Augmented Cocaine Seeking in Response to Stress or CRF Delivered into the Ventral Tegmental Area Following Long-Access Self-Administration Is Mediated by CRF Receptor Type 1 But Not CRF Receptor Type 2

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Stressful events are determinants of relapse in recovering cocaine addicts. Excessive cocaine use may increase susceptibility to stressor-induced relapse through alterations in brain corticotropin-releasing factor (CRF) regulation of neurocircuitry involved in drug seeking. We previously reported that the reinstatement of cocaine seeking by a stressor (footshock) is CRF dependent and is augmented in rats that self-administered cocaine under long-access (LgA; 6 h daily) conditions for 14 d when compared with rats provided shorter daily cocaine access [short access (ShA) rats; 2 h daily]. Further, we have demonstrated that reinstatement in response to intracerebroventricular CRF administration is heightened in LgA rats. This study examined the role of altered ventral tegmental area (VTA) responsiveness to CRF in intake-dependent increases in CRF- and stress-induced cocaine seeking. Bilateral intra-VTA administration of CRF (250 or 500 ng/side) produced reinstatement in LgA but not ShA rats. In LgA rats, intra-VTA CRF-induced reinstatement was blocked by administration of the CRF-receptor type 1 (CRF-R1) antagonist antalarmin (500 ng/side) or CP-376395 (500 ng/side), but not the CRF-R2 antagonist astressin-2B (500 ng or 1 μg/side) or antisauvagine-30 (ASV-30; 500 ng/side) into the VTA. Likewise, intra-VTA antalarmin, but not astressin-2B, blocked footshock-induced reinstatement in LgA rats. By contrast, neither intra-VTA antalarmin nor CP-376395 altered food-reinforced lever pressing. Intra-VTA injection of the CRF-R1-selective agonist cortagine (100 ng/side) but not the CRF-R2-selective agonist rat urocortin II (rUCN II; 250 ng/side) produced reinstatement. These findings reveal that excessive cocaine use increases susceptibility to stressor-induced relapse in part by augmenting CRF-R1-dependent regulation of addiction-related neurocircuitry in the VTA.

Introduction
Cocaine addiction is associated with a persistent susceptibility to drug relapse that emerges in an intake-dependent manner with repeated use. Understanding the neurobiological mechanisms that underlie drug relapse in cocaine addicts is critical to the development of effective treatment. Much evidence suggests that stress contributes to relapse. Stress promotes craving in abstinent human cocaine addicts (Sinha et al., 1999) and precipitates reinstatement in rodent relapse models (Erb et al., 1996; Ahmed and Koob, 1997).

The preclinical study of addiction-related, drug-induced neuroplasticity has involved the use of the long-access self-administration approach (Ahmed and Koob, 1998) in which rats provided repeated daily long access (LgA) to cocaine for self-administration (6–10 h/daily) are compared with rats provided shorter daily drug access [short access (ShA); 1–2 h/d]. We have reported that, compared with ShA rats, LgA rats are more susceptible to reinstatement by a stressor, footshock, suggesting that repeated cocaine use can produce intake-dependent alterations in how stress regulates neurocircuitry subserving motivation and relapse susceptibility (Mantsch et al., 2008). Our findings are consistent with reports that the magnitude of stress-induced craving is influenced by the amount of prior use in human addicts (Fox et al., 2005). The neuropeptide, corticotropin releasing factor (CRF), is a key mediator of stress-induced cocaine seeking (Shalev et al., 2010). CRF receptor antagonists block stress-induced reinstatement (Erb et al., 1998; Shaham et al., 1998; Graf et al., 2011), while delivery of CRF directly into the brain reinstates cocaine seeking (Erb et al., 2006; Mantsch et al., 2008). One key site at which CRF promotes reinstatement is the ventral tegmental area (VTA) (Wise and Morales, 2010) where CRF delivery precipitates reinstatement following self-administration (Wang et al., 2005, 2007). The VTA receives CRF-containing projections from a number of brain regions (Rodaros et al., 2007), and in vivo microdialysis studies have found that, during footshock-induced reinstatement, VTA extracellular CRF levels are elevated (Wang et al., 2005).
Although both CRF-R1 and CRF-R2 subtypes have been reported to be expressed in the VTA, the degree of expression, cellular localization, and function of these receptors in the VTA is controversial, and the receptor mechanism through which CRF regulates cocaine seeking remains unclear. Wang et al. (2005, 2007) have reported that reinstatement by centrally administered CRF is also augmented following LgA drug self-administration, suggesting that enhanced CRF responsiveness contributes to heightened stress-induced relapse (Mantsch et al., 2008; Graf et al., 2011), and with CRF binding protein, consistent with the mechanism through which CRF acutely enhances NMDA receptor-mediated currents which CRF-R2 (Shaham et al., 1998; Lu et al., 2001, 2003).

We have reported that reinstatement by centrally administered CRF is also augmented following LgA drug self-administration, suggesting that enhanced CRF responsiveness contributes to heightened stress-induced relapse (Mantsch et al., 2008). In this study, we examine intake-dependent augmentation of intra-VTA CRF-induced reinstatement and the role of VTA CRF-R1 versus CRF-R2 in reinstatement in response to CRF and stress.

Materials and Methods

Adult male Sprague Dawley rats (Harlan Laboratories) were housed individually under a 12 h reversed light/dark cycle (lights on at 7:00 P.M.) in a temperature and humidity controlled Association for Assessment and Accreditation of Laboratory Animal Care-accredited animal facility. All procedures were approved by the Marquette University Institutional Animal Care and Use Committee and performed in accordance with the NIH Guide for the Care and Use of Laboratory Animals.

Catheter and cannula implantation. For the reinstatement studies, rats were implanted with chronic indwelling jugular catheters under ketamine HCl (100 mg/kg, i.p.) and xylazine (2 mg/kg, i.p.) anesthesia, as previously described (Mantsch et al., 2008; Graf et al., 2011), and with bilateral 11 mm 26 gauge guide cannulae aimed at the VTA for intracranial injections. The tips of the guide cannulae were aimed 0.5 mm above the target injection site using the following coordinates determined from Paxinos and Watson (1998): 12° angle away from midline; anteroposterior −5.6 mm from bregma; mediolateral +2.2 mm from midline; and dorsoventral −6.7 mm from the skull surface. Placements for cannulae targeting the VTA for rats from each of the experiments are depicted in Figure 1.

Figure 1. A–C. Injection sites within the VTA for the following: LgA rats (triangles), ShA rats (squares), and Sal rats (circles) (A); LgA rats tested for CRF-induced reinstatement following CP-376395 (triangles), ASV-30 (circles), antalarmin (diamonds), and astressin-2B (squares), or LgA rats tested for footshock-induced reinstatement (stars) (B); and rats tested for effects on food-reinforced responding (open circles) and LgA rats tested for reinstatement by CRF receptor agonists (closed circles) (C).

Self-administration training. After recovery from surgery, rats were trained to self-administer cocaine (1.0 mg/kg/infusion, i.v.; NIDA Drug Supply Program) by pressing a lever under a fixed ratio 1 (FR1) schedule during daily 2 h sessions, within which the active (i.e., front) lever was extended into the chamber and the corresponding stimulus light was illuminated. Pressing the lever resulted in an intravenous infusion of drug or saline (Sal) solution (200 µl over 5 s) followed by a 25 s time-out period during which the stimulus light was extinguished but the lever remained extended. Responding on a second, inactive (i.e., back) lever was recorded but had no programmed consequences. Response requirements were gradually increased until rats displayed stable responding (within 10% of the three session mean) under an FR4 schedule, at which time they entered into a 14 day period of self-administration testing.

Effects of LgA self-administration on intra-VTA CRF-induced reinstatement. To examine intake-dependent effects of cocaine self-administration on later reinstatement by intra-VTA CRF, rats were assigned to ShA or LgA groups after self-administration training according to their access conditions for cocaine self-administration for the next 14 d. ShA rats (n = 6) continued to have access to cocaine for 2 h daily, as described above. LgA rats (n = 8) had access to the same cocaine dose for 6 h daily. Additionally, a third group of rats with no history of cocaine self-administration had access to infusions of saline during 14 daily 2 h sessions and served as a control to examine nonspecific effects of intra-VTA CRF on lever pressing (n = 6). Following 14 d of self-administration, rats underwent extinction during 10 consecutive 2 h sessions within which the cocaine solution was replaced by saline. After extinction, rats received a bilateral sham intra-VTA injection before testing for reinstatement in response to bilateral intra-VTA delivery of CRF (250 or 500 ng/side;
To examine the role of CRF-R1 and CRF-R2 in reinstatement by intra-VTA CRF administration, the ability of CRF (500 ng/side) to reinstate cocaine seeking following a 15 min bilateral intra-VTA pretreatment (0.25 μl/ side over 1 min) with the CRF-R1-selective antagonists antalarmin (500 ng/side; Sigma Aldrich; n = 7) (Webster et al., 1996) or CP-376395 (500 ng/side; Tocris Biosciences; n = 6) (Chen et al., 2008), or the CRF-R2-selective antagonists astressin-2B (500 ng and 1 μg/side; Sigma-Aldrich; n = 6) or antiauxagine-30 (ASV-30; 500 ng/side; Sigma-Aldrich; n = 6) was determined in separate groups of LGa rats. The astressin-2B and ASV-30 doses that were selected were based on those previously used for intracranial injections (Henry et al., 2006; Forster et al., 2008). After 14 d of LGa self-administration and extinction, rats were tested twice for CRF-induced reinstatement in counterbalanced sequence: once following intra-VTA pretreatment with CRF receptor antagonist; and once following pretreatment with vehicle. In the case of the CRF-R2 antagonist astressin-2B, rats were also treated with a second, higher antagonist dose (1 μg/side).

Effects of CRF receptor antagonists on stress-induced reinstatement. To examine the role of CRF-R1 and CRF-R2 activation in the VTA in stress-induced reinstatement, separate groups of rats were tested for the ability of electric footshock, delivered though the stainless steel grid floors of the self-administration chambers, to reinstate cocaine seeking following bilateral intra-VTA delivery of the CRF-R1-selective antagonist antalarmin (500 ng/side; n = 7) or the CRF-R2-selective antagonist astressin-2B (500 ng/side; n = 8). During the 15 min footshock period, the houselight was illuminated, the levers were retracted, and stimulus lights extinguished. Shocks (0.5 mA, 0.5 s duration) were delivered an average of every 40 s (range, 10–70 s). We have reported that these parameters produce robust reinstatement after long-access, but not short-access, self-administration (Mantsch et al., 2008). As was the case with intra-VTA CRF, rats were tested twice for footshock-induced reinstatement in counterbalanced fashion: once following intra-VTA pretreatment with CRF receptor antagonist; and once following pretreatment with vehicle.

Food self-administration. To confirm that the effects of intra-VTA antalarmin and CP-376395 on reinstatement were not attributable to nonspecific motor impairments, rats were tested for effects on sucrose pellet-reinforced lever pressing (data not shown). These rats were maintained at 90% of their free-feeding body weights and were trained to self-administer 45 mg sucrose-sweetened food pellets (BioServ) by pressing a response lever under an FR4 schedule of reinforcement during 30 min sessions. Once stable response patterns were obtained (responding within 10% of the mean over three sessions), separate groups of rats were tested for the effects of intra-VTA delivery of antalarmin (500 ng/side; n = 10) or CP-376395 (500 ng/side; n = 6), as described above, on responding. Each rat was tested twice with intra-VTA treatment in counterbalanced sequence: once with the CRF-R1 antagonist; and once with vehicle.

Reinstatement by intra-VTA administration of CRF receptor agonist drugs. To further examine the role of CRF-R1 and CRF-R2 in reinstatement, a separate group of LGa rats (n = 5) was tested for reinstatement following bilateral intra-VTA administration of the CRF-R1-selective agonist cortagine (100 ng/side) (Tezval et al., 2004), the CRF-R2-selective agonist rat urocortin II (250 ng/side) (Reyes et al., 2001), or vehicle (5 mM acetic acid in sterile saline). Following self-administration and extinction, each rat was tested with each intra-VTA treatment in counterbalanced sequence. After the first test, rats were only tested again for reinstatement if they emitted <20 responses during an intervening extinction session.

Histological confirmation of injection sites. The accuracy of cannula implantation was confirmed postmortem in each rat after cardiac perfusion with 60 ml of NaCl followed by 60 ml of 2.5% buffered neutral formalin under sodium barbital anesthesia (55 mg/kg). Brains were removed and stored in 2.5% buffered formalin for at least 1 d. Two hundred micrometer sections were cut using a vibratome, slide-mounted, and stained with cresyl violet for placement confirmation using a light microscope. Rats with injection sites outside of the VTA were excluded from data analysis. Notably, in cases where the injection sites were outside of the VTA (eight rats), bilateral intracranial delivery of the 500 ng/side CRF dose failed to produce significant reinstatement (mean responses, 20.38 ± 5.51 vs. 47.50 ± 7.20 for rats with injection sites within the VTA).

Statistical analyses. Statistical analyses were conducted using Predictive Analytics SoWfware statistics software (SPSS). Statistical significance was determined using ANOVA or Student’s t tests followed, when appropriate, by further analyses of main effects using ANOVA and/or post hoc testing using Bonferroni-corrected t tests.

Results

Effects of self-administration access condition on reinstatement by intra-VTA CRF self-administration

As expected, escalated self-administration was observed in rats provided daily long access (6 h; LGa rats, n = 8) but not short access (2 h; ShA rats, n = 6) to cocaine for self-administration and was not observed in rats provided access to saline (Sal rats, n = 6). For analysis, the mean daily hourly self-administration (infusions per hour) was compared across groups over the 14 d time period (Fig. 2A). Two-way ANOVA examining the effects of self-administration condition (ShA, LGa, and Sal) and day (1–14, repeated-measures) on the hourly number of self-administered infusions showed a significant overall main effect of self-administration condition (F(2,14) = 23.561; p < 0.001) but not day, and a significant condition × day interaction (F(26,208) = 2.708; p < 0.01). The daily number of self-administered infusions increased across the 14 d of self-administration in LGa but not ShA or Sal rats (one-way ANOVA: F(13,78) = 2.520; p < 0.01). Post hoc testing showed that self-administration was increased in LGa rats on days 5 and 9–14 compared with ShA rats (p < 0.05 for each comparison, Bonferroni-corrected t test). The mean net cocaine intake over the 14 sessions was markedly higher in LGa rats compared with ShA rats (t(11) = 5.889; p < 0.001) and is shown in the inset of Figure 2A.

Extinction and intra-VTA CRF-induced reinstatement

Responding did not differ between ShA and LGa rats during the 10 extinction sessions before reinstatement testing (Fig. 2B). However, differences in reinstatement in response to intra-VTA CRF (250 or 500 ng/side) were observed (Fig. 2C). Reinstatement by intra-VTA CRF was only observed in rats with a history of LGa self-administration. Two-way self-administration group (ShA, LGa, and Sal) × reinstatement condition (Veh: 250 ng/side CRF, 500 ng/side CRF) repeated-measures ANOVA showed significant main effects of self-administration group (F(2,17) = 16.774; p < 0.001) and reinstatement condition (F(2,34) = 7.934; p = 0.001), and a significant group × reinstatement condition interaction (F(4,68) = 5.969; p = 0.001). CRF-induced increases in responding were only found in rats with a history of LGa self-administration (one-way repeated-measures ANOVA; F(2,14) = 10.639; p < 0.01). Post hoc testing showed that responding following intra-VTA administration of the 500 ng/side CRF dose was significantly increased compared with vehicle (p < 0.05). Additionally, responding on the previously active lever was significantly increased in LGa rats compared with ShA and Sal rats.
followin intra-VTA delivery of either 250 ng/side CRF (one-way ANOVA, F_{2,19} = 4.179, p < 0.05; significant increase in LgA vs either ShA or Sal, p < 0.05, Bonferroni-corrected t test) or 500 ng/side CRF (one-way ANOVA, F_{2,19} = 23.103, p < 0.001; significant increase in LgA vs either ShA or Sal, p < 0.05). By contrast, we failed to find reinstatement effects or self-administration group effects on responding on the previously inactive lever or a reinstatement × group interaction (Fig. 2 D).

Effects of intra-VTA CRF-R1 antagonists on CRF-induced reinstatement

Bilateral intra-VTA pretreatment with the CRF-R1-selective antagonist antalarmin or CP-376395 blocked reinstatement by intra-VTA CRF (500 ng/side) in LgA rats (Fig. 3). The effect of antalarmin on CRF-induced reinstatement (n = 7) is shown in Figure 3A, and the effect of CP-376395 on CRF-induced reinstatement (n = 6) is shown in Figure 3B. In both cases, two-way repeated measures (CRF vs extinction; antagonist vs vehicle) ANOVA showed significant main effects of CRF delivery (F_{1,6} = 12.603, p < 0.05 for antalarmin; F_{1,5} = 6.920, p < 0.05 for CP-376395), but not antagonist treatment, and significant interactions between CRF-R1 antagonist pretreatment and CRF-induced reinstatement (F_{1,6} = 5.696, p = 0.05 for antalarmin; F_{1,5} = 9.120, p < 0.05 for CP-376395). Intra-VTA CRF produced significant reinstatement following pre-treatment with vehicle (p < 0.01 for each experiment), but not antalarmin or CP-376395, and responding following intra-VTA CRF administration was significantly lower following antalarmin or CP-376395 pretreatment compared with vehicle pretreatment (p < 0.01).

Effects of intra-VTA CRF-R1 antagonists on stress-induced reinstatement

Antagonism of CRF-R1 in the VTA also blocked footshock-induced reinstatement in LgA rats (n = 6) (Fig. 3C). Two-way (footshock vs extinction; antalarmin vs vehicle) repeated-measures ANOVA showed a significant overall effect of footshock (F_{1,5} = 28.817, p < 0.01), but not antalarmin pretreatment, and a significant antalarmin × footshock interaction (F_{1,5} = 6.294, p = 0.05). Intra-VTA CRF produced significant reinstatement following pre-treatment with vehicle (p < 0.01), but not antalarmin, and responding following EFS was significantly lower following antalarmin pretreatment compared with vehicle pretreatment (p < 0.01).

Effects of intra-VTA CRF-R1 antagonists on food-reinforced lever pressing

To confirm that the effects of intra-VTA antalarmin and CP-376395 on reinstatement were not attributable to nonspecific motor impairments, effects on sucrose pellet-reinforced lever pressing were examined. Neither intra-VTA antalarmin (185.83 ± 28.86 responses/30 min session vs 211.80 ± 13.41 responses/session for vehicle) nor intra-VTA CP-376395 (192.67 ± 23.73 responses/30 min session vs 203.10 ± 13.17 responses/session for vehicle) significantly decreased food-reinforced responding at doses that blocked CRF- or footshock-induced reinstatement. When interpreting these findings, it is important to note that in contrast to our self-administration rats, these rats had no history of cocaine intake and were food restricted, possibly altering their sensitivity to CRF-R1 antagonism.

Effects of intra-VTA CRF-R2 antagonists on CRF- and stress-induced reinstatement

The effects of bilateral intra-VTA pretreatment with CRF-R2-selective antagonists astressin-2B or ASV-30 on reinstatement by intra-VTA CRF (500 ng/side) are shown in Figure 4 A and B. The effects of the astressin-2B on CRF-induced reinstatement (n = 6) are shown in Figure 4 A. We initially tested rats for the effect of astressin-2B at the 500 ng/side astressin-2B dose. However, since this dose of astressin-2B did not attenuate reinstatement, we also tested rats with a twofold higher dose of astressin-2B (1 μg/side),
Two-way repeated-measures ANOVA examining the effects of both astressin-2B doses on reinstatement showed a significant overall effect of CRF-induced reinstatement ($F_{(1,5)} = 52.077, p = 0.001$), but not astressin-2B pretreatment, and no significant astressin-2B $\times$ CRF reinstatement interaction. Significant intra-VTA CRF-induced reinstatement was observed in rats pretreated with vehicle or either astressin-2B dose ($p < 0.01$ for each comparison), and reinstatement was not significantly different following astressin-2B delivery when compared with vehicle administration. Likewise, intra-VTA pretreatment with ASV-30 failed to alter reinstatement by intra-VTA CRF ($n = 6$) (Fig. 4B). Two-way repeated-measures ANOVA showed a significant overall effect of CRF-induced reinstatement ($F_{(1,5)} = 8.462, p < 0.05$), but not ASV-30 pretreatment, and no significant ASV-30 $\times$ CRF reinstatement interaction. Significant intra-VTA CRF-induced reinstatement was observed in rats pretreated with vehicle or ASV-30 ($p < 0.01$), and reinstatement was not different following ASV-30 delivery when compared with vehicle administration.

Effects of intra-VTA CRF-R2 antagonist on stress-induced reinstatement

Intra-VTA pretreatment with the CRF-R2-selective antagonist astressin-2B also failed to alter footshock-induced reinstatement ($n = 8$) (Fig. 4C). As was the case with CRF-induced reinstatement, ANOVA showed a significant overall effect of footshock reinstatement ($F_{(1,2)} = 20.587, p < 0.01$), but not astressin-2B pretreatment, and no significant interaction. Significant footshock-induced reinstatement was observed in rats pretreated with vehicle or astressin-2B ($p < 0.01$), and reinstatement was not different following astressin-2B delivery when compared with vehicle.

Reinstatement by intra-VTA administration of the CRF receptor agonists

To further examine the role of VTA CRF-R1 and CRF-R2 in reinstatement, the ability of bilateral intra-VTA injection of the selective CRF-R1 agonist cortagine (100 ng/side) or the selective CRF-R2 agonist rUCN II (250 ng/side) to reinstate cocaine seeking was examined in a separate group of LgA rats ($n = 5$) (Fig. 5). Reinstatement was observed following intra-VTA delivery of cortagine but not rUCN II. A two-way repeated-measures ANOVA with reinstatement condition (extinction vs intra-VTA treatment) and drug treatment (cortagine vs vehicle) as factors showed a significant overall main effect of reinstatement ($F_{(1,4)} = 17.316, p < 0.05$), but not cortagine treatment, and a significant reinstatement $\times$ cortagine treatment interaction ($F_{(1,4)} = 7.626, p = 0.05$). By contrast, an identical analysis examining rUCN II-induced reinstatement failed to show effects of rUCN II treatment or reinstatement testing or a rUCN II $\times$ reinstatement testing interaction. Intra-VTA cortagine treatment increased responding compared with the preceding extinction session ($p < 0.05$) and with vehicle-treated rats ($p < 0.05$).

Discussion

We have reported that self-administration under LgA/high-intake conditions augments later reinstatement by footshock stress and intracerebroventricular CRF (Mantsch et al., 2008), and that footshock-induced reinstatement following LgA self-administration is CRF dependent (Graf et al., 2011). These findings are consistent with prior reports that stress-induced reinstatement of drug seeking following self-administration of cocaine (Erb et al., 1998), heroin (Shaham et al., 1997), alcohol (Lé et al., 2000), or nicotine (Brujinzeel et al., 2009) involves CRF, as does stress-induced reinstatement of palatable food-seeking behavior (Ghitza et al., 2006). Our current findings demonstrate that heightened CRF responsiveness in the VTA likely contributes to intake-dependent increases in vulnerability to stress-
induced relapse that emerge with repeated drug use. Previous studies have reported that VTA CRF levels are increased during stress-induced reinstatement and that stress-induced cocaine seeking involves CRF actions in the VTA (Wang et al., 2005, 2007). Here we report that, similar to stress-induced reinstatement, reinstatement by intra-VTA CRF was augmented, and in fact was only observed, in rats with a history of self-administration under LgA/high-intake conditions. Further, we find that cocaine seeking induced by CRF delivery into the VTA or by footshock is mediated by VTA CRF-R1 and not CRF-R2, a finding that contrasts with previous reports implicating CRF-R2 in the VTA in stress-induced reinstatement (Wang et al., 2005, 2007).

A circuit involving the regulation of VTA dopaminergic neurons that project to the dorsal medial prefrontal cortex has been implicated in stress-induced relapse (McFarland et al., 2004). According to this model, inputs from several regions, including the ventral bed nucleus of the stria terminalis (vBNST), the central nucleus of the amygdala (CeA), and the NAc shell positively regulate mesocortical dopamine neurons in the VTA, thereby promoting cocaine seeking through activation of corticostratal glutamatergic inputs into the NAc core. It has been reported that the VTA receives CRF-containing afferents from two of these regions, the CeA and vBNST (Rodaros et al., 2007), and dual immunolabeling studies indicate that these CRF-containing projections are primarily glutamatergic (Tagliaferro and Morales, 2008). CRF-containing afferents form synapses on dopaminergic and GABAergic cells in the VTA (Tagliaferro and Morales, 2008), and the activity of both cell types is altered upon local CRF application (Korotkova et al., 2006). In the case of dopaminergic cells, CRF-positive synapses are primarily asymmetric (Tagliaferro and Morales, 2008), suggesting that CRF-releasing glutamatergic projections into the VTA, likely originating in the vBNST and CeA, positively regulate the activity of dopaminergic cells. Accordingly, delivery of CRF into the VTA stimulates local and terminal field dopamine release (Kalivas et al., 1987; Wang et al., 2005).

Although CRF-R1s are abundant in the VTA, their expression on dopaminergic cells appears to be somewhat limited (Korotkova et al., 2006), suggesting that actions of CRF on nondopaminergic cells may contribute to its effects on cocaine seeking. GABAergic cells in the VTA have also been reported to express CRF-R1s (Korotkova et al., 2006). The pattern of expression on other cell types in the VTA (glial cells, glutamatergic neurons) is unknown. The picture is less clear for CRF-R2s. Despite pharmacological evidence for CRF-R2 expression in the VTA (Wang et al., 2007), examination of the VTA using in situ hybridization suggests that there is little or no cell body expression (Van Pett et al., 2000). However, CRF-R2 expression has been reported as determined by RT-PCR (Korotkova et al., 2006). The purported ability of CRF-R2s to locally regulate glutamate release suggests a potential presynaptic localization of CRF-R2s on glutamate terminals (Wang et al., 2007). A clear understanding of the localization of CRF-R2s in the VTA awaits the availability of better antibodies for immunohistochemical characterization.

The precise mechanism of CRF regulation of dopaminergic cells in the VTA is also unclear and has been reported to involve both CRF-R1-mediated activation of protein kinase C (PKC) (Wanat et al., 2008) and possibly PKA (Riegel and Williams, 2008) signaling and/or CRF-R2 facilitation of NMDA receptor function (Angless et al., 2003). The CRF-R2 effects on signaling also appear to require CRF interaction with CRF binding protein (Angless et al., 2003; Wang et al., 2005) and may also involve presynaptic enhancement of glutamate release (Wang et al., 2005, 2007).

While our data implicate CRF actions in the VTA in stress-induced reinstatement, others have reported that CRF can act elsewhere within this circuit, including at sites upstream from the VTA, to regulate cocaine seeking, most notably in the vBNST, where CRF receptor antagonism blocks stress-induced cocaine seeking (Erb and Stewart, 1999). As the CRF actions in the vBNST involve projections from the CeA (Erb et al., 2001) and both regions send CRF-containing efferents to the VTA (Rodaros et al., 2007), it is likely that CRF exerts actions at multiple sites in the circuit that contribute to stress-induced cocaine seeking.

Similar to what we previously reported with stress-induced reinstatement (Mantsch et al., 2008), reinstatement by intra-VTA CRF appeared to represent an emergent phenomenon in that it was only observed following LgA self-administration. This finding is consistent with a report by Wang et al. (2005), who found that while footshock-induced increases in VTA CRF were observed in the VTA, the ability of CRF to regulate dopamine and glutamate levels in the VTA required a history of cocaine self-administration. The apparent augmentation of CRF responsiveness is also consistent with reports that CRF responsiveness in a number of brain regions is heightened following repeated cocaine administration (Erb et al., 2005; Liu et al., 2005; Pollandt et al., 2006; Orozco-Cabal et al., 2008; Francesconi et al., 2009; Guan et al., 2009). Although the effects of LgA cocaine self-administration on CRF-R1-mediated actions in the VTA have not been examined, some insight into potential mechanisms underlying heightened CRF regulation of cocaine seeking can be provided by studies involving repeated experimenter-delivered cocaine. In the VTA, CRF-R1 activation has been reported to produce both excitatory effects (via potentiation of NMDA and AMPA receptor-mediated neurotransmission) (Hahn et al., 2009) and possibly PKA (Riegel and Williams, 2008) and possibly PKA (Riegel and Williams, 2008) and possibly PKA (Riegel and Williams, 2008) and possibly PKA (Riegel and Williams, 2008) and possibly PKA (Riegel and Williams, 2008) and possibly PKA (Riegel and Williams, 2008) and possibly PKA (Riegel and Williams, 2008) and possibly PKA (Riegel and Williams, 2008) and possibly PKA (Riegel and Williams, 2008) and possibly PKA (Riegel and Williams, 2008) and possibly PKA (Riegel and Williams, 2008) and possibly PKA (Riegel and Williams, 2008) and possibly PKA (Riegel and Williams, 2008) and
wardly rectifying potassium channels) (Beckstead et al., 2009) on dopaminergic neurons. With repeated cocaine exposure, the excitatory effects of CRF-R1 activation are augmented (Hahn et al., 2009), while the inhibitory effects diminish (Beckstead et al., 2009), likely resulting in a net shift toward positive CRF-R1 regulation of dopaminergic cells in the VTA. Further, CRF binding in the VTA, as measured using autoradiography, is increased with repeated cocaine delivery (Goeders et al., 1990).

Our findings that VTA CRF-R1s mediate CRF- and stress-induced cocaine seeking are inconsistent with previous reports suggesting involvement of CRF-R2s in the VTA, apparently through a mechanism that also involves CRF binding protein (Wang et al., 2005, 2007). However, others have reported that stress-induced reinstatement is inhibited by systemic or intracerebroventricular delivery of CRF-R1, but not CRF-R2, antagonists (Shaham et al., 1998; Lu et al., 2001) and that CRF-R1, but not CRF-R2, antagonism can reduce evoked increases in VTA and NAc dopamine (Lu et al., 2003; Lodge and Grace, 2005). The reason for the discrepancy between our findings and those of Wang et al. (2005, 2007) is unclear but may involve different modes of CRF and receptor antagonist delivery (microinjection vs reverse dialysis), different CRF receptor antagonist and agonist doses, different rat strains, and differential experimental histories. Most importantly, it may be that the regulation of cocaine seeking by CRF-R1 in VTA requires a history of very high levels of cocaine intake [the daily cocaine intake reported by Wang et al. (2005, 2007) was ~33 mg/kg compared with >70 mg/kg in our LgA rats], which is consistent with reports that the selective CRF-R1 antagonists antalarmin and N,N-bis(2-methoxyethyl)-3-(4-methoxy-2-methylphenyl)-2,5-dimethyl-pyrazolo[1,5a] pyrimidin-7-amine (MPZP) only reduce cocaine self-administration following escalation in LgA rats (Specio et al., 2008).

The ability of stressful life events to precipitate drug use through actions involving CRF-R1s in the VTA may represent an emergent consequence of excessive cocaine use. Identification of the precise mechanisms through which VTA CRF-R1 activation produces cocaine seeking and the neuroadaptations that contribute to heightened susceptibility to CRF-dependent reinstatement should provide important insight into how stressor responsiveness changes in cocaine addicts in a way that promotes further use.

References