

Peripheral and central mechanisms of stress resilience



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ABSTRACT

Viable new treatments for depression and anxiety have been slow to emerge, likely owing to the complex and incompletely understood etiology of these disorders. A budding area of research with great therapeutic promise involves the study of resilience, the adaptive maintenance of normal physiology and behavior despite exposure to marked psychological stress. This phenomenon, documented in both humans and animal models, involves coordinated biological mechanisms in numerous bodily systems, both peripheral and central. In this review, we provide an overview of resilience mechanisms throughout the body, discussing current research in animal models investigating the roles of the neuroendocrine, immune, and central nervous systems in behavioral resilience to stress.

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1. Introduction

Decades of research on human stress resilience have followed its initial description in at risk children in the 1970s (Masten, 2001). Resilience is defined as the adaptive maintenance of normal physiology, development and behavior in the face of pronounced stress and adversity. Human resilience literature contains countless examples of adults and children who, despite marked psychological stress, display minimal adverse changes in emotional wellbeing or behavioral disturbances (Feder et al., 2009; Yehuda et al., 2006b; Alim et al., 2008; Fredrickson et al., 2003; Bonanno, 2004). Although it is tempting to attribute human resilience to the possession of exceptional abilities and coping mechanisms, both social and biological, most people do not develop anxiety and depression when faced with stress (Masten, 2001; Bonanno, 2004). Resilience is a common outcome that more likely involves the successful application of the body's adaptive stress response to maintaining the status quo. The biological processes underlying resilience are often collectively termed “allostasis” and constitute

variation in bodily systems that functions to maintain homeostasis in response to a stressor (McEwen, 2002). In some cases, allostasis is exaggerated or fails to cease along with the stressor, and mechanisms that were once protective can become pathological. This phenomenon—termed “allostatic load”—can potentially result in physiological and psychological damage, including enhanced susceptibility to disorders such as depression and anxiety (McEwen, 2002; Charney, 2004).

Mechanisms of resilience are of great interest due to the serious burdens imposed on patients and society by stress-related disorders including anxiety and depression. One in six Americans will develop Major Depressive Disorder (MDD) during their lifetime, a particularly alarming statistic as only 30% of patients achieve complete remission of symptoms following treatment with current first-line therapies, the monoamine-based antidepressants (Krishnan and Nestler, 2008; Kessler et al., 2005). When not adequately treated, MDD can become a chronic, recurrent condition characterized by escalating disability (Moussavi et al., 2007). Comprehensive knowledge of the etiology of depression is still lacking. Understanding the adaptive, allostatic mechanisms that protect most individuals against psychopathology can potentially inform therapeutic development and treatment strategies for more vulnerable individuals.

Depression and anxiety are increasingly considered to be “whole body” illnesses involving the dysregulation of multiple systems, both peripheral and central. Similarly, resilience likely results from successful allostatic mechanisms in the

Abbreviations: CUS, chronic unpredictable stress; SCUS, subchronic unpredictable stress; CSDS, chronic social defeat stress; LH, learned helplessness; LG, licking and grooming; OVX, ovariectomy; SDR, social disruption stress.

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hypothalamo–pituitary–adrenal (HPA) axis, autonomic nervous system, immune system and the brain (McEwen, 2002). In this review, we summarize recent research into the roles of the neuroendocrine, immune and central nervous systems in resilience to stress, focusing primarily on animal models. We describe both active, compensatory mechanisms as well as passive mechanisms in which the absence of a maladaptive stress response promotes resilience. We present resilience as an integrated process by which adaptive mechanisms in multiple systems promote psychological resilience to stress.

2. Animal models for the study of stress vulnerability and resilience

Research on human subjects has yielded important insights into the roles of various neurotransmitters, neuropeptides and hormones as well as genetic factors in the neurobiology of resilience (for comprehensive reviews, see Charney, 2004; Russo et al., 2012). For ethical and practical reasons, animal models are often employed to examine the causative effects of stress on biological processes in the brain and body. Resilience to stress has been documented and characterized in animal models throughout the lifespan. Below, we describe in detail several behavioral paradigms commonly used to elicit and study stress resilient phenotypes in juvenile and adult animals.

2.1. Early life stress

Models of early life stress have informed our understanding of a form of resilience called stress inoculation, whereby early stressful experience attenuates stress response in adulthood. In children, early stress can have a “steeling” effect, promoting subsequent stress resistance and successful psychological functioning (Rutter, 2006). Animal models of early life stress typically involve exposure to stressful stimuli during either the prenatal or postnatal periods. Prenatal stressors include maternal stress such as glucocorticoid administration or food deprivation while early postnatal stressors include brief bouts of maternal separation, altered maternal care behavior, or glucocorticoid administration (Lupien et al., 2009). Prolonged early life stress can cause programmed HPA axis overactivity, altered glucocorticoid response, structural changes in the brain, and deleterious effects on cognition, emotion and behavior (Lupien et al., 2009). These effects can be reconciled with the concept of stress inoculation by imagining adult outcomes of early life stress as a U-shaped curve—animals exposed to moderate stress in early life show better outcomes and more adaptive responses to stress in adulthood than do animals exposed to minimal or severe stress (Macri et al., 2011). Stress inoculation has been demonstrated in both primates and rodents. Infant squirrel monkeys separated from their mothers for brief, intermittent periods demonstrate reduced hormonal stress response in subsequent developmental stages (Lyons et al., 2010; Parker et al., 2005). They also demonstrate cognitive and emotional resilience across measures relevant to anxiety and depression, such as enhanced novelty tolerance, exploratory behavior and behavioral response inhibition (Lyons et al., 2010; Parker et al., 2004, 2005). There is a rich literature on stress inoculation in rodents demonstrating that rats exposed to early life stress, including brief maternal separations and neonatal corticosterone administration, display blunted HPA axis response to stress in adulthood as well as behavioral resilience in the form of reduced anxiety-like behavior and enhanced performance in cognitive tasks (Macri et al., 2009; Plotsky and Meaney, 1993; Macri and Wurbel, 2007; Levine, 1962; Levine et al., 1967). The relationship between early life stress exposure and subsequent resilience in both primates and

rodents follows the abovementioned U-shaped curve. Prolonged maternal separation and social isolation in infant rhesus monkeys produce an increased stress response and “despair-like” behavior in subsequent social separation tests (Young et al., 1973). Rats exposed to moderate early life stress show enhanced measures of resilience compared to both severely and minimally stressed rats (Macri and Wurbel, 2007). For example, early postnatal rats exposed to brief daily handling (a moderate stressor) subsequently show attenuated stress response compared to undisturbed pups and pups exposed to prolonged daily maternal separation (a more severe stressor) (Plotsky and Meaney, 1993; Macri et al., 2004).

2.2. Chronic unpredictable stress

Chronic unpredictable stress (CUS) is a useful model for examining stress vulnerability and resilience in rodents (Ricon et al., 2012; LaPlant et al., 2009). In CUS paradigms, animals are exposed to varying mild stressors sequentially for a period of 1–7 weeks (Krishnan and Nestler, 2011; Willner, 1997). Stressors can include mild foot shock, physical restraint, tail suspension, light/dark cycle disruption, food or water restriction, changes to cage mate, etc., and are changed after several hours to minimize habituation (LaPlant et al., 2009; Willner, 1997). CUS produces a range of depression and anxiety-like behaviors in rodents including anhedonia, measured as decreased sucrose preference, despair-like behavior, measured as increased immobility in the forced swim and tail suspension tests, and novelty suppressed feeding, measured as a decrease in approach to a novel food item (Krishnan and Nestler, 2011; Mineur et al., 2006; Feng et al., 2012). Mice exposed to CUS also display decreased grooming, aggression, and sexual behaviors. Certain CUS-induced behavioral changes, such as novelty suppressed feeding, can be reversed only by chronic antidepressant treatment (Willner, 1997), making CUS relevant to human antidepressant responses. Female mice display immobility in the forced swim test after just 6 days of subchronic unpredictable stress (SCUS) whereas males are generally resilient to SCUS and require 20–28 days of CUS exposure to elicit depression- and anxiety-like behavior (Hodes, G.E. et al., *Soc. Neurosci. Abstr.* 219.01, 2011). Interestingly, age is a factor in response to CUS—male rats exposed to 60 days of CUS in the juvenile period exhibit greater memory retention in a two-way shuttle avoidance task compared to rats exposed to the same stressor in adulthood, indicating enhanced cognitive resilience (Ricon et al., 2012). Sex differences and age effects in susceptibility to CUS-induced depression and anxiety-like behavior make this a powerful tool for investigating the hormonal and neural basis for stress vulnerability and resilience across the lifespan.

2.3. Chronic social defeat stress

Chronic social defeat stress (CSDS) is a prolonged social stress paradigm used to delineate stress resilient and susceptible phenotypes in adult rodents. In the CSDS model, a C57BL/6J mouse is repeatedly subordinated by a larger, aggressive CD-1 mouse for 10 consecutive days (Golden et al., 2011). Each physical bout is followed by overnight sensory contact with the aggressor through a plastic partition. Following CSDS, approximately 2/3 of experimental mice, termed “susceptible,” develop a constellation of depression-like behaviors including social avoidance and anhedonia (Krishnan et al., 2007; Donahue et al., 2014) as well as metabolic syndrome marked by dysregulated feeding peptides, weight gain and insulin insensitivity (Chuang et al., 2010; Lutter et al., 2008). Conversely, the remaining 1/3 of mice, termed “resilient,” develop a much milder phenotype, including elevated corticosterone and increased anxiety-like behavior (Krishnan et al., 2007).

Similar to human depression, CSDS-induced depression- and anxiety-like behavior can be reversed by chronic, but not acute, administration of antidepressants (Berton et al., 2006; Tsankova et al., 2006). Importantly, a number of biomarkers identified in humans with MDD are similarly disrupted in susceptible mice following CSDS, further highlighting its utility in studying depression mechanisms (Krishnan et al., 2007; Golden et al., 2013; Robison et al., 2014).

2.4. Learned helplessness

The learned helplessness (LH) model is an acute stress paradigm that, similar to CSDS, produces heterogeneous responses, enabling researchers to delineate stress susceptible and resilient animals (Krishnan and Nestler, 2011). The proportion of animals exposed to the LH paradigm that demonstrate phenotypic resilience ranges from 10% to 80% (Cryan and Mombereau, 2004). In this model, rodents are exposed to repeated inescapable foot shocks followed by a test period in which an easy escape mechanism is made available during shock exposure. Compared to control animals trained with escapable shocks and resilient animals, susceptible animals demonstrate “helplessness,” measured as longer escape latency or failure to escape (Seligman and Beagley, 1975). Like CSDS, the LH paradigm produces numerous behavioral and physiological changes including weight loss, HPA axis dysfunction, circadian alterations, and reductions in hippocampal synaptic spine number (Krishnan and Nestler, 2011). A weakness of the model is that LH-induced changes are short-lived, usually lasting only 2–3 days and can be reversed with acute antidepressant treatment (Cryan and Mombereau, 2004).

3. Neuroendocrine mechanisms of resilience

Appropriate response to stress involves the coordinated activity of the autonomic nervous system (ANS) and the HPA axis as well as the neural circuits in the hypothalamus, brainstem and forebrain that control their activity (for a comprehensive review, see Ulrich-Lai and Herman, 2009). Briefly, stress induces ANS processes by first activating preganglionic sympathetic neurons in the spinal cord that project to pre- or paravertebral ganglionic neurons. These neurons terminate on cardiovascular and visceral organs or on the adrenal medulla, and stimulate the release of adrenaline from the adrenal medulla and noradrenaline from sympathetic nerve fibers. Consequences of ANS activation by stress include changes in heart rate and vasoconstriction. In the HPA axis, stress activates neurons in the paraventricular nucleus (PVN) of the hypothalamus to secrete corticotropin releasing factor (CRF) and arginine vasopressin (AVP) into the portal circulation via the median eminence, which in turn stimulate the anterior pituitary gland to release adrenocorticotrophic hormone (ACTH). ACTH activates glucocorticoid synthesis and release from the adrenal cortex, which functions primarily to mobilize energy stores during stress. There is ample cross-talk between the ANS and the HPA axis—the adrenal cortex receives innervation from the sympathetic nervous system, regulating glucocorticoid release, and glucocorticoids mediate ANS-dependent stress responses including vasoconstriction. Modulation of these systems has been noted in cases of human resilience to MDD and post-traumatic stress disorder (PTSD), although results have been largely correlative (Russo et al., 2012). High dose glucocorticoid administration following traumatic stress exposure has emerged as a potential treatment for individuals vulnerable to PTSD, perhaps working by controlling hyperactive fear response and fear memory consolidation (Kearns et al., 2012). This strategy has yielded positive results in critically ill hospital patients and combat-exposed veterans (Schelling et al., 2006; Suris et al., 2010).

Additionally, dehydroepiandrosterone (DHEA) and neuropeptide Y (NPY) have emerged as potential pro-resilience biomarkers in humans. DHEA is released from the adrenal cortex with cortisol in response to stress and can counter the effects of glucocorticoids (Yehuda et al., 2006a). In combat-exposed veterans, both DHEA level and DHEA/cortisol ratio correlate negatively with PTSD symptom severity, suggesting that DHEA may serve a protective role in situations of extreme stress. NPY is co-released with noradrenaline from sympathetic nerves and has been shown to correlate positively with interrogation performance and negatively with dissociative symptoms in soldiers undergoing a U.S. Army survival training course (Morgan et al., 2000b).

The Hypothalamic Pituitary Gonadal (HPG) axis shares numerous component structures and neural circuitry with the HPA axis, and accordingly, reproductive hormones serve a prominent role in susceptibility and resilience to stress. Mood disorders including MDD and anxiety are about two times more prevalent in adult women than men, a sex difference that emerges in puberty and persists until menopause, suggesting a role for sex hormone fluctuations and activating effects of gonadal hormones on neural circuitry (Deecheer et al., 2008; Holden, 2005; Epperson et al., 2014). Estrogen supports cognitive processes, influences catecholamine and monoamine neurotransmission and metabolism, and regulates the expression of transcription factors and neurotrophins (Shanmugan and Epperson, 2014). Activation of brain stress response and reward circuitry depends on menstrual cycle stage in healthy adult women (Goldstein et al., 2010; Dreher et al., 2007). Women with a history of MDD display hypoactivation of brain stress response circuitry associated with lower serum estradiol levels and higher serum progesterone levels compared to healthy controls (Holsen et al., 2011). Mechanistically, perimenopause-associated estradiol fluctuations have been shown to contribute to vulnerability in part by increasing brain levels of monoamine oxidase A (MAO-A), an enzyme involved in apoptosis, oxidative stress, and monoamine metabolism (Rekkas et al., 2014). Conversely, testosterone has emerged as a potential pro-resilience factor in men (Russo et al., 2012). There is a strong positive correlation between testosterone and degree of social connectedness, feelings of personal success, and social dominance (Edwards et al., 2006). Given its role in social behavior and positive mood, it is not surprising that blood and saliva testosterone levels decrease following stress (Morgan et al., 2000a) and that low circulating levels are often found in individuals with PTSD or MDD (Mulchahey et al., 2001; Pope et al., 2003). Early studies in men suggest that testosterone may be effective in alleviating treatment resistant depression and as an adjunct to treatment with selective serotonin reuptake inhibitors (Pope et al., 2003). Although much future work is needed, together this work suggests that testosterone may serve as a pro-resilience factor by promoting positive mood and social connectedness.

3.1. The HPA axis

Animal studies investigating the mechanistic underpinnings of resilience related to the HPA axis largely focus on models of developmental stress. Adult rats that have undergone stress inoculation in the form of postnatal handling display lower basal levels of CRF, blunted stress-induced increases in ACTH, CRF and corticosterone secretion, and a more rapid post-stress recovery to basal stress hormone levels compared to unstressed rats or those that have undergone maternal separation (Plotsky and Meaney, 1993). Meaney and Szyf (2005) have identified maternal care behavior as a mediator of early life stress resilience that produces long lasting individual differences in gene expression and subsequent neuroendocrine stress response. In a study by Liu et al. (1997), they report that mothers of handled rats displayed more licking, grooming and

arched back nursing behaviors than mothers of nonhandled rats. The amount and frequency of these maternal behaviors correlated negatively with stress-induced plasma ACTH and corticosterone in adulthood (Liu et al., 1997). This pro-resilient neuroendocrine stress response in adult offspring of high maternal care mothers resulted from increased hippocampal expression of glucocorticoid receptors (GR), causing greater sensitivity to glucocorticoid negative feedback on CRF and AVP. The group has since identified a number of molecular mediators of enhanced GR expression in handled pups such as increased thyroid hormone secretion, serotonin turnover in the hippocampus, and hippocampal expression of nerve growth factor-inducible protein A (NGFI-A), a cAMP-inducible transcription factor that binds exon 1₇ of the *GR* promoter (Meaney and Szyf, 2005; Meaney et al., 2000; Weaver et al., 2004). In adult rats, epigenetic mechanisms maintain glucocorticoid receptor sensitivity in resilient animals. The 5' CpG dinucleotide site of the NGFI-A consensus sequence on *GR* is always methylated in offspring of low licking and grooming (LG) mothers whereas it is associated with acetylated H3 in the offspring of high LG mothers (Meaney and Szyf, 2005). Methylation of this site prevents the binding of NGFI-A to the *GR* promoter whereas acetylation has the opposite effect. In sum, high LG maternal care produces sustained epigenetic modifications that induce enhanced glucocorticoid receptor expression, enhanced sensitivity to glucocorticoid negative feedback, reduced hypothalamic release of AVP and CRF, and ultimately attenuated HPA axis response to subsequent stress (Kappeler and Meaney, 2010).

Although less is known about the HPA mechanisms underlying resilience to adulthood stress, two recent studies identify pro-resilience epigenetic modifications at the *CRF* gene in PVN neurons and CRF gating of brain-derived neurotrophic factor (BDNF) in the nucleus accumbens (NAc) as important mediators. Following CSDS exposure, Elliott et al. (2010) reported increased *CRF* mRNA expression in the PVN and decreased methylation at four CpG sites in the *CRF* promoter in susceptible, but not resilient, mice. Viral-mediated knockdown of *CRF* in the PVN after social defeat promoted resilient behavior in the social interaction test, suggesting that *CRF* promoter methylation in resilient animals underlies adaptive neuroendocrine and behavioral responses. Walsh et al. (2014) found that optogenetic induction of phasic firing in dopaminergic neurons of the ventral tegmental area (VTA) promoted social avoidance behavior in mice following subthreshold social defeat stress, an effect dependent upon CRF-gated induction of BDNF in the NAc, a structure in which VTA dopaminergic projections terminate. As CRF antagonist infusion blocked the effects of phasic stimulation on social avoidance behavior, CRF is likely an essential mediator of vulnerability and resilience to defeat stress. Future investigation of individual differences in CRF in the NAc will further elucidate CRF activity in resilient animals.

3.2. Gonadal sex hormones

The effects of sex hormones on resilience and vulnerability to stress are highly complicated and dependent upon the timing of stress (adulthood vs. developmental) and behavioral domain (cognitive vs. emotional resilience) (see Table 1). Adult female animals tend to show greater cognitive resilience to chronic stress than males, while males display enhanced emotional resilience. Following chronic restraint stress, male rats exhibit deficits in hippocampal-dependent memory tasks including the radial arm maze, object recognition test, Y maze, Morris water maze and object location task, whereas female rats are either unaffected or perform better on spatial and visual memory tasks (Luine, 2002; Conrad et al., 2003; Kittraki et al., 2004). Mechanisms underlying these differences in cognitive resilience include sex differences in stress-induced changes to hippocampal morphology and

corticosteroid receptor sensitivity. Chronic restraint stress induces atrophy of apical dendrites, measured as reduced dendritic length and branch points, in the CA3 region of the hippocampus in male, but not female, rats (Galea et al., 1997). Following 21 days of restraint stress, Kittraki et al. (2004) reported reduced GR immunoreactivity in the male rat hippocampus and impaired spatial learning and memory in the Morris water maze. In contrast, hippocampal GR and mineralocorticoid receptor immunoreactivity were enhanced in stressed females.

Our laboratory has identified nuclear factor κ B (NF κ B) signaling as a potential mediator of emotional resilience to CUS (LaPlant et al., 2009). NF κ B is a transcription factor that, although most commonly associated with immune function, can also be activated by stress (Christoffel et al., 2011a). The activation and nuclear translocation of NF κ B is regulated by the I κ B kinase complex (I κ K), which triggers the degradation of the cytoplasmic inhibitor of NF κ B, I κ B (Mohamed and McFadden, 2009). As mentioned earlier, gonadally intact females display depression-like behavior after SCUS, whereas males do not exhibit emotional dysregulation at this time point. Ovariectomy (OVX) abolished this enhanced behavioral susceptibility and blunted transcriptional response to stress—170 genes were regulated by SCUS in OVX mice vs. 619 in gonadally intact females, and many overlapping genes were regulated in opposite directions. Viral overexpression of I κ K (significantly upregulated in OVX mice) prevented SCUS-induced immobility in the forced swim test in gonadally intact female mice, whereas overexpression of a dominant negative form of I κ K increased immobility in OVX mice. These findings suggest that I κ K–NF κ B signaling is necessary and sufficient for the expression of resilience following SCUS in females. Additional findings suggest a role for DNA methyltransferase 3a (Dnmt3a) in enhanced male emotional resilience to SCUS. RNA sequencing analysis revealed that female mice exhibit higher expression of Dnmt3a mRNA than do males at baseline. Genetic deletion of Dnmt3a shifts female transcriptional profiles following SCUS exposure toward a more male-like pattern and promotes behavioral resilience to SCUS, whereas viral overexpression of Dnmt3a promotes vulnerability to SCUS. We consistently find that the SCUS-induced transcriptional profiles of male and female mice overlap minimally (Hodes, G.E. et al., *Soc. Neurosci. Abstr.* 219.01, 2011; Pfau, M.L. et al., *Soc. Neurosci. Abstr.* 541.26, 2013). Further mining of these data sets may reveal promising patterns and candidate genes for further understanding of sex-dependent stress resilience.

In addition to the activating effects of sex hormones on stress circuitry in adulthood, prenatal perturbations can exert organizational effects on the brain that dictate sex differences in adult stress response. Mueller and Bale (2008) reported increased depression-like behavior in male, but not female, mice whose mothers had been exposed to CUS during early pregnancy. Male mice displayed elevated amygdala CRF expression and decreased hippocampal GR expression that corresponded with epigenetic alterations—reduced methylation of the *CRF* promoter and enhanced methylation of the 1₇ exon of the *GR* promoter. The authors identified sex differences in prenatal stress-induced placental gene expression profiles, particularly differences in the methylation maintenance enzyme Dnmt1, as potential developmental mechanisms underlying adult phenotypes. Moreover, a recent study showed that stress-induced pro-inflammatory placental gene expression contributes to enhanced male susceptibility to prenatal stress (Bronson and Bale, 2014). Maternal nonsteroidal anti-inflammatory drug treatment reversed the stress-induced increase in placental Interleukin 6 (IL-6) expression and ameliorated locomotor hyperactivity (a behavioral indicator of dopaminergic dysfunction) in prenatally stressed adult male mice. While much work has focused on the maternal environment, an interesting study by Rodgers et al. (2013) demonstrated a role for paternal

Table 1
Sexual dimorphism in rodent behavioral response to stress.

| Stress paradigm | Age | Duration of stress | Male behavioral phenotype | Female behavioral phenotype |
|------------------------------|--|---|---|---|
| Paternal stress | Sires were exposed to stress one month prior to breeding (Dietz et al., 2011) | 10 Days of CSDS (Dietz et al., 2011) | <ul style="list-style-type: none"> • Social avoidance (Dietz et al., 2011) • Anxiety-like behavior (Dietz et al., 2011) • Increased locomotion in response to novelty (Dietz et al., 2011) • Decreased sucrose preference (Dietz et al., 2011) • Increased immobility in the Forced Swim Test (FST) (Dietz et al., 2011) | <ul style="list-style-type: none"> • Anxiety-like behavior (Dietz et al., 2011) • Increased immobility in the FST (Dietz et al., 2011) |
| Gestational stress | Dams were exposed to the stressor during early gestation (days 1–7) (Mueller and Bale, 2007, 2008; Bronson and Bale, 2014) | 7 Days of chronic variable stress (Mueller and Bale, 2007; 2008; Bronson and Bale, 2014) | <ul style="list-style-type: none"> • Impaired memory acquisition, enhanced recall in the Barnes maze (Mueller and Bale, 2007) • Increased immobility in the Tail Suspension Test (TST) and FST (Mueller and Bale, 2008) • Decreased sucrose preference (Mueller and Bale, 2008) • Locomotor hyperactivity (Bronson and Bale, 2014) | <ul style="list-style-type: none"> • Enhanced memory acquisition in the Barnes maze (Mueller and Bale, 2007) |
| Early life stress | P2–P10 (Macri et al., 2009) P1–P13 (Macri et al., 2004) P1–P14 (Caldji et al., 2000) | <p>Brief/mild: 9 days of low corticosterone administration to lactating mother (Macri et al., 2009), 13 (Macri et al., 2004) or 14 (Caldji et al., 2000) days of 15-min maternal separation</p> <p>Prolonged/severe: 9 days of high corticosterone administration to lactating mother (Macri et al., 2009), 13 days of 4 h maternal separation (Macri et al., 2004), 14 days of 3 h maternal separation (Caldji et al., 2000)</p> | <p>Brief/mild:</p> <ul style="list-style-type: none"> • Enhanced attentional set-shifting performance (cognitive flexibility) (Macri et al., 2009) • Reduced latency to approach food and water in a novel environment (Macri et al., 2004; Caldji et al., 2000) • Decreased startle response (Caldji et al., 2000) • Reduced anxiety-like behavior in the open field (Caldji et al., 2000) <p>Prolonged/severe:</p> <ul style="list-style-type: none"> • Similar in behavior to control animals (Macri et al., 2004; Caldji et al., 2000) | <ul style="list-style-type: none"> • The effects of early life stress are poorly understood in females, although recent evidence suggests differential effects of early life stress on exploratory behavior, social behavior and anhedonia (Kundakovic et al., 2013) |
| Chronic restraint stress | 80 Days old (Kitraki et al., 2004) 2–3 Months old (Luine, 2002) | 21 Days of 6 h restraint (Luine, 2002; Conrad et al., 2003; Kitraki et al., 2004) | <ul style="list-style-type: none"> • Spatial and visual memory deficits (Y maze, Morris water maze, radial arm maze, object recognition, object location) (Luine, 2002; Conrad et al., 2003; Kitraki et al., 2004) | <ul style="list-style-type: none"> • Normal or enhanced spatial and visual memory (Luine, 2002; Conrad et al., 2003; Kitraki et al., 2004) |
| Chronic social defeat stress | 7–8 Weeks old (Golden et al., 2011; Krishnan et al., 2007; Donahue et al., 2014) | 10 Days of 10-min defeat bouts (Golden et al., 2011; Krishnan et al., 2007; Donahue et al., 2014) | <ul style="list-style-type: none"> • “Susceptible” animals display social avoidance, increased anxiety-like behavior and decreased sucrose preference (anhedonia) (Krishnan et al., 2007; Donahue et al., 2014) • “Resilient” animals display increased anxiety-like behavior (Krishnan et al., 2007) | <ul style="list-style-type: none"> • The CSDS model has not been adequately adapted for use in females due to a lack of conspecific aggressive behavior |
| Unpredictable stress | Subchronic & Chronic: 7–8 weeks old (LaPlant et al., 2009; Hodes, 2013) | <p>Subchronic: 6 days of varied 1 h stressors (LaPlant et al., 2009; Hodes, 2013)</p> <p>Chronic: 21–28 days of varied 1 h stressors (Hodes, 2013)</p> | <p>Chronic:</p> <ul style="list-style-type: none"> • Decreased sucrose preference (Hodes, 2013) • Decreased grooming behavior (Hodes, 2013) • Increased latency to approach food in a novel environment (Hodes, 2013) • Increased immobility in the FST (Hodes, 2013) | <p>Subchronic & Chronic:</p> <ul style="list-style-type: none"> • Decreased sucrose preference (Hodes, 2013) • Decreased grooming behavior (Hodes, 2013) • Increased latency to approach food in a novel environment (Hodes, 2013) • Increased immobility in the FST (LaPlant et al., 2009; Hodes, 2013) |

stress in male offspring susceptibility. Adult male mice sired by fathers exposed to CUS in puberty or adulthood displayed HPA axis hypoactivity, which correlated with changes in paternal sperm microRNA expression profiles. Together these results highlight the complex interactions between genetics and environment in stress resilience.

4. Immunity & resilience

The interaction of stress and the immune system has become a major focus of psychiatric research since the introduction of the “cytokine hypothesis of depression” in the 1990s (Maes et al.,

2009). The hypothesis asserts that many of the central abnormalities observed in depression—enhanced HPA axis activity, neurodegeneration, decreased neurogenesis, oxidative stress, and serotonergic signaling dysfunction—are at least in part due to peripheral inflammatory cytokines released in response to external, psychological stressors and internal stressors such as chronic disease and “leaky gut.”

A growing literature explores the connection between stress, proinflammatory cytokines, and depression and anxiety-like behavior in both humans and animals. Cytokines are soluble proteins that are released at a site of infection by leukocytes. Upon release, they can readily cross the blood–brain barrier and act

within the central nervous system to mediate sickness behaviors. Sickness behaviors due to inflammation, such as social withdrawal and disinterest in food, overlap greatly with depression behaviors but are attenuated when infection is cleared (Dantzer et al., 2008). Altered regulation of this adaptive behavioral response to immune challenge by chronic illness or psychosocial stress contributes to depression (Maes et al., 2009; Dantzer et al., 2008). For example, patients with chronic inflammatory diseases such as multiple sclerosis, rheumatoid arthritis and asthma can be up to 6 times more likely to develop depression than healthy individuals (Moussavi et al., 2007). Depressed patients also show markers of inflammation, including elevated levels of cytokines and their soluble receptors in serum and cerebrospinal fluid, the most consistently elevated being IL-6 (Maes et al., 1997; Dowlati et al., 2010). Inflammatory markers are also elevated in rodent stress models—chronic stress causes an elevation in serum and brain cytokines including IL-6 and Interleukin-1 β (IL-1 β) (Sukoff Rizzo et al., 2012; Voorhees et al., 2013; Koo and Duman, 2008). In both humans receiving immunotherapy and animal models of inflammation, administration of pro-inflammatory cytokines produces depression and anxiety-like behaviors (Bonaccorso et al., 2001, 2002; Anisman et al., 2002; Sakic et al., 2001). While some studies have shown that antidepressant medications reduce peripheral inflammation (Kubera et al., 2001a,b), others suggest the opposite (Hannestad et al., 2011; Maes et al., 2012), resulting in a shift in drug development efforts that focus on the use of more direct anti-inflammatory agents to promote resilience.

Recent studies form a growing body of evidence that supports the existence of individual differences in inflammatory response to stress and subsequent physiological and behavioral vulnerability. Here, we will discuss peripheral markers characteristic of vulnerability and resilience to stress as well as central mechanisms that contribute to inflammation-mediated behavioral outcomes.

4.1. Heightened peripheral inflammation as a predictor and consequence of stress vulnerability and resilience

Several reports examine changes in immune cell localization and reactivity driven by stress exposure in rodents. Many of these studies utilize a social stress model similar to CSDS called social disruption stress (SDR). SDR involves chronic disruption of established social hierarchies in cages of male mice. Male cagemates establish a social hierarchy such that one mouse is the dominant, alpha male and the remaining males are codominant or subordinate (Avitsur et al., 2009). Once a day for a total of six days, a novel, dominant intruder mouse previously screened for aggressive behavior is placed into the housing cage for a period ranging from hours to overnight (Avitsur et al., 2001). The dominant intruder repeatedly attacks and defeats the resident mice, eliciting submissive behaviors. SDR has been shown to activate the HPA axis, enhance proinflammatory cytokine release, drive an increase in splenic and blood CD11b+ monocytes via influx from the bone marrow and stimulate glucocorticoid resistance in immune cells (Avitsur et al., 2009; Engler et al., 2008).

Following SDR, splenic leukocytes from stressed mice release more Tumor Necrosis Factor α (TNF- α) and IL-6 in response to stimulation with lipopolysaccharide (LPS), a bacterial endotoxin and toll-like receptor 4 agonist, compared to leukocytes from control mice, an effect that is driven both by increased number of leukocytes as well as enhanced release from each leukocyte (Avitsur et al., 2005). Enhanced cytokine release likely stems from the glucocorticoid resistance demonstrated by splenic macrophages and monocytes post-SDR, and indicates dysregulation of negative feedback mechanisms by which glucocorticoids and cytokines together self regulate stress-induced hyperinflammation (Stark et al., 2001). SDR-

induced glucocorticoid resistance in macrophages is at least partly due to a cytokine-mediated failure of corticosterone to stimulate nuclear translocation of glucocorticoid receptors and prevent NF κ B-induced proinflammatory transcription (Quan et al., 2003). Splenic macrophage enrichment and glucocorticoid resistance is dependent upon Interleukin-1 (IL-1)—mice lacking IL-1 receptor type 1 do not display these phenotypes (Engler et al., 2008). Interestingly, Avitsur et al. (2001) observed individual differences in macrophage glucocorticoid resistance based upon level of social subordination. Submissive mice were more likely to develop splenocyte corticosterone insensitivity following SDR than were control or dominant mice. Glucocorticoid resistance correlated negatively with time spent in social exploration and positively with time spent in submissive postures. Level of social exploration prior to SDR exposure was predictive of submissive behavior during the first session of SDR, suggesting that pre-existing differences in mouse behavior may predict response to SDR. Collectively, these results imply that the adaptive mechanism by which corticosterone represses the immune system in response to stress is compromised in susceptible (submissive) mice but maintained in resilient (dominant) mice. Further study is required to determine whether active molecular and cellular mechanisms maintain glucocorticoid sensitivity in resilient mice following SDR exposure and, similar to subordinate behavior, whether baseline differences in these mechanisms can predict ultimate behavioral response. As glucocorticoid resistance is a hallmark symptom of depression, further understanding of immune cell resilience to glucocorticoid insensitivity may prove particularly advantageous for therapeutics.

Recent findings by Hodes et al. (in press) suggest that pre-existing differences in IL-6 signaling from leukocytes also predict behavioral response to CSDS. Prior to any stress, leukocytes from peripheral blood of mice that ultimately became susceptible to CSDS released more IL-6 in response to *ex vivo* immune challenge with LPS. Pro-susceptible mice had higher numbers of circulating leukocytes, and leukocyte number and IL-6 release correlated negatively with social interaction ratio, indicating a predictive relationship. Increased leukocyte number is likely driven by an increase in blood CD11b+ monocytes, as significant differences in proportions of leukocyte subtypes between susceptible and resilient mice were observed only in monocytes. Generation of chimeric mice via transplantation of bone marrow hematopoietic progenitor cells from a susceptible donor produced a robust social avoidance phenotype compared to control bone marrow chimeras. In contrast, chimeras generated via transplantation of progenitors from an IL-6^{-/-} donor demonstrated behavioral resilience to CSDS, behaving similarly to IL-6^{-/-} mice and mice treated with an IL-6 antibody, which binds and neutralizes IL-6 in the peripheral circulation. These findings suggest that peripherally derived IL-6 drives susceptibility to CSDS, and that susceptible and resilient mice display baseline differences in leukocyte number and responsiveness. The mechanisms contributing to pre-existing differences in stress responsive IL-6 release and circulating leukocyte number are under investigation and will inform our understanding of immune regulation in resilience. Experiments investigating whether peripheral blood leukocytes of mice susceptible to CSDS, like splenic leukocytes in mice exposed to SDR, display glucocorticoid resistance may prove particularly fruitful.

As mentioned above, increased levels of CD11b+ monocytes in blood and spleen are both a risk factor for susceptibility to CSDS and a consequence of RSD in mice. A likely mechanism underlying this increased number of monocytes/macrophages is direct sympathetic nervous system innervation of bone marrow and control of bone marrow hematopoiesis via β -adrenergic signaling. Two recent studies propose that stress promotes proliferation and egress of immature, pro-inflammatory myeloid cells from the bone marrow.

Powell et al. (Powell et al., 2013) reported that in mice subjected to RSD, stress induces a transcriptional pattern that ultimately leads to myelopoiesis favoring immature, proinflammatory monocytes and granulocytes that express high and intermediate levels, respectively, of the surface marker Ly6c. RSD results in a 4-fold higher prevalence of monocytes in blood and spleen as well as a 50–70% increase in monocytic and granulocytic bone marrow progenitor cells. Post-RSD genome-wide analysis of the peripheral blood mononuclear cell transcriptome revealed a transcriptional mechanism underlying this phenomenon—enhanced expression of proinflammatory genes and genes related to myeloid cell lineage commitment accompanied by decreased expression of genes related to terminal myeloid differentiation. This transcriptional pattern was dependent upon β -adrenergic signaling as administration of propranolol, a β -adrenergic antagonist, prevented proinflammatory transcription in peripheral blood monocytes as well as the RSD-induced increase in Ly6c^{high} monocytes. A transcriptional profile favoring pro-inflammatory monocytes and β -adrenergic signaling was also identified in human subjects of low socioeconomic status, a form of chronic social stress. Further, Heidt et al. (2014) found that chronic variable stress increases numbers of monocytes and neutrophils in mouse blood and bone marrow due to proliferation of leukocyte progenitors. Stress-enhanced hematopoietic activity was accompanied by increased bone marrow noradrenaline levels and decreased expression of CXCL12, a negative regulator of hematopoietic stem and progenitor cell (HSPC) proliferation and migration that is in turn regulated by the β_3 -adrenergic receptor. Treatment of stressed mice with a β_3 -adrenergic receptor antagonist increased CXCL12 expression, reduced HSPC proliferation and attenuated the stress-induced increase in circulating neutrophils and Ly6c^{high} monocytes. Together, these studies provide compelling evidence in both humans and mice linking stress vulnerability to sympathetic nervous system mediated leukocytosis. Potentially informative future studies include an investigation of leukocyte population shifts and transcriptional profiles in blood and bone marrow of stress resilient subjects.

Many of the peripheral findings we've discussed focus primarily on stress susceptible animals and suggest immune mechanisms of passive resilience—resilient and control animals lack peripheral markers that are present and detrimental in susceptible animals. However, as research in the field shifts to focus more on pre-existing individual differences in inflammation as a proxy for vulnerability and resilience to depression and anxiety, we anticipate elucidation of active immune mechanisms of resilience, an exciting prospect due to the relative feasibility of therapeutically targeting peripheral systems with monoclonal antibodies, thus reducing off-target effects in the central nervous system.

4.2. Central immune regulation of physiology and behavior

Peripheral cytokine signals reach the central nervous system via two main pathways—stimulation of the vagal nerves and brain-stem nuclei (the neural pathway) and crossing of the blood–brain barrier (the humoral pathway, see Fig. 1) (Dantzer et al., 2008; Wohleb et al., 2013; Pavlov and Tracey, 2012; Quan, 2008). Centrally derived cytokine signals are produced by microglia, resident brain macrophages. Within the brain, inflammatory signals can influence behavior through mechanisms including activation of the HPA axis and glucocorticoid-induced neuronal atrophy (Iwata et al., 2013) as well as excitatory synaptic plasticity (see Fig. 2) (Christoffel et al., 2011a; Boersma et al., 2011).

Numerous studies investigating central stress-induced inflammatory processes have revealed a prominent role for IL-1 β . Iwata et al. (2013) propose that stress induces IL-1 β release in the brain primarily via the microglial NLRP3 inflammasome. NLRP3 is a

cytosolic pattern recognition receptor (PRR) that, when stimulated by toll-like receptor 4 (TLR4) activation or ATP, both of which are regulated by stress, binds to pro-caspase-1, forming the inflammasome complex. Pro-caspase-1 is cleaved and in turn cleaves pro-IL-1 β into IL-1 β , which is then released from the cell. Microglia constitutively express the components of the NLRP3 inflammasome, and acute restraint stress activates the NLRP3 inflammasome in the hippocampus, the brain region containing the highest concentration of microglia and IL-1 β receptors (Iwata et al., 2013; Farrar et al., 1987). Intracerebroventricular (i.c.v) administration of IL-1 results in increased anxiety-like behavior in the elevated plus maze and open field as well as spatial memory deficits in the Morris water maze (Song et al., 2006). In contrast, pharmacologic or genetic inhibition of Interleukin-1 Receptor 1 (IL-1R1) blocks anhedonia in rats exposed to CUS (Koo and Duman, 2008). Interestingly, i.c.v administration of an IL-1R1 antagonist prevented

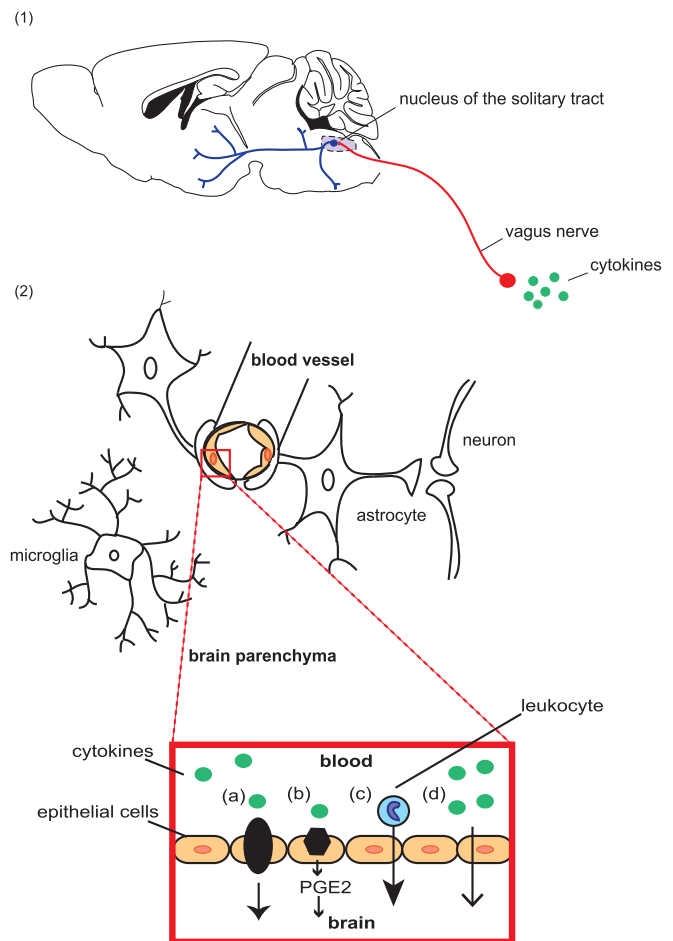


Fig. 1. The interface of the immune and central nervous systems. Peripheral immune cells and signals reach the CNS via two primary routes: the neural pathway and the humoral pathway. (1) In the neural pathway, peripheral cytokines activate primary afferent nerves, including the vagus nerve, which terminates in the nucleus of the solitary tract (NTS) in the medulla oblongata. Second order neurons project to brain-stem, hypothalamic and forebrain nuclei involved in behavioral stress response. (2) In the humoral pathway, cytokine signals cross the blood–brain barrier by transport, diffusion, or trafficking of immune cells. Cytokines in the peripheral circulation cross via cytokine transporters (a) or by activation of cytokine receptors on endothelial cells of the brain vasculature (b), leading to the production of prostaglandins E2 (PGE2), lipophilic molecules that can easily diffuse throughout the brain. Chronic social stress can also promote the recruitment of peripherally derived macrophages into the brain parenchyma of reward regions including the amygdala and prefrontal cortex (c). Cytokines produced locally by macrophage-like cells in the circumventricular organs and choroid plexus cross the BBB by volume diffusion (d). Once within the CNS, peripheral signals activate microglia that in turn produce pro-inflammatory cytokines.

shuttle box escape failure following pretreatment with repeated, inescapable tail shocks (Maier and Watkins, 1995). These results suggest that IL-1 β signaling is an important mediator of behavioral vulnerability and resilience to LH and CUS in rats, and that IL-1 β and its downstream effectors may be promising targets for promoting behavioral resilience to stress.

Downstream mechanisms by which IL-1 β influences behavioral outcomes to stress include HPA axis activation as well as modulation of hippocampal neurogenesis. Stress-induced IL-1 β modulates the HPA axis by stimulating release of CRF from the hypothalamus and subsequent downstream release of ACTH from the pituitary gland (Iwata et al., 2013; Sapolsky et al., 1987; Berkenbosch et al., 1987). Blockade of IL-1R1 via antagonist administration or null mutation prevents CUS-induced reductions in cells positive for BrdU (Bromodeoxyuridine, a marker of cell division) and DCX (doublecortin, a marker of immature neurons), indicating that chronic stress inhibits neurogenesis in an IL-1 β dependent fashion (Koo and Duman, 2008). In the same study, *in vitro* incubation with IL-1 β decreased the proliferation of adult hippocampal progenitor cells, an effect blocked by co-incubation with inhibitors of NF κ B signaling. As the I κ K–NF κ B signaling pathway is activated by IL-1 β and other pro-inflammatory cytokines, it is a promising candidate mediator of the downstream effects of IL-1 β . A follow-up study revealed that, indeed, exposure to an acute stressor activated NF κ B signaling in neural stem-like cells (NSCs), and NF κ B activation in NSCs was dependent upon IL-1 β signaling (Koo et al., 2010). Moreover, *i.c.v.* administration of an NF κ B inhibitor throughout CUS blocked the subsequent stress-induced decrease in BrdU⁺DCX⁺ cells as well as the expression of anxiety-like and anhedonic behaviors.

Our lab has reported I κ K–NF κ B dependent structural and functional plasticity in the mesolimbic dopaminergic reward pathway following CSDS (Christoffel et al., 2011a). I κ K and its downstream targets I κ B and phosphorylated I κ B were upregulated in the NAc of susceptible mice following CSDS. Interestingly, activation of I κ K–NF κ B signaling promotes susceptibility to CSDS by altering plasticity of glutamatergic synapses in the NAc. Strategies to blunt I κ K–NF κ B activation directly in NAc promote resilience. A subsequent study revealed that constitutive viral overexpression of I κ K promotes baseline anxiety and depression-

like behaviors in the open field and forced swim tests as well as social avoidance and anhedonic behavior in response to an acute social defeat stress (Christoffel et al., 2012). I κ K expression induced the formation of immature spines (primarily thin spines) in mice exposed to acute social defeat stress. Again, spine density correlated significantly with social avoidance behavior, suggesting that I κ K-dependent, stress-induced morphological changes may drive behavioral response to stress. Together, these data suggest a critical role for I κ K–NF κ B signaling in NAc in susceptibility vs. resilience to social stress. Future studies will be important to identify the upstream inflammatory signaling pathways responsible for such effects.

5. Molecular and cellular mechanisms of resilience in the central nervous system

Much of our current knowledge regarding central mechanisms of resilience centers on mesocorticolimbic reward circuitry. Brain reward circuitry serves the adaptive purpose of focusing one's attention on the acquisition of natural rewards to ultimately ensure survival (Russo and Nestler, 2013). Mesocorticolimbic circuitry comprises neurons from the medial prefrontal cortex (mPFC), hippocampus, NAc, amygdala, VTA, lateral hypothalamus, and lateral habenula, among other brain regions (see Fig. 3). Collectively, these brain regions are involved in numerous psychological and cognitive processes that are impacted by stress and compromised in patients with depression or anxiety (Christoffel et al., 2011b). Connections between mesocorticolimbic regions are dense and often complex. Here, we will focus primarily on the most well characterized connections, those of the VTA–NAc reward circuit. Dopaminergic neurons of the VTA project onto GABAergic medium spiny neurons (MSNs) of the NAc, a structure within the ventral striatum. VTA neurons release dopamine in response to reward-related stimuli to initiate consumption and sometimes also in response to aversive stimuli. The NAc sends reciprocal connections back to the VTA via two pathways—the direct pathway, via D1-type MSNs; and the indirect pathway, via D2-type MSNs, which innervate GABAergic interneurons in the ventral pallidum that in turn synapse onto VTA neurons. Although findings are variable, human imaging studies support a role for

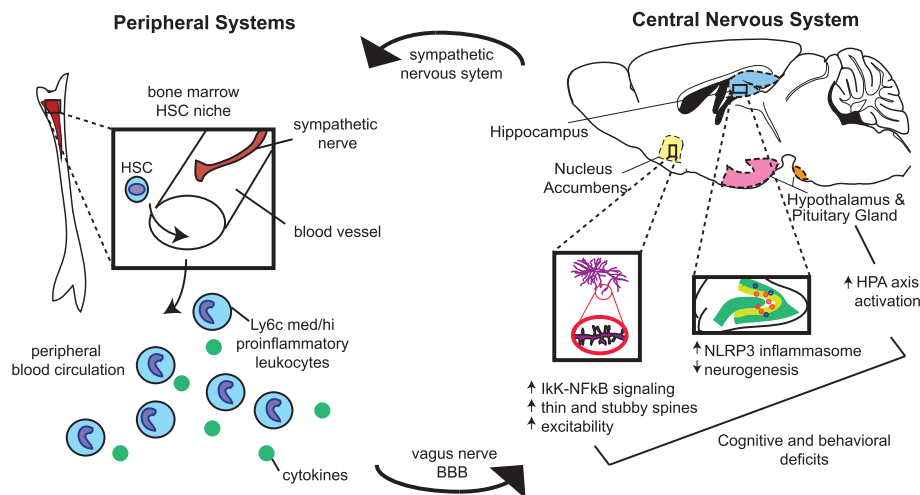


Fig. 2. Overview of immune activation in the peripheral and central nervous systems by stress. Chronic stress activates the sympathetic nervous system, precipitating a series of immune events in peripheral systems. Sympathetic nerves release noradrenaline in the bone marrow hematopoietic stem cell (HSC) niche, stimulating myelopoiesis favoring the release of immature, proinflammatory monocytes and neutrophils into the peripheral circulation. Cytokine signals derived from these leukocytes and, in some cases, the cells themselves, traffic to the brain by activating afferent nerves or by crossing the blood–brain barrier. Within the CNS, they produce lasting changes relevant to depression and anxiety-like behavior by activating the HPA axis, the microglial NLRP3 inflammasome in the hippocampus, and NF κ B–I κ K signaling in the ventral striatum.

the VTA–NAc pathway in human depression, demonstrating consistent reductions in NAc activity in adult patients as well as NAc volume reductions in elderly patient populations (Husain et al., 1991; Krishnan et al., 1992; Drevets et al., 1992; Mayberg et al., 2000). Deep brain stimulation procedures targeting the NAc and its efferent connections in the VTA have shown good therapeutic efficacy in treatment resistant depression (T. Schlaepfer, personal communication). However, it is currently unknown how these stimulation protocols affect NAc microcircuitry and whether they indirectly stimulate fibers of passages that synapse outside the NAc.

5.1. Epigenetic and transcriptional mechanisms

Numerous epigenetic and transcriptional mechanisms in mesocorticolimbic reward circuitry underlie antidepressant action and resilient behavioral responses to chronic stress. The transcription factor Δ FosB is upregulated in the NAc of resilient mice following CSDS in a serum response factor (SRF) dependent manner, and genetic overexpression or antagonism of Δ FosB expression promotes behavioral resilience or susceptibility, respectively (Vialou et al., 2010a,b). Furthermore, Δ FosB levels are reduced in post-mortem NAc samples of human depressed patients. Chronic

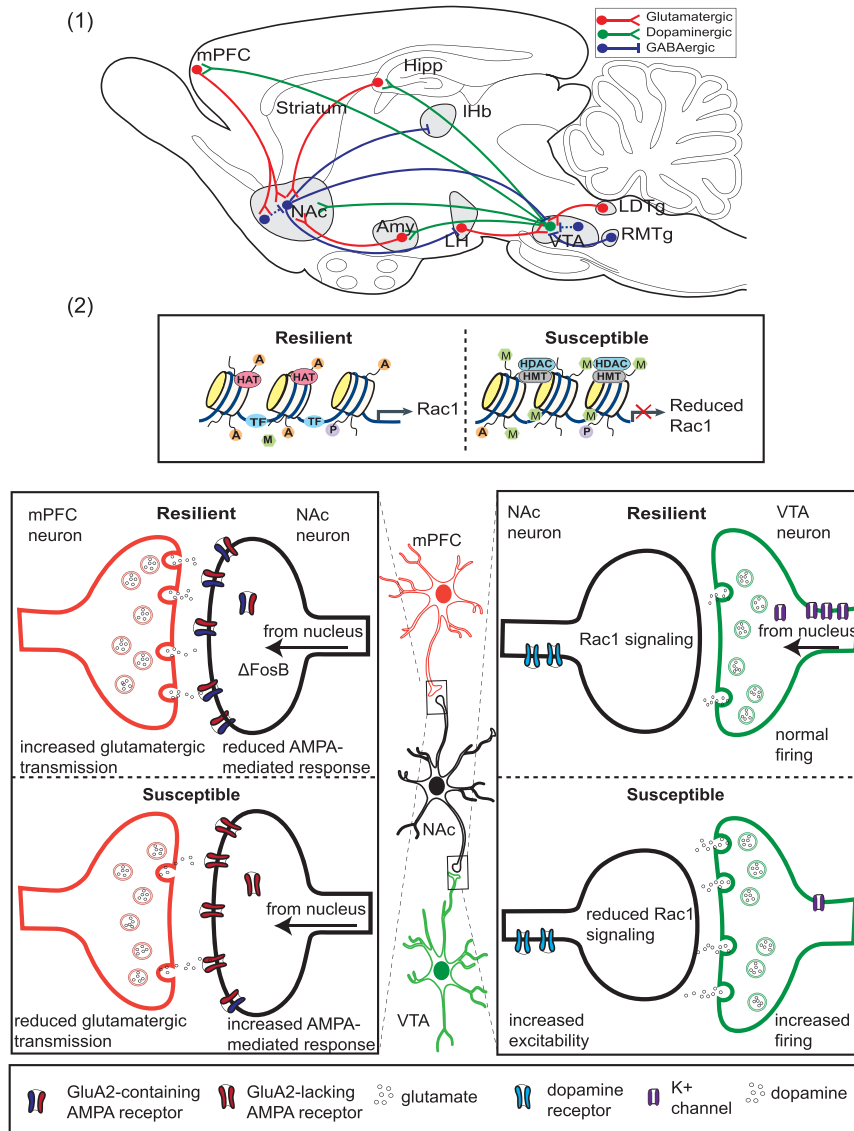


Fig. 3. Molecular and cellular mechanisms of resilience in the mesocorticolimbic reward circuitry. (1) A simplified schematic of the VTA–NAc circuit, showing the major glutamatergic, GABAergic, and dopaminergic projections to and from the ventral tegmental area (VTA) and nucleus accumbens (NAc) in the rodent. Dashed lines indicate internal, inhibitory connections mediated by interneurons. Hipp, hippocampus; mPFC, medial prefrontal cortex; Amy, amygdala; LH, lateral habenula, RTMg, rostromedial tegmentum; LDTg, lateral dorsal tegmentum. (2) Shown are examples of processes that contribute to resilience vs. susceptibility to chronic social defeat stress. Within the NAc, resilient mice lack the reduced Rac1 gene transcription characteristic of susceptible animals and attributed to reduced histone pan-acetylation (specified by A) and enhanced lysine 27 methylation (specified by M). In susceptible animals, reduced expression of Rac1 leads to actin cytoskeletal reorganization, and an increase in stubby, immature spines. Also in NAc neurons, Δ FosB activity promotes resilience by increasing expression of GluA2. Although shown here for simplicity in the synapse, Δ FosB is a transcription factor, and its activity occurs in the nucleus. GluA2-containing AMPA glutamate receptors are Ca²⁺ impermeable, reducing AMPA-mediated glutamate response and excitability. In the mPFC, increased glutamatergic transmission has been associated with resilience, and optogenetic stimulation of ChR2 in the mPFC has been shown to promote resilience. In the VTA, resilient animals homeostatically maintain normal firing rate through the induction of K⁺ channels. HAT, histone acetyltransferase; HDAC, histone deacetylase; HMT, histone methyltransferase; P, phosphorylation; TF, transcription factor.

fluoxetine treatment enhances Δ FosB concentration in the mouse NAc, and Δ FosB is required for fluoxetine-mediated antidepressant effects in susceptible mice. Δ FosB exerts its pro-resiliency effects through its transcriptional targets, including AMPA glutamate receptor subunit GluA2 and Sparc-like 1 (SC1). Following CSDS, resilient mice show greater NAc expression of GluA2 than do control or susceptible mice, an effect mediated by Δ FosB binding to the *GluA2* promoter. Δ FosB-mediated enhanced GluA2 expression promotes resilience by decreasing AMPA function—GluA2-containing AMPA receptors are Ca^{2+} impermeable with lower receptor conductance and reduced inwardly rectifying currents. In addition, SC1, a protein localized to the PSD and necessary for proper synapse assembly, is upregulated both in mice overexpressing Δ FosB and in mice resilient to CSDS. SC1 overexpression reverses social avoidance behavior following CSDS.

Epigenetic regulation of ras-related C3 botulinum toxin substrate 1 (*Rac1*) has been shown by our laboratory to mediate susceptibility vs. resilience to CSDS (Golden et al., 2013). *Rac1* is a Rho GTPase involved in the organization and maintenance of the actin cytoskeleton, largely through regulation of its downstream target cofilin, an actin severing protein critically involved in synaptic plasticity. Following CSDS, *Rac1* was downregulated in the NAc of susceptible, but not resilient, mice, and its expression correlated with social avoidance behavior. Viral-mediated overexpression and knockdown experiments demonstrated that *Rac1* is necessary and sufficient for the expression of resilient behavior following CSDS. ChIP analysis of permissive H3 acetylation (acH3) and repressive histone H3 Lys9 trimethylation (H3K9me3) at the *Rac1* promoter and its upstream regions revealed that, in both susceptible and resilient mice, *Rac1* expression was mediated by epigenetic mechanisms. Susceptible, but not resilient, mice exhibited reduced permissive acetylation at the *Rac1* promoter and its 2000-bp upstream region. Resilient mice showed reduced methylation within the *Rac1* promoter whereas susceptible mice showed enhanced methylation in the 1000-bp upstream region. Chronic intra-NAc administration of an HDAC inhibitor reversed social avoidance behavior and rescued *Rac1* expression in susceptible mice. Collectively, these results suggest that epigenetic mechanisms maintain *Rac1* expression in resilient mice, promoting adaptive behavioral response to stress, but have the opposite effect in susceptible mice. Analysis of human postmortem NAc tissue samples from depressed patients corroborated these animal findings. *Rac1* expression was strongly reduced in unmedicated patients compared to controls, and depressed patients showed decreased acetylation in regions ~200-bp upstream and downstream of the transcription start site (TSS) accompanied by increased methylation in the gene region ~200-bp upstream of the TSS. *Rac1* likely promotes resilient responses to CSDS via its effects on MSN spine structure. Viral-mediated overexpression of *Rac1* reduced the CSDS-induced enhancement in dendritic stubby (immature) spine density whereas Cre-mediated *Rac1* genetic deletion had the opposite effect.

5.2. Neurophysiology of susceptibility and resilience in the VTA

A robust neurophysiological correlate of susceptibility to CSDS is the enhanced excitability of VTA dopamine neurons following stress (Krishnan et al., 2008; Cao et al., 2010). CSDS increases the spontaneous firing rate of VTA dopamine neurons and the percentage of neurons demonstrating burst firing events in susceptible, but not resilient, mice (Cao et al., 2010). These physiological changes correlate inversely with social interaction score and can be reversed with chronic antidepressant treatment, suggesting an involvement of stress-induced changes to neuronal excitability in depression-like behavior. One mechanism underlying enhanced excitability in susceptible mice is the I_h (hyperpolarization-

activated cation) current (Cao et al., 2010). The I_h current regulates tonic firing of dopamine neurons as well as the transition from single-spike to burst firing, and is robustly increased only in susceptible mice following CSDS. I_h inhibitor infusion reverses social avoidance behavior, and chronic antidepressant treatment reduces the stress-induced increase in I_h current. Enhanced neuronal excitability is also mediated by reduced activation of AKT (thymoma-viral proto-oncogene) in the VTA, which likely produces a subsequent reduction in inhibitory tone (Krishnan et al., 2008). Phosphorylated AKT is reduced in the VTA of susceptible mice following CSDS, and this reduction is necessary and sufficient to produce social avoidance behavior. AKT targets the β_2 subunit of the GABA_A receptor, and *in vitro* pharmacological reduction of AKT reduces the amplitude and frequency of spontaneous GABA-mediated inhibitory postsynaptic currents (IPSCs) onto VTA dopaminergic neurons. Future *in vivo* studies are needed to causally link AKT-GABA changes to social avoidance behavior. Recently, Chaudhury et al. (2013) demonstrated that the CSDS-induced high frequency phasic firing in dopamine neurons of the VTA–NAc pathway is sufficient to functionally drive susceptible behavior. Optogenetic induction of phasic, but not tonic, firing in tyrosine hydroxylase positive (TH+) VTA neurons during or after exposure to subthreshold defeat rapidly produced robust social avoidance and anhedonia behaviors. Induction of phasic firing during the social interaction test following 10 days of CSDS was sufficient to reverse behavior in mice previously identified as resilient, generating social avoidance, and to produce long-lasting changes in excitability, as evidenced by maintenance of depression-like behavior (decreased sucrose preference) 8–12 h post-stimulation. These effects were VTA–NAc pathway specific, as selective optogenetic stimulation of VTA TH+ neurons projecting to the PFC did not induce social avoidance or anhedonia. Halorhodopsin inhibition of VTA firing reversed depression-like behavior in susceptible mice following CSDS exposure. These experiments demonstrate that stress-induced phasic firing in NAc-projecting VTA dopamine neurons is necessary and sufficient for the development of depression-like behavior.

Normal dopamine neuron firing rate, AKT activation and signaling, and I_h current dynamics are allostatically preserved in resilient mice during and after stress exposure, although the mechanisms underlying this allostasis are less understood than those driving susceptibility. A recent study by Friedman et al. (2014) identified an active mechanism by which normal dopamine neuron firing is maintained in resilient mice. Surprisingly, VTA dopamine neurons of resilient animals do not show a return to a normal I_h current comparable to that of controls following CSDS. Instead, they exhibit an I_h current increase that is much larger than that of susceptible mice. Underlying this phenomenon is a homeostatic enhancement in multiple K^+ channel currents—the potentiated I_h current augments neuronal firing to such an extent that K^+ channels are activated, returning firing rate to a normal level. Indeed, current injection in dopamine neurons of resilient mice produces a reduction in spike number, whereas current injection produces the opposite effect in susceptible mice. Repeated intra-VTA infusion of lamotrigine, an I_h potentiator, or VTA viral-mediated overexpression of hyperpolarization-activated and cyclic nucleotide-gated channel 2 (HCN2), a channel that regulates I_h current, reversed social avoidance and anhedonic behavior in susceptible mice. Both manipulations increased I_h and K^+ currents, and reduced neuronal excitability. Further, repeated optogenetic induction of hyperactivity in VTA dopamine neurons increased K^+ currents and reversed social avoidance behavior. This type of homeostatic plasticity was specific to VTA–NAc projecting neurons. These findings provide exciting insight into the biology of resilience as well as a potential therapeutic avenue.

5.3. Glutamatergic synapses in resilience and reward

In addition to dopaminergic innervation from the VTA, the NAC also receives glutamatergic innervation from the PFC, amygdala, thalamus and hippocampus. Decreased PFC activity, as measured by cerebral blood flow and glucose metabolism, is the most robust finding reported by human imaging studies of depressed patients (Mayberg, 2009). Findings from rodent models are generally consistent with those in humans and suggest that stress leads to hypofrontal function. First, chronic stress leads to significant atrophy and synapse loss on glutamatergic neurons in the PFC (Christoffel et al., 2011b; McEwen and Morrison, 2013; Duman and Li, 2012). Importantly, loss of synapses has also been observed in the PFC of humans with MDD (Kang et al., 2012). Covington et al. (2010) reported decreased expression of the immediate early genes (IEGs) *zif268* (also termed *egr1*) and *arc* in human postmortem prefrontal cortical tissue of unmedicated depressed patients. IEG expression was also reduced in the ventromedial PFC of susceptible mice, but was unchanged in resilient mice following CSDS. As IEG expression is considered a representation of brain activity, these results suggest that activity is reduced in susceptible mice and depressed patients, but maintained in resilient mice. Optogenetic stimulation of the mPFC of susceptible mice had an antidepressant effect, reversing social avoidance and anhedonic behavior, and indicating that burst firing in mPFC neurons promotes behavioral resilience. Optogenetic induction of burst firing also increased expression of the IEG *c-fos*. The NAC is another region of brain reward circuitry that undergoes significant stress-induced remodeling of glutamatergic synapses. Following CSDS, susceptible, but not resilient, mice have an increased density of glutamatergic synapses on NAC MSNs, which correlates with increased mini excitatory postsynaptic potential (mEPSP) frequency (indicative of more functional glutamatergic synapses or altered presynaptic release). Data from our lab using circuit specific optogenetic tools to stimulate glutamatergic neurons terminating in the ventral striatum (vStr), find that glutamatergic projections from the intralaminar thalamus (ILT) promote susceptibility to CSDS whereas stimulation of projections from the PFC exert opposite effects (Christoffel, D.J. et al., *Soc. Neurosci. Abstr.* 705.08, 2013). Both chronic, viral-mediated expression in the ILT of tethered toxins (tToxins, designed to inhibit excitatory transmission by selectively blocking calcium influx at the pre-synaptic voltage gated Ca^{2+} channels Cav2.1 and Cav2.2) and rapid optogenetic inhibition of ILT–vStr terminal projections prevented social avoidance and reduced MSN stubby spine density (a parameter that is known to positively correlate with social avoidance). Accordingly, rapid terminal stimulation of ILT projections to the vStr was pro-susceptible, inducing social avoidance in a subthreshold defeat. Together, these findings suggest that ILT–vStr projections are necessary and sufficient for the expression of social avoidance. In line with previous work on the role of PFC activity in depression-like behavior, chronic tToxin-mediated inhibition of PFC projections was pro-susceptible. Surprisingly, the more specific, rapid optogenetic inhibition of PFC–vStr glutamatergic terminals failed to induce social avoidance. This indicates that PFC-mediated resilience may require sustained activation of PFC–NAC terminals or the activity of other PFC terminal projections outside the striatum. While the PFC may provide a promising target for promoting resilience to stress, further research is needed to fully elucidate (1) the particular anatomical and physiological parameters of pro-resilient PFC activity, and (2) whether allostatic mechanisms maintain normal PFC–vStr firing patterns in resilient animals to prevent pathological changes in reward circuit activity.

6. Conclusion

With the exception of the rapidly acting antidepressant ketamine and the advent of deep brain stimulation paradigms to treat depression, both of which are limited to severe, treatment resistant cases of depression, there has been a decades long void of new treatment options for depression and anxiety. However, the future of treatment and research is hardly dire. Modern research on stress-related disorders has yielded numerous potential targets and biomarkers for diagnosis and treatment, largely due to an enhanced focus on alternatives to monoamine-based mechanisms, such as epigenetic mechanisms, immune-related factors, sex, and the biology of resilience. Stress-related disorders, and resilience to them, can be considered products of the coordinated activity of the brain and numerous bodily systems. The results of resilience research we've described here are particularly exciting as they offer an opportunity for personalized science and medicine. We've described potential targets and biomarkers specific to type of stress (developmental vs. adulthood), sex, and inflammatory state. As women are more likely to suffer from mood disorders, the continuing identification of sex-based, pro-resilience markers may enable the development of more effective, sex specific treatments. The NIH-mandated inclusion of female subjects in research studies will hopefully encourage further elucidation of sex-based resilience. We feel that immune mechanisms are particularly promising as many potential targets are peripheral, removing the blood–brain barrier as a therapeutic obstacle. Preclinical experiments in our lab indicate that peripherally targeting IL-6 with monoclonal antibodies is antidepressant in mice (Hodes, G.E. et al., *Soc. Neurosci. Abstr.* 542.10, 2013). Further research on active mechanisms that maintain normal immune function in resilient individuals, particularly the maintenance of glucocorticoid sensitivity and normal leukocyte differentiation and release from bone marrow, may yield additional therapeutic options for peripheral administration. An enhanced focus throughout the field on individual differences in response to stress and inclusion of resilient animals as research subjects is necessary, particularly in regard to studies of the immune system, where study of stress-resilient subjects has been minimal. Further interrogation of the mechanisms of what we've termed “passive” resilience will also be helpful. As described in this review, the adaptive failure of resilient animals to display the pathological markers seen in susceptible animals is often accomplished by active mechanisms. An enhanced focus on resilient subjects may enable us to harness mechanisms of resilience in the body and brain for the successful treatment of stress-related disorders.

Acknowledgments

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