Novelty-induced arousal enhances memory for cued classical fear conditioning: Interactions between peripheral adrenergic and brainstem glutamatergic systems

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Exposure to novel contexts produce heightened states of arousal and biochemical changes in the brain to consolidate memory. However, processes permitting simple exposure to unfamiliar contexts to elevate sympathetic output and to improve memory are poorly understood. This shortcoming was addressed by examining how novelty-induced changes in peripheral and/or central arousal modulates memory for Pavlovian fear conditioning. Male rats were either exposed to the conditioning chamber for 5-min or given no exposure 24 h before conditioning with five tone–shock (0.35 mA) pairings. Retention was assessed 48 h later in a different context. Non-pre-exposed animals exhibited significantly greater freezing during conditioned stimulus (CS) presentations than did pre-exposed animals (P < 0.05). The improvement in retention produced by novelty was attenuated by pretreating a blockade of peripheral β-adrenergic receptors with sotalol (6 mg/kg, i.p.). Study 2 revealed that novelty-induced increases in peripheral autonomic output are conveyed to the brain by visceral afferents that synapse upon brainstem neurons in the nucleus tractus solitarius (NTS). Blocking AMPA receptor activity in the NTS with CNQX (1.0 μg) significantly reduced freezing to the CS in non-pre-exposed animals (P < 0.01). Study 3 showed that elevating epinephrine levels in habituated animals influences learning through mechanisms similar to those produced by novelty-induced arousal. Pre-exposed animals given epinephrine (0.1 mg/kg) froze significantly more than saline controls (P < 0.01), and this effect was attenuated by intra-NTS infusion of CNQX. The findings demonstrate that novelty-induced arousal or increasing sympathetic activity with epinephrine in pre-exposed animals enhances memory through adrenergic mechanisms initiated in the periphery and transmitted centrally via the vagus/NTS complex.
forebrain and limbic structures display phasic bursts of activity upon initial exposure to a novel environment, but increased discharge does not occur in rats returned to a familiar context (Vankov et al. 1995). Other findings, reporting that norepinephrine concentrations in the frontal cortex and hypothalamus are significantly elevated following exposure to a novel illuminated environment or training context containing an unfamiliar rat (McQuade et al. 1999), provide more direct evidence that norepinephrine mediates central changes in response to novelty exposure. These collective findings demonstrate that novelty induced by subtle exposure to an unfamiliar context influences a number of neurochemical and synaptic changes that are required for new experiences to be encoded effectively into long-term memory.

The consequences of brief exposure to unfamiliar environments are not limited to the well-documented biochemical changes observed in the brain. Autonomic indices of sympathetic activity, including skin conductance, cardiac output, and circulating concentrations of the adrenal hormones corticosterone and epinephrine, are all elevated by presenting humans or animals with novel stimuli or after allowing free exploration in an unfamiliar environment (De Boer et al. 1990; Bradley et al. 1993; Handa et al. 1994; Gerra et al. 1996; Codispoti et al. 2006). These findings reveal important parallels between the class of physiological changes that emerge as a direct result of exposure to novel environment and those elicited by emotionally arousing events. Although both conditions induce changes that modulate peripheral visceral activity and brain limbic output to encode new events into memory, the mechanism by which novelty-induced peripheral and/or central arousal may influence memory formation is not completely understood.

Several lines of evidence suggest that the arousal-related hormone epinephrine plays complementary roles in both processes. For example, systemic injection of epinephrine in a range of doses that improve memory in laboratory rats (Williams and McGaugh 1993; Clayton and Williams 2000; Nordby et al. 2006; Dornelles et al. 2007) increases the firing rate of noradrenergic LC neurons (Holdefer and Jensen 1987) that display high levels of discharge following exposure to novel contexts (Vankov et al. 1995). As with novelty, epinephrine administration facilitates LTP (Korol and Gold 2008) and reverses deficits in retention for contextual fear conditioning displayed by mice with the transcription factor CREB genetically disrupted (Frankland et al. 2004). Presentation of novel visual slides to humans improves memory (Fonlupt et al. 2004) and initiates epinephrine secretion from the adrenals (Gerra et al. 1996), and this change in arousal is sufficient to improve later retention performance (Cahill et al. 1994) comparable to that produced by direct administration of this hormone (Cahill and Alkire 2003). The arousal-induced enhancement in human memory with novel visual slides (Strange and Dolan 2004) and the novelty-induced facilitation of LTP discussed above (Li et al. 2003; Straube et al. 2003a,b) are both attenuated by blocking noradrenergic receptor transmission with the β-adrenergic receptor antagonist propranolol. These types of findings provide the foundation for determining whether novelty-induced arousal and the subsequent physiological changes that assist in encoding features of new experiences into memory are mediated through interactions involving peripheral hormonal systems that influence noradrenergic activity in the brain.

If brief periods of novelty exposure induce arousal through this mechanism, then it is plausible that one means by which arousal impacts the strength that emotional episodes are stored into memory is by activating neural pathways that transmit the sympathomimetic actions of epinephrine mediated in the peripheral to brain systems that influence norepinephrine output in the CNS. Peripheral branches of the vagus serve a key role in this process as ascending fibers of the vagus are densely embedded with β-adrenergic receptors that bind epinephrine (Schreurs et al. 1986; Lawrance et al. 1995), and peripheral endings of the vagus innervate sensory organs that are highly responsive to sympathetic arousal produced by epinephrine release or novelty, including the heart, liver, stomach, and lungs (Shapiro and Miselis 1985; Coupland et al. 1989; Paton 1998a, b). Moreover, electrical stimulation of ascending vagal fibers produce significant burst firing in LC neurons (Groves et al. 2005; Dorr and Debonnel 2006) and lead to long-lasting elevations in norepinephrine concentrations collected from the amygdala (Hassett et al. 2004) and hippocampus (Miyashita and Williams 2002).

Information regarding heightened activity in peripheral sensory organs is transmitted by ascending vagal fibers to a specific cluster of cells in the brainstem known as the nucleus of the solitary tract (NTS) (Kalida and Sullivan 1982; Sumal et al. 1983). In response to these changes, NTS neurons influence central noradrenergic activity through direct synapses on LC neurons (Van Bockstaele et al. 1999) that not only become active in the presence of novel stimuli (Vankov et al. 1995) but also modulate norepinephrine release in structures that play important roles in encoding new experiences into long-term memory such as medial prefrontal cortex, hippocampus, and amygdala (Ricardo and Koh 1978; Loughlin et al. 1986; Florin-Lechner et al. 1996).

If novelty-induced arousal increases epinephrine secretion, then it is plausible that one means by which arousal impacts the strength in which emotional episodes are stored into long-term memory is by activating this vagal/NTS pathway. The present study tests this hypothesis by using the “familiarity” versus “novelty” of the training context as a manipulation to increase physiological arousal prior to learning and examine whether storage of emotionally laden memories are influenced by peripheral adrenergic activation. Pavlovian fear conditioning is frequently used to understand the neural circuits involved in forming memories for emotionally arousing experiences (Kim and Jung 2006), although the effects of manipulating physiological arousal during the formation of fear conditioned memory has not been widely explored.

Given this shortcoming, these studies examined how changes in peripheral physiological activity are transmitted by the vagus/NTS complex to reveal the mechanisms by which novelty-induced arousal influences memory for fear conditioning. The objective of experiment 1 was to assess the contribution of peripheral adrenergic activity in mediating novelty-induced arousal and its subsequent effects on mnemonic processing. In this study, novelty was induced in separate groups by withholding habituation and waiting until the day of conditioning to introduce subjects to the training context for the first time. The consequences of blocking peripheral adrenergic receptors prior to Pavlovian conditioning was examined in groups in which the training context represented novelty exposure and compared to groups that were familiarized with the fear conditioning chamber through previous habituation. Study 2 examined whether the pathway between peripheral vagal afferents and brainstem nuclei in the NTS mediate the mnemonic consequences of novelty-induced increases in sympathetic activity during fear conditioning. The amino acid glutamate is the primary transmitter mediating synaptic communication between vagal afferents and NTS neurons since vagal terminals contain glutamate (Sykes et al. 1997) and glutamate receptors are localized on NTS dendrites (Aicher et al. 1999, 2002). In addition, intra-NTS infusion of the AMPA glutamatergic receptor antagonists CNQX (6-cyano-7-nitroquinoxaline-2,3-dione) suppresses excitatory burst firing in NTS neurons activated by stimulating the vagus nerve (Granata and Reis 1983a; Andresen and Yang 1990) with a range of currents that increase LC discharge (Groves et al. 2005; Dorr and Debonnel
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2006) or potentiates norepinephrine release in the amygdala or hippocampus (Miyashita and Williams 2002; Hassert et al. 2004). To this end, the AMPA receptor antagonist CNQX was used to block postsynaptic glutamate receptors in the region of the NTS that receives input from vagal terminals. The location of cannulae and injection needle tips aimed at the NTS is depicted in Figure 1.

Study 3 investigated whether the poor memory exhibited by habituated control groups familiar with the training context could be enhanced by increasing peripheral activity after Pavlovian conditioning with systemic injection of epinephrine. This study also determined if glutamatergic transmission between vagal afferents and NTS neurons plays a critical role in mediating the direct changes on memory produced by elevated concentrations of epinephrine. Findings emerging from these studies reveal that arousal induced by environmental novelty or by exogenously amplifying sympathetic activity with epinephrine enhances Pavlovian fear conditioned memory through adrenergic mechanisms initiated in the periphery and transmitted centrally via the vago/NTS complex.

Results

Experiment 1

Fear conditioned training

This study determined if the improvement in memory produced by novelty exposure and subsequent Pavlovian fear conditioning training is mediated by activating peripheral adrenergic systems. It was hypothesized that the secretion of epinephrine would be a necessary component for novelty-induced arousal to improve memory. This hypothesis was examined by using the peripheral β-adrenergic receptor antagonist sotalol to block epinephrine binding to peripheral β-adrenergic receptors in rats exposed to the novel conditioning context.

A two-way factorial ANOVA on the mean percentage of freezing, exhibited to the final presentation of the conditioned stimulus (CS; tone) during acquisition with five CS-unconditioned stimulus (US) pairings, revealed no statistical differences between the treatment groups in their capacity to learn that the CS tone is a reliable predicator of the US footshock and elicits freezing, \( F_{(1,20)} = 1.48, P = \text{NS} \) (pre-exposed/saline 88.38 ± 7.3, pre-exposed/sotalol 90.68 ± 4.0, non-pre-exposed/saline 97.28 ± 2.0, non-pre-exposed/sotalol 84.16 ± 6.9).

Retention test

A two-way ANOVA indicated a significant overall effect of treatment on the mean percentage of freezing exhibited during three presentations of the CS during retention testing in a completely different Pavlovian chamber (\( F_{(1,20)} = 21.26, P < 0.01 \); Fig. 2A). Post-hoc tests revealed that non-pre-exposed animals exhibited significantly more freezing during CS presentations than did the habituated animals that were pre-exposed to the conditioning chamber 24 h prior to training (\( P < 0.05 \)). Additionally, non-pre-exposed animals, administered the peripherally-acting β-adrenergic receptor antagonist sotalol, exhibited significantly less overall freezing during the three CS presentations relative to non-pre-exposed animals given saline injections (\( P < 0.01 \)). Tone-by-tone analysis of freezing with factorial ANOVAs indicated that non-pre-exposed subjects displayed a significantly higher level of freezing to each individual tone presentation relative to all other groups (see Fig. 2B). Thus, the reported arousal associated with placing organisms in a novel context (De Boer et al. 1990; Handa et al. 1994) contributes to enhanced encoding of emotional learning. Moreover, the beneficial consequences of arousal on memory for CS–US pairings are contingent upon activation of peripheral hormonal systems that bind to β-adrenergic receptors.

Experiment 2

Fear conditioned training

The second study examined whether physiological changes induced in the periphery by exposure to a novel environment affect memory for fear conditioning by activating neurons in the NTS. It was predicted that novelty-induced arousal reflected by autonomic activation enhances memory via epinephrine binding to β-adrenergic receptors along ascending fibers of the vagus nerve. Increased transmission along the vagus would in turn excite neurons in the NTS that are innervated by vagal terminals that release glutamate. Given this assumption, blocking AMPA receptor activity associated with glutamate release in the NTS immediately post-conditioning for non-pre-exposed animals should attenuate the memory improvement from novelty-induced arousal. The initial findings from this study indicated no differences between the treatment groups in their capacity to learn the CS–US associations during training. All groups demonstrated comparable levels of freezing to the final CS presentation during conditioning, \( F_{(1,25)} = 0.670, P = \text{ns} \) (pre-exposed/PBS 92.0 ± 5.0, pre-exposed/CNQX 86.1 ± 5.2, non-pre-exposed/PBS 96.0 ± 2.2, non-pre-exposed/CNQX 96.3 ± 1.1).

Retention test

A two-way ANOVA revealed significant overall effects on mean percentage of freezing to the three CSs presented during retention testing, \( F_{(1,25)} = 9.60, P < 0.01 \). As with experiment 1, non-pre-exposed animals given vehicle injections into the NTS froze for a significantly higher percentage of time when the CS was presented compared with pre-exposed controls and pre-exposed animals administered CNQX into the NTS (\( P < 0.01 \); Fig. 3A). Post-hoc results indicated that a bilateral infusion of CNQX into the NTS significantly reduced the high percentage of freezing observed in non-pre-exposed animals to levels that were comparable to that of

![Figure 1](image-url) Location of needle tip placements in the nucleus of the solitary tract (NTS) overlaid onto a representative photomicrograph from animals trained and tested in experiments 2 and 3. (●) Indicates correct cannula and needle tip placements within the caudal NTS; X, incorrect cannula and needle tip placements; 4V, fourth ventricle; ECu, external cuneate nucleus; and Sp5n, spinal trigeminal nucleus.
pre-exposed controls ($P < 0.01$). Figure 3B shows the percentage of freezing during each of the three CS tone presentations. On only the first presentation of the CS, non-pre-exposed saline treated animals froze significantly more than non-pre-exposed CNQX treated animals ($P < 0.02$), but not the pre-exposed groups. The non-pre-exposed group froze significantly more than all groups during the second and third presentations of the CS ($P < 0.01$). These findings demonstrate that novelty-induced enhancement in memory for Pavlovian fear conditioning is attenuated by blocking access to postsynaptic glutamate receptors in the NTS with bilateral infusions of the AMPA receptor antagonist CNQX.

**Experiment 3**

**Fear conditioned training**

The final study examined whether increasing circulating concentrations of epinephrine improves Pavlovian conditioning through mechanisms similar to those produced by novelty-induced arousal. If the two types of manipulations share similar pathways, then any changes in conditioning mediated by epinephrine should be attenuated by disrupting the same NTS mechanism shown to be critical for novelty-induced arousal to affect memory. This hypothesis was examined by infusing the AMPA receptor antagonist CNQX in the NTS 2 min before systemic administration of epinephrine (0.1 mg/kg). Both treatments were given after conditioning with the five CS-US pairings. All pre-exposed treatment groups exhibited comparable percentages of freezing to the final presentation of the CS during conditioning, $F_{(1,26)} = 0.057$, $P = \text{NS}$ (saline/PBS 94.6 ± 3.1, saline/CNQX 92.5 ± 4.0, epinephrine/PBS 97.9 ± 2.1, epinephrine/CNQX 94.3 ± 4.0).

**Retention test**

A two-way ANOVA indicated a significant overall effect on the mean percentage of freezing demonstrated to three presentations of the CS during retention testing, $F_{(1,26)} = 12.13, P < 0.01$. Post-hoc tests revealed that pre-exposed animals given an intra-NTS infusion of PBS and a systemic injection of epinephrine after bilateral infusions of CNQX into the NTS were indistinguishable from PBS injected controls ($P = \text{NS}$; Fig. 4A). Figure 4B shows the percentage of freezing during each of the three CS tone presentations. The epinephrine group (0.1 mg/kg) froze significantly more than did saline controls during the initial presentation of the CS ($P < 0.05$), but their percentage of freezing...
produced by emotionally laden experiences to brain systems that encode and store memory for fear conditioning.

Discussion

These experiments examined whether the intensity of arousal induced by the novelty of a learning context influences memory for Pavlovian fear conditioning. Findings from the three experiments reveal that memory for tone–shock pairings is enhanced in groups conditioned in a completely novel context relative to groups that were previously exposed by habituation in the training context 24 h before fear conditioning. Study 1 also examined whether the beneficial actions of arousal produced by exposure to the novel training context involves activation of peripheral sympathetic hormones. To this end, sotalol was administered preconditioning to block peripheral β-adrenergic receptors that bind the arousal related hormone epinephrine. The higher percentage of freezing behavior observed in groups conditioned in a novel context was attenuated by blocking these receptors before conditioning with sotalol. The results from study 1 suggest that arousal-induced changes in memory produced by the novelty of a conditioning context involve secretion of adrenal hormones and subsequent actions of these hormones on peripheral β-adrenergic receptors.

The interpretation of this finding was extended in study 2 by determining whether the mnemonic consequences of novelty are mediated in part by activation of brainstem neurons that are responsive to arousal-induced fluctuations in peripheral hormonal and sympathetic output. The adrenal hormone epinephrine binds to β-adrenergic receptors along vagal nerve fibers (Lawrence et al. 1995) that ascend to the brainstem and synapse upon neurons in the NTS (Kalia and Sullivan 1982). Arousal-induced changes in adrenal hormonal secretion increase discharge along vagal afferent fibers (Miyashita and Williams 2006) that, in turn, excite NTS neurons by releasing glutamate from its terminals (Granata and Reis 1983b; Allchin et al. 1994). Study 2 assessed the functional significance of glutamate release from excited vagal afferents onto NTS neurons in mediating the effects of novelty on memory. Findings from this study demonstrated that increased freezing observed during the tone retention test in subjects trained in a novel conditioning chamber was attenuated by blocking AMPA receptors in the NTS with CNQX.

It is important to note that although the experimental conditions used to induce novelty in experiments 1 and 2 facilitate later retention of cue–shock associative learning, novelty exposure with more intense environmental stimuli has been observed to produce opposite effects on mnemonic processing. For example, placement in an unfamiliar context in conjunction with restraint, restraint plus intermittent tail shock, in the presence of a freely moving cat or in an elevated platform that is brightly illuminated, disrupts induction of LTP, primed burst potentiation, and memory for spatial learning (Diamond et al. 1990, 1994; Xu et al. 1997; Akirav and Richter-Levin 1999; Diamond and Park 2000). The differences in memory and synaptic plasticity observed in these studies relative to those

Figure 3. (A) Groups: CNQX (1.0 μg) blockade of glutamatergic transmission in the nucleus of the solitary tract (NTS) attenuates novelty-induced memory enhancement. The non-pre-exposed group given vehicle into the NTS exhibited a significantly higher percentage of freezing than all experimental groups during three CS presentations given during a 48 h retention test (**P < 0.01). The enhancement in memory produced by novelty at the time of conditioning was attenuated by blocking AMPA receptors in the NTS with CNQX. The non-pre-exposed-CNQX group showed significantly poorer memory for the CS reflected in reduced freezing to the CS relative to the non-pre-exposed group given PBS into the NTS (*P < 0.05). Twenty-nine subjects were divided into the following treatment groups: pre-exposed PBS, n = 8; non-pre-exposed PBS, n = 8; pre-exposed CNQX, n = 6; and non-pre-exposed CNQX, n = 7. (B) Retention trials: Line graph depicting trial-by-trial freezing to CS tone presentations during retention testing. Subjects in the non-pre-exposed saline group showed a significantly higher level of freezing than all other groups during the second and third presentation of the tone CS (**P < 0.01). The high level of freezing exhibited by non-pre-exposed subjects was attenuated by blocking AMPA receptors in the NTS with CNQX.
reporting memory enhancement by using brief periods of non-stressful novelty exposure (Kinney and Routtenberg 1993; Vankov et al. 1995; Izquierdo et al. 2000, 2001, 2003; Viola et al. 2000; Li et al. 2003; Straube et al. 2003a,b; Davis et al. 2004; Moncada and Viola 2007; Sierra-Mercado et al. 2008) may be related to the magnitude of arousal and subsequent levels of stress induced by the respective training conditions.

If, however, brief exposure to novel contexts creates a moderate level of arousal through secretion of adrenal hormones, then administration of epinephrine to habituated subjects should increase arousal to a level comparable to that produced by Pavlovian conditioning in a context that was completely new. This premise was tested in the final study by examining whether more intensive levels of fear-induced freezing are exhibited on a 48-h retention test in pre-exposed subjects given post-training epinephrine (0.1 mg/kg) relative to saline-treated pre-exposed controls that showed only mild levels of freezing behavior in experiments 1 and 2. Findings from study 3 revealed that pre-exposed subjects given epinephrine post-conditioning exhibited a significantly greater percentage of freezing behavior during tone-only presentations on a 48-h retention test than did pre-exposed controls. The epinephrine-induced memory enhancement reflected in a higher percentage of freezing behavior was attenuated by interrupting impulse flow between the vagus nerve and the brainstem by blocking postsynaptic glutamate receptors in the NTS. There were no differences in the percentage of CS-induced freezing between controls and the group given epinephrine systemically and the glutamate receptor antagonist CNQX in the NTS. The overall findings suggest that exposure to a novel context increases physiological arousal, and these changes impact the strength of Pavlovian conditioning by influencing peripheral hormonal systems.

Previous studies demonstrate that several physiological indices of arousal such as heart rate and blood pressure are increased after the initial exposure to a new context (Carrive 2000). For instance, exposure to unfamiliar stimuli, such as water-immersion, handling, or placement into a new cage, results in increased activation of the sympathetic-adrenal system that is reflected by elevated concentrations of epinephrine in plasma (De Boer et al. 1990). Research also indicates that these exaggerated hormonal responses to novelty are suppressed by familiarizing subjects to a novel context by either repetitive or prolonged exposure to arousing stimuli (De Boer et al. 1988; Konarska et al. 1989, 1990). Based on these physiological findings, the present studies were conducted to examine the mechanism by which novelty produced physiological arousal impacts memory.

Findings from experiment 1 indicate epinephrine is involved in the capacity for novelty-induced arousal to impact the strength new events are encoded into memory. The level of freezing exhibited by non-pre-exposed animals given the peripheral
β-adrenergic receptor antagonist sotalol were comparable to the levels of freezing demonstrated when the CS was presented for saline-injected pre-exposed animals. This view is also supported by studies showing that arousal-induced changes in peripheral autonomic functioning involving elevated heart rate, increased discharge along vagal nerve fibers, and blood pressure are significantly reduced by blocking peripheral β-adrenergic receptors (van den Buuse et al. 2001; van den Buuse 2002; Carriere 2006; Miyashita and Williams 2006). The findings from experiment 1 demonstrate arousal states can be regulated by the novelty of stimuli, and novelty-induced arousal affects the formation of memory. It is important to note that the dose of sotalol selected for the present study did not in itself impair memory for pre-exposed control animals, suggesting this dose of sotalol was low enough to only partially saturate β-adrenergic receptors (Nattel et al. 1989). The absence of any observable impairment in pre-exposed sotalol-treated subjects may be related to floor effects of freezing performance produced by the mild training footshock. For example, the mild intensity 0.35-mA footshock used in this study was identified as the lowest level of shock intensity that is capable of eliciting cued-conditioned learning (Phillips and LeDoux 1992; Baldi et al. 2004). Therefore, this intensity was utilized to produce mild levels of freezing in controls to better examine whether the arousal produced by novelty improves overall fear conditioned learning in the non-pre-exposed groups. It is plausible that training parameters that produce higher percentages of freezing behavior in controls would, in fact, demonstrate that blocking peripheral β-adrenergic receptors with sotalol produces learning deficits. However, this type of training regimen will obscure the changes in learning and memory formation produced by novelty-induced increases in arousal.

Numerous studies indicate the adrenergic stress hormone epinephrine modulates memory formation for emotional events experienced by humans or animals. These effects are attributed to epinephrine acting directly on peripheral β-adrenergic receptors (Sternberg et al. 1986; Introni-Collison et al. 1992) and indirectly on NTS and LC neurons to potentiate noradrenergic activation of the amygdala and hippocampus (Williams et al. 1998, 2000; Miyashita and Williams 2004). Experiment 2 examined whether novelty-induced increases in peripheral autonomic and hormonal output affect central mnemonic processing by increasing synaptic transmission between peripheral vagal fibers and the neurons they synapse upon in the NTS. The vagus nerve was targeted as a putative pathway since the peripheral endings of the vagus nerve innervate a broad spectrum of sensory organs that show heightened activity in response to epinephrine secretion (Shapiro and Misels 1985; Coupland et al. 1989; Paton 1998a, b), and systemic administration of epinephrine increases neural impulses propagated along the vagus nerve as well as the firing rates in NTS neurons (Papas et al. 1990; Miyashita and Williams 2004). These findings suggest that the vagus nerve is capable of relaying peripheral physiological changes following the secretion of epinephrine to the brain in response to highly arousing experiences.

Arousal-induced increases in autonomic activity that are conveyed to the brain by the vagus nerve play an important role in producing functional and structural changes in brainstem neurons that are conducive to learning. For example, synaptic modifications increasing the efficiency of glutamatergic signaling in the NTS, such as increases in AMPA receptor subunit expression and structural changes at the synapse, occur from increased and sustained ascending peripheral signals such as hypertension and vagus nerve stimulation (for review, see Kline 2008). Furthermore, selective breeds with heightened states of autonomic activity such as spontaneously hypertensive rats show a number of synaptic modifications in the NTS such as a greater number of dendritic spines, an increase in the proportion of those spines that contain the GluR1 subunit of AMPA receptors, and an increase in the total AMPA receptor mRNA expression within the NTS compared with normotensive rats (Aicher et al. 2003; Saha et al. 2004; Hermes et al. 2008). Likewise, brief changes in blood pressure comparable to acute episodes of heightened arousal from an emotional event induce structural changes in neurons that indicate increased transcription at glutamatergic synapses in the NTS. As such, the collective findings suggest that highly arousing experiences that produce structural adaptations in the NTS through the release of glutamate may represent one mechanism by which emotional events are initially encoded and later processed by other limbic structures into long term memory.

Several behavioral studies demonstrate increased glutamate transmission in the NTS enhances memory for emotionally arousing experiences. For instance, microinjecting glutamate into the NTS where its neurons synapse with vagal afferents improves memory for the context where laboratory animals were last shocked in a water-motivated inhibitory avoidance task (Miyashita and Williams 2002; Kerfoot et al. 2008). The present study demonstrated that antagonizing glutamatergic transmission in the NTS with the selective AMPA receptor antagonist CNQX blocks the memory enhancing effects of heightened arousal from non-pre-exposure to the conditioning chamber. This study extends our understanding of the consequences of arousal on cognitive processes by revealing that postsynaptic AMPA receptors in the NTS transmit the physiological changes from novelty-induced arousal that enhance cue-fear conditioned memory.

Overall, findings from experiments 1 and 2 suggest novelty-induced arousal affects mnemonic processes by influencing peripheral hormone release and subsequent activation of the vagal/NTS complex. The final experiment was conducted to directly address interactions between peripheral hormones that are released following novelty-induced arousal and their impact on NTS neurons in the brain stem that are sensitive to fluctuations in peripheral autonomic functioning. To this end pre-exposed (non-aroused) animals were trained in the Pavlovian fear conditioning task with procedures identical to that employed in experiments 1 and 2 with the exception that separate groups received post-conditioning injections of saline or epinephrine. Findings from experiment 3 suggest that memory enhancement observed in response to novelty-induced arousal may involve peripheral hormonal secretion. This study demonstrated that increasing peripheral sympathetic output with epinephrine injections significantly enhanced the marginal levels of fear conditioning normally observed in groups pre-exposed to the conditioning chamber through habituation. Changes in physiological arousal produced by epinephrine in pre-exposed animals resulted in very high rates of freezing to the CS that were quite similar to that observed in non-pre-exposed animals tested in experiments 1 and
2. In addition, when physiological arousal is increased by handling following cued fear conditioning (Hui et al. 2006), administering epinephrine or corticosterone after learning tasks such as object recognition (Roosendaal et al. 2006; Donnelles et al. 2007), viewing a series of neutral slides (Cahill and Alkire 2003), or receiving a footshock in a distinct context (Introvini-Collison and McGaugh 1988), memory for the CS, the location of objects, slides viewed, or the context where given a footshock is improved. The findings from experiment 3 further suggest heightened states of arousal impact memory. Moreover, this study demonstrates that peripheral epinephrine is involved in the novelty-induced arousal memory enhancement in that it requires the same glutamatergic mechanism in the NTS. Given the mounting evidence that novelty and peripheral adrenergic mechanisms work in concert to strengthen synaptic connections, the current findings underscore the importance of signaling between the vagus and NTS complex in mediating the beneficial consequences of emotional arousal on memory.

Materials and Methods

Subjects

Eighty-three male Sprague–Dawley rats (275–300 g) obtained from Charles River Laboratories (Wilmington, MA) were used in experiments 1 (n = 24), 2 (n = 29), and 3 (n = 30). Rats were individually housed in plastic cages and maintained on a standard 12:12-h light–dark cycle with lights on at 7:00 a.m. Food and water were available ad libitum during the 7-d undisturbed adaptation period to the vivarium. All experiments were conducted in accordance to the policies and guidelines of the University of Virginia’s Animal Care and Use Committee.

Surgery

Each rat received an injection of atropine sulfate (0.1 mg/kg, i.p., American Pharmaceutical Partners, Inc.) followed 10 min later by an injection of the anesthetic sodium pentobarbital (50 mg/kg, i.p., Abbot Laboratories). A midline scalp incision was made, and 15-mm-long, extra-thin-wall stainless steel guide cannulae (25.0 gauge, Small Parts) were implanted bilaterally 2 mm above the NTS (AP: –13.3; ML: ±1.0 from bregma; DV: –5.6 from the skull surface) according to coordinates adapted from the atlas of Paxinos and Watson (1986). Guide cannulae and skull screws were anchored to the skull with dental cement, and the scalp was closed with sutures. Stylets (15 mm, 00 insect dissection pins) were inserted into the injection cannulae to maintain cannula patency. Penicillin (0.1 mL, i.m., Fort Dodge Animal Health) was administered immediately after surgery along with the analgesic buprenex (0.05 mL s.c., Hospira, Inc.) to alleviate postsurgical discomfort. The rats remained in a temperature-controlled chamber for at least 1 h following surgery and were given 7 d to recover before the start of each study.

Microinjection procedure

Each rat was restrained by hand in the experimenter’s lap, stylets were removed, and 17-mm-long, 30-gauge injection needles were inserted bilaterally into the NTS guide cannulae. The tip of the injection needle extended 2 mm beyond the base of the guide cannulae. The needles were connected to 10-μL Hamilton syringes via PE-20 (polyethylene) tubing. An automated syringe pump (Sage-Orion) delivered 0.5 μL of PBS or the AMPA receptor antagonist CNQX (1.0 μg; Sigma Aldrich) into the NTS over a period of 60 sec. The dose of CNQX used in this study was selected from a range of doses that effectively reduce NTS neural activity (Andresen and Yang 1990). The injection needles were retained in the guide cannulae for an additional 60 sec following infusion to ensure complete delivery of drugs. The stylets were then reinserted into the cannulae, and each rat received an i.p. injection of either saline or epinephrine (0.1 mg/kg).

Systemic injections

Subjects in the first experiment received preconditioning systemic (i.p.) injections of saline or sotalol (4 mg/kg), 5 min before placement into the conditioning chambers.

Behavioral apparatus

The apparatus used for Pavlovian fear conditioning consisted of a Coulbourn behavioral chamber (12 inches width × 10 inches depth × 12 inches height, model no. H13-16) that was enclosed in a larger sound-attenuating box (28 inches width × 16 inches depth × 16 inches height). The front and back walls of the chamber were made of clear plastic with stainless steel sides and a removable stainless steel grid floor. Freezing behavior was recorded during behavioral testing with an infrared activity monitor (model no. H24-61) that samples movement every 400 msec. The chambers used to assess retention for tone–shock pairings were identical in dimensions to the training apparatus but modified to be contextually different from the conditioning chambers and were located in a different room separate from the laboratory. The conditioning chambers were cleaned with a 10% alcohol solution after training and retention testing. All materials for the behavioral test apparatus were obtained from Coulbourn Instruments.

Behavioral procedures

Fear conditioning

Rats were transported from the vivarium into the laboratory 1 h before behavioral testing. One day prior to conditioning, the rats were habituated to the conditioning chamber with 5 min of free exploration. Animals assigned to the non-pre-exposure condition were also transported to the laboratory but remained in their home cage during the period that the pre-exposed group was habituated to the conditioning chamber. Twenty-four hours later animals in the pre-exposed or non-pre-exposed groups were placed into the chamber for conditioning. Three minutes after the rats were in the context, a 30-sec tone (5 kHz, 75 db) CS was presented and coterminated with a 1-sec, 0.35-mA footshock US. A 60-sec intertrial interval separated the footshock from the presentation of the next tone. Conditioning consisted of five tone–shock pairings.

Retention testing

Animals were transported in pairs to a completely different testing room and behavioral chamber to assess memory for the CS tone 48 h following conditioning. Each animal was given an initial 3-min period of exploration in the new chamber. Afterward, a CS tone (5 kHz, 75 db) was presented for 30 sec in the absence of the US footshock. A 30-sec intertrial interval separated the end of one tone and the presentation of the next. Three presentations of the CS tone were given during the retention test. The percentage of time subjects displayed a freezing response during presentation of the CS tone that was previously paired with footshocks was used as an index of retention.

Statistical analysis

Behavioral measures from the fear conditioning task are expressed as the mean percentage of time ± SE rats spent immobile during the presentation of the tone. Between-group comparisons for the freezing behavior measured during retention testing were made with a two-way ANOVA followed by Fisher’s post-hoc tests. Differences less than P < 0.05 were considered statistically significant.

Histology

To verify correct placement of injection needle tips and guide cannulae in the NTS after the completion of the experiment, each animal was anesthetized with the euthanasia solution Euthasil (0.5 mL, Virbac Corporation) and perfused intracardially with 0.9% saline followed by 10% formalin. The brains were stored in
10% formalin until sectioned on a vibratome. Sections were cut 60 μm thick, mounted on glass slides, subbed with chromium-aluminum, and stained with cresyl violet. The locations of the cannulae and injection needle tips were verified by examining enlarged projections of the slides (Fig. 1). The data from five animals were excluded from statistical analysis because of incorrect cannula placement.

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