Cocaine Dose and Self-Administration History, but Not Initial Cocaine Locomotor Responsiveness, Affects Sensitization to the Motivational Effects of Cocaine in Rats

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ABSTRACT

Cocaine addiction is a significant and complex disease. Part of this complexity is caused by the variability of the drug experience early in drug use (initial responsiveness, amount of use, etc.). In rats, individual differences in initial cocaine responsiveness and cocaine self-administration history both predict the development of cocaine sensitization, a putative mechanism contributing to the development of cocaine addiction. Here, we sought to determine the role of these factors and cocaine dose on the development of sensitization to cocaine’s motivational effects during the earliest stages of self-administration. Rats were classified as either low or high cocaine responders (LCRs or HCRs, respectively) based on acute cocaine-induced locomotor activity (10 mg/kg i.p.) before learning to self-administer cocaine (0.6 mg/kg/infusion i.v.) under a fixed ratio 1 (FR1) schedule of reinforcement. After acquisition, rats self-administered cocaine (0.6 or 1.2 mg/kg/infusion) under a progressive ratio (PR) schedule of reinforcement either immediately or after an additional five FR1 sessions (0.6 or 1.2 mg/kg/infusion). No LCR/HCR differences in sensitization were observed. However, regardless of LCR/HCR classification, exposure to the higher dose of cocaine produced sensitization to cocaine’s motivational effects on the PR schedule (i.e., increased break points) and an escalation of consumption on the FR schedule. Thus, our results reveal a novel model for studying escalation and sensitization very early after acquisition and suggest that sensitization may be important in the earliest stages of the cocaine addiction process.

Introduction

Cocaine addiction remains a significant public health issue. Recent statistics from the United States indicate that as of 2010 37.2 million people aged 12 and over had tried cocaine at least once in their lifetimes, with approximately 1.5 million current cocaine users (Substance Abuse and Mental Health Services Administration, 2011). These statistics are particularly troubling considering an estimated 10 to 15% of all initial intranasal cocaine users (Substance Abuse and Mental Health Services Administration, 2011). These statistics are particularly troubling considering an estimated 10 to 15% of all initial intranasal cocaine users progress to addiction (Gavin, 1991). For a subset of these individuals, the transition from recreational use to addiction occurs very rapidly and is associated with routes of administration that result in more rapid and higher brain cocaine levels (Wagner and Anthony, 2002; O’Brien and Anthony, 2005). Furthermore, individual differences in the initial subjective effects of cocaine have been found to correlate with long-term use and dependence (Lambert et al., 2006). Thus, understanding factors that contribute to the rapid development of this disease in susceptible individuals remains an important and significant challenge.

Addiction has been defined as “a chronic disorder characterized by the compulsive use of a substance resulting in physical, psychological, or social harm to the user and continued use despite that harm” (Rinaldi et al., 1988). Because of the complex etiology of addiction, animal models tend to focus on one aspect of the human condition (but see Deroche-Gamonet et al., 2004). For example, the long access model of cocaine self-administration and extinction/reinstatement paradigms model escalation in cocaine consumption and “relapse” to cocaine use, respectively (Ahmed and Koob, 1998; Shaham et al., 2003), but do so after stable patterns of...
self-administration have been established. We are interested in factors that occur earlier in the drug-taking process, and we have recently established a method for measuring cocaine consumption during the earliest stages of self-administration. This method defines acquisition based on intake criteria and then measures each subject's postacquisition intake relative to the day of acquisition (day x), allowing for experimental manipulations after very limited initial cocaine exposure (Mandt et al., 2012).

One factor postulated to be important for the transition from recreational drug use to addiction is sensitization (Robinson and Berridge, 1993, 2001). In animals, sensitization to the locomotor-stimulating effects of drugs such as cocaine has been demonstrated for years (Post, 1980; Zahniser et al., 1988; Kalivas and Duffy, 1993); however, whether sensitization is seen in humans and nonhuman primates is an area of active debate (Bradberry, 2007). One possibility for this discrepancy is that sensitization in humans may occur while addiction is developing, before the individual has entered treatment and subsequent clinical studies. Likewise, sensitization may also be more difficult to demonstrate during contingent (i.e., drug self-administration) than noncontingent (i.e., experimenter administered) cocaine exposure, because it occurs very early in the drug-taking process while animals are being trained on the operant. Nonetheless, sensitization during paradigms of cocaine self-administration has been demonstrated. For example, Roberts and colleagues have reported environmental conditions that result in sensitization to the reinforcing effects of cocaine measured with a progressive ratio (PR) schedule of reinforcement (Morgan and Roberts, 2004; Liu et al., 2005; Morgan et al., 2006).

Over the past 10 years, we have worked with a model of individual differences in rats that distinguishes animals based on the magnitude of their acute locomotor response to a single cocaine injection (10 mg/kg i.p.; Sabeti et al., 2002). One of the most consistent behavioral findings has been that rats with lower, but not higher, initial responsiveness to cocaine [low cocaine responders (LCRs), but not high cocaine responders (HCRs)] more readily develop cocaine-induced locomotor sensitization after repeated noncontingent cocaine (10 mg/kg i.p.) (Sabeti et al., 2003; Allen et al., 2007; Mandt et al., 2008, 2009; Nelson et al., 2009). It is noteworthy that the LCR/HCR difference in cocaine-induced locomotor sensitization was not found to predict acquisition of low-dose cocaine self-administration (Mandt et al., 2008). However, whether LCRs and HCRs differ in the development of other forms of sensitization (e.g., during contingent cocaine self-administration) remains unknown.

Thus, the present study sought to investigate the role of self-administration history and initial cocaine responsiveness on the development of sensitization to the motivational effects of cocaine. Specifically, we used our new method of acquisition analysis combined with PR schedules of cocaine reinforcement to determine whether: 1) we could establish conditions that reveal sensitization to the motivational effects of contingent cocaine administration and 2) LCRs and HCRs differ in the development of this form of sensitization.

Materials and Methods

Animals. Four groups of outbred male Sprague-Dawley rats (n = 70; group A, n = 16; group B, n = 14; group C, n = 20; group D, n = 20), weighing between 225 and 250 g (~8 weeks of age), were purchased from Charles River Breeding Laboratories (Portage, MI) and used in this study. All rats were housed individually with ad libitum access to food and water in an animal care facility at the University of Colorado, Denver, CO. To remain consistent with all previous LCR/HCR studies, rats were housed on a 12-h light/dark cycle (lights on at 6:00 AM), and all testing was conducted during the light cycle. All animal care and use procedures were in strict accordance with the National Institutes of Health Guide for the Care and Use of Laboratory Animals (Institute of Laboratory Animal Resources, 1986) and approved by the University of Colorado Denver Institutional Animal Care and Use Committee.

Catheter Construction and Placement. Intravenous catheters were constructed in the laboratory and surgically implanted into the right jugular vein under ketamine (100 mg/kg i.m.) and xylazine (10 mg/kg i.m.) anesthesia by using established procedures (Thomasen and Caine, 2005). Rats received acetaminophen as an analgesic in their drinking water (20 mg/ml) for 48 h before and after surgery. Rats recovered from surgery for at least 1 week before self-administration training began. Catheters were flushed with 0.3 ml of bacteroiostatic 0.9% sodium chloride containing heparin (30 U/ml) before and after each self-administration session. Sodium thiopental (20 mg/kg i.v.) was administered at the conclusion of the experiment to verify catheter patency.

Locomotor Activity. Locomotor activity testing was conducted as described previously (Sabeti et al., 2002; Mandt and Zahniser, 2010) and consisted of a single cocaine exposure 48 to 72 h before self-administration training. In brief, rats were taken to the behavioral testing room in their home cages and allowed to habituate for 45 to 60 min. At the start of the behavioral recording session, rats were placed in open field activity chambers consisting of Plexiglas boxes (43.2 × 43.2 cm) fitted with a photobeam frame (16 beams per horizontal photobeam interruptions converted to distance traveled (centimeters) per 10-min bin. The sum of distance traveled over the 30 min postcocaine was used to determine the median split for all rats within each of the four groups, which was then used to classify rats as either LCRs or HCRs.

Self-Administration Training. Rats self-administered cocaine during the light cycle in 16 Plexiglas and metal operant conditioning chambers (29 × 24 × 21 cm; MED Associates) that were housed within sound-attenuating cabinets. The chambers had two retractable levers on the front wall with stimulus lights positioned 6 cm above each lever. A tone presentation speaker (Sonalert Tone Generator, 2900 Hz; MED Associates) and a white noise speaker (90 dB) were mounted 12 cm above the floor on the wall opposite the levers. A house light (100 mA) was mounted 6 cm above the tone speaker, and a computer-controlled syringe pump delivered cocaine infusions. All behavioral events were monitored and controlled by a personal computer using MED-PC for Windows software (MED Associates). All self-administration sessions began with the extension of the retractable levers, white-noise activation, and illumination of the stimulus light on the right side of the chamber. For fixed ratio 1 (FR1) sessions, 1 s after session initiation, a cocaine priming infusion was delivered (0.6 or 1.2 mg/kg/infusion in a volume of 0.2 ml over 5–7 s, based on the weight of the rat). During this priming infusion and all subsequent self-administered infusions, the stimulus light over the active lever was turned off, and a tone-house light stimulus complex was activated for 15 s coinciding with a “timeout” period.

Acquisition of cocaine self-administration was measured in 2-h sessions, and testing was conducted 5 days a week. Responses on the right lever were reinforced with a cocaine infusion (0.6 mg/kg) according to a FR1 schedule of reinforcement. All groups (A–D) acquired cocaine self-administration with this training dose. Re-
sponses emitted on the right lever during cocaine infusion and stimulus complex (i.e., timeout) were not reinforced and were recorded separately from reinforced responses. Responses on the left lever were recorded but had no programmed consequence. Acquisition in these experiments was defined as the first of three consecutive sessions during which a rat consumed at least 4 mg/kg cocaine (Mandt et al., 2012). In our work, rats that meet these criteria continue to reliably self-administer cocaine; and these criteria are similar to intake-based acquisition criterion used by other laboratories (Carroll and Lac, 1997; Mantsch et al., 2001).

After acquisition, rats that met criterion were advanced to PR testing either immediately (groups A and B) or after an additional five FR1 sessions (groups C and D). Similar to FR testing, PR testing was conducted 5 days a week, and responding was reinforced with either 1.2 mg/kg/infusion cocaine (groups A and C) or 0.6 mg/kg/infusion cocaine (groups B and D). PR sessions were 5 h in duration, and the response requirement increased progressively according to the following schedule: 1, 2, 4, 6, 9, 12, 15, 20, 25, 32, 40, 50, 62, 77, 95, 118, 145, 178, 216, 288, 328, 402, 492, 603, and 737 for cocaine infusions 1 to 25, respectively (Richardson and Roberts, 1996). Breakpoint was defined as the last response ratio completed before 1 h without earned reinforcement or the end of 5 h. As with the FR sessions, left lever responses were recorded but had no programmed consequences. The details of experimental conditions for each group are shown in Table 1.

Exclusions and Exceptions. In total, 20 of the 70 rats used in this study were excluded from final PR analysis. One rat in group A, two rats in group B, four rats in group C, and six rats in group D did not acquire cocaine self-administration. An additional three rats (one in group B and two in group D) had catheter patency failure before the end of the acquisition phase of the study. Two rats (one in group A and one in group C) had catheter patency failure after acquisition, but before the end of PR testing, and as such, these rats were included in acquisition, but not PR analysis. One rat in group C underwent locomotor classification, but was not advanced to self-administration testing because of health concerns. Finally, one rat in group D was advanced prematurely to PR testing and thus was not included in group D intake or PR analysis.

Data Analysis. All statistical analyses were conducted by using PASW Statistics, version 18.0 (IBM, White Plains, NY). For acquisition analysis, cocaine intake on the session before acquisition and three postacquisition sessions was analyzed with three-way repeated-measures analysis of variance (RMANOVA). Classification (LCR or HCR), group (A and B or C and D), and session (within-subjects variable) were treated as independent variables, and intake was treated as the dependent measure. Postacquisition intake analysis in groups C and D was analyzed with three-way RMANOVA. Classification, dose (0.6 or 1.2 mg/kg/infusion), and session were treated as independent variables, and cocaine intake was treated as the dependent measure. The PR experiments were analyzed with three-way RMANOVA. Classification, dose (0.6 or 1.2 mg/kg/infusion), and session were treated as independent variables, and break-point-associated infusions were treated as the dependent measure. When main or interaction effects were revealed, independent samples t tests or one-way RMANOVA were used for post hoc analyses. When the assumption of sphericity was violated for a particular repeated-measures analysis, as revealed by Mauchly’s test statistic, tests of significance were based on the more conservative Huynh-Feldt corrected degrees of freedom. Superscript a (*) indicates Huynh-Feldt corrected values throughout the text. Between-groups analysis of break point-associated infusions on the first PR session were analyzed with independent samples t tests (group A versus C or group B versus D).

Drugs. The National Institute on Drug Abuse (Bethesda, MD) generously provided the (−)-cocaine hydrochloride used in these studies. For intraperitoneal injections, cocaine was dissolved in sterile saline (9.9% sodium chloride) at a concentration of 10 mg/ml and administered in a volume of 1 ml/kg. For intravenous infusions, cocaine was dissolved in sterile saline containing 1.7 USP units/ml heparin. To check catheter patency, sodium thiopental (Sigma-Aldrich, St. Louis, MO) was dissolved in saline and administered intravenously at 20 mg/kg. Drug weights refer to the salt.

Results

Groups A and B: Cocaine Self-Administration on a FR1 Schedule of Reinforcement. Groups A and B were given 20 sessions to acquire cocaine self-administration (0.6 mg/kg/infusion) by using a FR1 schedule of reinforcement. Overall, 15/16 rats in group A and 11/13 rats in group B met criteria within this timeframe with average times to acquisition of 6.5 ± 1.3 and 6.9 ± 1.2 days, respectively. Cocaine intake over the two sessions before acquisition and three acquisition sessions are shown in Fig. 1A. It should be noted that three rats in group A met criterion starting on the first day of the experiment and, consequently, do not have x − 2 and x − 1 values. Thus, n values are decreased for those time points in both the data analysis and Fig. 1, A and B. Analysis of cocaine intake over the session before acquisition (x − 1) and three acquisition sessions (x to x + 2) with three-way RMANOVA revealed a significant effect of session (F 3, 86 = 70.7; p < 0.001), but no other significant effects or interactions. Compared with sessions x − 1 and x, rats in both groups consumed significantly more cocaine on sessions x + 1 and x + 2, but intake was not significantly different between the groups.

Active and inactive lever responding in groups A and B over the two sessions before acquisition and three acquisition sessions are shown in Fig. 1B. Although the ratio of active to inactive lever responding was not part of the acquisition criterion, rats in both groups displayed clear lever discrimination upon meeting our intake criterion. By the third acquisition session, rats in groups A and B displayed a more than 10:1 ratio of active to inactive lever responding (Fig. 1B).

Groups A and B: Postacquisition Cocaine Self-Administration on a PR Schedule of Reinforcement. After acquisition, rats were immediately advanced to PR testing reinforced by either a higher dose (1.2 mg/kg/infusion; group A)

<table>
<thead>
<tr>
<th>TABLE 1</th>
<th>Experimental groups</th>
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<tbody>
<tr>
<td>Group</td>
<td>Acquisition Sessions and Cocaine Doses</td>
</tr>
<tr>
<td>A, n = 15*</td>
<td>3 × FR1, 0.6 mg/kg/inf</td>
</tr>
<tr>
<td>B, n = 11</td>
<td>3 × FR1, 0.6 mg/kg/inf</td>
</tr>
<tr>
<td>C, n = 15*</td>
<td>3 × FR1, 0.6 mg/kg/inf</td>
</tr>
<tr>
<td>D, n = 12</td>
<td>3 × FR1, 0.6 mg/kg/inf</td>
</tr>
</tbody>
</table>

* One rat in group A and one rat in group C were excluded after the acquisition phase and, as such, n values for PR testing were 14 for both groups A and C (see Materials and Methods: Exclusions and Exceptions).
or the training dose (0.6 mg/kg/infusion; group B) of cocaine. Analysis with three-way RMANOVA revealed a significant effect of session ($F_{4, 92} = 26.3; p < 0.001$), dose ($F_{1, 23} = 15.4; p = 0.001$), and a session × dose interaction ($F_{4, 92} = 5.7; p < 0.001$) interactions. Separate analysis of intake with one-way RMANOVA revealed a significant effect of group in C ($F_{2, 58} = 24.6; p < 0.001$), but no other significant effects or interactions. Relative to session $x - 1$, rats in both groups C and D consumed significantly more cocaine in sessions $x$ to $x + 2$.

In contrast to the acquisition sessions, analysis of the five additional FR1 sessions revealed significant effects of session ($F_{4, 92} = 21.6; p < 0.001$), dose ($F_{1, 23} = 15.4; p = 0.001$), classification × session ($F_{4, 92} = 2.9; p = 0.026$), and dose × session ($F_{4, 92} = 2.9; p < 0.001$) interactions. Separate analysis of intake with one-way RMANOVA revealed a significant effect of session in group C ($F_{2, 58} = 26.3; p < 0.001$), but not group D. Relative to the first session of 1.2 mg/kg/infusion cocaine self-administration (x + 3), rats in group C consumed significantly more cocaine in sessions 3 to 5 ($x + 3$ to $x + 7$; Fig 2A). Furthermore, pairwise comparisons revealed that whereas intake on session $x + 3$ was not significantly different between groups C and D, rats in group C consumed significantly more cocaine than rats in group D on each of the next four sessions (Fig 2A). Analysis of classification on each of the five additional FR1 sessions did not reveal significant differences in intake between LCRs and HCRs at any point (Table 2).

After the five additional FR1 sessions, groups C and D were advanced to PR testing reinforced by either 1.2 or 0.6...
TABLE 2
Cocaine self-administration in rats classified as LCRs or HCRs

<table>
<thead>
<tr>
<th>Group</th>
<th>Cocaine-Induced Locomotor Activity (cm/30 min)</th>
<th>Acquisition %</th>
<th>Average Intake (x to x + 2, mg/kg)</th>
<th>Average Intake (x + 3 to x + 7, mg/kg)</th>
<th>Average Break Point-Associated Infusions</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Median</td>
<td>Mean</td>
<td>(x to x + 2, mg/kg)</td>
<td>(x + 3 to x + 7, mg/kg)</td>
<td></td>
</tr>
<tr>
<td>A</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>LCR (n = 8)</td>
<td>2382</td>
<td>1331 ± 221</td>
<td>88 (7/8)</td>
<td>10.3 ± 1.4</td>
<td>15.5 ± 1.6</td>
</tr>
<tr>
<td>HCR (n = 8)</td>
<td>4204</td>
<td>532</td>
<td>100 (8/8)</td>
<td>11.3 ± 1.5</td>
<td>17.0 ± 1.8</td>
</tr>
<tr>
<td>B</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>LCR (n = 7)</td>
<td>2772</td>
<td>2065 ± 175</td>
<td>100 (7/7)</td>
<td>9.5 ± 1.8</td>
<td>11.3 ± 1.6</td>
</tr>
<tr>
<td>HCR (n = 7)</td>
<td>4914</td>
<td>749</td>
<td>67 (4/6)</td>
<td>8.9 ± 1.6</td>
<td>13.6 ± 1.8</td>
</tr>
<tr>
<td>C</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>LCR (n = 10)</td>
<td>2750</td>
<td>1322 ± 275</td>
<td>89 (7/9)</td>
<td>9.7 ± 1.5</td>
<td>15.4 ± 1.4</td>
</tr>
<tr>
<td>HCR (n = 10)</td>
<td>4272</td>
<td>349</td>
<td>70 (7/10)</td>
<td>9.5 ± 2.6</td>
<td>16.6 ± 1.6</td>
</tr>
<tr>
<td>D</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>LCR (n = 10)</td>
<td>2363</td>
<td>1271 ± 144</td>
<td>88 (7/8)</td>
<td>7.4 ± 0.8</td>
<td>9.1 ± 1.2</td>
</tr>
<tr>
<td>HCR (n = 10)</td>
<td>4375</td>
<td>644</td>
<td>56 (5/9)</td>
<td>10.0 ± 1.5</td>
<td>12.6 ± 2.2</td>
</tr>
</tbody>
</table>

The current study had two aims. First, we sought to identify conditions that reveal sensitization to the motivational effects of cocaine. Second, we sought to determine whether individual differences in acute cocaine-induced locomotor activity (i.e., LCRs and HCRs) predict this form of sensitization. Although this study did not reveal differences in this form of sensitization between LCRs and HCRs, we did identify conditions that reveal sensitization to the motivational effects of cocaine. Similar to other reports (Liu et al., 2005; Morgan et al., 2006), cocaine dose and exposure history were important for sensitization on the FR schedule. Thus, this study replicates previous findings on the important role of cocaine dose and exposure history for sensitization to the effects of contingent cocaine and provides new evidence that sensitization may occur very early during the drug-taking process.
Sensitization has been suggested to be important during the transition from recreational to compulsive cocaine use, and this idea has been argued most formally by proponents of the incentive-sensitization theory of addiction (Robinson and Berridge, 1993, 2001). In animals, sensitization to the locomotor-stimulating effects of drugs such as cocaine has been demonstrated for years and a great deal is known about its neurobiological mechanisms (Post, 1980; Zahniser et al., 1988; Kalivas and Duffy, 1993; Churchill et al., 1999; Grignaschi et al., 2004). However, it is not clear whether sensitization is seen in humans and nonhuman primates (Bradberry, 2007). One likely factor in this discrepancy is that humans actively take drugs, whereas animals passively receive drugs during locomotor sensitization paradigms. Consequently, demonstrating sensitization under paradigms of cocaine self-administration becomes extremely important for understanding the contribution of this phenomenon to complex emitted behaviors.

Roberts and colleagues have reported conditions that produce sensitization to the reinforcing effects of cocaine in rats evident by increasing break points on a PR schedule of reinforcement (Liu et al., 2005; Morgan et al., 2006). They found that a dose of 1.5 mg/kg/infusion cocaine produced the most robust increase in break points and limited initial exposure (e.g., 60–160 mg/kg total exposure before PR) was necessary to produce this effect. However, these studies used very specific conditions (animals housed in self-administration chambers with 12-h access to cocaine) and whether this type of sensitization could be produced under more widely used conditions (animals housed in separate rooms and 2-h sessions) remained unknown.

We have recently established a method for determining acquisition that allows for experimental manipulation of self-administration behavior during the earliest stages of the drug-taking process (Mandt et al., 2012). This method defines acquisition as the first of three consecutive sessions when a rat consumes ≥4 mg/kg cocaine during a 2-h FR1 session and aligns intake to that session (i.e., day x; see Fig. 1A). Previously, we found that rats consuming 1.2 mg/kg/infusion cocaine rapidly increased consumption from this point, whereas rats self-administering a range of lower doses exhibited more stable intake (Mandt et al., 2012). However, this new method of analysis was a product of that study and not part of the design. Thus, the present study used this method of analysis to assess sensitization to the motivational effects of cocaine after very limited initial cocaine exposure, similar to other reports (Liu et al., 2005; Morgan et al., 2006).

It is noteworthy that when rats were given limited exposure before PR testing (groups A and B), we found a robust replication of the Liu et al. (2005) study. In both that study and ours, break points did not differ on the first PR session regardless of cocaine dose. However, rats given the higher cocaine doses (1.5 mg/kg/infusion in our study or 1.2 mg/kg/infusion in ours) progressively increased break points over the next 5 to 10 sessions, whereas rats given the lower doses exhibited stable break points. It was possible that in our study the initial low break points for 1.2 mg/kg/infusion cocaine were an artifact of the switch in cocaine dose and reinforcement schedule, caused either by learning or dose-induced rate-limiting effects. Indeed, it has been reported that stability of responding on a PR schedule can take time to develop (Depoortere et al., 1993). However, given that rats exposed to five additional FR1 sessions at 1.2 mg/kg/infusion cocaine before PR testing (group C) exhibited significantly greater rates of responding (i.e., break points) than the limited exposure group on the first session, these differences are not likely explained by the need to learn to respond at higher rates. Furthermore, in the Liu et al. (2005) study, rats were trained with 1.5 mg/kg/infusion cocaine before PR testing with that same dose and still exhibited break points similar to those in our study on the first session. Thus, it also seems unlikely that the switch in cocaine dose produced competitive behaviors on the first session, preventing higher break points.

Rather, we believe the dose-dependent change in break points seen in the limited-exposure group reflects sensitization to the motivational effects of cocaine, similar to previous reports (Liu et al., 2005; Morgan et al., 2006). In addition, our results revealed a dose-dependent early escalation in cocaine consumption that occurs during 2-h FR1 sessions. Rats given limited initial exposure to a moderate dose of cocaine (0.6 mg/kg/infusion) during acquisition and then switched to a higher dose (1.2 mg/kg/infusion) increased consumption 45% over five additional sessions (see Fig. 2A). Although at first this may be surprising given reports in the literature of well trained animals’ ability to regulate intake when cocaine dose is varied (Pickens and Thompson, 1968; Gerber and Wise, 1989; Lynch et al., 1998; Panlilio et al., 2003), our animals received minimal training before the switch in cocaine dose. Furthermore, there are now many paradigms that model an escalation of consumption, and they all demonstrate conditions that produce instability of intake (i.e., escalation) (Carroll et al., 1989; Fitch and Roberts, 1993; Ahmed and Koob, 1998; Tornatzky and Miczek, 2000). Thus, we believe we have established an exciting new model of escalation in cocaine consumption that can be revealed under standard self-administration conditions (FR1 schedules, 2-h sessions, separate housing rooms, etc.).

It is tempting to conclude that the increase in consumption on the FR1 schedule is related to sensitization, as was revealed on the PR schedule, given the similar time course of the two effects (first five sessions at the higher dose; see Figs. 1C and 2A). In addition, rats that escalated consumption (group C) displayed significantly greater break points than rats in the limited-exposure group on the first PR session (Fig. 2C), possibly because some change in the motivational effects of cocaine had occurred as a result. However, until we are able to fully assess the reinforcing effectiveness of cocaine after this escalation on the FR schedule (e.g., with dose-consumption analysis), we are unable to know whether this truly is a sensitization effect revealed under the FR schedule.

Although we were able to identify conditions that produced sensitization to the motivational effects of cocaine, we did not observe differences between LCRs and HCRs in this form of sensitization. Thus, under conditions that produced sensitization, break points increased for both LCRs and HCRs. It should be noted that the relationship between locomotor sensitization and sensitization to the effects of self-administered cocaine is unclear. Although some studies have found that prior noncontingent stimulant exposure decreases latency to acquisition and increases break points for cocaine (Schenk and Partridge, 2000; Suto et al., 2002), other studies...
(including our own LCR/HCR study) have found that cocaine-induced locomotor sensitization does not predict acquisition or break points (Lack et al., 2008; Mandt et al., 2008) and is dissociable from the motivational effects after compulsive cocaine consumption (Ahmed and Cador, 2006). Furthermore, noncontingent cocaine administration, such as that used to induce locomotor sensitization, is known under some conditions to produce different neurobiological effects than contingent cocaine administration (Chen et al., 2008; Migueús et al., 2008). Finally, the amount of cocaine exposure during self-administration far exceeds that of multiple intermittent injections, making direct comparison of these findings difficult.

The current study replicated previous findings on the important role of cocaine dose and exposure history for sensitization to the effects of self-administered cocaine. In addition, we extended these findings to show that this form of sensitization can be revealed under standard training conditions and occurs very early in the drug-taking process. Furthermore, we revealed conditions (switch from a moderate to a high dose of cocaine) that lead to an early escalation of cocaine consumption on a FR1 schedule of reinforcement during 2-h sessions. Thus, this study provides important evidence that sensitization may be a process involved during the earliest stages of drug taking when experience is limited and may help explain the contribution of this phenomenon to the rapid progression of cocaine addiction in some people.

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Authorship Contributions

Participated in research design: Mandt, Gomez, Johnston, Zahnisner, and Allen.

Conducted experiments: Mandt, Gomez, and Johnston.

Performed data analysis: Mandt and Allen.

Wrote or contributed to the writing of the manuscript: Mandt, Zahnisner, and Allen.

References


Substance Abuse and Mental Health Services Administration (2011) Results from...
Cocaine Self-Administration History and Sensitization


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