, and would appear to provide the appropriate setting for teasing apart the role of the HPA axis in the pathogenesis of depression. However, progress has been hampered by the fact that the neural circuitry and mechanisms accounting for limbic forebrain control over the HPA axis have proven difficult to unravel. While a number of these candidate regions have been implicated in HPA axis inhibition during emotional stress (Herman et al., 2003; Radley and Sawchenko, 2011), none of these cell groups provide any appreciable direct innervation of the PVH. Combined pathway tracing and immediate-early gene mapping studies have helped to identify a number of candidate cell groups that could serve as dysynaptic relays to interface between forebrain regulators and the PVH. The picture that emerges is one involving a complex network of higher-order
structures interconnected in a parallel and multisynaptic manner with the PVH (Cullinan et al., 1993; Roland and Sawchenko, 1993; van de Kar and Blair, 1999; Herman et al., 2003).

Here we highlight recent advances in our research suggesting an entirely different organization for limbic forebrain control over the stress axis: one involving convergence onto a circumscribed cluster of GABAergic neurons within the anterior bed nuclei of the stria terminalis (aBST), that, in turn, directly inhibits the PVH and HPA activation. This model has several implications for neural circuits and mechanisms underlying HPA axis control and GC-dependent negative feedback. An unforeseen but not incidental feature is that this model helps to clarify the sequelae of chronic stress-induced HPA axis hyperactivity, whereby structural reorganization within limbic forebrain cell groups (i.e., synapse loss/gain) throughout the network leads to an attrition of HPA axis control.

**EMOTIONAL STRESS CIRCUITRY: A SEARCH FOR THE MISSING LINK**

Over the years, attempts to organize stressors into a taxonomical framework have resulted in two major groupings, physiological (a.k.a., systemic), and emotional (a.k.a., neurogenic, psychogenic) (Fortier, 1951; Allen et al., 1971). More recent immediate-early gene mapping as a generic index of cellular activation in stress-related circuits has helped to provide a considerable degree of face validity for these distinctions (Cullinan et al., 1995; Li and Sawchenko, 1998; Dayas et al., 2001). Physiological stressors are generally considered to involve more targeted challenges that overwhelm selective homeostatic systems, such as hemorrhage, hypoxia, or immunogenic stimuli. Emotional stressors require interpretation by exteroceptive sensory modalities and integration with distinct cognitive (comparison with past experience) and affective information processing systems in the brain (Herman and Cullinan, 1997; Sawchenko et al., 2000; Dayas et al., 2001). Commonly employed animal models of emotional stress are restraint, immobilization, and footshock. Whereas each class of stressor enlists brainstem and hypothalamic effectors for activation of the sympathoadrenal and HPA axis output, emotional stressors manifest widespread activation in the limbic forebrain, and correspond to a broad array of behavioral changes (e.g., vigilance, fear, anxiety) that help to facilitate adaptive coping as required by the specific environmental demand (Cullinan et al., 1995; Campeau et al., 1997; Li and Sawchenko, 1998; Dayas et al., 2001).

Functional and lesion studies implicate a network of limbic forebrain cell groups in the inhibitory control of HPA activation during emotional stress (Cullinan et al., 1995; Herman and Cullinan, 1997; Akana et al., 2001; Jaferi and Bhatnagar, 2006). Noteworthy examples of regions implicated in HPA axis inhibition are the septum (Feldman and Conforti, 1980b), posterior paraventricular nucleus of the thalamus (PVTP; Jaferi and Bhatnagar, 2006), ventral subiculum (vSUB, the region issuing extrinsic projections of hippocampal formation involved in stress regulation; Herman et al., 1995), and mPFC (Diorio et al., 1993). These cell groups are conspicuously lacking in any direct innervation of HPA effector neurons within the PVH, instead issuing projections throughout numerous basal forebrain and hypothalamic structures (Sesack et al., 1989; Cullinan et al., 1993; Herman et al., 2003). Many of these regions (notably, mPFC and hippocampal outputs) give rise to predominantly excitatory projections, utilizing the neurotransmitter glutamate (Malthes-Sorensen et al., 1980; Walaas and Fonnum, 1980; Ottersen et al., 1995), implicating a hitherto unknown, GABAergic relay. Previous work has identified candidate GABAergic cell groups (i.e., preoptic area, aBST, posterior BST, dorsomedial hypothalamic nucleus, PVH-surround regions) between vSUB and PVH (Cullinan et al., 1993), laying a foundation for understanding how controls over the axis may be organized. Nonetheless, whether influences from vSUB, and other HPA-inhibitory cell groups, are mediated via several disynaptic relays arranged in parallel to each other, and which relays are capable of integrating inhibitory signals from the limbic forebrain during emotional stress, has remained elusive.

Our starting point into this problem was to first address the nature of mPFC involvement in acute emotional stress-induced HPA activation, and we have shown that distinct subregions of mPFC differentially modulate the stress axis (Radley et al., 2006a). These studies were inspired from the idea that a variety of other functions subserved by mPFC are differentiaeted in a dorsal-to-ventral manner (Morgan and LeDoux, 1995; Heidbreder and Groenewegen, 2003). Indeed, previous reports in the stress literature tended to treat the mPFC as a homogeneous structure, and discrepancies remained concerning the nature of mPFC’s influence (excitatory or inhibitory) on HPA output (Sullivan and Gratton, 1999; Akana et al., 2001; Figueiredo et al., 2003; Spencer et al., 2005). Through a series of experiments employing discrete excitotoxin lesions in cortical subfields of mPFC, we found that lesions of dorsal mPFC (encompassing prelimbic cortex, PL, and portions of dorsal anterior cingulate cortex, ACd) enhanced, whereas ventral mPFC (infralimbic cortex, IL) lesions inhibited HPA activation in response to acute restraint stress (Radley et al., 2006a). Furthermore, dorsal mPFC lesions resulted in a prolonged elevation of plasma corticosterone after the cessation of restraint, which is consistent with its role as a target site for GC negative feedback under normal conditions (Diorio et al., 1993) (Figure 1).

Follow-up work has shown that a discrete cluster of GABAergic neurons in aBST forms the missing link in a circuit conveying HPA-inhibitory influences of PL during emotional stress (Radley et al., 2009). First, functional neuroanatomical experiments assayed for sources of GABAergic input to PVH whose sensitivity (i.e., as measured with Fos activation) to an acute stress or (restraint) was diminished by dorsal mPFC lesions. Of the stress-sensitive, GABAergic, PVH-projecting regions analyzed, a circumscribed region in the aBST (corresponding to the dorsomedial and fusiform subdivisions of Dong et al., 2001) was exclusive in showing a decrement in Fos activation following PL lesions (Radley et al., 2009). By contrast, IL lesions were noted to attenuate Fos activation in PVH-projecting neurons in the same region, albeit in a subpopulation of non-GABAergic neurons (Figure 2).

In a second series of experiments, functional ablation of GABAergic neurons in aBST recapitulated the effects of PL lesions on acute stress-induced HPA activation (Radley et al., 2006b, 2009). These studies were performed by focally administering...
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**FIGURE 1** | Top: Darkfield photomicrographs showing corticotropin-releasing factor (CRF) mRNA expression in the paraventricular nucleus of the hypothalamus (PVH) as a function of treatment condition. Restraint stress results in a marked increase in CRF mRNA expression in the PVH dorsal medial parvicellular subdivision of intact animals, which is enhanced in prefrontal (PL)-lesioned rats. No such enhancement of stress-induced CRF mRNA expression was seen in infralimbic (IL)-lesioned animals.

Bottom: Stress exposure also significantly increases plasma corticosterone (CORT) levels in sham-lesioned animals. PL lesions result in a prolonged increase in stress-induced plasma CORT, while IL-lesioned animals show a more rapid recovery to pre-stress levels. ∗Differs significantly from basal (0 min) values from within each group, p < 0.05; †Differs significantly from sham-lesioned animals for a given time point, p < 0.01. Data are from Radley et al. (2006a).

an immunotoxin in aBST that preferentially ablates GABAergic, while sparing non-GABAergic, neurons (Radley et al., 2009). Ablation of GABAergic cell groups in aBST enhanced activation of PVH and hormonal indices of HPA axis output in response to acute restraint. Previous reports that indiscriminate lesions to aBST attenuate stress-induced HPA output (Choi et al., 2007), whereas stimulation of aBST may either facilitate or inhibit HPA activity (Dunn, 1987), are consistent with the idea that distinct HPA-regulatory influences arise from neurochemically heterogeneous subpopulations. Thus, opposing influences of the dorsal and ventral mPFC may commingle within the same region of aBST onto separate populations of PVH-projecting GABAergic and non-GABAergic neurons, respectively, to modulate emotional stress-induced HPA output (Figure 3).

Subsequent anatomical pathway tracing studies have that PL is the cortical subfield that provides the source of HPA-inhibitory influences that emanate from the mPFC (Radley et al., 2006b, 2008b, 2009). Whereas the subcortical projections of dorsal and ventral mPFC are considered to be highly divergent (e.g., Vertes, 2004), their projections to aBST distribute in a topographically
also interceded for by aBST, similar in nature to PL (Radley and Sawchenko, 2011). First, GABAergic PVH-afferent cell groups in aBST showed a diminished functional activation in animals bearing excitotoxin lesions of vSUB. In these experiments, vSUB lesions were also noted to increase multiple indices of acute restraint-induced HPA activation, as previously reported (Herman et al., 1995). Although HF is not typically regarded as one of the more stress-responsive regions in the limbic forebrain (Li and Sawchenko, 1998), vSUB, particularly its aBST-projecting neurons, does in fact display a moderate degree of engagement following acute restraint stress (Radley and Sawchenko, 2011). Finally, animals bearing dual tracer deposits show that vSUB projections overlap extensively, and make appositions with, stress-sensitive PVH-projecting cell groups in aBST. Collectively, these studies highlight a neural circuitry from vSUB → aBST (GABA) → PVH, with each node in the pathway showing functional activation in response to acute restraint stress, and lesions of vSUB resulting in corresponding alterations in output (i.e., decreased aBST, increased PVH/HPA activation).

A key feature of aBST, in addition to its role as a site of convergence, is that it appears to integrate limbic cortical influences (Radley and Sawchenko, 2011). For instance, animals bearing excitotoxin lesions of both PL and vSUB were found to exhibit more exaggerated central indices of stress-induced HPA responses as compared to lesions of either alone. Furthermore, ablation of GABAergic cell groups in aBST produced a greater enhancement of hormonal indices of HPA activation in response to acute restraint, as compared to animals with vSUB lesions alone. Given that dual lesions of PL and vSUB, or separately, by disruption of their interceding inhibitory relay, result in a greater overall effect on stress-induced HPA output than lesions of either, implicates aBST as a key integrator of stress-inhibitory influences emanating from the limbic cortex. Indeed, our examination of the projections of PL and vSUB reveal extensive overlap in their terminal innervation of PVH-projecting neurons within aBST, with evidence of some convergence onto single neurons (Figure 4).

IMPLICATIONS OF AN HPA-INHIBITORY NETWORK

The elucidation of this network should help to address some of the lingering questions concerning the central organization of HPA control. First is the generality of aBST as a site of convergence and integration of additional forebrain limbic influences on emotional stress-induced HPA output (Figure 5). None of the other forebrain cell groups implicated in the inhibition of emotional stress-induced HPA activity (i.e., septum, posterior paraventricular thalamic nucleus) provides any substantial direct innervation of PVH, although each projects to aBST (Shin et al., 2008). Thus, aBST GABAergic neurons are poised to receive and integrate these along with prefrontal and hippocampal influences. The amygdala is generally considered to exert an excitatory influence on HPA axis activation (Prewitt and Herman, 1997; Sullivan et al., 2004), however, the circuits and mechanisms accounting for this are poorly understood. Both CeA and MeA issue a massive GABAergic input into GABAergic regions of aBST (Sun and Cassell, 1993; Tsubouchi et al., 2007), particularly the aforementioned PVH-projecting population. This suggests that excitatory effects on HPA output may be mediated via disinhibition of this modulatory pathway. The basolateral amygdala (BLA), which
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FIGURE 4 | Top: Overlap of retrograde tracer injections in PVH (Fluoro-gold, FG; cyan), and anterograde tracers in PL (BDA; green) and vSUB (FlouroRuby, FR; red) was evaluated in fluorescence preparations using confocal microscopy. Instances of BDA- (arrows) and FR-labeled (arrowhead) terminals were found to make appositions onto single PVH-projecting neurons in aBST, by analysis of single optical planes containing fluorescence labeling for all three markers. Bottom row: After a single stress exposure, numerous instances of Fos-labeled nuclei are evident in PVH-projecting neurons containing appositions from BDA- (left) and FR-labeled (right) terminals. Data are based upon Radley and Sawchenko (2011).

FIGURE 5 | Proposed role of anterior bed nuclei of the stria terminalis (aBST) as an integrator of limbic forebrain influences on emotional stress-induced HPA output. Previous work of ours supports the pathways highlighted in red, with aBST providing an important source of GABAergic innervation of PVH, and relaying limbic cortical influences. Other forebrain cell groups known to influence HPA output (highlighted in black), notably via GC receptor-mediated negative feedback, also project to aBST, whose integrated output targets PVH directly. Like ventral subiculum (vSUB) and prelimbic cortex (PL), these regions do not provide any appreciable innervation of PVH, but do issue projections to the aBST. ACTH, adrenocorticotropic hormone; Amyg, amygdala; cc, corpus callosum; CRF, corticotropin-releasing factor; Glu, glutamate; HF, hippocampal formation; LS, lateral septum; ot, optic tract; Pit., pituitary gland; PVH, paraventricular nucleus of the hypothalamus; PVT, paraventricular thalamic nucleus.

Consists predominantly of pyramidal-like glutamatergic neurons, is also implicated in stimulating emotional stress-induced HPA activation (Bhatnagar et al., 2004). One likely scenario is for BLA to access the PVH via the BLA-to-CeA pathway widely implicated as the direction of information flow for the genesis of fear-related autonomic and behavioral responses (Pitkanen et al., 1997; LeDoux, 2000), then proceeding via a CeA (GABA) → aBST (GABA) → PVH pathway.

The model as proposed may also help to advance our understanding of the circuits and mechanisms accounting for GC receptor-mediated negative feedback. A number of cell groups implicated in inhibiting emotional stress-induced HPA activation (i.e., PL, HF, PVTp, lateral septum) are also capable of imparting GC receptor-mediated negative feedback on the axis (Feldman and Conforti, 1980a; Jacobson and Sapolsky, 1991; Diorio et al., 1993; Jaferi and Bhatnagar, 2006). This raises the possibility that aBST may integrate steroid-dependent feedback information from the limbic forebrain for conveyance to the PVH. Evidence increasingly suggests that GC negative feedback in the limbic forebrain may be mediated via an endocannabinoid signaling mechanism. For example, Hill and colleagues (2011) recently reported that GC receptor activation in mPFC neurons mobilizes the release of endocannabinoids and increases excitatory outflow from principal neurons via the presynaptic inhibition of GABA release from local interneurons (Hill et al., 2011). Understanding of a broader circuitry for imparting inhibitory influences over the stress axis should allow for assessment of whether GC negative feedback is restricted to upstream mediators, or whether aBST can intercede for these influences as a proximate source of steroid-mediated feedback, and, the generality of endocannabinoid signaling in relaying GC-dependent feedback in other components of the network.

Finally, if the proposed framework is inhibitory in nature, this helps to clarify a fundamentally important question of what drives the initial activation of PVH and HPA output during stress. As previously noted, IL appears to exert an excitatory influence on HPA output via a distinct relay in aBST, and may comprise one of the upstream components for an activating network. The idea that a non-GABAergic subpopulation of aBST neurons relays excitatory influences from IL to the PVH is consistent with evidence that indiscriminate excitotoxin lesions in aBST reliably attenuate acute emotional stress-induced HPA output (Choi et al., 2007). One proposal from Choi and colleagues (2007) is that the non-neuroendocrine CRF-expressing subpopulation within aBST (corresponding to the fusiform subdivision of Dong et al., 2001)
may provide a source of excitatory input into PVH, and future studies will help to clarify this relationship further. At least with regard to physiological stress, HPA output appears to be mediated predominantly via medullary aminergic inputs to PVH, as ablation of this pathway completely blocks central and peripheral indices of HPA activation under exposure to these challenges (Ritter et al., 2003; Schiltz and Sawchenko, 2007). By contrast, this pathway does not mediate HPA activation during acute emotional stress (Ritter et al., 2003; Schiltz and Sawchenko, 2007), and evidence for an equivalent activating system under this category of challenges remains elusive.

**CHRONIC STRESS-INDUCED NETWORK REORGANIZATION**

Chronic stress induces profound structural and synaptic changes in a variety of limbic forebrain regions. mPFC (ACd, PL, and IL) and CA3 hippocampal neurons show regressive alterations in apical dendritic and synapse morphology (Watanabe et al., 1992; Magarinos and McEwen, 1995a; Cook and Wellman, 2004; Radley et al., 2004, 2006b; Stewart et al., 2005; Hajszan et al., 2009), whereas amygdala (BLA) neurons show increases in these indices (Vyas et al., 2002, 2006). These changes are paralleled by reductions in gray matter volume and functional impairments in mPFC and HF of depressed individuals (Sheline et al., 1996, 2003; Drevets et al., 1997). Generally speaking, stress-induced structural plasticity is dependent on elevated GCs and excitatory glutamatergic signaling (Liu et al., 2008; Magarinos and McEwen, 1995b). Such structural alterations have been linked with disruptions in learning and memory (Luine et al., 1994; Stewart et al., 2005; Liston et al., 2006; Dias-Ferreira et al., 2009; Holmes and Wellman, 2009; cf. Conrad, 2010), and increases in anxiety-like behaviors (e.g., elevated plus maze performance; Mitra et al., 2005). HPA axis hyperactivity (i.e., sensitization, facilitation) is also widely documented to result from chronic stress (Ottenweller et al., 1989; Dallman et al., 1992; Willner, 1997; Bhatnagar and Dallman, 1998; Figueiredo et al., 2003a; Weinberg et al., 2010), although its relation to structural plasticity in the limbic forebrain remains to be thoroughly examined.

Much of the previous literature relevant to studying the effects of chronic stress on structural plasticity entailed examination of dendritic branching patterns. This is likely due to the fact that stress and GCs produce robust changes on neurons that are readily manifest at the morphological level, and that changes in dendritic branching patterns (i.e., complexity, length, branch numbers) generally were thought to correlate with changes in synaptic connectivity. Nonetheless, increasing attention has been given to more detailed analyses of synaptic alterations in neural circuits following chronic stress, given their critical role as junctional points of connectivity that mediate information flow between neurons. Dendritic spines represent sites of postsynaptic contact for the majority of excitatory synaptic input in cortical structures. Spines are highly motile and dynamic structures that exhibit a wide degree of morphological diversity, with alterations in shape and number providing a cellular correlate for learning capacity, learning, and memory (Bailey and Kandel, 1993; Sorra and Harris, 2000; Kasai et al., 2003; Wilbrecht et al., 2010). Despite the heterogeneity of spine morphology, their classification into broad categories has proven useful. For instance, long and thin spines tend to be regarded as immature, and are more abundant during development, whereas mushroom-shaped spines (large diameter head, small diameter neck) represent stronger, more well-established excitatory synapses.

A number of studies suggest that chronic stress leads to a net loss of excitatory synapses in PL neurons (Radley et al., 2006b; Michelsen et al., 2007; Liu and Aghajanian, 2008; Arnsten, 2009). Notably, dendritic spines in the distal portions of the apical dendritic tree appear to be most profoundly impacted by chronic stress, inclusive of retraction of distal processes and decreases in spine density (Figure 6) (Radley et al., 2006b; Liu and Aghajanian, 2008). We conducted a high-throughput analysis of over 17,000 dendritic spine morphologies in PL and ACd pyramidal neurons (Radley et al., 2008a), and found that chronic stress resulted in an overall decrease in apical dendritic spine density, manifested by a loss of mushroom-shaped spines, and an increased frequency of long and thin spines. Another recent report employing two-photon microscopic *in vivo* imaging of spines in has provided the most compelling evidence to date for the capacity of GCs to mediate stress-induced spine alterations in the cortex (i.e., primary motor, secondary motor, somatosensory; Liston and Gan, 2011). Whereas acute GC exposure increased the rate of spine turnover (elimination and formation), prolonged GC exposure selectively increased the elimination of spines, particularly ones that were older and more stable. These, and other studies (Michelsen et al., 2007; Liu and Aghajanian, 2008), support the idea that chronic stress, via increases in GC levels, may selectively target the mature, stable population of excitatory synapses throughout cortical structures.

Chronic stress has also been shown to decrease synapse and spine density in hippocampal neurons (Sousa et al., 2000; Sandi et al., 2003; Stewart et al., 2005; Hajszan et al., 2009). In one of the more rigorous demonstrations of this phenomenon, one study employed electron microscopy (EM) and stereological 3-D reconstructions in, finding that chronic stress induced significant decreases in dendritic spine density and synapse number, and was reversible following a stress-free recovery period (Sandi et al., 2003). In another study employing EM, Magarinos and colleagues (1997) reported ultrastructural differences in presynaptic terminals of synapses in the mossy fiber pathway in CA3 neurons following chronic stress, indicative of an up-regulation of presynaptic activity and release of glutamate. Taken together, decreases in density and in overall numbers of postsynaptic excitatory contacts may help to limit the extent of excitotoxic damage that would otherwise result from prolonged activation of glutamatergic synapses under chronic stress.

A number of studies have begun to identify the cellular mechanisms underlying chronic stress-induced spine synapse loss (for reviews, see Arnsten, 2009; Duman and Voleti, 2012). For example, reduced expression of certain neurotrophic/growth factors (notably, brain-derived neurotrophic factor) in the hippocampus, and more recently in mPFC, may contribute to dendritic spine synaptic compromise in these regions (Liu et al., 2012; Nibuya et al., 1995; Kuipers et al., 2003). Alterations in protein kinase C signaling have also been shown to underlie dendritic spine loss in mPFC (Hains et al., 2009). Finally, the mammalian target of rapamycin (mTOR) signaling pathway has recently been...
implicated in synaptic deficits that result from excessive glutamatergic stimulation, such as which ensues under chronic stress (Magarinos and McEwen, 1995b; Li et al., 2011). These studies highlight potentially important cellular mechanisms for investigating their role in the circuit alterations underlying neuroendocrine adjustments following chronic stress.

From the network perspective, large-scale decreases and destabilization of the excitatory synapse population in mPFC and HF could uncouple excitatory afferent input from excitatory outflow in PL and/or HF, resulting in their diminished influence over PVH-projecting GABAergic neurons in aBST. Concurrent increases in BLA neuronal dendritic branching and spine densities could also drive disinhibition of PVH-projecting GABAergic neurons, via increasing activation in the extrinsic GABAergic projections from CeA. One challenge concerns whether changes throughout the entire network are necessary for HPA axis hyperactivity following chronic stress, or whether this phenotype is regulated by a distinct pathway or mechanism. As many of the stress-related changes in HF have been demonstrated more dorsally in CA3, and to some extent in DG and CA1 (e.g., Sousa et al., 2000; Snyder et al., 2011) it is unclear whether vSUB serves as a way station, or whether stress effects within vSUB proper (or ventral hippocampus) account for alterations in excitatory outflow to aBST. Another issue concerns the fact that little is known about how chronic stress impacts other cell groups implicated in the stress-inhibitory network, such as PVTp and lateral septum. The fact that GCs appear capable to exert widespread effects throughout the cortex, inclusive of sensorimotor regions (Liston and Gan, 2011), poses additional challenges in teasing apart neural circuits that underlie stress-related behavioral and physiological alterations.

SUMMARY

It has been previously established that the HPA axis response to emotional stress involves a network of limbic forebrain afferents that exert their effects on the PVH via multisynaptic and parallel pathways. Recent evidence lends support for at least two limbic cortical regions, mPFC and HF, that impart their inhibitory influences over the stress axis by converging on a discrete target, the aBST, that in turn inhibits the PVH and HPA activity. Importantly, GABAergic neurons in the aBST exhibit the capacity to integrate the inhibitory prefrontal and hippocampal influences that they impart on the stress axis. There are a number of hypotheses that derive from this model that should help to inform future work. One idea is that aBST serves as a neural hub for receiving and integrating stress-modulatory influences from other limbic forebrain regions (i.e., PVTp, septum, amygdala). Another implication is that this network, notably via GABAergic relays in aBST, may serve to integrate GC receptor-mediated negative feedback signals from some, if not all of, these limbic forebrain regions via a presynaptic endocannabinoid signaling mechanism. As this network is inhibitory, this may help to inform the search for HPA-activating networks; i.e., something akin to the medullary aminergic inputs to PVH that are known to drive HPA output in response to physiological stressors.

Chronic stress-induced neuroplasticity throughout the limbic cortex, or within key regions, may lead to an attrition of their excitatory influence on the PVH-projecting GABAergic cell population in aBST, producing HPA axis hyperactivity. A key feature of this hypothesis is that regressive changes are evident in regions that normally serve to inhibit HPA axis activation during emotionally stressful experiences, and hypertrophic

FIGURE 6 | Summary of effects of chronic stress on structural plasticity in mPFC pyramidal neurons. In these studies (Radley et al., 2004, 2006b, 2008a), high resolution analyses were performed in digitally reconstructed dendritic segments from fluorescent dye-injected pyramidal neurons in dorsal anterior cingulate (ACd) and prelimbic (PL) areas. An atlas plate (lower left) depicts the approximate region within mPFC that neurons were filled for morphologic analyses. Distance in millimeters relative to bregma is indicated. Arrows highlight the fact that dendritic atrophy and spine/excitatory synapse loss is most prominent on distal apical dendrites (right). Spine morphologic analyses reveal fewer spines with mature (stubby, mushroom-shaped), and a greater number with immature (long and thin) phenotype.
changes (i.e., increased branching and synapse number) are evident in regions that contribute an excitatory influence on HPA output. Whereas considerable gains have been made in understanding the cellular mechanisms underlying dendritic spine dynamics, much of this work has not been applied to enhance our knowledge of how chronic stress leads to long-term changes in neuroendocrine function. Moreover, examination of why stress and GCs have bidirectional effects on excitatory synapse plasticity in BLA, relative to HF and mPFC, may shed light on what are likely to be categorically distinct effects on the regulation of gene expression in these cell types that are imparted by GCs. Finally, susceptibility to stress-related psychiatric illnesses depends on a number of factors (e.g., genetics, early-life experiences, previous stress exposure) that may help to explain why some individuals go on to develop stress-related disorders while others do not. The extent to which structural plasticity in limbic cortical regions is predictive of adaptation or failure of stress/HPA control systems is of fundamental importance for informing the issue of how these more complex hereditary and environmental factors may tip the balance between stress resilience and pathology.

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A novel HPA-inhibitory network in stress regulation

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