Behavioral/Systems/Cognitive

Involvement of the Olfactory Tubercle in Cocaine Reward: Intracranial Self-Administration Studies

Satoshi Ikemoto
Behavioral Neuroscience Branch, National Institute on Drug Abuse, National Institutes of Health, Department of Health and Human Services, Baltimore, Maryland 21224

Cocaine has multiple actions and multiple sites of action in the brain. Evidence from pharmacological studies indicates that it is the ability of cocaine to block dopamine uptake and elevate extracellular dopamine concentrations, and thus increase dopaminergic receptor activation, that makes cocaine rewarding. Lesion studies have implicated the nucleus accumbens (the dorsal portion of the “ventral striatum”) as the probable site of the rewarding action of the drug. However, the drug is only marginally self-administered into this site. We now report that cocaine (60 or 200 mM in 75 nl/infusion) is readily self-administered into the olfactory tubercle, the most ventral portion of the ventral striatum. Cocaine (200 mM) was self-administered marginally into the accumbens shell but not into the core, dorsal striatum, or ventral pallidum. In addition, cocaine injections (200 mM in 300 nl) into the tubercle but not the shell or ventral pallidum induced conditioned place preference. Rewarding effects of cocaine in the tubercle were blocked by coadministration of dopamine D₁ or D₂ antagonists (1 mM SCH 23390 or 3 mM raclopride) and were not mimicked by injections of the local anesthetic procaine (800 mM). In conclusion, the tubercle plays a critical role in mediating rewarding action of cocaine.

Key words: reinforcement; conditioned place preference; nucleus accumbens shell; core; ventral pallidum; dorsal striatum; procaine; dopamine D₁ and D₂ receptor antagonists; SCH 23390; raclopride

Introduction

It is widely held that the rewarding effects of psychomotor stimulants such as amphetamine and cocaine are mediated by the nucleus accumbens (United States Department of Health and Human Services, 1999). Injections of amphetamine into the accumbens readily support robust self-administration (Hoebel et al., 1983; Phillips et al., 1994; Ikemoto and Sharpe, 2001) and induce conditioned place preference (Carr and White, 1986). Specific lesions of dopamine terminals in the accumbens disrupt self-administration of systemic amphetamine (Lyness et al., 1979) as well as conditioned place preference (Spyraki et al., 1982) induced by the drug. Although these findings suggest that amphetamine is mediated by the nucleus accumbens, evidence that cocaine is mediated in the same site is unconvincing.

Facilitation of dopamine transmission by blocking dopamine uptake (Ritz et al., 1987) plays a primary role in the rewarding effects of cocaine because cocaine is not rewarding to animals with selective pharmacological blockade of dopamine systems (De Wit and Wise, 1977; Ettenberg et al., 1982). One of the regions that receive dense dopaminergic projections is the nucleus accumbens. Several lines of evidence have suggested that the accumbens plays an important role in the rewarding effects of cocaine. Rewarding effects of systemic cocaine administration are decreased markedly by specific lesions of dopaminergic terminals in the accumbens (Roberts et al., 1977, 1979) or injections of dopamine antagonists into the accumbens (Maldonado et al., 1993; McGregor and Roberts, 1993; Baker et al., 1998). Systemic administration of cocaine increases extracellular dopamine in the accumbens (Imperato and Di Chiara, 1986; Pettit and Justice, 1989), among other regions (Hurd et al., 1988; Maisonneuve et al., 1990; Carboni et al., 2000; Sizemore et al., 2000). However, unlike amphetamine injections, direct injections of cocaine into the accumbens are only marginally rewarding (Carlezon et al., 1995; Liao et al., 2000; Rodd-Henricks et al., 2002) or are not rewarding at all (Goeders and Smith, 1983; Hemby et al., 1992).

Dopaminergic neurons from the ventral tegmental area also innervate densely to the olfactory tubercle (Voorn et al., 1986), localized just ventral to the accumbens. The olfactory tubercle is not merely an olfactory structure. With respect to its morphology, chemistry, and connections with other structures (Heimer, 1978), the tubercle has more in common with the accumbens and dorsal striatum than the primary olfactory cortex. It has been suggested that the tubercle and the accumbens should be conceptualized jointly as the ventral striatum (Heimer, 1978; Heimer et al., 1995). Thus, it is not surprising to see that dopaminergic manipulations of the tubercle have somewhat similar effects as those of the accumbens; it was recently found that injections of cocaine into the tubercle induce robust locomotion and rearing (Ikemoto, 2002; Ikemoto and Witkin, 2003). Because these behaviors may indicate the activation of a brain system that also mediates the rewarding effects of drugs and brain stimulation (Panksepp, 1982; Wise and Bozarth, 1987), we sought evidence of
the rewarding effects of cocaine in the subregions of the ventral striatum and related regions. Here I report that cocaine injections into the anteromedial portion of the tubercle are robustly rewarding. Cocaine injections into other regions, including the accumbens, dorsal striatum, and ventral pallidum, were either marginally rewarding or not rewarding at all.

Materials and Methods

Animals. One hundred eleven male Wistar rats (Harlan, Dublin, VA; 250–350 gm at the time of surgery) were used. Food and water were available ad libitum except during testing. The procedures were approved by the Animal Care and Use Committee of the National Institute on Drug Abuse Intramural Research Program and were in accordance with National Institutes of Health guidelines.

Surgery. Each animal was implanted with a unilateral guide cannula (24 gauge) that ended 1.0 mm above one of seven target sites (Ikemoto, 2002; Ikemoto and Wise, 2002). Efflux from central injections tends to follow a pressure gradient up the cannula shaft (Routtenberg, 1972). Accordingly, the cannulas for medial shell and anteromedial tubercle sites were inserted at a 20° angle from the other hemisphere through the midline (Fig. 1A) to minimize diffusion of drug solution to the core or shell, respectively. The cannulas were inserted at a 10° angle toward the midline for the site in the core, whereas guides were implanted vertically for injections in the anterolateral and posteromedial tubercle, ventral pallidum, and dorsal striatum. The incisor bar was set at 3.3 mm below the interaural line. The stereotaxic coordinates were 2.0 mm anterior to bregma (A), 2.0 mm lateral to the midline (L), and 8.5 mm ventral to the skull surface (V) (measured along the trajectory of the angled cannula) for anteromedial tubercle placements; A 2.0, L 1.7, and V 7.2 for shell placements; A 2.0, L 2.0, and V 8.2 for anterolateral tubercle placements; A 2.0, L 3.5, and V 6.7 for core placements; A 0.6, L 2.0, and V 8.7 for posteromedial tubercle placements; A 0.6, L 2.0, and V 7.7 for ventral pallidum placements; and A 1.0, L 3.0, and V 5.0 for dorsal striatum placements.

Drugs. (--)-Cocaine HCl (Research Triangle Institute, Research Triangle Park, NC), procaine HCl (Sigma, St. Louis, MO), the dopamine D1 antagonist R (+)-SCH 23390 HCl (Sigma), and the dopamine D2 antagonist S (--)-raclopride 1-tartarate (Sigma) were dissolved in artificial CSF consisting of (in mM): 148 NaCl, 2.7 KCl, 1.2 CaCl2, and 0.85 MgCl2, pH adjusted to 6.5–7.8.

Operant conditioning procedure. For operant testing, each animal was placed in a 30 × 22 × 24 cm chamber equipped with a lever (45 mm wide, made of thin metal protruding 20 mm from the wall) and a cue light just above the lever. Each rat’s 31-gauge injection cannula was connected by polyethylene tubing to a micropump (Ikemoto and Sharpe, 2001) hanging a few millimeters above the rat’s head. A lever press turned on the cue light for 1 sec and the micropump for 5 sec, dispensing a 75 nl infusion; additional lever presses were not rewarded until an additional 5 sec passed after the completion of the infusion. The maximum number of infusions available per session was limited to 60 to minimize the possibility of tissue damage. Sessions lasted 90 min or until the rats received a total of 60 infusions.

To determine whether cocaine would be self-administered into specific regions, 68 rats with no previous operant training received 60 mM cocaine injections in each of seven sessions 2–4, 200 mM in sessions 6 and 7, and vehicle in sessions 1, 5, and 8. To determine whether cocaine reward was mediated by dopaminergic mechanisms, the effects of dopamine D1 and D2 antagonists SCH 23390 and raclopride and the local anesthetic procaine on intratubercular self-administration of cocaine were evaluated. Six rats with anteromedial tubercle guides were trained initially to lever press for 200 mM cocaine infusions in two or three sessions. In the D1 antagonist experiment, the rats received 200 mM cocaine plus 1 mM SCH 23390, 200 mM cocaine, and vehicle over three sessions. In the D2 antagonist experiment, the animals received 200 mM cocaine plus 3 mM raclopride, 200 mM cocaine, and vehicle over three sessions. In the local anesthesia experiment, they received 800 mM procaine, 200 mM cocaine, and vehicle over three sessions. The order of testing the injection treatments in each experiment was counterbalanced among the subjects.

Two-lever discrimination procedure. Six rats with no previous operant training were placed in operant chambers with two retractable levers. The chambers were identical to the one-lever chambers described above, except that two levers were placed on a chamber wall. A response on the “active” lever resulted in a 5 sec infusion (75 nl in volume), turned on a cue light above the lever, and retracted both levers for 30 sec. A response on the “inactive” lever did not deliver infusions but retracted both the active and inactive levers for 30 sec. The left lever was designated the active lever for three rats and the inactive lever for the other three. Responding on the active lever produced vehicle infusions in session 1 and 200 mM cocaine infusions in sessions 2–4. Sessions lasted 90 min or until the rats received a total of 60 infusions. Numbers of responses on each lever were recorded.

Conditioned place preference procedure. The place-conditioning chambers were configured as previously described (Ikemoto and Wise, 2002). In session 1, experimentally naïve rats were placed individually in the place-conditioning chamber for 15 min without any treatment; they had access to both compartments, and the time spent in each compartment was recorded. In sessions 2–5, they were placed individually in one of the compartments and received an intracranial injection (300 nl delivered over 60 sec) (Ikemoto, 2002; Ikemoto and Wise, 2002) of cocaine (200 mM) or vehicle. An additional 30 sec period elapsed before the injection cannula was removed. The animals remained in the compartment for an additional 5 min after the injection. Infusions of cocaine and vehicle were alternated over four sessions, as was the placement of each rat in one of the two compartments. The order of injection treatments and the assignment of the compartments with injection treatments were counterbalanced among the subjects. In session 6, the untreated rats were placed individually in the chamber and given access to both compartments; the time spent in each compartment was recorded for 15 min. Sessions were separated by 24 hr.

Histology. When each rat completed the experimental procedure, the brain was removed and processed as described previously (Ikemoto, 2002; Ikemoto and Wise, 2002).

Statistical analyses. Effects of 60 mM cocaine on infusion rates were analyzed with 2 × 2 within-subjects ANOVAs with session and treatment (vehicle sessions 1 and 5 vs cocaine sessions 3 and 4). Effects of 200 mM cocaine were also analyzed with 2 × 2 within-subjects ANOVAs with session and treatment (vehicle sessions 5 and 8 vs cocaine sessions 6 and 7). The data on two-lever responses were analyzed with a 2 × 4 within-subjects ANOVA with lever (active vs inactive lever) and session (1–4). Conditioned place preference data (times spent in compartments) were analyzed with 2 × 2 within-subjects ANOVAs with treatment (before vs after conditioning) and compartment (cocaine - vs vehicle-associated for each injection site separately. Effects of dopamine antagonists or procaine on infusion rates were analyzed with one-way within-subjects ANOVAs over three treatments (experimental treatment, cocaine, and vehicle).

Results

Cocaine self-administration

We used a self-administration procedure to determine which region around the ventral striatum mediates the most robust rewarding effects of cocaine. Rats with indwelling cannulas aimed at one of seven regions (the anteromedial, anterolateral, and posteromedial tubercle, core, dorsal striatum, and ventral pallidum; Fig. 1A–C) were placed in operant chambers and were trained to lever press for cocaine infusions. Rats learned to self-administer 60 mM cocaine into the anteromedial tubercle (Fig. 2A; F1,15 = 10.31; p < 0.01). More robust self-administration was induced by 200 mM cocaine into the same region (F1,15 = 9.22; p < 0.01). Both 60 and 200 mM cocaine supported persistent self-administration lasting the entire session or until the maximum number of infusions was obtained. However, vehicle infusions did not sustain self-administration (Fig. 2B). In the medial shell, 60 mM cocaine was not self-administered, whereas 200 mM cocaine was self-administered moderately (F1,9 = 8.15; p <
Similar concentration-dependent effects of cocaine were found in the anterolateral tubercle ($F_{(1,5)} = 9.03; p < 0.05$). In the posteromedial tubercle, the low concentration of cocaine induced persistent self-administration ($F_{(1,9)} = 20.54; p < 0.005$), whereas the high concentration did not. Cocaine did not support self-administration in the core, dorsal striatum, or ventral pallidum.

0.05). Similar concentration-dependent effects of cocaine were found in the anterolateral tubercle ($F_{(1,5)} = 9.03; p < 0.05$). In the posteromedial tubercle, the low concentration of cocaine induced persistent self-administration ($F_{(1,9)} = 20.54; p < 0.005$), whereas the high concentration did not. Cocaine did not support self-administration in the core, dorsal striatum, or ventral pallidum.
Roles of local anesthesia and dopamine receptors in cocaine reward

To examine the role of the local anesthetic effect of cocaine in the reward, we tested the drug procaine, which has similar anesthetic properties, to find out whether it would mimic cocaine reward. Although procaine has a chemical structure similar to that of cocaine, the drug is much less potent as a dopamine uptake blocker (Ritz et al., 1987; Woodward et al., 1995). Rats receiving 800 mM procaine, which is a concentration equipotent to 200 mM cocaine in blocking Na \(^+\) channels (Creveling et al., 1983; Reith et al., 1986; Wilcox et al., 1999), into the anteromedial tubercle decreased self-administration rates (Fig. 5A; \(F_{(2,10)} = 5.02; p < 0.05\)). The temporal pattern of procaine self-administration resembled that of vehicle self-administration (Fig. 5B); that is, cocaine injections induced extinction-like self-administration. To examine whether the rewarding effects of cocaine are mediated by dopaminergic mechanisms, the effects of dopamine receptor antagonists on cocaine self-administration were determined. Co-administration of the dopamine D\(_1\) or D\(_2\) receptor antagonists SCH 23390 (1 mM) and raclopride (3 mM) with cocaine decreased self-administration rates significantly (Fig. 5A; \(F_{(2,10)} = 8.95; p < 0.01\); \(F_{(2,10)} = 4.98; p < 0.05\), respectively). When the cocaine solution containing the D\(_1\) or D\(_2\) antagonist was given, self-administration did not typically persist throughout the session (Fig. 5B).

Two-lever discrimination

When active lever responding triggered vehicle infusions in session 1, the rats did not respond on the active lever more than the inactive lever (Fig. 3). When active lever responding triggered cocaine infusions in sessions 2–4, the animals responded on the active lever significantly more than the inactive lever. These observations are confirmed by a significant interaction between lever and session (\(F_{(3,15)} = 4.92; p < 0.05\)) followed by simple effects comparisons between the two levers.

Conditioned place preference

To provide independent evidence for the apparent rewarding action of cocaine in the tubercle, we used a conditioned place preference procedure to assess the conditioned effects of cocaine in the absence of the unconditional effects of the drug. After the association of one test chamber compartment with cocaine injection and the other with vehicle, we measured the animals’ preference for each compartment under a cocaine-free state. Rats receiving injections into the anteromedial tubercle spent more time in the cocaine-associated compartment (Fig. 4; a significant preference procedure to assess the conditioned effects of cocaine. This conclusion is elaborated below with considerations of the issues concerning behavioral assay and pharmacology and the anatomy of cocaine reward.

Arousing effects of cocaine

Injections of cocaine into the tubercle induce marked, unconditional locomotion and rearing (Ikemoto, 2002). The effective trigger zones of such responses parallel those in which cocaine induced rewarding effects. Significant locomotion and rearing are induced after cocaine administration into the anteromedial, anterolateral, or posterior tubercle or medial shell but not the lateral shell, core, or ventral pallidum (Ikemoto, 2002). Concentrations of cocaine that induce such responses are similar to the concentrations that induce rewarding effects. These findings arguably suggest that cocaine-induced arousal alone could be responsible for sustained lever press responding.

Two lines of evidence suggest that cocaine administration into the tubercle is reinforcing. When given a choice between two levers, rats responded selectively to the lever that delivered co-
The rewarding effects of intratubercle cocaine administration appear to depend on dopamine transmission, requiring both D₁ and D₂-like receptors. Coadministration of the D₁ and D₂ antagonists with cocaine did not sustain self-administration throughout the session. The pattern of the self-administration resembled that of vehicle self-administration.

Neuroanatomy of cocaine reward

Comparisons of the effects of cocaine among injection sites indicate that the tubercle, particularly the anteromedial portion, mediates the most robust rewarding effects of the drug. Cocaine administration into the medial shell was less rewarding than administration into the tubercle. The present results are consistent with recent findings in that cocaine injections into the medial shell but not the core are moderately rewarding, as reflected in rates of self-administration (Carlezon et al., 1995; Roddis-Henricks et al., 2002). Liao et al. (2000) reported that cocaine injections at 600 mM (bilaterally) but not 300 mM into the shell induced conditioned place preference (CPP). The present study found that 200 mM cocaine injections into the tubercle but not the shell induced CPP. The fact that a higher cocaine concentration was needed to induce CPP via shell injections (Liao et al., 2000) than tubercle injections (present study) is consistent with the present self-administration findings.

A previous study (Gong et al., 1996) suggests that bilateral injections of cocaine into the ventral pallidum induce CPP. The present study, on the other hand, found that injections of cocaine into the ventral pallidum did not induce CPP or support self-administration. Because there were no injection site controls in the study by Gong et al. (1996), the exact action site of cocaine studied is unclear. It is possible that ventral pallidal CPP (Gong et al., 1996) resulted from the diffusion of cocaine into the tubercle, which is located just ventral to the ventral pallidum. Indeed, the study by Gong et al. (1996) used a higher concentration (300 mM) and a larger volume (500 nl/hemisphere × 2) of cocaine with a longer conditioning trial duration (30 min) than those used in the present study (200 mM concentration, a single 300 nl volume, 5 min conditioning trial duration for place conditioning: 60 and 200 mM concentrations and 75 nl volume/infusion for self-administration).

Goeders and Smith (1983, 1986) and Goeders et al. (1986) showed that rats self-administer cocaine into the medial prefrontal cortex. Cocaine self-administration into the medial prefrontal cortex differs markedly from self-administration into the tubercle (examined in the present study) with respect to effective concentrations and rates of self-administration. Although much lower concentrations (between 0.5 and 1 mM) of cocaine are self-administered into the prefrontal cortex than into the tubercle, rates of prefrontal cortex self-administration are remarkably low, at ~0.05 infusions/min (Goeders et al., 1986). This low rate may reflect a small rewarding effect of cocaine in the medial prefrontal cortex. Indeed, subsequent studies suggest that lesions of dopaminergic terminals or cell bodies in the medial prefrontal cortex do not disrupt or even facilitate the acquisition and maintenance of intravenous self-administration of cocaine (Martin-Iverson et al., 1986; Schenk et al., 1991; McGregor et al., 1996; Weissenborn et al., 1997). The medial prefrontal cortex does not appear to mediate the rewarding effects of systemic cocaine administration.

Role of the nucleus accumbens in cocaine reward

There have been many studies devoted to the investigation of the role of the accumbens in cocaine reward and addiction, based on...
the assumption that the accumbens plays a major role in these processes. In the present study, cocaine administration was more rewarding in the tubercule than in the accumbens. In light of this finding, it is necessary to reconsider the exact role of the accumbens in cocaine reinforcement. We will consider possible roles of the accumbens in the rewarding effects of cocaine in terms of an integrative zone and a trigger zone. It is important to recognize the difference between the anatomical substrates initiating reward process after direct interaction with cocaine (trigger zones) and the anatomical substrates integrating reward signals initiated somewhere else (integrative zones). The accumbens appears to play an important role in incentive learning involving a number of reinforcers, such as food, water, and sexual intercourse (Robbins and Everitt, 1996; Ikemoto and Panksepp, 1999). Therefore, although the accumbens may or may not be a major trigger zone for the rewarding effects of cocaine, it is likely that the accumbens is important for integrating the reward signals of cocaine, which may be triggered elsewhere. Such integrative zones likely include many brain regions, for example, the ventral pallidum.

The accumbens may still be a major trigger zone of cocaine after systemic treatment despite marginal rewarding effects after direct injections. Two factors should be considered: the local anesthetic action of the drug and delivery method used to administer cocaine. The local anesthetic action of cocaine may have hindered the role of the accumbens as a trigger zone for the rewarding effects of cocaine. A recent study (Ikemoto and Witkin, 2003) suggests that core neurons are more vulnerable to the local anesthetic effects of cocaine than tubercule neurons. Injections of the local anesthetic procaine into the core inhibit spontaneous locomotion as well as amphetamine-induced locomotion, whereas injections of the same procaine concentrations into the anteromedial tubercule do not disrupt spontaneous or amphetamine-induced locomotion (Ikemoto and Witkin, 2003). The concentrations of procaine inducing such locomotor disruption in the core are equipotent in inhibiting sodium channels to the cocaine concentrations that are rewarding in the tubercule.

The manner in which cocaine affects accumbens neurons via systemic administration likely differs from the effect of the drug when it is administered centrally. Systemic administration, which saturates the entire body with cocaine, makes the drug available for much longer periods, with a more gradual concentration change than intracranial administration, which diffuses rapidly to an insignificant concentration. Lower concentrations (at which cocaine has no local anesthetic action) may be rewarding after prolonged exposure to the accumbens. Thus, it is quite possible that the accumbens plays a more important role in mediating the rewarding effects of systemic cocaine than that found in the present intracranial administration study.

Future investigation

Only a handful of studies have examined the role of the tubercule in reward-related functions (Prado-Alcala and Wise, 1984; Fibiger et al., 1987; Clarke et al., 1990; Kornetsky et al., 1991; Stein and Fuller, 1992; Porrino et al., 2002). Because the tubercule appears to be involved in positive reinforcement of cocaine, the roles of the tubercule in the rewarding effects of other drugs of abuse, as well as learning and motivation involving natural rewards such as food and sex, should be examined. Further studies are needed to characterize the role of the tubercule in cocaine reward and addiction (e.g., molecular mechanisms). It is particularly important to examine the contribution of the tubercule to the rewarding effects of systemic cocaine administration. Although the lesion study done by Roberts et al. (1979) has been taken to support the role of the accumbens in cocaine reward, the lesions in question were quite extensive, affecting the tubercule as well as the accumbens (Roberts et al., 1979). It is necessary to conduct studies examining selective lesions in the accumbens or tubercule to evaluate the contribution of these regions in cocaine reward.

References


Ikemoto S, Sharpe LG (2001) A head-attachable device for injecting nano-


