Paradoxical effects of injection stress and nicotine exposure experienced during adolescence on learning in a serial multiple choice (SMC) task in adult female rats

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ABSTRACT

Nicotine exposure in adolescent rats has been shown to cause learning impairments that persist into adulthood long after nicotine exposure has ended. This study was designed to assess the extent to which the effects of adolescent nicotine exposure on learning in adulthood can be accounted for by adolescent injection stress experienced concurrently with adolescent nicotine exposure. Female rats received either 0.033 mg/h nicotine (expressed as the weight of the free base) or bacteriostatic water vehicle by osmotic pump infusion on postnatal days 25–53 (P25–53). Half of the nicotine-exposed rats and half of the vehicle rats also received twice-daily injection stress consisting of intraperitoneal saline injections on P26–53. Together these procedures produced 4 groups: No Nicotine/No Stress, Nicotine/No Stress, No Nicotine/Stress, and Nicotine/Stress. On P65–99, rats were trained to perform a structurally complex 24-element serial pattern of responses in the serial multiple choice (SMC) task. Four general results were obtained in the current study. First, learning for within-chunk elements was not affected by either adolescent nicotine exposure, consistent with past work (Pickens, Rowan, Bevins, and Fountain, 2013), or adolescent injection stress. Thus, there were no effects of adolescent nicotine exposure or injection stress on adult within-chunk learning typically attributed to rule learning in the SMC task. Second, adolescent injection stress alone (i.e., without concurrent nicotine exposure) caused transient but significant facilitation of adult learning restricted to a single element of the 24-element pattern, namely, the "violation element," that was the only element of the pattern that was inconsistent with pattern structure. Thus, adolescent injection stress alone facilitated violation element acquisition in adulthood. Third, adolescent nicotine exposure, in this case both with and without adolescent injection stress, caused a learning impairment in adulthood for the violation element in female rats. Thus, adolescent nicotine impaired adult violation element learning typically attributed to multiple-item learning in the SMC task. Fourth, a paradoxical interaction of injection stress and nicotine exposure in acquisition was observed. In the same female rats in which violation-element learning was impaired by adolescent nicotine exposure, adolescent nicotine experienced without concurrent injection stress produced better learning for chunk-boundary elements in adulthood compared to all other conditions. Thus, adolescent nicotine without concurrent injection stress facilitated adult chunk-boundary element learning typically attributed to concurrent stimulus–response discrimination learning and serial-position learning in the SMC task. To the best of our knowledge, the current study is the first to demonstrate facilitation of adult learning caused by adolescent nicotine exposure.

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1. Introduction

Survey research has shown that cigarette smoking has been on a decreasing trend among adolescents since the late 1990s though 40% of 12th graders still admit to have smoked at least once in their life (Johnston et al., 2011). However, from 2011 to 2012, electronic cigarette use doubled for middle and high school students and every day, more than 3200 U.S. adolescents smoke their first cigarette with an estimated 2100 becoming daily smokers (Center for Disease Control and Prevention, 2014). Though smoking in adolescence may not be as prevalent as it was in the early 1990s, adolescents continue to be exposed to nicotine, perhaps in growing numbers.

Research in rats has shown that nicotine exposure during adolescence can cause long-lasting physiological and morphological changes to the brain that cause persistent changes in adult neural and behavioral
function (Abreu-Villaça et al., 2003a, 2003b). Nicotine exposure in adolescence has also been shown to cause cognitive deficits in adults such as decreased attentional performance, impairments of stimulus-response (S-R) learning, and impairments of memory in several behavioral paradigms (Counotte et al., 2009; Fountain et al., 2008; Jacobsen et al., 2005; Schochet et al., 2004; Slawecki et al., 2004; Spaeth et al., 2010).

For experimental purposes in animal models, controlled nicotine exposure in adolescence is typically achieved by subcutaneous or intraperitoneal injections, by transdermal patch, or by implantable osmotic pump. Administration can thus be intermittent via single or multiple bolus injections distributed through time, or continuous via chronic absorption or infusion. Very few indications are given by experimenters as to why one procedure is chosen over another. However, all of the foregoing methods of adolescent nicotine exposure in rats have been shown to cause neural and behavioral changes that last into adulthood (Abreu-Villaça et al., 2003a; Abreu-Villaça et al., 2003b; Adriani et al., 2003; Barron et al., 2005; Belluzzi et al., 2004; Brielmaier et al., 2007; Counotte et al., 2009; Dwyer et al., 2009; Fountain et al., 2008; McDonald et al., 2005; Natividad et al., 2013; Philpot et al., 2014; Pickens et al., 2013; Polesskaya et al., 2007; Quick et al., 2014; Schochet et al., 2004; Slawecki et al., 2004; Slotkin, 2002; Slotkin et al., 2008, 2007; Spaeth et al., 2010; Trauth et al., 2000c, 1999, 2000a, 2000b; Wheeler et al., 2013). Furthermore, recent research has found that various forms of stress in adolescence can have long-term effects on cognition (Green and McCormick, 2013; Ignor et al., 2004; Morrissey et al., 2011; Torregrossa et al., 2012). Since daily injections and other stress-inducing activities related to injections such as handling and weighing have been shown to be a source of stress in rats (Dilsaver and Majchrzak, 1990; Sharp et al., 2003), it is possible that the behavioral effects that have been observed in adult rats after adolescent nicotine exposure are a result of chronic injection stress experienced during adolescence. Injection stress alone, or through an interaction with nicotine, may result in cognitive impairments. The current study assessed the effects of injection stress and adolescent nicotine exposure on adult rat learning.

Prior research has shown that serial pattern learning in a serial multiple choice (SMC) task in rats is sensitive to learning impairments in adulthood caused by earlier adolescent nicotine exposure (Fountain et al., 2008; Pickens et al., 2013). The SMC task is modeled after a non-verbal task that has been used to study human associative versus rule learning (Fountain and Rowan, 1995; Kundey et al., 2013; Restle and Brown, 1970) and we have used it to study complex cognitive processes in rats (Fountain et al., 2012). When rats learn sufficiently complex serial patterns in this task, they have been shown to employ multiple cognitive systems concurrently, including simple S–R discrimination learning, multiple-item memory, and abstract rule learning processes (for a review, see Fountain et al., 2012). In addition, sex differences in acquisition rates have been observed in this paradigm for adult vehicle and nicotine-exposed rats that were not subject to chronic injection stress should make no more errors than vehicle rats.

### 2. Materials and methods

#### 2.1. Animal care and nicotine treatment

Sixty female Long Evans rats (Rattus norvegicus) that were bred in-house served as subjects for this experiment. They were randomly assigned to one of four conditions. Rats were received on postnatal day 21 (P21) and housed in groups of 3 in plastic shoe-box cages (40 cm wide × 85 cm long × 40 cm high) and given free access to water and food (LabDiet P5000 – ProLab® RMH 3000). Rats were housed so that each individual in a cage was from a different litter but part of the same experimental condition. Rats caged together were differentiated from one another by tail markings.

Delivery of nicotine (0.033 mg/h expressed as the weight of the free base) or vehicle occurred from P25 to P53 via osmotic pumps (Alzet®, model 2004). This dose was chosen based on previous work reported by Trauth et al. (1999), though their dose was based on the weight of the rat at the onset of exposure. Because adolescence is a period of rapid growth in rats, the concentration of nicotine we used was determined by estimating the average weight of female rats during the entire period of exposure (P25–P53) based on a Long Evans rat growth chart (Long Evans (Blue Spruce), 2006) with a target average exposure of 6.0 mg/kg/day over the entire period of exposure. Because osmotic pumps deliver contents at a constant rate, rats received 0.033 mg/h of nicotine or vehicle throughout the period of exposure. Weight data for nicotine-exposed rats during the period of exposure closely approximated the expected weights from the growth chart (Long Evans (Blue Spruce), 2006) for the days in the growth chart closest to the beginning and end of the exposure period. The group mean weight of nicotine-exposed rats near the beginning of exposure on P28 was 61.1 g (ranging from 34–75 g), which was comparable to a mean weight of 62.8 g from the growth chart for the same day. The group mean weight of nicotine-exposed rats near the end of exposure on P49 was 160.9 g (ranging from 141–181 g), which was comparable to a mean weight of 156.7 g from the growth chart for the same day. The overall mean nicotine dose for these rats based on constant nicotine exposure of 0.033 mg/h throughout the period of exposure and the overall mean body weight for the period of exposure was 6.79 mg/kg/day of nicotine expressed as the weight of the free base.

On P24 pumps were prepared with either nicotine bitartrate dissolved in bacteriostatic water (Sigma Chemical, Saint Louis, MO, expressed as the weight of the free base) or bacteriostatic water alone and left to prime in a saline solution for 24 h. On P25, rats were

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anesthetized under isoflurane and a pump was implanted subcutaneously in the mid-scapular region of the animal's back. Subjects were then allowed to recover from surgery in their home cages. Beginning the following day, P26, rats in the injection stress condition were weighed daily and received twice-daily intraperitoneal injections of 1.0 ml/kg of saline until P53 12 h apart. Rats in the No Stress condition were left undisturbed until P53 except for regular cage maintenance. On P53 the surgical procedure was repeated for all rats to remove their osmotic pumps. It should be noted that the osmotic pump, marketed as a four-week infusion device, actually takes approximately 33.3 days to be exhausted completely (information supplied by the manufacturer). After the osmotic pumps were removed, rats were housed individually (and remained so for the rest of the experiment) and allowed to recover in their home cages. This resulted in 4 groups of rats in this experiment: No Nicotine/No Stress, Nicotine/No Stress, No Nicotine/Stress, and Nicotine/Stress. Two rats were removed from the study prior to water restriction due to illness. Therefore, the current study used only 58 subjects: 15 subjects in the No Nicotine/No Stress group, 14 subjects in the Nicotine/No Stress group, 14 subjects in the No Nicotine/Stress group, and 15 in the Nicotine/Stress group.

At P61, access to water was restricted to prepare for shaping and experimental testing. Rats continued to receive free access to food in their home cage and after training received 5 min of access to water daily throughout the experiment. Rats were monitored for signs of dehydration, such as yellowing of the belly, loss of skin elasticity, and lethargy. If any rats displayed signs of dehydration their participation in the experiment was paused and they were given supplemental water. No rats in the current experiment displayed signs of dehydration, such as yellowing of the belly, loss of skin elasticity, and lethargy. If any rats displayed signs of dehydration their participation in the experiment was paused and they were given supplemental water. All rats in the current experiment displayed signs of dehydration, therefore supplemental water was not necessary. All rats were kept on a 15:9-h light–dark cycle, which is the standardized light–dark cycle for our facility, with testing occurring during the light portion of the cycle. A timeline of experimental procedures is depicted in Fig. 1.

2.2. Apparatus

Three clear Plexiglas® shaping chambers (15 cm wide × 30 cm long × 30 cm high) with stainless steel wire mesh flooring and a single nose poke receptacle (2.5-cm diameter PVC pipe end caps painted flat black) centered on one end wall 5.0 cm above the floor were used in this study. This receptacle contained an infra-red emitter and detector which were located on the left and right sides as well as a white LED cue light positioned on the back of the receptacle.

Six clear 1/4-inch Plexiglas® octagonal test chambers (15 cm wide × 30 cm tall walls with 40 cm between opposite walls) with wire mesh floors were used as the experimental apparatus. Each of the eight walls was equipped with a nose-poke receptacle described above centered 5.0 cm above the floor.

An opening located at the bottom of each receptacle, connected to a solenoid and syringe by plastic tubing, served to deliver water to the chamber. All chambers were enclosed within sound attenuating chambers with 10-ml syringes attached to an internal wall of the enclosure that served as water reservoirs. Syringes were connected by Tygon tubing (VWR Scientific, Performance Plastics 1/32-inch, #R-3603) to solenoids (General Valve Corp. Vac. 20 psig. 24 V) and then to the receptacles. The solenoid controlled the delivery of water droplets to the nose-poke receptacles.

Background white noise masked extraneous noise. All chambers were controlled by a computer running the MedPC interface (Med Associates interface; Grayson Stadler power supply Model E 783 DA) which was located in a separate room of the laboratory. Rats were monitored from the computer room via closed circuit cameras mounted inside the enclosures.

2.3. Procedure

2.3.1. Shaping procedure

On P63 and P64, all rats received two consecutive days of nose-poke shaping in shaping chambers. On each trial, the receptacle light was illuminated. After each nose-poke response, a reinforcer consisting of a 0.025-ml droplet of water was delivered through the bottom of the receptacle and the receptacle light was extinguished for the intertrial interval. Intertrial intervals were 1 s on P63 and 2 s on P64. Criterion for being included in the study was set at 240 responses within one hour on each of these two consecutive days. All rats met criterion on both shaping days.

2.3.2. Testing procedure

Testing in the SMC task began on P65, the day after the rats completed nose-poke shaping. All testing was conducted in our experimental 8-walled chamber. At the beginning of each trial, all 8 nose poke receptacles were illuminated and the rat was allowed to make a response at one of the 8 receptacles. If the rat's response was correct, all lights were extinguished and reinforcement consisting of a 0.025-ml droplet of water was delivered to the correct receptacle. If an incorrect choice was made at any time during experimental sessions, the correction procedure was initiated where all lights were extinguished except for the light of the correct receptacle and the rat was reinforced only after a nose poke response in the correct receptacle. After the correction procedure resulted in the rat's response to the indicated correct receptacle, the sequence continued as if a correct response had been produced on the trial. The computer recorded number of correct and incorrect responses as well as the location of the rats' response on each trial. Rats had to perform the following pattern:


where digits represent the clockwise position of the correct receptacles in the octagonal chamber on successive trials. Dashes indicate the location of 3-s pauses that served as phrasing cues separating structural chunks of the pattern. All other intertrial intervals were 1 s. The location of the receptacle designated as “I” was constant throughout the experiment for each rat but counterbalanced between rats. The first digit of each chunk is called a chunk-boundary element, the two digits following the chunk boundary are called within-chunk elements, and the last digit “8” in the pattern (underlined) is called the violation element because it violates pattern structure. Rats performed the

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Fig. 1. Timeline of experimental methods and procedures. Rats were weaned on postnatal day 21 (P21). Osmotic pumps containing either nicotine or vehicle were implanted on P26 and rats in the Stress groups began receiving twice-daily injections of saline on P26. Osmotic pumps were removed and stress injections were discontinued on P53. Starting on P63, all rats completed 2 days of shaping followed by 35 days of serial pattern learning training (ending on P99).
pattern 20 times a day without interruption, 7 days a week, for 35 days. Rats were allowed unlimited time to complete 20 pattern repetitions daily, resulting in sessions that lasted as long as approximately 2 h early in training, but average session length quickly decreased to approximately 30 min.

2.4. Statistical analysis

Analysis of variance (ANOVA) was used to examine the effects of adolescent exposure to nicotine and injection stress on rats’ acquisition for each element type (within-chunk, chunk-boundary, and violation elements) across the 35 days of the experiment. Main effects and interactions were considered significant if \( p < .05 \). To assess differences in acquisition of pattern elements, a 2 (drug condition) × 2 (stress condition) × 35 (days) repeated measures ANOVA was conducted on rats’ daily total correct responses for each element type. When significant effects were observed, planned comparisons based on the appropriate error term of the ANOVA (Fisher’s Least Significant Difference tests) were conducted to determine the direction of the effect as well as specific days within acquisition when groups differed. Multiple regression analysis was used to assess differences in asymptotic levels of performance over the last 10 days of acquisition. Lastly, “intrusion analysis” examined the types of errors committed by element type to assess possible changes in learning strategies caused by adolescent exposure to nicotine or injection stress.

3. Results

Generally, the results show that adolescent injection stress and adolescent nicotine exposure interacted in producing somewhat paradoxical effects in adulthood on some, but not all, aspects of serial pattern learning in adult female rats. Correct choice data showed that adolescent exposure to 0.033 mg/h of nicotine while not receiving stress from the injection procedure resulted in faster learning of the chunk-boundary element. However this exposure also resulted in impaired learning of the violation element on the last 10 days of acquisition.

Acquisition curves for within-chunk elements, which are the second and third elements of each chunk, are shown in Fig. 2. A drug × stress × day repeated measures ANOVA conducted on rats’ daily mean correct response rates on within-chunk elements revealed a significant main effect of days, \( F(34,1839) = 184.04, \ p < .001 \), but all other main effects and interactions were not significant (\( p > .05 \)), indicating that there were no significant differences in learning rate for within-chunk elements between groups.

Acquisition curves for chunk-boundary elements, that is, the first element of chunks that always immediately followed phrasing cues, are shown in Fig. 3. A drug × stress × day repeated measures ANOVA conducted on rats’ daily mean correct response rates on chunk-boundary elements revealed a significant main effect of day of the experiment, \( F(34,1836) = 415.11, \ p < .001 \). The ANOVA also revealed significant interactions for stress × day, \( F(34,1839) = 2.21, \ p < .001 \), and a drug × stress × day, \( F(34,1836) = 1.75, \ p < .005 \). The drug and stress main effects, however, were not significant (\( p > .05 \)). Planned comparisons based on the appropriate error term from the ANOVA showed that female rats in the Nicotine/No Stress group made significantly more correct responses than rats in all other groups, “A” indicates 1 day when Nicotine/No Stress rats made significantly more correct responses than rats in all other groups, “B” indicates 1 day when Nicotine/No Stress rats made more correct responses than the No Nicotine/No Stress and the Nicotine/Stress groups, and “C” indicates 1 day when Nicotine/No Stress rats made more correct responses than No Nicotine/Stress and the Nicotine/Stress rats (\( p < .05 \)). Error bars: ± SEM.

First, planned comparisons were performed to examine the effects of adolescent injection stress on violation element learning in adulthood independent of nicotine exposure. Planned comparisons revealed that adolescent injection stress alone caused a transient but significant
facilitation of learning to anticipate the violation element in adulthood in both No Nicotine and Nicotine groups. As shown in Fig. 5A, adolescent injection stress facilitated learning of the violation element for rats in No nicotine groups that did not receive adolescent nicotine exposure. Rats in the No Nicotine/Stress group made significantly more correct responses than rats in the No Nicotine/No Stress group on days 14, 16, and 18–20. It should be noted, however, that the opposite was true on days 8 and 9, that is, rats in the Nicotine/No Stress group made significantly more correct responses than rats in the Nicotine/Stress group. Overall, these results indicate that adolescent injection stress caused a transient but significant facilitation of violation element learning in adulthood, the opposite of what was found for chunk-boundary element acquisition.

Second, planned comparisons were performed to examine the effects of adolescent nicotine exposure on violation element learning in adulthood independent of adolescent injection stress. Planned comparisons showed that adolescent nicotine caused a transient but significant impairment of violation element acquisition in adulthood in rats that did not receive adolescent injection stress, but not in rats

Fig. 4. Acquisition curves for the violation element of the pattern showing daily mean total correct responses over 35 days of training beginning on P65 for all four groups (N = 14–15 per group). Rats received prior adolescent exposure to either 0.033 mg/h nicotine or an equivalent volume of saline via implanted osmotic pumps from P25–53. From P26–53 rats in the Stress conditions also received twice daily injections of 1.0 ml/kg saline. Error bars: ± SEM.

Fig. 5. Acquisition curves for the violation element of the pattern showing daily mean total correct responses over 35 days of training beginning on P65 replotted from Fig. 4 to assess the effects of adolescent nicotine for rats in the No Stress condition that received no injection stress (Panel A) or for rats that received stress via twice daily injections during adolescence (Panel B) (N = 14–15 per group). Error bars: ± SEM. *p < 0.05.
that did receive adolescent injection stress. As shown in Fig. 6A comparing data for No Stress groups, rats in the Nicotine/No Stress group made significantly fewer correct responses than rats in the No Nicotine/No Stress group on days 20, 25, 29, and 30. However, as shown in Fig. 6B comparing data for Stress groups, no differences were observed in violation element acquisition in adulthood between the Nicotine and No Nicotine groups in rats that received adolescent injection stress. Overall, these results indicate that adolescent nicotine exposure caused a transient but significant impairment of violation element learning in adulthood, but only in rats that did not experience adolescent injection stress.

In the earlier Pickens et al. (2013) study using the same serial pattern, in female rats the most profound learning impairment caused by adolescent nicotine exposure was observed for the violation element late in training. In order to determine if adolescent exposure to nicotine had an effect on violation element performance late in training, a multiple regression was performed for the last 10 days of training. This set of days was chosen because planned comparisons detected no significant differences in daily mean performance by rats in the No Nicotine/No Stress control group during this period. The multiple regression found that only the original violation error model (which did not include the interaction term) was significant. It accounted for 32% of the variance in violation errors. According to the regression model, only nicotine was a significant predictor of violation errors ($b = .32, t(57) = 2.53, p = .014$). This variable also accounted for a significant amount of the variance in the outcome ($R^2 = .105, F(2, 55) = 3.23, p = .047$). As shown in Fig. 7, further analysis of the data, determined that rats not exposed to nicotine made fewer errors on the violation element (Mean $M = 1.09$, Standard Deviation $SD = 1.01$) than rats that were exposed to nicotine ($M = 3.32$, $SD = 3.09$). No other significant main effects or interactions were found for the models ($p > .05$).

As shown in Fig. 8, error data were further analyzed with an intrusion analysis to determine types of errors committed for the different element types. For chunk-boundary elements, the most frequent type of error for all conditions was a perseveration response, that is, a repetition of the last correct response. An example of such an error after a 2-3-4 chunk would be a “4” response on the chunk boundary trial following the phrasing cue when a “3” response was correct. Across all groups, this type of error accounted for 36–42% of all errors made on chunk-boundary trials. For the violation element, rats in all conditions responded similarly, with all groups producing more than 80% of their errors as overextensions. This error occurs when the rats extrapolate the within-chunk “+1” rule; that is, on the third element of the violation chunk, 8-1-8, all rats tended to respond with a rule-consistent but incorrect “2”—rather than “8”—to produce an 8-1-2 chunk that was structurally consistent with the rest of the chunks of the pattern. These results suggest that adolescent injection stress and nicotine exposure do not create changes in behavioral strategy. Number of errors varied between groups but the cognitive strategies employed remained the same.

4. Discussion

This study was designed to examine the effects of chronic injection stress and concurrent chronic nicotine exposure experienced during adolescence on adult learning in female rats. Four general results were obtained in the current study. First, learning for within-chunk elements was not affected by either adolescent nicotine exposure, consistent with past work (Pickens et al., 2013), or adolescent injection stress. Thus, there were no effects of adolescent nicotine exposure or injection stress on adult within-chunk learning typically attributed to rule learning in the SMC task, as shown in Fig. 2. Second, adolescent injection stress alone (i.e., without concurrent nicotine exposure) caused transient but significant facilitation of adult learning restricted to a single element of the 24-element pattern, namely, the “violation element,” that was the only element of the pattern that was inconsistent with pattern structure. Thus, adolescent injection stress alone facilitated violation element acquisition in adulthood, as shown in Fig. 6A. Third, also consistent with past work (Pickens et al., 2013), adolescent nicotine exposure, in this case both with and without adolescent injection stress, caused a learning impairment in adulthood for the violation element in female rats, as shown in Fig. 7. Thus, adolescent nicotine impaired adult violation element learning typically attributed to multiple-item learning in the SMC task. Fourth, a paradoxical interaction of injection stress and nicotine exposure in acquisition was observed. In the same female rats in which violation-element learning was impaired (Fig. 7), adolescent nicotine experienced without adolescent injection stress produced better learning for chunk-boundary elements in adulthood compared to all other conditions (Fig. 3).

Previous research has shown that serial pattern learning in the SMC task recruits multiple cognitive systems concurrently, including associative S–R learning, serial position learning involving timing or counting processes, and rule abstraction processes (Fountain and Benson, 2006; Fountain et al., 2008; Fountain et al., 2012; Kundey and Fountain,
Learning to anticipate chunk-boundary elements has been shown to depend on both associative stimulus response (S–R) learning and serial-position learning concurrently (Muller and Fountain, 2010; Stempowski et al., 1999). Learning to anticipate the violation element has been shown to depend on associative multiple-item learning involving cues from several preceding trials and “intra-box” apparatus cues that signal the upcoming violation trial (Kundey and Fountain, 2010; Muller and Fountain, 2010). Learning to anticipate within-chunk elements, on the other hand, has been shown to depend on learning a motor program or abstract rules that are independent of external stimuli (Muller and Fountain, 2010). Since the different pattern element types are learned using distinct cognitive mechanisms, it is not surprising to find that the same drug or toxic agent can result in differential facilitation of learning, impairment of learning, and no effect on learning for different element types in individual rats in the SMC task. Dissociations in learning and performance consistent with the foregoing behavioral and cognitive distinctions have been observed in rats following acute systemic treatment with MK-801, an N-methyl-D-aspartate receptor (NMDAr) antagonist, and with atropine, a muscarinic cholinergic antagonist (Fountain and Rowan, 2000; Fountain et al., 2013).

Similar dissociations have also been observed in adolescent nicotine effects on adult learning. Pickens et al. (2013) demonstrated sex-specific impairments of discrimination learning for chunk-boundary elements in male rats, but not female rats, and impairments of multiple-item discrimination learning for violation elements in female rats, but not male rats. Neither adult male nor female rats were impaired in rule-based learning for within-chunk elements after adolescent nicotine exposure. The current study replicated the dissociation of effects observed by Pickens et al. (2013) where female rats exposed to nicotine in adolescence had impaired learning of multiple-item learning for violation elements whereas S–R and serial-position learning for chunk-boundary elements was not impaired. Thus, based on the results of Pickens et al. (2013), a dissociation of nicotine effects on multiple-item learning for violation elements versus S–R and serial-position learning of chunk-boundary elements in female rats in the current study was expected. However, facilitated learning for chunk-boundary elements in adulthood after adolescent nicotine exposure was unanticipated and, given the growing literature on learning impairments caused by adolescent nicotine exposure in rodent models, it was indeed surprising.

The current study found that when nicotine was experienced without concurrently experiencing injection stress, nicotine in adolescence facilitated learning in adulthood long after chronic adolescent nicotine exposure ceased. The critical behavioral effect supporting this claim is the fact that the group receiving adolescent nicotine without stress (i.e., the Nicotine/No Stress group) learned chunk-boundary elements faster during adulthood than all other groups including its control group (the No Nicotine/No Stress group), as shown in Fig. 3. Taken together, the evidence indicates that adolescent nicotine exposure without accompanying adolescent injection stress caused facilitation of S–R and/or serial position learning involved in learning to anticipate chunk-boundary elements in the SMC task. It should be noted that, although facilitated learning or performance is commonly observed after acute nicotine exposure in adulthood (Attaway et al., 1999; Barron et al., 2005; Levin and Torry, 1996; Semenova et al., 2007) evidence of facilitated learning in adulthood after adolescent nicotine exposure appears to be unprecedented in the literature on adolescent nicotine exposure effects on adult learning.

It is possible that the unusual pattern of behavioral effects observed in this study may have been related to the method used for adolescent nicotine exposure. Whereas earlier studies employing the SMC task used “bolus intraperitoneal” (c.f. Silverstone et al., 2008) injections of nicotine (Fountain et al., 2008; Pickens et al., 2013), the current study used continuous subcutaneous infusion of nicotine during adolescence. Intraperitoneal injections of nicotine cause a rapid surge in the concentration of nicotine followed by a gradual return to baseline levels and since injections were adjusted as a function of rats’ daily weight in prior studies, the concentration of nicotine at the time of injections was a constant dose throughout the experiment. In contrast, continuous subcutaneous infusion of nicotine over many days via osmotic pump, as used in the present study, delivered a steady infusion of a nicotine solution of consistent concentration throughout the exposure period of many days so that as a rat’s body weight increased during adolescence, the daily nicotine dose decreased as a proportion of body weight over the period of infusion. It is also important to note that the rats in the present study received a nicotine dose of 0.033 mg/h, which is an average over the course of the entire experiment of approximately 6.79 mg/kg nicotine daily via infusion. This dose was on average six times higher than the 1.0 mg/kg daily dose delivered via bolus intraperitoneal injection in our previous research (Fountain et al., 2008; Pickens et al., 2013). However, because continuous infusion by osmotic pump would deliver nicotine at only 0.033 mg/h, the highest tissue concentration of nicotine actually experienced at any time in the present study would have been much lower than the peak concentration experienced in earlier studies employing 1.0 mg/kg intraperitoneal injections. Perhaps a higher daily dose of nicotine or the fact that it was delivered via continuous subcutaneous infusion rather than bolus intraperitoneal injections contributed to the interaction of adolescent nicotine and adolescent injection stress that produced facilitated learning for chunk-boundary elements.

It should also be noted that the current experiment found that in the absence of adolescent nicotine exposure, there was no effect of adolescent injection stress on adult learning for within-chunk and chunk-boundary elements. However, injection stress during adolescence did cause transient but significant facilitation of violation element learning for both Nicotine and No Nicotine groups (Fig. 5A and 5B). No Nicotine/Stress rats made significantly more correct violation element responses for 3 days of acquisition compared to No Nicotine/No Stress rats. Furthermore, Nicotine/No Stress rats made significantly more correct violation responses on 6 days of acquisition compared to Nicotine/Stress rats. One question raised by the current experiment is how best to study the effects of adolescent nicotine while controlling stress as a factor. Methods for exposing rats to nicotine while minimizing stress due to handling, surgery, or injection—such as by adding nicotine to drinking water or food—also have methodological drawbacks.

The results of the current study demonstrated that injection stress did interact with adolescent nicotine exposure to produce paradoxic effects on adult learning. Adolescent nicotine exposure in the absence of concurrent adolescent injection stress resulted in facilitated S–R discrimination learning for chunk-boundary elements while concurrently impairing multiple-item memory of cues important for anticipating the violation element in adulthood in female rats. Previous behavioral and neurobiological work has shown that learning to anticipate chunk-boundary elements depends on different cognitive/behavioral mechanisms than learning to anticipate violation elements, and that these different behavioral mechanisms depend on dissociable neural mechanisms (for a summary, see Fountain et al., 2012). For that reason it is not surprising that the same manipulation—in this case, adolescent nicotine exposure—might affect multiple brain/behavioral mechanisms differentially in the same animal, perhaps even in opposite directions as observed in the current study in chunk-boundary element versus violation element learning. One reviewer of this paper suggested that perhaps adolescent injection stress may have one or more general effects, for example, that adolescent injection stress reverses or interferes with adolescent nicotine effects. Two pieces of evidence from the current study are consistent with this idea in the context of our multiple-process view. First, adolescent injection stress appeared to reduce nicotine-induced facilitation of learning for chunk-boundary elements in Fig. 3. Second, adolescent injection stress reduced the nicotine-induced impairment of learning the violation element in Fig. 6. One hypothesis is that adolescent nicotine exposure can produce
differential effects—facilitation or impairment—depending on the system, but that adolescent stress always interferes with the expression of adult sensitive responses. Although this view of adolescent injection stress has its appeal, data such as those of Fig. 7, where adolescent nicotine effects appear to be the same across different levels of adolescent injection stress, are not consistent with this hypothesis. Interestingly, adolescent injection stress when considered alone appears to facilitate learning of violation elements, as shown in both Fig. 5A and 5B, but has no such effect on learning chunk-boundary elements, as shown in Fig. 3. One conclusion is that like the effects of adolescent nicotine exposure, the effects of adolescent injection stress may vary across different neural, behavioral, and cognitive systems.

How to interpret the results of the current study from a public health perspective is an open question. However, adolescent smokers may encounter a wide variety of acute and chronic stressors as they go about their lives. Thus, the effects of adolescent nicotine exposure under different conditions of concurrent adolescent stress deserve further scrutiny, especially with regard to possible effects of both of these factors on cognitive capacity in adulthood. Furthermore, other studies have shown that exposure to nicotine can cause cognitive enhancement when nicotine is administered in adulthood and acutely just prior to behavioral assessment (Attaway et al., 1999; Barron et al., 2005; Levin and Torry, 1996; Semenova et al., 2007). Numerous studies with rodent models, including our own demonstrating sex differences in adolescent nicotine effects on adult learning in rats, have already shown that adolescent exposure to nicotine causes cognitive impairments in adulthood. To the best of our knowledge, the current study is the first to demonstrate facilitation of learning in adulthood caused by adolescent exposure to nicotine. Following on this discovery, it will be able to properly predict the sequelae of adolescent nicotine exposure it will be of great importance to identify the conditions under which adolescent nicotine causes cognitive facilitation or enhancement in adulthood versus cognitive impairment. This, in turn, will provide a necessary foundation for identifying the relevant neurobiological mechanisms responsible for both these effects.

Transparency document

The Transparency document associated with this article can be found, in the online version.

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