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## RAPID COMMUNICATION

# Real Time Detection of Acute (IP) Cocaine-Enhanced Dopamine and Serotonin Release in Ventrolateral Nucleus Accumbens of the Behaving Norway Rat

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BRODERICK, P. A., E. P. KORNAK, JR., F. ENG AND R. T. WECHSLER. Real time detection of acute (IP) cocaine-enhanced dopamine and serotonin release in ventrolateral nucleus accumbens of the behaving Norway rat. PHARMA-COL BIOCHEM BEHAV 46(3) 715-722, 1993. - Cocaine (10 mg/kg), administered intraperitoneal (IP), was studied for its effects on dopamine (DA) and serotonin (5-HT) release in ventrolateral nucleus accumbens (vlNAcc) of conscious and behaving male, virus-free, Sprague-Dawley rats with in vivo electrochemistry (voltammetry). Miniature stearate probes detected DA and 5-HT release, on line and within a temporal resolution of seconds. Psychostimulant behaviors, in the form of four behavioral components (i.e., the classically DA-dependent behaviors of locomotor activity [ambulations], rearing, and stereotypy, and a 5-HT-ergic behavior, central ambulations) were studied concurrently with infrared photobeam detection. The results show that (IP) cocaine significantly increased vlNAcc DA release (p < 0.0001) and 5-HT release (p < 0.0001) 0.0012). Each of the four parameters of cocaine-induced psychostimulant behavior was concurrently and significantly increased as well (ambulations: p < 0.0001; rearing: p < 0.0008; stereotypy: p < 0.0004; central ambulations: p < 0.0082). Moreover, exactly coincident data points for DA and 5-HT release occurred 10 and 40 min after (IP) cocaine administration. Cocaine-induced DA and 5-HT release were highly and positively correlated during the first hour of study (p < 0.01). As expected, increased DA release in vINAcc after cocaine administration was significantly and positively correlated with classically DA-dependent behaviors (first- and second-hour effects) (p < 0.01) and with the 5-HT-ergic behavior, central ambulations (p < 0.01). Also, cocaine-induced 5-HT release was significantly and positively correlated with 5-HT behavior (p < 0.01). However, not as expected, classically DA-dependent behaviors were more positively correlated with cocaineinduced 5-HT release in vINAcc throughout the two-hour period of study. Thus, the present findings show that 5-HT is a comediator with DA in the cocaine response in vINAcc. Importantly, 5-HT may signal the known DA response to cocaine.

Cocaine	Psychostimulant behavior	Dopamine	Serotonin	Ventrolateral nucleus accumbens (vlNAcc)
In vivo electrochemistry (voltammetry)		Anxiety	Agoraphobia	

THE relevance of  $A_{10}$  nucleus accumbens (NAcc) dopaminergic (DAergic) function to brain reward seems sufficiently explicit. Particularly, in paradigms of classical self-stimulation reinforcement phenomena two distinct models for the neuronal *modus operandi* of DA in the acute reinforcement process have been proposed. The first has been called "the two neuron" model, in which descending medial forebrain bundle (MFB) fibres synapse in  $A_{10}$  somatodendrites, ventral tegmental area (VTA), and subsequently give rise to ascending projections to NAcc. In this model, DA fibres may *directly carry* 

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the reward signal (56,64). In the second model, the "modulatory model," the reward signal may be *carried* by non-DAergic fibres via an *oblique* rather than direct  $A_{10}$  neuronal pathway from  $A_{10}$  somatodendritic VTA to nerve terminal NAcc (23, 65; cf. 58 for review).

Interestingly, the relevance of DAergic functionality in  $A_{10}$  circuits to *cocaine-induced brain reward* also seems explicit. The first studies to describe this cocaine- $A_{10}$ -DA relationship (52,53) precede several studies of supporting evidence, predicated on the acute (single dose) effects of cocaine in NAcc (6,7,29,34,62; cf. 33 for review), and also precede evidence for an acute cocaine-DA functionality in  $A_{10}$  somato-dendrites, VTA as well (3,8,11,21,46).

At first glance, since only the neurotransmitter DA is involved in these previous studies, acute cocaine brain reward could be analogous to the "two neuron" model of brain reward. New neurochemical data, though, implicate serotonin (5-HT) as a co-modulator mechanism for the SC and IV cocaine-induced DAergic response in A<sub>10</sub> neuronal circuits (6-10). Moreover, electrophysiological studies show a role for a 5-HT-ergic mediation in the mechanism of action of acutely administered cocaine (16,17,20,42,62). Other studies have assayed acute cocaine-induced 5-HT-ergic effects on reuptake processes, in vitro (22,31,54,55); release processes, in vivo (7); behavior, in vivo (18,47,48); and synthesis processes post mortem (24). Furthermore, there is evidence that acute iontophoretic applications of DA and 5-HT in combination more efficiently mimic the effect of cocaine on NAcc neurons than application of either DA or 5-HT alone (63). Thus, a concept of acute cocaine-induced brain reward encompassing a 5-HTergic component is unfolding. Such a concept could interestingly be analogous to the "modulatory model" of brain reward.

Therefore, we examine here the effect of acute cocaine on 5-HT release concurrently with DA release in ventrolateral (vl)NAcc, on line with cocaine-induced psychostimulant behavior, when cocaine is administered by the intraperitoneal (IP) route. Importantly, the route of administration of cocaine has been shown to cause important differences in the determination of its consequent acute behavioral (37) and neurochemical effects (9). Moreover, release mechanisms are primarily addressed. Previous studies have shown that cocaine's neurochemical effects are dependent on impulse flow (7, 12,24). Cocaine utilizes release (41) as well as reuptake inhibitory processes (50) presynaptically.

#### MATERIALS AND METHODS

Animals

The studies were done in unrestrained freely moving male Sprague-Dawley rats (Charles River, Kingston, NY) (weight range 362-446 g at the time of the in vivo electrochemical and behavioral studies). The animals were fed Purina Rat Chow and water ad lib and were group housed before surgery and individually housed after surgery. A 12-h dark/light cycle was maintained both during the housing of the laboratory rats and throughout the experimental studies. The rats were tested free from the following viruses: Sendai Virus, Kilham Rat Virus, Reo Virus Type 3, Sialodacryoadenitis Virus, Rat Corona Virus, Toolan's H1 Virus, Micro Plasma Pulmonis Virus, Lymphocytic Choriomeningitis Virus, Hantaan Virus, and Encephalitozoon Cuniculi Virus.

#### Surgery

Pentobarbital Na (50 mg/kg IP) was the general anesthetic employed to produce surgical anesthesia. A booster injection of pentobarbital Na (0.10 cc of the same solution) was administered once after the first two hours of surgery, and another booster (0.05 cc) was administered each of the two subsequent hours of surgery to maintain adequate anesthesia. Rats were tested for an absence of corneal, pinnal, and leg flexion responses. Body temperature was continuously monitored with a rectal probe thermometer (Fisher Scientific, Fadem, NJ) and was maintained at 37  $\pm$  0.5°C with an aquamatic K module heating pad (American Hospital Supply, Edison, NJ). Rats were stereotaxically implanted with stearate working microelectrodes in vlNAcc (anterior-posterior [AP] = +2.6, medial-lateral [ML] = +2.5, dorsal-ventral [DV] = -7.3) (44). Stereotaxic equipment was purchased from Kopf Stereotaxic (Tujunga, CA). Ag/AgCl reference microelectrodes and stainless steel auxiliary microelectrodes were placed in contact with cortex. The working (indicator), reference, and auxiliary microelectrodes were held in place with dental acrylic (Kadon Cavity Liner, Caulk, Becker Parkin Dental Supply Co. Inc., NY). Animals recovered in an appropriately bedded Plexiglas chamber  $(12'' \times 12'' \times 18'')$ . Animals were treated with physiological saline immediately and for two days after surgery. Each animal was treated with care throughout the surgical procedures and the studies.

#### In Vivo Electrochemical (Voltammetric) Biotechnology

The methods for the manufacture of each of the three in vivo electrochemical microelectrodes have been published by this laboratory (4). The methodology previously described includes the conditioning or preconcentration steps for the working microelectrode and the specifications for the formulation and synthesis of the stearic acid carbon paste. A review of the historical and technical aspects of the field of in vivo electrochemistry is referenced (5). Electrocatalytic interactions between DA and ascorbic acid (AA) have been reported with a stearic acid macroelectrode in vitro (25), but more recent reports show that these interactions are insignificant in neuronal tissue in vivo when a stearic acid microelectrode is used (2). Precalibration and postcalibration procedures were done as previously described (6).

In vivo voltammetric (semiderivative) studies on conscious rats were begun approximately 9 to 14 days after the aseptic surgical procedures were performed. On each experimental day, an animal was placed in a faradaic, Plexiglas chamber  $(24'' \times 18'' \times 23.5'')$ . The three-microelectrode assembly, enclosed within the animal's prosthetic acrylic cap, was connected to a CV37 detector (BAS, West Lafayette, IN) by means of a mercury commutator (Brain Research Instruments, Princeton, NJ), a flexible cable, and a mating connector (BJM Electronics, Staten Island, NY). The CV37 was electrically connected to a Minigard surge suppressor (Jefferson Electric, Magnetek, NY) which was then connected to an isolated electrical ground. Stable in vivo electrochemical signals for DA and 5-HT were evident before cocaine (10 mg/kg IP) was administered. Cocaine (Sigma, St. Louis) was dissolved in doubly distilled water, and solutions were made fresh on the day of each study.

#### Behavior

On each day of the cocaine study, each animal was placed in the faradaic copper-enclosed Plexiglas chamber described above. The behavioral chamber was novel to each animal, although each animal was habituated (i.e., had essentially completed exploratory behavior) before cocaine injection. Moreover, the behavioral chamber was equipped with side by side double doors (W 15.75"  $\times$  H 16") to enable a facile injection procedure. A series of infrared photobeams was encased in aluminum frames around the chamber's perimeter. When activated with an IBM computerized circuit, these infrared photobeams detected the animal's position in the behavioral chamber on an X-Y axes positional basis. Thus, multiple concurrent measures of the animal's activity were simultaneously assayed. The specific activities of each animal assayed were the "classically DA-dependent" behaviors-that is, 1) ambulations (locomotor activity or running ["running" is forward locomotion interacting with the maintenance of a horizontal position of the head, without lateral turning (61)]), 2) rearing behavior [maximal upward vertical movement of the head involving recruitment of the body, without any forward or lateral movement (61)], 3) fine movements (combined stereotypic movements of head bob, sniffing, and grooming)-in addition to a 5-HT-ergic behavior, and 4) central ambulations (locomotor activity into the central part of the chamber). [Central ambulatory behavior is called agoraphobic (thigmotactic) inhibition and indicates reduced fear on the part of the animal (26)]. The status of the infrared photobeams was sampled every 100 ms. The system is a modified version of an Activity Pattern Monitor (San Diego Instruments, San Diego). Data were collected as measures of concurrent and separate activities for 10-min time periods.

#### Confirmation of Microelectrode Placement

Following the completion of the study, the prosthetic acrylic cap was removed from the skull while the animal was under Na pentobarbital anesthesia. Placement of working microelectrodes in vlNAcc was confirmed by the potassium ferrocyanide in 10% formalin blue dot method with transcardial perfusion (80 ml saline). The precise electrical specifications for deposition of the blue dot in vlNAcc was 50  $\mu$ A current in a 30-s time period. Virtually no damage to brain tissue occurred. The working microelectrode was postcalibrated for in vitro electrochemical detection of DA and 5-HT.

#### Statistics

Each component of the psychostimulant behavior monitored, in addition to DA and 5-HT release assayed, was tested for statistically significant differences between pre- and postcocaine (10 mg/kg IP) (same animal control) by standard repeated-measures analysis of variance (ANOVA) (Statview, Brain Power Inc., Calabasas, CA). ANOVAs were followed by post hoc tests, Fisher PLSDs (least square differences), and the Scheffe F test (Statview, Brain Power Inc.) to determine hourly statistically significant differences. Statistically significant differences were also calculated on the individual time course data points by 95% confidence limits (95% CL), setting the p value at p < 0.05. Changes in DA and 5-HT values after (IP) cocaine treatment vis-à-vis untreated (same animal) controls are presented as percent change, whereas behavioral data are presented as frequency or number of behavioral events. Control is represented as 100%.

Since the actual detection time for DA is 10-15 s, the percent change in synaptic concentrations of DA at each data point, post-cocaine, represents a 10-15-s current change (pA) from baseline (i.e., actual current detected within a discrete synaptic environment of vlNAcc at the working [indicator] microelectrode surface within a 10-15-s time period is measured). The same principle of in vivo electrochemical detection applies for 5-HT. Since the actual detection time for 5-HT at each data point is 10-13 s, the percent change in synaptic concentrations of 5-HT at each data point post-cocaine represents a current measurement within the same discrete synaptic environment as that for DA, within vlNAcc within a 10-13-s time period.

Finally, cocaine-induced DA and 5-HT release in vlNAcc and consequent psychostimulant behavior were studied for statistically significant correlative value by the Pearson product-moment coefficient of correlation (r), simple and polynomial regression (Statview, Brain Power Inc.); corresponding  $z_f$  values were derived from the table of z for values of r from 0.0 to 1.0.

#### RESULTS

Figure 1 shows the effect of cocaine (10 mg/kg IP) on synaptic concentrations of DA and 5-HT in the vlNAcc. Cocaine (10 mg/kg IP) significantly increased the in vivo electrochemical signal for DA, F(2, 10) = 96.604, p < 0.0001, N = 6. Post hoc analysis further shows that there were statistically significant differences from baseline in each hour of the two hours tested (Fisher PLSD = 4.852, Scheffe F = 33.062and 95.616, first and second hours respectively). Dopamine release was significantly increased 110% (p < 0.05, 95% CL) over baseline within 10 min and was maximally increased 136% (p < 0.05, 95% CL) over baseline within 90 min after cocaine administration (baseline = 100%).

Also in Fig. 1, cocaine (10 mg/kg IP) simultaneously and significantly increased the in vivo electrochemical signal for 5-HT, F(2, 10) = 14.135, p < 0.0012, N = 6. Post hoc analysis further shows that there were statistically significant differences from baseline in the first hour of study (Fisher PLSD = 6.048, Scheffe F = 13.125 and 0.886, first and second hours respectively). 5-HT was significantly increased to 108% (p < 0.05, 95% CL) over baseline (100%) within 10 min and was maximally increased to 123% (p < 0.05, 95% CL) over baseline within 40 min after cocaine administration.

Moreover, cocaine's colocalized effects on DA and 5-HT release in vlNAcc were significantly and positively correlated in the first hour of study (Pearson product:  $r_{(a)} = 0.833$ , z = 1.1881, p < 0.01). Interestingly, exact coincident points occurred at the 10-min and 40-min marks of the time course study after (IP) cocaine administration.

Figure 2 shows the concurrent effect of cocaine (10 mg/kg IP) on the frequency (number) of ambulations (locomotor activity) in the same group of animals in which neurochemistry was assayed. Cocaine (10 mg/kg IP) significantly increased the frequency of ambulations, F(2, 10) = 27.502, p < 100.0001, N = 6). Post hoc analysis further shows that there were statistically significant differences from baseline in the first hour of the two hours tested (Fisher PLSD = 193.583. Scheffe F = 26.852 and 3.581, first and second hours respectively). Thus, cocaine's effects on hyperactive behavior progressively declined in the second hour of study with the exception of the 90-min mark of the time course, at which time ambulatory behavior abruptly rose and then fell. Frequency of ambulations was significantly increased to 943  $\pm$  130 photobeam interruptions (p < 0.05, 95% CL) from a baseline of 187  $\pm$  35 within 10 min, and maximally increased to 1070  $\pm$ 124 photobeam interruptions (p < 0.05, 95% CL) within 20 min after cocaine administration.

Figure 3 shows the concurrent effect of cocaine (10 mg/kg IP) on the frequency (number) of rearings. Cocaine (10 mg/ kg IP) significantly increased the rearing frequency, F(2, 10) = 15.749, p < 0.0008, N = 6. Furthermore, post hoc analysis shows that there were statistically significant differences

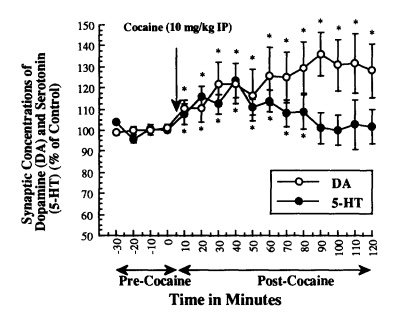


FIG. 1. The effects of cocaine (10 mg/kg IP) on concurrent DA and 5-HT release in vlNAcc in male, virus-free, Sprague-Dawley rats (N = 6) (cf. text for analysis of variation statistics). Detection limits for synaptic concentrations of DA and 5-HT as low as 5 nmol and 1 nmol, respectively, are currently possible with this biotechnology. \*p < 0.05 (95% confidence limits).

from baseline in the first hour of the two hours tested (Fisher PLSD = 9.056, Scheffe F = 15.592 and 2.661, first and second hours respectively). Cocaine's effects on rearing were dissipated in the second hour except at the 90-min mark of the time course, at which time behavior abruptly increased and

subsequently decreased. Rearing frequency was significantly increased to  $28 \pm 5$  photobeam interruptions (p < 0.05, 95% CL) from a baseline of  $3 \pm 0.8$  within 10 min and maximally increased to  $31 \pm 3$  (p < 0.05, 95% CL) within 50 min after cocaine administration.

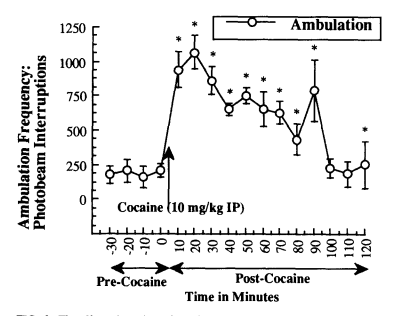


FIG. 2. The effect of cocaine (10 mg/kg IP) on the frequency of ambulations (locomotor activity or running behavior) in the same group of Norway rats (cf. text for analysis of variation statistics). Baseline photobeam interruptions were 187  $\pm$  35 (representing habituated behavior). \*p < 0.05 (95% confidence limits).

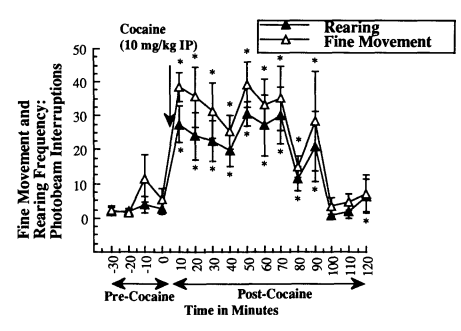


FIG. 3. The effect of cocaine (10 mg/kg IP) on frequency of rearing behavior and fine movement behavior of head bob, sniff, and groom in the same group of Norway rats (cf. text for analysis of variation statistics). Baseline photobeam interruptions were  $3 \pm 0.8$  and  $5 \pm 2$ , respectively (representing habituated behavior). \*p < 0.05 (95% confidence limits).

Figure 3 also shows the concurrent effect of cocaine (10 mg/kg IP) on the frequency (number) of fine movements. Cocaine (10 mg/kg IP) significantly increased the frequency of fine movements, F(2, 10) = 18.833, p < 0.0004, N = 6. Post hoc analysis further shows that there were statistically significant differences from baseline in the first hour of the two hours tested (Fisher PLSD = 10.532, Scheffe F =18.402 and 2.486, first and second hours respectively). Cocaine's effects on stereotypy dissipated in the second hour of study with the exception of the behavior seen at the 90-min mark which abruptly rose and momentarily fell. Frequency of fine movements was significantly increased to  $38 \pm 4$ photobeam interruptions (p < 0.05, 95% CL) from a baseline of  $5 \pm 2$  within 10 min and was maximally increased to  $39 \pm 6$  (p < 0.05, 95% CL) within 50 min after cocaine administration.

Figure 4 shows the concurrent effect of cocaine (10 mg/kg IP) on the frequency (number) of central ambulations. Cocaine (10 mg/kg IP) significantly increased the frequency of central ambulations, F(2, 10) = 8.074, p < 0.0082, N = 6). Post hoc analysis further shows that there were statistically significant differences from baseline in the first hour of the two hours tested (Fisher PLSD = 2.751, Scheffe F = 7.949and 1.216, first and second hours respectively). Cocaine's effects on central ambulations were completed during the second hour. However, at the 90-min point of the time course, central ambulatory behavior underwent a transient rise and fall, not unlike its previous pattern but very similar to the pattern of the ambulatory, rearing, and stereotypic fine movement behavior seen in Figs. 2 and 3. Frequency of central ambulations was significantly increased to  $3 \pm 1$  photobeam interruptions (p < 0.05, 95% CL) from a baseline of 0 ± 0.04 within 10 min and maximally increased to 9  $\pm$  5 (p < 0.05, 95% CL) within 50 min after cocaine administration.

Increased DA release in vINAcc after cocaine administra-

tion was significantly and positively correlated with classically DA-dependent behaviors (first- and second-hour effects) (Pearson product:  $r_{(a)} > 0.651$ ,  $z_f > 0.7753$ , p < 0.01) and with the 5-HT-ergic behavior, central ambulations (Pearson product:  $r_{(a)} = 0.606$ ,  $z_f < 0.7089$ , p < 0.01). Cocaine-induced 5-HT release was significantly and positively correlated with the 5-HT behavior (Pearson product:  $r_{(a)} = 0.595$ ,  $z_f < 0.6931$ , p < 0.01). However, classically DA-dependent behaviors were significantly and more positively correlated with cocaine-induced 5-HT release in vlNAcc than with DA release throughout the two-hour period of study (Pearson product:  $r_{(a)} > 0.732$ ,  $z_f > 0.9287$ , p < 0.01).

Provocatively, the noted abrupt rise and fall in each of the cocaine-induced psychostimulant behaviors occurred when 5-HT release underwent a divergence in direction from concurrent DA release. Interestingly, the Pearson product-moment coefficient of correlation tests show that DA and 5-HT release were highly and positively correlated with classically DA-dependent behaviors up to the 90-min mark,  $r_{(a)} > 0.697$ ,  $z_f > 0.8673$ , p < 0.01.

Preliminary results from studies of the immediate aftereffects of acute (IP) cocaine show that 5-HT release continues to increase after the two-hour period of study at a time during which DA release begins to decrease and cocaine-induced psychostimulant behaviors have begun to reach completion.

#### DISCUSSION

These data demonstrate that acute (IP) cocaine increases both DA and 5-HT release in vlNAcc concurrently and in vivo in the freely moving and behaving animal. The DAergic elements of the cocaine effect seen here are consistent with the body of evidence already presented. Moreover, new findings show that (IP) cocaine increased 5-HT release in vlNAcc. Thus, these data show that the effects of cocaine on DA neu-

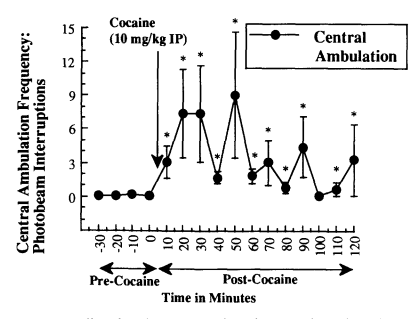


FIG. 4. The effect of cocaine (10 mg/kg IP) on frequency of central ambulation behavior (agoraphobic inhibition) in the same group of Norway rats (cf. text for analysis of variation statistics). Baseline photobeam interruptions were  $0 \pm 0.04$  (representing habituated behavior). \*p < 0.05 (95% confidence limits).

rons in  $A_{10}$  circuits are co-mediated by 5-HT. The data are consistent with a proposed 5-HT involvement in drug discrimination (15), in self-administration (13,39,40,49), and in the endocrine effects of cocaine (38).

Importantly, vINAcc is a neuroanatomically specific site of NAcc, for which a reciprocal connection with the midlateral (i.e., the middle rostro-caudal VTA) has been shown with anterograde and retrograde tract tracing studies (19,59,60). Using light microscopic immunocytochemistry and silver intensification procedures, with anterograde and retrograde tracing tract studies, we have found in vINAcc a core which contains a dense terminal field of tyrosine hydroxylase (TH)that is, TH-IR axons that have an extensive overlap with 5-HT-IR axons in the periphery within the core (45). Thus, a DA-5-HT-ergic interaction may play a critical role in cocaine's manipulation of the compensatory negative feedback pathway between vlNAcc and VTA in the A<sub>10</sub> circuit. A likely scenario mechanistically for enhanced 5-HT release in vlNAcc after (IP) cocaine, may be compensatory negative feedback due to decreased 5-HT cell firing in DR (16,17). Alternatively, a cocaine-induced 5-HT reuptake inhibition, combined with the increased release of 5-HT at vINAcc, may stimulate autoreceptors that act presynaptically at 5-HT somatodendrites to decrease DR cell firing.

Interestingly, studies which have not tested cocaine effects on biogenic amines but have tested 5-HT-ergic effects on DA neurons per se have shown that 5-HT stimulates DA neurons in NAcc (14,32,43) and in VTA (1). In addition, interactive effects by 5-HT on NAcc DA neurons by VTA (28) and by dorsal raphe (DR) (27,30) have been reported. Taken together, the present data suggest that cocaine reward and/or dysfunction may derive from a malfunctioning of this dual biogenic amine system. Whether or not reward-relevant synaptic contacts are made may actually be a separate consideration.

Notably, the present results are different from the effects of (SC) cocaine on 5-HT release in vlNAcc, in the same paradigm (6), and are similar to the effect of (IV) cocaine on 5-HT-ergic release in vlNAcc, in the chloral hydrate-anesthetized rat paradigm (9). Thus, the present data demonstrate, consistent with others (37), that the route of administration for cocaine administered acutely is a crucial factor in its consequent effects. Also, the data demonstrate that anesthesia does not significantly influence the 5-HT-ergic response to cocaine (9).

Real time detection of DA and 5-HT release in vINAcc in the conscious animal provides an excellent tool for studying the nature of the "classically DA-dependent" and 5-HT-ergic psychostimulant behaviors induced by cocaine. Psychostimulant behaviors have been termed dysfunctional, nonadaptive, composite aggregates of subsystems which mediate movement along independent spatial dimensions (61). In this view, psychostimulant behaviors occur as a result of the initial activation and then deactivation of DAergic systems. However, in another view, the neurotransmitter DA functions in mesolimbic and nigrostriatal neuronal circuitry differentially; locomotor activity is primarily controlled by NAcc (ventral striatum) and stereotypic fine movements are primarily controlled by dorsal striatum (51). Placing the present findings within the latter framework, cocaine induced an increase in DA release in vlNAcc that was correlated as expected with cocaine-induced increased locomotor activity. Correspondingly, cocaine-induced maladaptive rearing behavior and stereotypic fine movement behavior were correlated with increased DA release as well. This response too was expected (i.e., based on the postulated partial mediation of these behaviors by NAcc). Also, predictably, the behavioral profiles for the classically DA-dependent behaviors (running, rearing, and fine movement behaviors) are remarkably similar. Indeed, the behavioral profiles for rearing and fine movement are superimposable, differing only in degree, thereby supporting the concept of rearing as a simple vertical extension of a maladaptive head movement (61).

Nonetheless, the present data show that the ebb and flow of each of the cocaine-induced classically DA-dependent behaviors were also dramatically correlated with the neurotransmitter 5-HT. Therefore, the present behavioral data further support a contributory role for 5-HT in the underlying mechanism of action of cocaine. That cocaine has the capability of showing anti-agoraphobic inhibitory characteristics in the "central ambulations" paradigm is also consistent with its 5-HT-ergic effects.

In conclusion, the present studies show that 5-HT may signal or precede the DAergic events associated with the well-known acute cocaine-induced DAergic dysfunction in  $A_{10}$ 

neuronal circuits. Interpretation of these results appears to parallel the "modulatory model" of brain reward. More importantly though, the data may bear relevance to aspects of chronic cocaine abuse such as those described in the Opponent Process Theory (35,36,57). Perhaps 5-HT may serve as a putative regulator during addictive and withdrawal processes.

#### ACKNOWLEDGEMENTS

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#### REFERENCES

- 1. Beart, P. M.; McDonald, D. 5-Hydroxytryptamine and 5hydroxytryptaminergic-dopaminergic interactions in the ventral tegmental area of rat brain. J. Pharm. Pharmacol. 34:591-593; 1982.
- Blaha, C. D.; Jung, M. E. Electrochemical evaluation of stearate modified graphite paste electrodes: Selective detection of dopamine maintained after exposure to brain tissue. J. Electroanal. Chem. 310:317-334, 1991.
- Bradberry, C. W.; Roth, R. H. Cocaine increases extracellular dopamine in rat nucleus accumbens and ventral tegmental areas as shown by *in vivo* microdialysis. Neurosci. Lett. 103:97-102; 1989.
- 4. Broderick, P. A. Characterizing stearate probes in vitro for the electrochemical detection of dopamine and serotonin. Brain Res. 495:115-121; 1989.
- Broderick, P. A. State-of-the-Art microelectrodes for in vivo voltammetry. Electroanalysis 2:241-251; 1990.
- Broderick, P. A. Cocaine-on-line analysis of an accumbens amine neuronal basis for psychomotor behavior. Pharmacol. Biochem. Behav. 40:959-968; 1991.
- Broderick, P. A. In vivo voltammetric studies on release mechanisms for cocaine with γ-butyrolactone. Pharmacol. Biochem. Behav. 40:969-975; 1991.
- Broderick, P. A. Cocaine's colocalized effects on synaptic serotonin and dopamine in ventral tegmentum in a reinforcement paradigm. Pharmacol. Biochem. Behav. 42:889-898; 1992.
- 9. Broderick, P. A. Distinguishing effects of cocaine IV and SC on mesoaccumbens dopamine and serotonin release with chloral hydrate anesthesia. Pharmacol. Biochem. Behav. 43:929-937; 1992.
- Broderick, P. A.; Wechsler, R. T.; Phelan, F. T.; Eng, F. Cocaine increases accumbens serotonin release concurrently with dopamine release in psychostimulant reinforcement. Soc. Neurosci. Abstr. 18:1456; 1992.
- 11. Brodie, M. S.; Dunwiddie, T. V. Cocaine effects in the ventral tegmental area: Evidence for an indirect dopaminergic mechanism of action. Naunyn Schmiedebergs Arch. Pharmacol. 342: 660-665; 1990.
- Carboni, E.; Imperato, A.; Perezzani, L.; DiChiara, G. Amphetamine, cocaine, phencyclidine and nomifensine increase extracellular dopamine concentrations preferentially in the nucleus accumbens of freely moving rats. Neuroscience 28:653-661; 1989.
- Carrol, M. E.; Lac, S. T.; Asencio, M.; Kragh, R. Fluoxetine reduces intravenous cocaine self-administration in rats. Pharmacol. Biochem. Behav. 35:237-244; 1990.
- Chen, J.; van Praag, H. M.; Gardner, E. L. Activation of 5-HT<sub>3</sub> receptor by l-phenylbiguanide increases dopamine release in the rat nucleus accumbens. Brain Res. 543:354-357; 1991.
- 15. Cunningham, K. A.; Callahan, P. M. Monoamine reuptake inhibitors enhance the discriminative state induced by cocaine in the rat. Psychopharmacology 104:177-180; 1991.
- Cunningham, K. A.; Lakoski, J. M. Electrophysiological effects of cocaine and procaine on dorsal raphe serotonin neurons. Eur. J. Pharmacol. 148:457-462; 1988.
- 17. Cunningham, K. A.; Lakoski, J. M. The interaction of cocaine

with serotonin dorsal raphe neurons. Neuropsychopharmacology 3:41-50; 1990.

- Darmani, N. A.; Martin, B. R.; Pandey, U.; Glennon, R. A. Inhibition of 5-HT<sub>2</sub> receptor-mediated head twitch response by cocaine via indirect stimulation of adrenergic alpha<sub>2</sub> and serotonergic 5-HT<sub>1A</sub> receptors. Pharmacol. Biochem. Behav. 38:353-357; 1991.
- Domesick, V. B. Neuroanatomical organization of dopamine neurons in the ventral tegmental area. Ann. N. Y. Acad. Sci. 537: 10-26; 1988.
- Drescher, K.; Hetey, L. Influence of antipsychotics and serotonin antagonists on presynaptic receptors modulating the release of serotonin in synaptosomes of the nucleus accumbens of rats. Neuropharmacology 27:31-36; 1988.
- Einhorn, L. C.; Johansen, P. A.; White, F. J. Electrophysiological effects of cocaine in the mesoaccumbens dopamine system: Studies in the ventral tegmental area. J. Neurosci. 8:100-112, 1988.
- Friedman, E.; Gershen, S.; Rotrosen, J. Effect of acute cocaine treatment on the turnover of 5-hydroxytryptamine in the rat brain. Br. J. Pharmacol. 54:61-64; 1975.
- Gallistel, C. R. The role of dopaminergic projections in MFB self-stimulation. Behav. Br. Res. 20:313-321; 1986.
- Galloway, M. P. Regulation of dopamine and serotonin synthesis by acute administration of cocaine. Synapse 6:63-72; 1990.
- Gelbert, M. B.; Curran, D. J. Alternating current voltammetry of dopamine and ascorbic acid at carbon paste and stearic acid modified carbon paste electrodes. Anal. Chem. 58:1028-1032; 1986.
- Geyer, M. A. Approaches to the characterization of drug effects on locomotor activity in rodents. In: Adler, M. W.; Cowan, A., eds. Testing and evaluation of drugs of abuse. New York: A.R. Liss; 1990:81-99.
- Glowinski, J.; Tassin, J. P. Increased utilization of dopamine in the nucleus accumbens but not in the cerebral cortex after dorsal raphe lesion in the rat. Neurosci. Lett. 15:127-133; 1979.
- Guan, X. M.; McBride, W. J. Serotonin microinfusion into the ventral tegmental area increases accumbens dopamine metabolites. Brain Res. Bull. 23:541-547; 1989.
- Hernandez, L.; Guzman, N. A.; Hoebel, B. G. Bidirectional microdialysis *in vivo* shows differential dopaminergic potency of cocaine, procaine and lidocaine in the nucleus accumbens using capillary electrophoresis for calibration of drug outward diffusion. Psychopharmacology 105:264-268; 1991.
- Herve, D.; Simon, H.; Blanc, G.; LeMoal, M.; Glowinski, J.; Tassin, J. P. Opposite changes in dopamine utilization in the nucleus accumbens and the frontal cortex after electrolytic lesion of the median raphe in the rat. Brain Res. 216:422-428; 1981.
- Izenwasser, S.; Rosenberger, J. G.; Cox, B. M. Inhibition of [<sup>3</sup>H] dopamine and [<sup>3</sup>H] serotonin uptake by cocaine: Comparison between chopped tissue slices and synaptosomes. Life Sci. 50:541-547; 1992.
- 32. Jiang, L. H.; Ashby, C. R.; Kasser, R. J.; Wang, R. Y. The effect of intraventricular administration of the 5-HT<sub>3</sub> receptor agonist 2 methyl-serotonin on the release of dopamine in the

nucleus accumbens: An in vivo coulometric study. Brain Res. 513:156-160; 1990.

- 33. Johanson, C. E.; Fischman, M. W. The pharmacology of cocaine related to its abuse. Pharmacol. Rev. 41:3-52; 1989.
- Kalivas, P. W.; Duffy, P. Effect of acute and daily cocaine treatment on extracellular dopamine in the nucleus accumbens. Synapse 5:48-58; 1990.
- Koob, G. F.; Markou, A.; Heinrichs, S.; Schulteis, G.; Weiss, F. Motivational effects of drug withdrawal: Neuronal substrates. Int. Behav. Neuro. Soc. Abstr. 2:20; 1993.
- Koob, G. F.; Stinus, L.; LeMoal, M.; Bloom, F. E. Opponent process theory of motivation: Neurobiological evidence from studies of opiate dependence. Neurosci. Biochem. Rev. 13:135-140; 1989.
- 37. Lau, C. E.; Imam, A.; Ma, F.; Falk, J. L. Acute effects of cocaine on spontaneous and discriminative motor functions: Relation to route of administration and pharmacokinetics. J. Pharmacol. Exp. Ther. 257:444-456; 1991.
- Levy, A. D.; Li, Q.; Kerr, J. E.; Rittenhouse, P. A.; Milonas, G.; Cabrera, T. M.; Battaglia, G.; Alvarez Sanz, M. C.; van deKar, L. D. Cocaine-induced elevation of plasma adrenocorticotropin hormone and corticosterone is mediated by serotonergic neurons. J. Pharmacol. Exp. Ther. 259:495-500; 1991.
- Loh, E. A.; Roberts, D. C. Break-points on a progressive ratio schedule reinforced by intravenous cocaine increase following depletion of forebrain serotonin. Psychopharmacology (Berl.) 101:262-266; 1990.
- Meert, T. F.; Awouters, F.; Niemegeers, C. J. E.; Schellekens, K. H. L.; Janssen, P. A. J. Ritanserin reduces abuse of alcohol, cocaine and fentanyl in rats. Pharmacopsychiatry 24:159-163; 1991.
- Ng, J. P.; Hubert, G. W.; Justice, J. B., Jr. Increased stimulated release and uptake of dopamine in nucleus accumbens after repeated cocaine administration as measured by *in vivo* voltammetry. J. Neurochem. 56:1485-1492; 1991.
- 42. Pan, Z. Z.; Williams, J. T. Differential actions of cocaine and amphetamine on dorsal raphe neurons in vitro. J. Pharmacol. Exp. Ther. 251:56-62; 1989.
- 43. Parsons, L. H.; Justice, J. B. J. Perfusate serotonin increases extracellular dopamine in the nucleus accumbens as measured by in vivo microdialysis. Brain Res. 606:195-199; 1993.
- Pellegrino, L. J.; Pellegrino, A. S.; Cushman, A. J. A stereotaxic atlas of the rat brain. New York: Plenum Press; 1979.
- Phelix, C. F.; Tshoepe, L.; Broderick, P. A. Convergence of serotonin and dopamine in ventrolateral nucleus accumbens: Anatomical and *in vivo* electrochemical analysis. Soc. Neurosci. Abstr. 19; 1993.
- 46. Piercey, M. F.; Lum, J. T.; Hoffmann, W. E.; Carlsson, A.; Ljung, E.; Svensson, K. Antagonism of cocaine's pharmacological effects by the stimulant dopaminergic antagonists, (+)-AJ76 and (+)-UH232. Brain Res. 588:217-222; 1992.
- Reith, M. E. A. 5-HT<sub>3</sub> receptor antagonists attenuate cocaine-induced locomotion in mice. Eur. J. Pharmacol. 186:327-330; 1990.
- Reith, M. E. A.; Wiener, H. L.; Fischette, C. T. Sertraline and cocaine-induced locomotion in mice. I. Acute studies. Psychopharmacology 103:297-305; 1991.
- Richardson, N. R.; Roberts, D. C. S. Fluoxetine pretreatment reduces breaking points on a progressive ratio schedule reinforced by intravenous cocaine self-administration in the rat. Life Sci. 49: 833-840; 1991.

- Ritz, M. C.; Kuhar, M. J. Relationship between self-administration of amphetamines and monoamine receptors in brain: Comparison with cocaine. J. Pharmacol. Exp. Ther. 248:1010-1017; 1989.
- Robbins, T. W.; Mittleman, G.; O'Brien, J.; Winn, P. The neuropsychological significance of stereotypy induced by stimulant drugs. In: Cooper, S. J.; Dourish, C. T., eds. Neurobiology of stereotyped behavior. New York: Oxford Science Publishers; 1990:25-63.
- Roberts, D. C. S.; Koob, G. F. Disruption of cocaine self-administration following 6-hydroxydopamine lesions of the ventral tegmental area in rats. Pharmacol. Biochem. Behav. 17:901-904; 1982.
- Roberts, D. C. S.; Koob, G. F.; Klonoff, P.; Fibiger, H. C. Extinction and recovery of cocaine self-administration following 6-hydroxydopamine lesions of the nucleus accumbens. Pharmacol. Biochem. Behav. 12:781-787; 1980.
- Ross, S. B.; Renyi, A. L. Inhibition of the uptake of tritiated 5-hydroxytryptamine in brain tissue. Eur. J. Pharmacol. 7:270– 277; 1969.
- Rudnick, G.; Wall, S. C. Binding of the cocaine analog 2-Beta-(<sup>3</sup>H) carboxymethoxy-3-Beta-(4-Fluorophenyl)tropane to the serotonin transporter. Mol. Pharmacol. 40:421-426; 1991.
- Shizgal, P.; Bielajew, C.; Corbett, D.; Skelton, R.; Yeomans, J. Behavioral methods for inferring anatomical linkage between rewarding brain stimulation sites. J. Comp. Physiol. Psychol. 94: 227-237; 1980.
- Solomon, R. L.; Corbit, J. D. An opponent process theory of motivation. I. Temporal dynamics of affect. Psychol. Rev. 81: 119-45; 1974.
- Stellar, J. R.; Rice, M. B. Pharmacological basis of intracranial self-stimulation reward. In: Liebman, J. M.; Cooper, S. J., eds. The neuropharmacological basis of reward. New York: Oxford University Press; 1989:14-65.
- Swanson L. W. The projections of the ventral tegmental area and adjacent regions: A combined fluorescent retrograde tracer and immunofluorescence study in the rat. Brain Res. Bull. 9:321-353; 1982.
- Swanson, L. W.; Cowan, W. M. A note on the connections and development of the nucleus accumbens. Brain Res. 92:324-330; 1975.
- Teitelbaum, P.; Pellis, S. M.; DeVietti, T. L. Disintegration into stereotypy induced by drugs or brain damage: A microdescriptive behavioral analysis. In: Cooper, S. J.; Dourish, C. T., eds. Neurobiology of stereotyped behavior. New York: Oxford University Press; 1990:169-199.
- Uchimura, N.; North, R. A. Actions of cocaine on rat nucleus accumbens neurons *in vitro*. Br. J. Pharmacol. 99:736-740; 1990.
- 63. White, F. J.; Wachtel, S. R.; Johansen, P. A.; Einhorn, L. C. Electrophysiological studies of the rat mesoaccumbens dopamine system: Focus on dopamine receptor subtypes, interactions and the effects of cocaine. In: Chiodo, L. A.; Freeman, A. S., eds. Neurophysiology of dopaminergic systems. Grosse Pointe, MI: Lakeshore Publishing Co.; 1987:317-365; 1987.
- 64. Wise, R. A.; Bozarth, M. A. Brain reward circuitry, four circuit elements "wired" in apparent series. Brain Res. Bull. 12:203-208; 1984.
- Yim, Y.; Mogenson, G. J. Response of nucleus accumbens neurons to amygdala stimulation and its modification by dopamine. Brain Res. 239:401-415; 1982.