Dopamine Uptake Inhibitors but Not Dopamine Releasers Induce Greater Increases in Motor Behavior and Extracellular Dopamine in Adolescent Rats Than in Adult Male Rats

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ABSTRACT

Most life-long drug addiction begins during adolescence. Important structural and functional changes in brain occur during adolescence and developmental differences in forebrain dopamine systems could mediate a biologic vulnerability to drug addiction during adolescence. Studies investigating age differences in psychostimulant responses have yielded mixed results, possibly because of different mechanisms for increasing extracellular dopamine. Recent research from our laboratory suggests that adolescent dopamine systems may be most affected by selective dopamine uptake inhibitors. We investigated age-related behavioral responses to acute administration of several dopamine uptake inhibitors [methylphenidate, 1-{2-[bis-(4-fluorophenyl)methoxy]ethyl}-4-(3-phenylpropyl)piperazine (GBR12909), and nomifensine] and releasing agents [amphetamine and methylenedioxymethylamphetamine (MDMA)] in adolescent and adult male rats. Methylphenidate and amphetamine effects on stimulated dopamine efflux were determined using fast-scan cyclic voltammetry in vivo. Dopamine uptake inhibitors but not dopamine releasing agents induced more locomotion and/or stereotypy in adolescent relative to adult rats. MDMA effects were greater in adults at early time points after dosing. Methylphenidate but not amphetamine induced much greater dopamine efflux in periadolescent relative to adult rats. Periadolescent male rats are particularly sensitive to psychostimulants that are DAT inhibitors but are not internalized and do not release dopamine. Immaturity of DAT and/or DAT associated signaling systems in adolescence specifically enhances behavioral and dopaminergic responses in adolescence.

Introduction

Lifelong drug addiction usually begins with drug use during adolescence or young adulthood (Spear, 2000; Schramm-Sapyta et al., 2009). Longitudinal and retrospective studies consistently demonstrate that early exposure to drugs and alcohol is one of the strongest predictors of adult substance abuse (Spear, 2000; Chambers et al., 2003). The onset of drug addiction during adolescence is correlated with an increased severity of addiction including higher rates of morbidity and mortality (for reviews, see Spear, 2000; Schramm-Sapyta et al., 2009). Finally, the progression from initial drug use to the expression of addictive behaviors occurs more rapidly during adolescence than in adulthood. Although such studies demonstrate the importance of adolescence in human drug use, the biological basis for these vulnerabilities is not fully understood.

Adolescence is a time of both sexual maturation and attainment of adult nervous system function. Neurobiologic changes during this phase of development contribute to age-related differences in drug sensitivity (Andersen, 2003; McCutcheon and Marinelli, 2009). Dopamine systems, which mediate the rewarding effects of addictive drugs, undergo significant development and reorganization during adolescence, and may explain, in part, why this period is so important for the development of drug addiction (Spear, 2000; Andersen, 2003; Chambers et al., 2003).

Psychostimulants increase extracellular dopamine by mobilizing different storage pools of transmitter (McMillen, 1983) and so might be predicted to exhibit age-related behavioral effects to the extent that there are age-related differences in the dopamine pools. However, the pharmacological literature exploring behavioral sensitivity to psychomotor stimulants across adolescence is mixed. Several groups have

ABBREVIATIONS: PN, postnatal day; GBR12909, 1-{2-[bis-(4-fluorophenyl)methoxy]ethyl}-4-(3-phenylpropyl)piperazine; MDMA, methylenedioxymethylamphetamine; DAT, dopamine transporter; ANOVA, analysis of variance; DA, dopamine.
reported that rats in the periadolescent period [postnatal day (PN) 30–40] are hyperactive at baseline but have smaller increases in locomotion and stereotyped behaviors than younger or older cohorts after a single dose of amphetamine or cocaine (Bolanos et al., 1998). In contrast, our laboratory has reported that cocaine induces more acute locomotor behavior and stereotypy in periadolescent than adult male rats (Caster et al., 2005; Parylak et al., 2008) and that adolescent rats will consume more of a sweetened cocaine solution (Walker et al., 2009). Periadolescent rats exhibit an “intra-binge” sensitization (Caster et al., 2005) and greater sensitization than adults 24 h after a single high dose of cocaine (Caster et al., 2007). Fast-scan cyclic voltammetry showed that 15 mg/kg cocaine enhanced dopamine efflux in dorsal striatum nearly 3-fold more in adolescent than in adult males, suggesting that greater cocaine-stimulated dopamine efflux might mediate the greater behavioral responses of periadolescents (Walker and Kuhn, 2008).

The present study compared the behavioral and neurochemical effects of methylphenidate, nominifensine, GBR12909, methylenedioxyamphetamine (MDMA), and amphetamine. Psychostimulants can be classified as either dopamine transporter (DAT) inhibitors or amphetamine-like dopamine releasers (McMillen, 1983). This classification is based, in part, on the observation that the ability of amphetamine-like drugs to stimulate behavior is antagonized by the dopamine synthesis inhibitor α-methyl-p-tyrosine, whereas the action of DAT inhibitors is not affected (Carlsson et al., 1966; Weissman et al., 1966). In contrast, behavioral effects of DAT inhibitors, but not amphetamine-like drugs, are antagonized by reserpine, a drug that depletes catecholamine storage vesicles (Weissman et al., 1966). The ability of the DAT inhibitors but not dopamine releasers to increase extracellular dopamine in dialysis experiments is blocked by tetrodotoxin, showing that impulse flow or neuronal activity is necessary for the DAT inhibitors to exert their effects (Westerink et al., 1987; Carboni et al., 1989; Nomikos et al., 1990). The current study seek to determine whether the enhanced behavioral and neurochemical responses induced by cocaine in adolescents are induced by other stimulant drugs from both classes of psychostimulants. To accomplish this goal, we have contrasted the behavioral and neurochemical responses to representative drugs from each class in adolescents and adults. Differences in developmental effects of each drug class could identify age-related differences in DAT function and dopamine neurotransmission.

Fast-scan cyclic voltammetry in anesthetized rats was used to measure electrically stimulated dopamine efflux in dorsal striatum. There is good correlation between the effects of cocaine on electrically stimulated dopamine efflux in anesthetized rats and spontaneous release and uptake events in awake rats (Greco and Garris, 2003; España et al., 2008). Michaelis-Menten parameters for uptake are not different in anesthetized and awake rats (Garris et al., 2003). A single compound was chosen from each category, DAT inhibitors and dopamine releasers. We chose methylphenidate because it effectively increased locomotor behavior and behavioral rating in all ages. We chose amphetamine (1 mg/kg) because our initial observations suggested a trend for PN28 rats to be more activated than adults in the earliest intervals. Neurochemical experiments were performed only in the youngest and oldest age groups because behavioral stimulation was most disparate in these groups.

Materials and Methods

Subjects

Male Sprague-Dawley rats were acquired from Charles River Laboratories (Raleigh, NC) and housed in self-ventilated cages by age. Animals were housed in a vivarium with a 12-h light/dark cycle and given ad libitum access to food and water. Rats PN28, 42, and 65 (≥1 day) were used to correspond to early adolescence, midadolescence, and adulthood, respectively (Spear, 2000). These animals were shipped and received on PN21, 35, and 58 and given 1 week to acclimate in our facility. All experiments were approved by the Duke University Institutional Animal Care and Use Committee.

Drugs

Methylphenidate, nominifensine, GBR12909 [1-[2-bis-(4-fluorophenyl)methoxyethyl]-4-(3-phenylpropyl)piperazine], urethane, and amphetamine were purchased from Sigma-Aldrich (St. Louis, MO), and solutions were made fresh in saline and injected intraperitoneally at 1 ml/kg. MDMA was obtained from RTI International (Research Triangle Park, NC), courtesy of the National Institute on Drug Abuse.

Locomotor Activity

Motor activity was determined in eight open-field photocell devices (Kinder Scientific, Inc., Poway, CA). The devices consisted of a Plexiglas arena (40 cm for each dimension) with corn cob bedding on the floor. Computer software supplied by the manufacturer recorded interruptions of photobeams spaced 2.54 cm (1 inch) apart and reported distance traveled. Assignment to test chambers was counterbalanced across testing days with respect to age. Habituation test sessions began when rats were placed in the open arena without injection. After this 1-h session, all rats were injected with one of the drugs, and data recording was started immediately.

Observational Behavioral Measurements

The topography of behavior was assessed simultaneously with locomotor activity by recording the occurrence of inactivity, rearing, grooming, locomotion, sniffing, continuous sniffing, and behavioral rating during three observation periods consisting of 15 s each, every 5 min, beginning 5 min after dosing. This approach ensured that the automated locomotor behavior measurements were not confounded by a greater stereotypy response in a particular age group and not another. Observations after injection of 1 mg/kg amphetamine were done at 10-min intervals for 1 h. Stereotypy included head weaving or bobbing, patterned locomotion, paw treading, and dyskinesia. A single observer, blinded to the drug treatment, watched all the rats in individual experiments. For each of the three 15-s observation periods, a summed behavioral rating score according to a noncontinuous 6-point behavioral rating scale that has been described previously (Walker et al., 2001). This scale provides a relative measure of behavioral activity with higher numbers denoting more intense behavioral activity than lower numbers. These three scores were then averaged to obtain a score for that minute. The scoring system was as follows: 1, inactive; 2, grooming or locomotion or sniffing or rearing; 3, sniffing with locomotion and/or rearing, or continuous sniffing; 4, continuous sniffing with continuous motion; 5, frequent stereotyped movements with locomotion; and 6, almost continuous stereotyped movements, restricted to one place in the cage.

In Vivo Electrochemistry

In Vivo Methods. Rats were anesthetized with urethane (1.5 g/kg i.p.) and positioned in a stereotaxic apparatus (David Kopf Instruments, Tujunga, CA). Body temperature was maintained at 37°C with a Deltapath Isothermal Pad (Braintree Scientific, Braintree, MA). A bipolar stimulating electrode (Plastics One Inc., Roanoke, VA) was positioned in the medial forebrain bundle, and biphasic stimulation parameters were 300 μA, 2 ms each phase. The
sere stereotactic coordinates (in millimeters) anteroposterior (AP) and mediolateral (ML) from bregma and dorsoventral (DV) from dura follow: the stimulating electrode was placed at −4.6 AP, +1.4 ML, and −7.5 to −9.0 DV. The carbon-fiber microelectrode was directed at the center of the caudate (+1.2 AP, 2.0 ML, and −4.5 to −5.6 DV). To compensate for the smaller size of the PN28 rats, the ML placement of the stimulating electrode was +1.35.

The locations of the stimulating and working electrodes were optimized to give maximal dopamine responses. Extracellular dopamine concentrations resulting from 60 pulse stimulation trains at frequencies from 10, 20, 30, 40, 50, to 60 Hz were recorded. Immediately after the final baseline data collection, the rat was administered 10 mg/kg methylphenidate or 1 mg/kg amphetamine intraperitoneally. These doses were used in behavioral experiments. The time course of drug effects on extracellular dopamine was monitored at 20 Hz because the effect of uptake inhibition is frequency-dependent and most robust at this frequency. Twenty-Hertz stimulations commenced immediately after drug injection (approximately 1 min) and were repeated at 2.5, 5, 7.5, 10, 15, and 30 min after drug. Drug responses to stimulations at the other frequencies were recorded between 20 and 40 min after drug administration.

**Electrochemistry.** Voltammetry procedures were similar to our previously published methods (Walker and Kuhn, 2008). Fast-scan cyclic voltammetry was conducted with an EI-400 potentiostat (Ensnman Instrumentation, Bloomington, IN). The potential at carbon fiber electrodes was held at −400 mV, ramped to 1 V, and then back to −400 mV at 300 V/s. Cyclic voltammograms were recorded at 10 Hz. Carbon-fiber microcylinder electrodes, prepared from 7-µm-diameter T-300 fibers with approximately 50 to 100 µm of exposed carbon fiber (Amoco, Greenville, SC), were used in the in vivo experiments along with a silver/silver chloride reference wire.

Changes in extracellular dopamine were determined by monitoring the current over a 100-mV window at the peak oxidation potential for dopamine. The electroactive substance was identified as dopamine by comparing background subtracted cyclic voltammograms from the in vivo stimulations with those collected at the same electrode in vitro after the experiment. Oxidation currents in vivo were converted to dopamine concentrations by calibrating the electrodes with dopamine standard solutions in a flow injection system after experimental use.

**Data Analysis**

Group averages are expressed as the mean ± S.E.M., and n is the number of rats. Effects of age and time after injection on locomotor behavior and behavioral rating were determined using two-way ANOVA with repeated measures on time. Drug-induced changes in dopamine efflux were expressed relative to the baseline in each rat

**Methylphenidate.** The age-dependent effects of DAT inhibitors methylphenidate, nomifensine, and GBR12909 on horizontal activity and experimenter-observed behaviors were determined. Figure 2 shows time courses of effects for each drug, and Fig. 4 shows session totals for locomotion and session means for behavioral rating. The locomotor stimulatory effect of 10 mg/kg methylphenidate (n = 20 for PN28 and PN65 and 19 for PN42) was greatest in the youngest rats.

![Fig. 1. Horizontal activity during habituation to a novel open-field device and after saline injection intraperitoneally. Initial habituation data from all male rats used in subsequent drug experiments were combined in the left graph: PN28 (n = 109), PN42 (n = 110), and PN65 (n = 111). There was no main effect of age but a significant age by time interaction. *, PN42 > PN28 at 40 min. A subset of these rats was injected with saline intraperitoneally after habituation, and the results are shown in the right graph: PN28 (n = 17), PN42 (n = 18), and PN65 (n = 15). No overall age difference was observed after saline injection, although the interaction of age and time was significant. *, PN42 < PN28 and PN65 at 10 min. Group means ± S.E. are shown in this and all figures. Error bars are smaller than the symbols in some cases.]
ANOVA indicated an effect of age ($F_{2,56} = 5.25; p = 0.008$), and post hoc analysis indicated that PN28 rats exhibited more locomotion than PN42 and PN65 rats. Effects of interval after methylphenidate injection ($F_{8,448} = 101, p < 0.001$) and an interaction of interval and age also were found ($F_{16,896} = 2.63; p < 0.001$). Activity in PN42 rats was high in early intervals relative to adults, similar to PN28, but then it fell to lower levels similar to PN65.

Observer-rated behavioral activation in a subset of these animals ($n$ values: PN28 and PN65 = 16 and PN42 = 15) mirrored those for locomotion. Behavioral rating was highest in the youngest rats and ANOVA indicated a significant effect of age ($F_{2,44} = 6.35; p = 0.004$). Post hoc analysis showed that PN28 rats had significantly higher ratings than PN42 and PN65 rats ($p < 0.05$). Time after methylphenidate varied significantly ($F_{8,352} = 7.83; p < 0.001$) and time and age significantly interacted ($F_{16,352} = 3.10; p < 0.001$). Behavioral rating was higher in PN28 than PN65 in early intervals and waned in PN42 in late intervals.

**Nomifensine.** Figures 2 and 4 display the effects of 5 mg/kg nomifensine on behavior of PN28, PN42, and PN65 male rats ($n = 10$ or $11/age$). Nomifensine-induced locomotor behavior was inversely related to age. ANOVA indicated an overall effect of age ($F_{2,28} = 6.57; p = 0.005$). Post hoc analysis showed that PN28 rats ambulated more than PN65 rats ($p < 0.05$). Activity in PN42 rats was intermediate but not statistically different from other ages by post hoc analysis.

Activity varied with time after injection ($F_{11,308} = 11.6; p < 0.001$), and there was no interaction with age ($p = 0.13$).

The effect of nomifensine on behavioral rating exhibited a similar age dependence. Age significantly affected behavioral rating ($F_{2,28} = 5.82; p = 0.008$). Newman-Keuls test indicated that nomifensine increased behavioral rating in both adolescent age groups more than in adult rats ($p < 0.05$). Behavioral rating varied with time after injection ($F_{11,308} = 2.43; p = 0.007$). Time did not interact with age ($p = 0.81$).

**GBR12909.** GBR12909 (5 mg/kg) increased locomotor behavior and behavioral rating but only the effect on locomotion was age-related. GBR12909 effects on locomotion varied significantly across development ($F_{2,37} = 3.92; p = 0.03; n = 13$ or 14/age). Post hoc analysis showed that PN28 rats exhibited more locomotor behavior than PN65 rats ($p < 0.05$). GBR12909 effects varied with time after injection ($F_{5,185} = 9.59; p < 0.001$). The interaction of age and time was not significant ($p = 0.07$).

In contrast to the locomotor effects, GBR12909 did not affect behavioral rating differently across the age groups ($n = 8$). ANOVA reported a significant effect of time after injection ($F_{2,104} = 6.63; p < 0.001$), no age effect ($p = 0.61$) and no age × time interaction ($p = 0.15$).

**Amphetamine.** Figures 3 and 4 show the effects of dopamine releasing drugs on spontaneous behavior. Multiple doses of amphetamine were tested because it is the prototypical drug in this class, and we wanted to span the dose range...
from low to high induction of stereotypy. A single dose of another amphetamine, MDMA, also was investigated.

The effect of 1 mg/kg amphetamine on ambulatory behavior did not differ by age ($F_{2,42} = 0.30; p = 0.74; n = 14–16/group$). Time after injection significantly affected ambulations ($F_{11,462} = 15.3; p < 0.001$), and the interaction with age was not significant ($p = 0.22$).

The effect of 1 mg/kg amphetamine on behavioral rating was also not age-dependent ($p = 0.33; n = 15 or 16/group$). Time after injection was significant ($F_{5,215} = 44; p < 0.001$), but its interaction with age did not quite reach significance ($F_{10,215} = 1.80; p = 0.062$).

The effect of 2 mg/kg amphetamine on ambulatory behavior did not exhibit an overall effect of age ($F_{2,42} = 0.67; p = 0.52; n = 9–11/group$). Time after injection significantly affected ambulations ($F_{11,297} = 12.4; p < 0.001$), but the interaction with age was not significant ($p = 0.086$).

The effect of 2 mg/kg amphetamine on behavioral rating in the same rats was also not age-dependent ($F_{2,27} = 0.22; p = 0.80$). Time after injection was significant ($F_{11,297} = 49.6; p < 0.001$), but its interaction with age did not reach significance ($F_{10,297} = 2.17; p = 0.045$).
The effect of 5 mg/kg amphetamine on ambulatory behavior did not exhibit an overall effect of age ($F_{2.24} = 0.22; p = 0.80; n = 8–10/group). Time after injection significantly affected ambulations ($F_{11.297} = 10.8; p < 0.001$), but its interaction with age was not significant ($p = 0.67$).

The effect of 5 mg/kg amphetamine on behavioral rating in the same rats also was not age-dependent ($F_{2.24} = 2.54; p = 0.10$). Time after injection was significant ($F_{11.297} = 6.31; p < 0.001$). The interaction of time and age was significant ($F_{22,294} = 1.63; p = 0.04$) because in mid-to-late intervals, ratings tended to be highest in PN28 and lowest in PN42 rats.

**Methylenedioxymethamphetamine.** Figures 3 and 4 show the effect of 5 mg/kg MDMA on locomotor behavior ($n = 15$ or 17/group). Unlike the DAT inhibitors tested, locomotor effects of MDMA were not enhanced in peradolescents. In fact, the trend was the opposite. The effect of age on MDMA effects were not enhanced in periadolescents. In adolescents more than adults particularly at the lowest frequency.

**Amphetamine.** Amphetamine effects on stimulated extracellular dopamine levels were determined using the methods described for methylphenidate in PN28 and PN65 male rats. Amphetamine effects were analyzed relative to predrug baseline levels (percentage of baseline) at each frequency tested. Amphetamine (1 mg/kg) increased stimulated extracellular dopamine in a frequency-dependent manner ($F_{5,59} = 12.9; p = 0.007$), time ($F_{6,48} = 23.4; p < 0.001$), and the interaction of the two ($F_{6,48} = 3.60; p = 0.005$). Age differences in dopamine concentrations were greatest between 7.5 to 10 min after intraperitoneal injection.

**Effects of Psychostimulants in Adolescence**

![Graphs showing distance traveled and stereotypy score for different groups](image-url)
indicated by main effect of age ($F_{1,11} = 8.23; p = 0.015; n = 6$ for PN28 and $n = 7$ for PN65). Amphetamine effects varied with time after administration ($F_{8,82} = 24.7; p < 0.001$), and age and time significantly interacted ($F_{8,82} = 6.36; p < 0.001$). This interaction reflects the results showing that peak amphetamine effects occurred between 10 to 30 min for adolescents and between 30 to 60 min for adults.

Amphetamine enhanced extracellular dopamine more in adolescent rats at 20 Hz, and the peak increases were earlier in adolescents than adults. Relative to methylphenidate effects however, age differences caused by amphetamine were more modest.

Discussion

The present study demonstrates that selective DAT inhibitors induce more spontaneous motor behavior in adolescent than adult rats. In contrast, dopamine releasing drugs did not. The early increase in locomotor behavior induced by MDMA was greater in adult than adolescent rats. The age differences spanned a large range of maximal locomotor stimulation, suggesting that the phenomenon is robust and consistent. Effects on stimulated dopamine efflux reflected the behavioral differences: methylphenidate stimulated dopamine efflux more in adolescents, but amphetamine did not. Thus, greater relative stimulation of dopamine by selective DAT inhibitors in periadolescents partly explains the distinct age differences.

Reported effects of amphetamine vary across laboratories. Vasilev et al. (2003) reported that amphetamine (1.5 mg/kg) induced less locomotion and stereotyped behavior in PN28 to 30 than 90-day-old adult hooded males. Bolanos et al. (1998) also found lower locomotor effects of 0.5 and 1.5 mg/kg amphetamine in PN35 rats than in PN80 male rats. Although these results differed from the present results no age differences in the effects of 1, 2 or 5 mg/kg amphetamine, they support our conclusion that developmental effects of amphetamine differ from those of dopamine uptake inhibitors. Wooters et al. (2006) showed that the acute locomotor effects of methylphenidate in periadolescent males were approximately double those in the adults. These reports generally agree with the present findings.

Age differences in pharmacokinetics could contribute to age differences in the behavioral effects of psychostimulants. Determining the kinetics of each of these drugs across development was beyond the scope of the present study. Unfortunately, the literature provides few answers for the issue. In general, blood levels of these psychostimulants may be slightly lower in adolescents than adults (Spear, 2007). We have previously examined brain cocaine levels in adolescent and adult males using a repeated dose model and found no significant differences across age (Caster et al., 2005). Age-related differences in acute cocaine metabolism have not been identified, which would explain greater the behavioral responses of adolescents. A review article mentions that brain amphetamine levels are lower in adolescent (PN25) than adult rats (Spear and Brake, 1983). This age-related pharmacokinetic difference did not correlate with the age-related behavioral differences reported in that article. If amphetamine concentrations are in fact lower in the adolescent brain, this would confound the present results. However, the weight of the evidence from this and our other studies with cocaine showing that four DAT inhibitors are more effective in adolescents and two dopamine releasers are not suggests that it is unlikely that age differences in pharmacokinetics is a sufficient explanation.

The present study confirms and extends other dopamine work. Stamford (1989) showed that nomifensine (10 mg/kg i.p.) increased electrically stimulated dopamine efflux more
in the striatum of young (30-day-old) rats relative to adults, using very similar voltammetry methods to those in the current study. In addition, our laboratory showed previously that cocaine increased extracellular dopamine more in PN28 than PN65 rats (Walker and Kuhn, 2008). Thus, three studies using electrically stimulated dopamine efflux have reported that effects of cocaine, nomifensine and now methylphenidate are enhanced in dorsal striatum of periadolescents relative to adults. Other reports of age-related effects of these compounds on dopamine include microdialysis studies in nucleus accumbens. A low dose of cocaine increased extracellular dopamine more rapidly in nucleus accumbens of PN35 rats (Badanich et al., 2006). Frantz et al. (2007) did not find age differences in the nucleus accumbens shell at baseline or after 20 mg/kg cocaine i.p. This agrees with our previous report finding greater electrically stimulated dopamine efflux in dorsal striatum not nucleus accumbens core (Walker and Kuhn, 2008). Cao et al. (2007) found lower basal dopamine in ventral but not dorsal caudate putamen of PN29 rats relative to adults but did not report whether cocaine induced a significant effect. One important caveat with the current results is that our technique might not have captured spontaneous increases in basal dopamine that might have been induced by the low dose used here.

Differences in the mechanism of DAT inhibition might explain the present behavioral results. Amphetamine-like compounds are fundamentally different from cocaine and other DAT inhibitors because they are a substrate for DAT and are transported into the cell. Slightly different binding sites on DAT or interactions with DAT between these classes of stimulants may explain their differing functional effects. Some studies show that the binding sites of dopamine, cocaine, and amphetamine overlap (Beuming et al., 2008), whereas others demonstrate differences between amphetamine and other psychostimulants in their binding or inhibition of DAT (Dersch et al., 1994; Wayment et al., 1998). Differences in DAT inhibition have functional implications. For example, high-cocaine-responding rats were found to have greater dopamine uptake than low responders, and uptake in individual rats was correlated with cocaine-stimulated behavior (Briegleb et al., 2004). However, none of these relationships existed for amphetamine-stimulated behavior, suggesting that functional DAT expression on the cell surface is related to cocaine- but not amphetamine-stimulated behavioral activation (Briegleb et al., 2004). Furthermore, a novel benztrapine analog that occupies DAT completely blocks behavioral and conditioned effects of cocaine but not amphetamines (Velazquez-Sanchez et al., 2009). Carboni et al. (1989) showed that the dopamine-elevating effect of amphetamine is independent of impulse flow because γ- butyrolactone blocked the dopamine increases induced by cocaine and nomifensine but not amphetamine-induced increases. Similarly, using tetrodotoxin to inhibit action potential propagation, nomifensine, cocaine, GBR12909, and methylphenidate but not amphetamine were shown to be dependent on impulse flow to increase extracellular dopamine in dialysis (Nomikos et al., 1990). In this context, the present results suggest that developmental differences in DAT structure–function are related to the cocaine but not the amphetamine binding site on DAT.

Forebrain dopamine systems continue to mature and reorganize across adolescence (for reviews, see Andersen, 2003; Kuhn et al., 2010). Dopamine innervation of the dorsal and ventral striatum is incomplete during early adolescence, and most presynaptic markers including DAT expression and dopamine stores have not yet attained adult levels. There is also an overproduction followed by regressive “pruning” of striatal dopamine receptors during adolescence. Such maturational events in dopamine systems could probably affect the behavioral responsiveness to stimulants across adolescence independent of sex hormones. However, the parallel increase in DAT expression and DA stores that have been observed do not suggest an obvious explanation for differences in the actions of DAT inhibitors and DA-releasing drugs.

That adolescent dopaminergic transmission is more regulated by DAT than adults would contribute to the present findings. We have postulated that uptake inhibition by cocaine enhances extracellular dopamine more in the adolescent striatum because at baseline, the ratio of uptake to release is greater in adolescent striatum (Walker and Kuhn, 2008). Release capacity is lower in periadolescents than adult striatum. The lesser relative effects of amphetamine could be related to the limited stores available for release in early adolescence as dopamine content and release capacity is less in dorsal striatum of adolescents than adults (Kuhn et al., 2010). Serotonin innervation is also immature in the periadolescents (Moll et al., 2000; Galineau et al., 2004), which is significant because the two dopamine releasers used in these studies would have elevated extracellular serotonin levels more than the DAT inhibitors (Kuczenski and Segal, 1997). Increased serotonin levels should attenuate the hyperlocomotion induced by hyperdopaminergia (Gaineddinov et al., 1999), an effect that should be greater in the adults. The enhanced serotonin elicited by releasers would be expected to decrease locomotor behavior preferentially in the adults, serving to attenuate the presently observed age by psychostimulant category difference.

Putative developmental differences in DAT glyclosylation represent one potential mechanism for the greater sensitivity of adolescents to DAT inhibitors. Patel et al. (1994) found that DAT from the striatum of adult rats had a higher molecular weight than DAT from rats at 0, 4, and 14 days of age, and they showed that this size difference is due to adult DAT being more glycosylated. Cocaine was more potent for inhibition of dopamine uptake into cells expressing the least glycosylated mutant because nonglycosylated DAT has greater affinity for dopamine than normal DAT (Li et al., 2004). Thus, specific DAT inhibition should induce a greater relative change in extracellular dopamine in the adolescents and presumably induce more behavioral effects.

We have shown previously that cocaine effects on behavior and stimulated dopamine efflux are greater in adult female than adult male rats and that psychostimulant responses fall across development in males (Parylak et al., 2008). The present results suggest that this latter effect exists for a broad array of DAT inhibitors. Thus, the behavioral response to clinically used psychostimulants might be expected to change across development, depending on the mechanism of action. This phenomenon could have implications for pharmacotherapy of attention-deficit hyperactivity disorder for children, adolescents, and adults.
References


