Neonatal Amygdala Lesions: Co-Occurring Impact on Social/Fear-Related Behavior and Cocaine Sensitization in Adult Rats

R. Andrew Chambers, Tammy J. Sajdyk, Susan K. Conroy, Joan E. Lafuze, Stephanie D. Fitz, and Anantha Shekhar
Indiana University School of Medicine

Neurodevelopmental abnormalities of temporal–limbic structures may underlie both adult psychiatric syndromes and increased addiction vulnerability, leading to high frequencies of “dual diagnosis” disorders. Although the amygdala is implicated in various mental disorders and drug addiction, no studies have explored the impact of early developmental damage to the amygdala on phenotypes relating to mental illness and addictions as co-occurring processes. We tested rats with neonatal amygdala lesions (NAML) vs. SHAM-operated controls in a battery of tests—novel field activity, elevated plus maze (EPM), and social interaction (SI) at baseline and after odor and restraint stress—followed by measures of cocaine sensitization (15 mg/kg vs. saline × 5 days + challenge session 2 weeks later) and remeasurement of SI. NAMLs showed increased novelty-related locomotion, less fear responding in the EPM, and resistance to predator-odor- but not to restraint-induced suppression of SI. NAMLs also had elevated cocaine sensitization profiles, and cocaine history differentially affected subsequent SI in NAMLs compared with SHAMS. NAMLs may provide models for understanding a shared neurobiological basis for and complex interactions among psychiatric symptoms, drug exposure history, and addiction vulnerability.

Keywords: amygdala, cocaine, neurodevelopment, dual diagnosis, social interaction.

Substance use disorder comorbidity in mental illness spans differential psychiatric diagnoses and drugs of abuse (Kessler, 2004; Regier et al., 1990). In many treatment settings, these “dual diagnosis” presentations are the majority of cases, associated with increased medical and psychiatric morbidity and mortality, financial destitution, and criminal incarceration (Dickey, Normand, Weiss, Drake, & Azeni, 2002; Dixon, 1999; RachBeisel, Scott, & Dixon, 1999). Emerging clinical and basic research suggests that varieties of mental illness and the addiction process involve integrated neurocircuits leading to increased addiction vulnerability regardless of or despite psychoactive consequences of drug use (Chambers, Krystal, & Self, 2001; Volkow, 2004). Functional and/or developmental abnormalities of the hippocampal formation and other temporal–limbic circuits are associated with a range of major neuropsychiatric and personality disorders that are frequently dually diagnosed (Blumberg et al., 2003; Bremner et al., 2000, 1995; Driessen et al., 2000; Weinberger, 1999). Meanwhile, these temporal–limbic regions are also directly connected with and modulate functional and neuroadaptive processes of frontal cortical–striatal motivational circuits impacted by addictive drugs (Chambers et al., 2001; Goto & O’Donnell, 2004; Kelley & Domesick, 1982; O’Donnell, Lewis, Weinberger, & Lipska, 2002; Pennartz, Groenewegen, & Lopez da Silva, 1994).

Animal modeling work shows that adult-onset or developmental disruptions to temporal–limbic structures produce not only clusters of behavioral abnormalities that model affective or schizophrenic syndromes but also multiple behavioral markers of enhanced addiction vulnerability. For example, adult-onset olfactory bulbectomy in rats produces distributed neural systems abnormalities involving components of the amygdala, hippocampal formation, and bed nucleus of the stria terminalis and a behavioral syndrome encompassing features of affective disorders, heightened behavioral sensitization to cocaine, and self-administration of amphetamine (Chambers, Sheehan & Taylor, 2004; Holmes et al., 2002; Kelly, Wrynn, & Leonard, 1997). Rats with neonatal ventral hippocampal lesions (NVHL) show a developmental behavioral and neurobiological marker syndrome consistent with features of schizophrenia (Flores et al., 2005; Lipska & Weinberger, 2000), increased cocaine self-administration behavior across several stages of the addiction process (Chambers & Self, 2002), impulsivity in natural-reward learning worsened by cocaine history (Chambers, Jones, Brown, & Taylor, 2005), and enhanced long-term sensitization to cocaine and ethanol (Chambers & Taylor, 2004; Conroy, Rodd, & Chambers, 2007).

The amygdala is another key brain region implicated in multiple neuropsychiatric disorders spanning psychotic, mood, anxiety, personality, and pervasive developmental disorder classifications.
(Anand & Shekhar, 2003; Hull, 2002; Joyal et al., 2003; Rusch et al., 2003; Shekhar, Truitt, Rainnie, & Sajdhyk, 2005; Sheline, Sanghavi, Mintun, & Gado, 1999; Sweeten, Posey, Shekhar, & McDougle, 2002). It is also involved in reinforcement learning and drug taking (Baxter & Murray, 2002; Carelli, Williams, & Holander, 2003; Everitt, Cardinal, Parkinsson, & Robbins, 2003; Kilts, 2001), and its basolateral regions have direct connectivity with frontal cortical–striatal circuits implicated in addiction (Baxter & Murray, 2002; Ishikawa & Nakamura, 2003; Kita & Kitai, 1990; Sah, Faber, Lopez De Armenta, & Power, 2003).

Although a growing literature has examined the role of neonatal amygdala lesions (NAML) in developmental aspects of autism, schizophrenia, and other disorders of abnormal social and affective behavior (Daenen, Van der Heyden, Kruse, Wolterink, & Van Ree, 2001; Diergaarde, Gerrits, Stuy, Spruijt, & Van Ree, 2004), no researchers have investigated the impact of NAML from an addiction-model perspective. We examined the effects of repeated cocaine injections in shaping the psychomotor behavior of NAML rats as a possible indirect measure of addiction vulnerability. For characterizing the impact of NAML and drug history on psychiatrically relevant behavioral phenotypes, the sensitization paradigm was embedded within a larger series of tests measuring aspects of novelty responsivity, anxiety, and social interaction (SI). This study design attempted to model complex dual diagnosis phenomena in which psychiatric syndromes and heightened drug responsivity may not only emerge in parallel from the same neurobiological abnormalities but also in which drug history might differentially influence neuropsychiatric phenotypes.

Method

Subjects and Surgery

Pregnant Wistar rats (Harlan, Indianapolis, IN) arrived in the laboratory at 14–15 days' gestation and were individually housed on a 12-hr light–dark cycle (lights on 7 a.m.) and given free access to food and water. Seven days after birth (Postnatal Day [PD] 7), male pups weighing 15–18 g were removed for 1–2 hr for surgery based on the protocol developed by Lipska, Jaskiw, and Weinberger (1993) and modified for NAML by Daenen et al (2001). Pups were randomly assigned SHAM (sham-operated) or NAML status, with balancing of lesion assignments within litters. For surgery, subjects were anesthetized by hypothermia on ice for 5 min. Time spent in each set of arms was scored from videotape review.

Phase 1: Socio–Affective Behavior

Open field habituation and social interaction (SI). All sessions occurred between 8 a.m. and 1 p.m. in a 91 cm x 91 cm x 30 cm SI box and were videorecorded from above under low red light (40 W). On the first day of testing (PD 59), novel open field activity was measured during a 5-min habituation session in the SI box. Activity was scored after dividing the box into nine 30 cm x 30 cm zones and counting entries into corner and center zones. SI sessions were conducted based on the protocol devised by File (1980). During each 5-min session, the experimental rat was placed into the box with a healthy partner rat with which it had been matched by sex, weight, and housing conditions and with which it had had no previous contact. Total time interacting was scored as the duration of physical contacts initiated by the experimental rat, including nonaggressive interactions such as sniffing, grooming, and playing. SI was measured for each rat in three independent sessions: as a baseline measure (second day after initial novel field assessment) and immediately after exposure to predator odor and restraint stressors (beginning second day after elevated plus maze assessment with a rest day between odor and restraint sessions). In the predator odor condition, animals were placed for 5 min into an airtight Plexiglass (Rohm & Haas, Philadelphia, PA) chamber (30 cm x 30 cm x 50 cm) in which a cat-odor-impregnated zeolite charcoal air filter (12 cm x 20 cm x 1.5 cm) had been taped to the inside of the lid. The filter had been previously placed in an enclosed litter box (Van Ness Plastics, Clifton, NJ; model CP-7) for 90 days of continuous use by a domesticated cat and then fixed to the odor chamber for 1 hr prior to rat entry. In the restraint stress condition, rats were placed in a disposable DecapiCone (Braintree Scientific, Braintree, MA) for 30 min.

Elevated plus maze (EPM). The EPM, standing 50 cm above the ground, consisted of two open arms (50 cm x 10 cm) and two enclosed arms (50 cm x 10 cm x 30 cm) with an open roof. The open and closed arms extended perpendicularly from a common center. Rats were placed in the center and exploration of the maze was videorecorded from above for 5 min. Time spent in each set of arms was scored from videotape review.
Phase 2: Cocaine Sensitization

Locomotor activity was measured under red light in 43.2 cm $\times$ 43.2 cm $\times$ 30.5 cm arenas that were made of clear plastic and equipped with three sets of infrared beam transmitters and detectors (Med Associates, St. Albans, VT). Horizontal motion ($x$, $y$ dimensions) and rearing ($z$ dimension) beam arrays were located 5 cm and 17.5 cm above the chamber floor, respectively. Each array consisted of 16 beams located 2.5 cm apart. Data were collected using Activity Monitor Version 5.0 software (Med Associates, St. Albans, VT) configured to separate stereotypic movements from gross horizontal locomotion. The dependent measure of stereotypic activity counted beam breaks occurring when the animal was not demonstrating forward horizontal locomotion, whereas the dependent measure of distance traveled was derived from beam breaks associated with displacement of the rat’s estimated center of mass approximately 5 cm in horizontal locomotion. During the sensitization paradigm, activity was assessed over five 2-hr sessions (one session/day), each of which consisted of a 60-min pre-injection phase and a 60-min post-injection phase. After the pre-injection phase, rats received either 15 mg/kg cocaine (15 mg cocaine HCl/1.0 ml saline) or (1.0 ml/kg) saline according to initial random assignments. Two weeks later, in another 2-hr session, all rats received a challenge dose of 15 mg/kg cocaine after the pre-injection hour.

Phase 3: Impact of Cocaine History on Social Interaction (SI)

Four days after all rats had received a cocaine injection in the challenge session and 18 days after the initial injection series, the rats were again tested in the SI paradigm with a novel partner as described above.

Lesion Verification

After behavioral testing, animals were decapitated under isoflurane anesthesia. Brains were removed and immediately frozen in isopentane. Coronal sections (20 µm) were cut on a cryostat at 300-µm intervals from approximately −0.11 mm to −5.25 mm relative to bregma (Swanson, 2004), corresponding to a range that included the entire amygdala, beginning just before the medial crossing of the anterior commissure and ending in the caudal hippocampus. Sections were placed on slides, dehydrated, stained with thionin, and coverslipped. Subjects were included in the behavioral analysis if, upon microscopic examination, they showed evidence of bilateral damage to the greater amygdala in at least two consecutive sections. Subjects with notable damage to basal ganglia, piriform cortex, or hippocampal formation were excluded.
**Statistical Analysis**

Differences between NAML and SHAM groups in the unitary measure pre-drug behavioral assays (novel SI field habituation, EPM, and pre-drug locomotor activity from Sensitization Day 1) were analyzed using independent samples t tests. The pre-drug SI series was tested using a repeated measures analysis of variance (ANOVA) (between: lesion status; within: pre-test condition). Subsequent testing examined independent factors of lesion status and drug exposure with two-way ANOVAs for unitary measures or repeated measures ANOVAs in which serial dependent measures were assessed. As indicated by significant main effects or interactions, post hoc testing was conducted with t tests (between: lesion) or least significant difference (LSD; between: lesion and drug history). Statistical significance was assumed at p < .05. Informative significant and nonsignificant (ns; p > .05) statistics are presented throughout. Data are presented as means ± standard errors of the means.

**Results**

**Lesion Analysis**

NAMLs were attempted on a total of 38 rats, of which 18 were assigned to receive saline and 20, cocaine during sensitization. From these groups, 3 saline- and 10 cocaine-assigned rats received inappropriate lesions, leaving 15 saline-assigned (NAML-SAL) and 10 cocaine-assigned (NAML-COC) rats for behavioral analysis. Appropriate amygdala damage was scored upon evidence in serial sections of cellular disarray, gliosis, atrophy, or absence of one or more major amygdalar nuclei (central, basolateral, basomedial, medial, anterior, intercalated), with or without extension or hypertrophy of the lateral ventricle into the tissue-poor amygdalar region. Figure 2 shows the minimal and maximal extents of amygdalar damage in the included set. In nearly all cases, damage encompassed both the central and basolateral nuclei; in no cases were one or the other of these nuclei undamaged on both sides of the brain. Damage to the dorsal or ventral endopiriform nuclei was deemed acceptable as long as it occurred at their boundaries with the amygdala proper within which the bulk of damage had occurred. The 13 subjects excluded from analysis showed highly heterogeneous patterns of inappropriate damage. The 3 excluded saline-assigned rats showed mutually exclusive unilateral extension of damage on the right side into the lateral hypothalamus versus hippocampus versus globus pallidus/substantia innominata. Among the 10 excluded cocaine-assigned rats, 2 showed insufficient damage to the amygdala bilaterally; 1, unilateral amygdalar damage only; 1, extension into right ventral hippocampus; 2, extension into right globus pallidus/substantia innominata; 1 bilateral extension into globus pallidus/substantia innominata; 2, extension into right lateral hypothalamus; and 1, extension into left lateral hypothalamus. Due to the pronounced heterogeneity of these inappropriate lesions, these rats were excluded from further analysis. None of the 24 SHAM rats showed evidence of excitotoxic damage.

**Behavioral Analysis**

**Phase 1: Socio-Affective Behavior**

_Habituation to novel social interaction (SI) arena._ Behavioral testing began for all animals in early adulthood (PD 59), beginning with this assessment of activity in the novel SI arena. During the initial 5-min exposure to the SI arena, NAML rats (N = 25) produced 48.2 ± 2.1 line crossings compared with 41.1 ± 1.4 produced by SHAM rats (N = 24). These differences were significant in two-way t testing, t(47) = -2.8, p < .01, indicating greater novel-field-induced locomotor activation in NAMLs.

_Elevated plus maze (EPM)._ Over the 5-min period, NAML rats spent 16.3% ± 2.3% of their time in the open arms of the EPM compared with 11.6% ± 1.6% for SHAM rats. These differences were indicative of a trend analogous to group differences in locomotor activity in the novel SI arena. Given an a priori hypothesis of impaired stimulus-induced fear responding associated with amygdala lesions (Kalin, Shelton, & Davidson, 2004; Takahashi, Hubbard, Lee, Dar, & Sipes, 2007), one-tailed t testing revealed significant group differences, t(47) = -1.7, p < .05.

_Social interaction (SI) after stressors._ During the 5-min baseline test, NAML and SHAM rats showed similar times in SI (23.7 ± 3.0 s and 23.1 ± 1.8 s, respectively). Repetition of the SI test with pre-conditioned stimuli of predator odor and restraint stress produced overall decreases in SI, with pre-condition, F(2, 94) = 53.9, p < .001, interacting with lesion status—Pre-Condition × Lesion Status, F(2, 94) = 3.7, p < .05 (Figure 3). Post hoc analysis revealed that SI was significantly suppressed after exposure to the predator odor to a greater extent in SHAM (12.1 ± 1.1 s) compared with NAML rats (19.6 ± 2.2 s), t(31) = 3.7, p < .01, but SI was further suppressed after restraint stress to similar levels in SHAM (8.0 ± 1.0 s) and NAML (8.8 ± 1.3 s) rats.

**Phase 2: Cocaine Sensitization**

_Pre-injection activation._ During the pre-injection hour on the first day of cocaine sensitization regimen, in the novel context, NAML rats (N = 25) again showed significantly greater locomotor activity compared with SHAM rats (N = 24): 10,250 ± 3,290 cm

---

_Figure 2 (follows)._ Histological verification mapping (upper panel, left) showed minimal (dark gray) and maximal (light gray) extents of lesions in brains of subjects included in the study: 25 rats with neonatal amygdala lesions (NAML) vs. 24 SHAM-operated controls. Coordinates shown are relative to bregma based on work by Swanson, 2004. Damage was usually centered at the boundary of the central (CEA) and basolateral (BLA) nuclei but often extended into the basomedial (BMA) and medial (MEA) nuclei. Exemplary whole coronal sections from a SHAM brain and a NAML brain with bilateral CEA/BLA damage (upper panel, right) is denoted by arrows. Exemplary sections from a SHAM and 3 different NAML brains featured at 25x magnification (lower panel) show variation in lesion damage with differential extents of cellular change and frank vacuolization in the amygdala. All sections were taken from approximately the same level relative to bregma (e.g. ±2.5).
main drug effect, a significant main effect of lesion, similarly progressive activity declines across all groups. However, differences in pre-injection activity (activity minus pre-injection activity) as a means to correct for baseline terms of injection-induced change in activity (post-injection activity) were therefore measured in analyses of drug-induced activation were therefore measured in terms of injection-induced change in activity (post-injection activity minus pre-injection activity) as a means to correct for baseline differences in pre-injection activity.

**Figure 3.** Social interaction times of experimental rats over 5-min encounters with novel healthy partners at baseline and immediately after exposure to predator odor and restraint stress. Repeated measures analysis of variance showed significant interactions between pre-test condition and lesion (p < .05) that was carried by significant differences between 25 rats with neonatal amygdala lesions (NAML) versus 24 sham-operated controls (SHAM) after the predator odor pre-condition (p < .01, post hoc t testing). Error bars indicate ±1 SEM.

Table 1

<table>
<thead>
<tr>
<th>Daily Pre-Injection Locomotion (cm/hr) in Rats by Experimental Group</th>
<th>Sensitization session (Day)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>1</td>
</tr>
<tr>
<td>SHAM-SAL</td>
<td>7664 ± 449</td>
</tr>
<tr>
<td>NAML-SAL</td>
<td>9961 ± 631</td>
</tr>
<tr>
<td>SHAM-COC</td>
<td>8339 ± 617</td>
</tr>
<tr>
<td>NAML-COC</td>
<td>10683 ± 1924</td>
</tr>
</tbody>
</table>

**Note.** SHAM-SAL = sham-operated saline-assigned rats; NAML-SAL = neonatal-amygdala-lesioned saline-assigned rats; SHAM-COC = sham-operated cocaine-assigned rats; NAML-COC = neonatal-amygdala-lesioned cocaine-assigned rats.

**Short-term sensitization.** Daily cocaine injections produced locomotor activation in NAML (N = 10) compared with SHAM (N = 13) and saline-receiving NAML (N = 15) and SHAM (N = 11) rats on the first day as exemplified in Figure 4. Over the 5 days of injections, drug-induced sensitization was confirmed in both NAML and SHAM groups as indicated by significant interactions of Day × Drug, F(4, 180) = 6.1, p < .001, but not Day × Lesion interactions, F(4, 180) = 1.0, ns. Significant main effects of drug, F(1, 45) = 88.0, p < .001, and Lesion × Drug interactions, F(1, 45) = 5.3, p < .05, but not of lesion, F(1, 45) = 2.1, ns, indicated that NAML rats demonstrated significant overall elevations in cocaine sensitization patterns. This was confirmed by post hoc testing (LSD) in which both NAML-COC and SHAM-COC rats showed greater activation than saline-receiving NAML or SHAM rats (p < .001 for all comparisons), while NAML-COC rats also activated to a greater extent than SHAM-COC rats (p < .05). NAML-SAL and SHAM-SAL groups were not significantly different.

Post-injection stereotypic movement (defined as beam breaks not counted as components of horizontal locomotion) over the 5 injection days is shown in Figure 5. Cocaine produced progressive daily—Day × Drug: F(4, 180) = 10.7, p < .001—and overall—drug: F(1, 45) = 172, p < .001—increases in stereotypic movement; however, there were no significant main effects of lesion, F(1, 45) = 1.1, ns, or Lesion × Drug interactions, F(1, 45) = 1.0, ns, indicating that stereotypic movement was not differentially sensitized by cocaine according to lesion status.

**Long-term sensitization.** Cocaine injections (15 mg/kg) were delivered to all rats 2 weeks after the initial short-term series of injections (15 mg/kg cocaine vs. saline × 5 days) (Figure 6). A two way ANOVA revealed a marginally nonsignificant main effect of drug history, F(1, 45) = 3.7, p = .059, on injection-induced change in activity. However, NAML rats showed significant increases in cocaine-induced change in activity—lesion: F(1, 45) = 4.6, p < .05—in which NAML rats with cocaine history showed significantly greater activation than SHAM rats with cocaine history and the NAML and SHAM rats with saline histories (p < .01 for all comparisons by LSD post hoc).

**Phase 3: Impact of Cocaine History on Social Interaction (SI)**

Four days after the cocaine challenge session, NAML versus SHAM rats showed differential levels of SI based on history of repeated cocaine versus saline injections in the short-term sensitization series (Figure 7, upper panel). While neither the main
Effects of lesion status, $F(1, 45) = 0.6, \text{ns}$, nor history of repeated cocaine injections, $F(1, 45) = 1.7, \text{ns}$, affected SI time, Lesion $\times$ Drug History interactions, $F(1, 45) = 6.4, p < .5$, were significant. Post hoc analysis (LSD) revealed that SHAM rats with histories of repeated cocaine injections showed greater SI time compared with SHAM rats with histories of only single cocaine injections or NAML rats with histories of repeated cocaine injections, $p < .05$; all other comparisons were not significant. A secondary analysis of the entire series of SI tests occurring both pre- and post-cocaine sensitization for only those rats receiving repeated cocaine injections (Figure 7, lower panel), showed general effects of pre-condition, $F(3, 63) = 14.9, p < .001$, that were differentially expressed based on lesion status: Pre-Condition $\times$ Lesion, $F(3, 63) = 3.4, p < .05$. While the main effect of lesion alone did not alter overall social behavior across these tests, $F(1, 21) = 0.07, \text{ns}$, post hoc testing ($t$ tests) specified that NAML rats were refractory to odor-induced suppression of SI before serial cocaine injections ($p < .05$) and refractory to cocaine history–related elevation in SI after cocaine sensitization seen in SHAM rats ($p < .05$).

Discussion

Young adult NAML rats show persistent elevations in locomotor behavior to novel conditions, reduced fear-related behavior suggested by EPM testing, and resistance to suppression of SI produced by pre-exposure to predator odor. Although generally consistent with deficits in stimuli-related discrimination and behavioral disinhibition long attributed to temporal-limbic and/or amygdalar damage (S. Brown & Schafer, 1887; Kluver & Bucy, 1937; Weiskrantz, 1956), this syndrome also encompassed elevated short- and long-term sensitization patterns with repeated cocaine injections. Finally, history of repeated versus single cocaine injections differentially impacted subsequent SI in NAML compared with SHAM rats.

Altered responses to novel objects or settings have been characterized as typical of bilateral amygdala damage in rats and monkeys whether induced neonatally or in adulthood (Burns, Annett, Kelley, Everitt, & Robbins, 1996; Daenen et al., 2001; Diergaarde et al., 2004; Kalin et al., 2004; Nachman & Ashe, 1987; S. Brown & Schafer, 1887; Kluver & Bucy, 1937; Weiskrantz, 1956).
Since normal upper ranges of exploratory behavior may be bounded by threat-awareness (e.g., novelty neophobia), these findings may reflect a fundamental deficit of amygdala-mediated cautionary behavior normally provoked by ambiguous contexts or stimuli (Bauman, Lavenex, Mason, Capitanio, & Amaral, 2004a; Kalin et al., 2004; Kalin & Shelton, 2003). In this study, locomotor reactivity to novel contexts was tested twice: first, at the beginning of experimental series in the SI arena, and second, during the initial pre-injection hour of the cocaine sensitization series. Both of these tests identified elevated activation to novel environments in NAML rats, suggesting the robustness and persistence of this trait. Consistent with distributed roles of temporal–limbic structures (e.g., both amygdala and ventral hippocampus) in modulating novelty reactivity, increased responsiveness to contextual novelty has also been identified in adult rats with neonatal ventral hippocampus lesions (NVHLs; Lipska et al., 1993). However, this effect is often minimal or inconsistent in NVHLs (Chambers & Taylor, 2004; Conroy et al., 2007), while NAMLs more powerfully augment novel open field activity when directly compared with NVHLs (Daenen et al., 2001).

That NAMLs produce reductions in fear-related behaviors was also suggested by elevations in time spent in the anxiogenic open arm of the EPM and failure of predator odor pre-exposure to suppress SI to the extent observed in SHAMs. In contrast, adult-onset lesions to the central nucleus of the amygdala do not appear to produce differential EPM responding in rats with such lesions compared with controls at baseline, although pre-exposure to restraint stress suppresses open arm entries in control but not adult-lesioned rats (McHugh, Deacon, Rawlins, & Bannerman, 2004; Moller, Wiklund, Sommer, Thorsell, & Heilig, 1997). The SI series demonstrated that neither gross deficits in SI per se nor generalized impairments in the capacity of all pre-test stressors to suppress SI can account for the failure of predator odor to suppress SI in NAML rats. Differential forms and severities of psychological stress produce differential patterns and extremes of neural activation within the amygdala and other stress-responsive circuits (Crane, French, & Buller, 2005; Takahashi, Nakashima, Hong, & Watanabe, 2005). Our data suggest that NAMLs produce deficits in the generation of fear-responses that are most pronounced under mild to moderately stressful conditions and/or are encoded to some extent by olfactory processing channels. Although the present study did not rule out whether NAMLs produce anosmia, olfactory sensation and some capacity for olfactory-based discrimination and learning are maintained with both neonatal (Sullivan & Wilson, 1993) and adult-onset amygdala lesions (Shoenbaum, Setlow, Nugent, Saddoris, & Gallagher, 2003). In general, our results are in agreement with work showing that unconditioned fear responding to predator odor is reduced in rats with adult-onset ibotenic acid lesions to the basolateral amygdala (Takahashi et al., 2007).

Elevated short-term sensitization profiles to repeated cocaine injections in NAML rats suggests persistence of hypersensitivity to activating effects of cocaine superimposed on an intact sensitization process across injection days. Although NAML rats showed elevated daily pre-injection activity compared with SHAMs, this activity could not have accounted for elevations in post–cocaine injection activity, given our analysis of daily change in activity post-injection (i.e., post-injection activity minus pre-injection activity), which rendered the injection-induced activation curves of saline-receiving NAML and SHAM rats identical. Also, lesion-based differences in nonhorizontal behavior (i.e., stereotypic behavior) could not have accounted for differences in locomotor sensitization by interfering with locomotion, because cocaine-

![Figure 6.](image_url)
receiving NAML and SHAM rats showed similar patterns of stereotypic behavior.

Effects of single or repeated injections of cocaine or other psychostimulants in NAML rats have not been previously reported, although increased locomotor responses to single injections of phencyclidine (Daenen, Wolterink, & Van Ree, 2002a, 2002b). However, future studies involving differential age testing of sensitization or head-to-head comparisons between subjects with neonatal and subjects with adult-onset amygdala lesions are needed to accurately characterize developmentally specific effects of NAMLs on cocaine sensitization.

In the cocaine challenge session, NAML rats with prior cocaine history showed the greatest activation to a single injection of cocaine compared with their cocaine-history SHAM and saline-history NAML counterparts, demonstrating that NAMLs augment long-term cocaine sensitization. Sensitization has been demonstrated in rodents using a variety of drugs (cocaine, amphetamine, nicotine, opiates, alcohol) that are pharmacologically and psychoactively divergent but share common capacities to produce dopamine efflux into the ventral striatum and ultimately, addictive behavior (Di Chiara & Imperato, 1988; Wise & Bozarth, 1987). Behavioral sensitization may be analogous to processes of motivational sensitization as a core feature of addictive disease (Robinson & Berridge, 1993), and neuroplastic changes underlying behavioral sensitization occur in cortical–striatal circuits also implicated in reward-motivated operant behavior (Vanderschuren & Kalivas, 2000). Given previous modeling of dual diagnosis syndromes involving temporal–limbic disruptions via the NVHL or olfactory bulbectomy, in which enhancement of novelty reactivity, drug sensitization and self-administration have all been identified (Chambers & Self, 2002; Chambers et al., 2004; Chambers & Taylor, 2002; Holmes et al., 2002), the present results are suggestive of NAML-enhancement of addiction vulnerability that requires confirmation in self-administration studies.

By placing the cocaine injection series between two phases of SI testing, the present study allowed assessment of the contribution of drug history toward altering social behavior as a general component of psychiatric illness. SHAMs with repeated cocaine injection histories showed increased SI compared with their single-cocaine-injection SHAM counterparts, while repeated cocaine injection histories nonsignificantly reduced SI in NAML rats compared with their single injection counterparts. These results suggest that recent cocaine sensitization history can have differential effects on subsequent social behavior, based on underlying neurodevelopmental amygdalar damage. In lesioned rats, recent cocaine history can impair SI (Costall et al., 1990) and alter levels of stress-related neuropeptide neurotransmitters within the amygdala (Richter-Weiss, 1999; Zhou, Spangler, Ho, & Kreek, 2003). Whether our results reflect NAML-based alterations in neural mechanisms mediating the expression of cocaine withdrawal, tolerance, or sensitization within the domain of social behavior requires further investigation. Examination of the entire sequence of SI tests in only those rats that received repeated cocaine injections does suggest that across pre-conditions that alter social behavior, NAML rats were relatively refractory to pre-conditions that decrease (odor pre-condition) or increase (repeated cocaine injection history) SI in SHAM rats. This pattern is consistent with the putative role of the intact amygdala in flexibly configuring social and other forms of motivated behavior, according to current affective states or learned (e.g. sensitized) affective associations.

Further work is needed to better characterize how NAMLs model aspects of one or more human psychiatric disorders. Cur-
rently, only a few studies have examined the NAML syndrome from behavioral (e.g., Bauman, Lavenex, Mason, Capitanio, & Amaral, 2004b; Daenen et al., 2001; Diergaard et al., 2004) or neurobiological perspectives (Bouwmeester, Snapper, Ronken, Kruse, & Van Ree, 2003; Gerrits, Wolterink, & Van Ree, 2006; Van den Buse, Garner, & Koch, 2003) with many investigators suggesting NAMLs model social developmental deficits of schizophrenia and/or autism (Daenen et al., 2002b). However, our results and other lines of evidence suggest the NAML syndrome could more directly relate to forms of developmental psychopathology marked most prominently by impulsive behavior rather than by the social deficits and stereotypic behaviors associated with autism. For instance, although our procedural methods likely lacked sensitivity for detecting relatively fine or qualitative differences between groups, we observed neither gross deficits in baseline social behavior and interest nor abnormalities in saline- or psychostimulant-induced rates of stereotypic behavior in NAMLs. Similarly, in NAML monkeys that grow up in groups, social interest and ranges of social behavioral repertoires are not altered per se, but the timing of particular social behaviors are disordered such that either affiliative–approach behavior or fear-related social withdrawal occur at inappropriate junctures (Bauman et al., 2004b). Humans with progressive bilateral amygdalar degeneration can also show a full range of social behaviors and interest but demonstrate alterations in socio-affective perception, poor social decision making, and susceptibility to predatory or unstable relationships (Adolphs, Tranel, & Damasio, 1998; Adolphs, Tranel, Damasio, & Damasio, 1995). Given the similarity of these traits with features of the borderline personality syndrome and recent neuroimaging evidence identifying amygdalar abnormalities in patients with borderline personality syndrome (Rusch et al., 2003; Tebartz van Elst et al., 2003), NAMLs may also model aspects of certain personality disorders, which, like schizophrenia, entail high rates of dual diagnosis.

Regardless of the psychiatric diagnostic specificity of the NAML syndrome, we speculate that it entails mechanisms of addiction vulnerability similar to those implicated in the NVHL model of schizophrenia. Like the ventral hippocampus, the basolateral amygdala sends direct glutamatergic fibers into both the prefrontal cortex and the nucleus accumbens (Baxter & Murray, 2002; Ishikawa & Naka- mura, 2003; Kita & Kitai, 1990; Sah et al., 2003). The basolateral nucleus also has reciprocal connectivity with the ventral hippocampus (Pitkanen, Pikkarainen, Nurminen, & Ylinen, 2000), and neural recordings suggest the hippocampus and the amygdala work interactively to modulate frontal–cortical–ventral striatal information processing and learning (Finch, 1996). On the basis of this functional anatomy and the fact that NVHLs produce developmental alterations in frontal–cortical–striatal cellular markers and physiology (O’Donnell et al., 2002), further studies should test whether NAMLs also introduce analogous cortical–striatal alterations leading to enhanced response profiles to addictive drugs.

In summary, these findings contribute to growing data that developmental disturbance of temporal–limbic regions—that may instantiate a neurobiological basis component of several major adult psychiatric syndromes—also confers increased short- and long-term responsiveness to addictive drugs and differential effects on psychiatric-related phenotypes after drug exposure. Further studies using the NAML may be useful for understanding addiction vulnerability in the mentally ill or interpreting clinical data suggesting that history of chronic substance use alters neuropsychiatric illness trajectories and treatment responsiveness (Le Fauve et al., 2004).

References


