Iptakalim attenuates self-administration and acquired goal-tracking behavior controlled by nicotine

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IPTAKALIM ATTENUATES SELF-ADMINISTRATION AND ACQUIRED GOAL-TRACKING BEHAVIOR CONTROLLED BY NICOTINE

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Abstract

Iptakalim is an ATP-sensitive potassium channel opener, as well as an \( \alpha_4\beta_2 \)-containing nicotinic acetylcholine receptor (nAChR) antagonist. Pretreatment with iptakalim diminishes nicotine-induced dopamine (DA) and glutamate release in the nucleus accumbens. This neuropharmacological profile suggests that iptakalim may be useful for treatment of nicotine dependence. Thus, we examined the effects of iptakalim in two preclinical models. First, the impact of iptakalim on the interoceptive stimulus effect of nicotine was evaluated by training rats in a discriminated goal-tracking task that included intermixed nicotine (0.4 mg/kg, SC) and saline sessions. Sucrose was intermittently presented in a response-independent manner only on nicotine sessions. Sucrose was intermittently presented in a response-independent manner only on nicotine sessions. On intervening test days, rats were pretreated with iptakalim (10, 30, 60 mg/kg, IP). Results revealed that iptakalim attenuated nicotine-evoked responding controlled by the nicotine stimulus in a dose-dependent manner. In a separate study, the impact of iptakalim on the reinforcing effects of nicotine was investigated by training rats to lever-press to self-administer nicotine (0.03 mg/kg/infusion). Results revealed that pretreatment with iptakalim (1, 3, 6 mg/kg, IV) decreased nicotine intake (i.e., less active lever responding). Neither behavioral effect was due to a non-specific motor effect of iptakalim, nor to an ability of iptakalim to inhibit DA transporter (DAT) or serotonin transporter (SERT) function. Together, these finding support the notion that iptakalim may be an effective pharmacotherapy for increasing smoking cessation and better understanding its action could contribute medication development.
INTRODUCTION

Over 400,000 people in the United States die annually from tobacco-related diseases (CDC, 2008). It is estimated that $193 billion per year of expenditures are associated with health care cost and loss of productivity related to the consumption of tobacco products (CDC, 2008). Even with a number of smoking cessation treatments available (e.g., Zyban®, Chantix®), only about 20% of users remain abstinent for over a year (Schnoll and Lerman, 2006). Clearly there is still great need for new and effective treatments for nicotine dependence.

Iptakalim is an adenosine triphosphate-sensitive potassium channel opener that is approved for use in treating hypertension in China (Wang et al., 2005a; Wang et al., 2005b). Along with its action on potassium channels, iptakalim has several other neuropharmacological effects that suggest that it could be used to treat nicotine dependence. For example, iptakalim is an α4β2 nicotinic acetylcholine receptor antagonist (Hu et al., 2006). Nicotine increases dopamine and glutamate release in the nucleus accumbens. Pretreatment with iptakalim reduces this nicotine-induced intracumbal dopamine and glutamate release (Hu et al., 2006; Liu et al., 2006). Behaviorally, iptakalim under some conditions can decreases nicotine-induced hyperactivity (Hu et al., 2006; Liu et al., 2006; Volf et al., 2012). Given these findings, the first goal of the current research was to examine the effect of iptakalim on nicotine in two separate animal models: the discriminated goal-tracking task that assesses the interoceptive stimulus effects of nicotine and the intravenous (IV) drug self-administration task that assess the positive reinforcing effects of nicotine.

The interoceptive stimulus effects of nicotine and the associated learning processes are thought to be important to chronic tobacco use and nicotine dependence (Bevins and Palmatier, 2004). Indeed, the pharmacological effects of nicotine that generate a perceptible internal stimulus (cf. drug intoxication) can come to control reinforced behaviors in rodents and primates—including humans (for reviews see Smith and Stolerman, 2009; Wooters et al., 2009). In the present study, we used the drug discriminated goal-tracking (DGT) task with rats to investigate the effects of iptakalim on nicotine in two separate animal models: the discriminated goal-tracking task that assesses the interoceptive stimulus effects of nicotine and the intravenous (IV) drug self-administration task that assess the positive reinforcing effects of nicotine.

Another factor contributing to chronic tobacco use is the reinforcing effect of nicotine (Henningfield and Goldberg, 1983; Le Foll and Goldberg, 2009; Stolerman and Shoaib, 1991). Intravenous nicotine self-administration in rats has been commonly used to assess the reinforcing effects of nicotine. In this task, rats have a catheter implanted into the jugular vein. Rats are then given an opportunity to make one of two responses (e.g., press on the right or left lever). Responding as prescribed by the experimenter on one of the two levers will produce an infusion of nicotine; responding on the other inactive lever has no programmed consequence. Nicotine self-administration, as indicated by greater responding
on the drug lever, has been demonstrated in many laboratories (Caggiula et al., 2002; Corrigall and Coen, 1989; DeNoble and Mele, 2006; Donny et al., 1995; Feltenstein et al., 2012; Neugebauer et al., 2006). The α₄β₂ antagonist DHβE attenuates nicotine self-administration (Grottick et al., 2000; Tobey et al., 2012; Watkins et al., 1999).

We found that iptakalim attenuated responding in the DGT task and decreased the intake of nicotine in the self-administration task. Monoamine transporter inhibition can likewise diminish these behaviors. For example, pretreatment with the selective serotonin reuptake inhibitor citalopram blocks nicotine-evoked responding in the DGT task (Dion et al., 2012). Further, the dopamine reuptake inhibitor bupropion substitutes for the nicotine stimulus (Wilkinson et al., 2010). Given these findings, a second goal of the present research was to examine the effects of iptakalim on DAT and SERT to determine if these transporters contribute to iptakalim’s mechanism of action.

**MATERIALS**

**Animals**

Thirty male Sprague-Dawley rats weighing 275–299 grams upon arrival from Harlan (Indianapolis, IN, USA) were housed individually in clear polycarbonate cages (48.3x26.7x20.3 cm) lined with wood shavings. The temperature- and humidity-controlled colony was on a 12-h light/dark schedule (lights on at 0600); experiments were conducted during the light cycle. Water was freely available in the home cage. Rats’ weight in the DGT task was maintained at 85% of their free-feeding body weight. For rats in the self-administration study, chow was available ad libitum for the day before surgery and the 6 days following surgery. Otherwise, rats were maintained at 90% of their free-feeding body weight. Four weeks into the self-administration study, the target weight was increased by 2 g to adjust for average growth pattern provided by the supplier. Experimental protocols were approved by the University of Nebraska-Lincoln Institutional Animal Care and Use Committee.

**Apparatus**

The conditioning chambers (ENV-008CT; Med Associates, Inc.; St. Albans, VT, USA) measuring 30.5x24.1x21.0 cm (l×w×h), were enclosed in a sound- and light-attenuating cubicle equipped with an exhaust fan. Each chamber had aluminum sidewalls, metal rod floors with polycarbonate front, back, and ceiling. A recessed receptacle (5.2x5.2x3.8 cm; l×w×d) was centered on one sidewall. A dipper arm, when raised, provided access to 0.1 ml of 26% (w/v) sucrose solution in the receptacle. Access to the dipper was monitored by an infrared beam mounted 1.2 cm into the receptacle and 3 cm above the floor. A second infrared beam that monitored chamber activity was located 4 cm above the floor and 14.5 cm from the side wall containing the receptacle. Two retractable levers (147 nN required for micro-switch closure) were mounted on each side of the receptacle. A white cue-light (2.54 cm diameter; 28V, 100 mA) was mounted 7 cm above each lever. A house light (two white 28V, 100 mA lamps) was located 10 cm above the conditioning chamber. The infusion pump (PMH-100VS; Med Associates; St. Albans, VT, USA) for each chamber was located outside the sound-attenuating cubicle. A 5-ml syringe mounted on the infusion pump was connected with Tygon® tubing (AAQ04103; VWR; West Chester, PA, USA). The tubing was attached to a swivel coupled with a spring leash (C313C; Plastics One; Roanoke, VA, USA) which were suspended over the ceiling of the chamber on a balanced metal arm. Med Associates interface and software (Med-PC for Windows, version IV) were used to collect data and present programmed events.
Drugs—(-)-Nicotine hydrogen tartrate (Sigma; St. Louis, MO, USA) and iptakalim hydrochloride (99.9%; synthesized and provided by the Institute of Pharmacology and Toxicology, Academy of Military Medical Sciences of China) were dissolved in 0.9% saline. The pH of the nicotine solution was adjusted to 7.0±0.2 with a dilute NaOH solution. Nicotine doses (reported as base), iptakalim doses (reported as salt), route of injection, and injection-to-placement intervals (i.e., time between the injection and the onset of behavioral assessment) were selected based on previous research (Charntikov et al., 2012; Murray and Bevins, 2007a, b; Sun et al., 2010). In the DGT study, nicotine was administered subcutaneously (SC) and iptakalim was administered intraperitoneally (IP) at a volume of 1 ml/kg. In the self-administration study, nicotine was infused intravenously (IV) 35.74 μl per infusion across 1 sec; iptakalim was administered IV at 1 ml/kg (see later for more detail).

METHODS

Discriminated goal-tracking task

Rats in this study (n=10) were trained on our standard DGT protocol (as previously described in Bevins et al., 2012; Wilkinson et al., 2010). Before this study, these rats served in a separate experiment investigating how drug combinations, some of which bind to nAChRs (e.g., sazetidine A, nornicotine, bupropion), affected responding. This use of trained animals in the drug discrimination field, and in our lab specifically, is common given that performance is reliable and repeatable across time (Polewan et al., 2013; Smith et al., 2009; Struthers et al., 2009; Wooters et al., 2009). For the present study, all rats were tested on all doses of iptakalim using a repeated 5-day testing cycle. The first 4 days of this standardized testing protocol were training sessions to insure the nicotine/saline discrimination was maintained. For each daily training session, rats were injected SC with either nicotine (0.4 mg/kg) or saline 5 min before placement into the chamber for a 20-min session. There were 2 nicotine and 2 sucrose sessions intermixed per 4-day training cycle. Access to sucrose was initiated between 124 to 152 s from the start of the nicotine session. There were 36 separate 4-sec deliveries of sucrose per nicotine session. Time between sucrose deliveries ranged from 4 to 80 s (mean = 25 s). Sucrose was unavailable during the saline sessions. On day 5 of the test cycle, rats were injected with a randomly assigned dose of iptakalim (0, 10, 30, or 60 mg/kg; IP) 10 min before the start of a 4-min test session; nicotine was administered 5 min before the test. During test sessions, head entries into the dipper receptacle and general chamber activity were recorded, but sucrose was not available. Once all doses of iptakalim in combination with nicotine were tested, we noted that the combination with 60 mg/kg iptakalim decreased chamber activity. To determine if iptakalim impaired spontaneous locomotion in a manner that could account for the decrease in goal-tracking behavior evoked by the nicotine stimulus, we then tested all doses of iptakalim alone (no nicotine). These tests proceeded similar to those previously described except saline replaced nicotine as the solution injected 5 min before the 4-min test session. For all DGT sessions, the house light remained off and both levers were retracted.

IV nicotine self-administration task

Preliminary training—Before the start of this study, a separate set of rats (n=20) were handled for a minimum of 2 min per each of three consecutive days. Rats were then trained to lever press. The start of each lever-press session was signaled by illumination of the house light and insertion of a randomly selected lever (right or left). A lever press or a lapse of 15 sec resulted in 4-sec access to sucrose, retraction of the lever, and commencement of a timeout that lasted on average 60 s (range=30 to 89 s). Following the timeout, a randomly selected lever was again inserted into the chamber with the condition that the same lever could not be presented more than 2 times in a row. This protocol was repeated for 60 sucrose deliveries. Daily sessions ranged from 65–80 min, depending on individual
performance and were continued until a lever press was made on at least 80% of the lever insertions for two consecutive days (i.e., 3 to 5 training sessions).

**Catheter surgery**—Following at least 24 h after preliminary training, rats were anesthetized with 1 ml/kg ketamine (100 mg/ml) and xylazine (20 mg/ml) mixture (2:1 ratio; administered intramuscularly; Sigma; St. Louis, MO, USA). Polyurethane catheter (RJVR-23; Strategic Applications Inc.; Lake Villa, IL, USA) with rounded tip and double suture beads (one secured internally and other externally) was implanted into the right external jugular vein. The other end of the catheter was subcutaneously placed around the shoulder and exited below the scapula via subcutaneously implanted polycarbonate back-mount access port (313-000BM; Plastics One Inc.; Roanoke, VA, USA). Immediately following surgery, catheters were flushed with 0.2 ml of streptokinase (2 mg/ml; Sigma; St. Louis, MO, USA) diluted in sterile heparinized saline (30 U/ml; Midwest Veterinary Supply; Lincoln, NE, USA). Atipamezole hydrochloride (0.5 mg/kg; IM; Sigma; St. Louis, MO, USA) diluted in saline was used to terminate anesthesia (Wee et al., 2006). To manage post-surgical pain, buprenorphine hydrochloride (0.1 mg/kg; SC) was administered immediately after the surgery and daily for the next two recovery days. Starting from the day after surgery, catheters were flushed daily with heparinized saline (30 U/ml). Catheter patency was assessed when patency loss was suspected or upon completion of the self-administration study using an infusion of 0.05 ml xylazine (20 mg/ml; IV). This xylazine concentration produces motor ataxia within 5 sec (cf. Bevins, 2005; Reichel et al., 2008). Only the 13 rats with patent catheters were included in the analyses.

**Continued preliminary training**—Following 7 days of recovery from surgery in the home cage, rats were trained for 3 consecutive daily sessions to lever press for liquid sucrose on a variable ratio (VR3) schedule of reinforcement (i.e., on average every third response was followed by access to sucrose; range=1 to 6 presses). This training was similar to the pre-surgery training except lever pressing was now required to access sucrose. Across the 3 daily sessions, all rats earned at least 80% of the 60 available sucrose deliveries and were thus moved to the self-administration phase (Nb. this protocol insures a high baseline level of lever pressing with both levers having a similar reinforcement history).

**Nicotine self-administration**—Before the start of each 60-min self-administration session, catheters were flushed with 0.2 ml of heparinized saline. The start of each session was signaled by turning the house light off, priming the catheter with nicotine (ca. 31 μl or 90% of internal catheter volume), and insertion of both levers. Which lever served as the active lever was balanced across the rats. The active lever was reinforced on aVR3 schedule of reinforcement. A VR schedule was used because in preliminary studies in our laboratory, this reinforcement schedule produced more reliable self-administration and lever discrimination than other schedules (e.g., fixed ratio [FR] 1 or FR5). Further, VR schedules typically control high and steady rates of behavior which may provide enhanced sensitivity to experimental manipulations (Williams, 1988). Upon completion of the VR3, there was a 1-sec infusion of nicotine (0.01 mg/kg/infusion), retraction of both levers, and illumination of the house light for a 20-sec timeout. Following the timeout, the house light was turned off and levers were inserted back into the chamber. Inactive lever responding was recorded but there was no programmed consequence. Immediately after each self-administration session, catheters were flushed with the antibiotic cefazolin (10 mg) diluted in 0.2 ml of heparinized saline (30 U/ml). After rats had reached a criterion of 80% discrimination between the active and inactive lever, iptakalim testing commenced.

**Iptakalim dose response**—Iptakalim (0, 1, 3, or 6 mg/kg) was administered IV 5 min before the start of the self-administration session. The IV route was adopted because of...
limited drug availability at the time of the experiment. The 10 fold decrease in doses for iptakalim for IV administration was adapted from Bardo et al (1999). All doses were tested in a randomly selected order for each rat. Following each test day there were at least 2 iptakalim-free self-administration sessions. At least 80% responding had to be on the active lever before being tested with the next randomly selected iptakalim dose.

\[ ^3\text{H} \]dopamine and \[ ^3\text{H} \]serotonin uptake

Inhibition of \[ ^3\text{H} \]dopamine and \[ ^3\text{H} \]serotonin uptake by iptakalim was determined using a rat striatal (dopamine) or hippocampal (serotonin) synaptosomal preparation previously described (Hadlock et al, 2011). Briefly, synaptosomes were prepared by homogenizing freshly dissected striatal or hippocampal tissue in ice-cold 0.32 M sucrose pH 7.4, and centrifuged (800 × g, 12 min; 4°C). The supernatants were centrifuged (22,000 × g, 15 min; 4°C) and the resulting pellets were re-suspended in ice-cold assay buffer (in mM: 126 NaCl, 4.8 KCl, 1.3 CaCl2, 16 sodium phosphate, 1.4 MgSO4, 11 glucose and 1 ascobic acid; pH 7.4) and 1 μM pargyline. Iptakalim (1 nM–1 mM) was present in the assay tubes. Samples were incubated for 10 min at 37°C and the assays initiated by the addition of \[ ^3\text{H} \]dopamine or \[ ^3\text{H} \]serotonin (0.5 nM or 5 nM final concentration, respectively). Following incubation for 3 min, samples were placed on ice to stop the reaction and were filtered through GF/B filter paper (Whatman; Florham Park, NJ, USA) soaked previously in 0.05% polyethylenimine. The filter paper was rapidly washed three times with 3 ml of ice-cold 0.32 M sucrose buffer using a filtering manifold (Brandel; Gaithersburg, MD, USA). For \[ ^3\text{H} \]dopamine uptake, nonspecific values were determined in the presence of 50 μM cocaine. For \[ ^3\text{H} \]serotonin uptake, nonspecific values were determined in the presence of 10 μM fluoxetine. Radioactivity trapped in the filter paper was counted using a liquid scintillation counter.

STATISTICAL ANALYSES

An omnibus analysis of variance (ANOVA) preceded all planned comparisons. Higher-order interactions were further analyzed by two-way ANOVAs or ANCOVA and followed, if necessary, by Tukey’s HSD post-hoc tests. Statistical significance was set at p<0.05. To examine the effect of iptakalim on dipper entry rates and chamber activity, a separate one-way ANOVA with iptakalim dose as the within-subjects factor was performed. To fully assess the potential locomotor effects of iptakalim on nicotine-associated activity, a separate LSD comparisons to a saline control (0 mg/kg iptakalim + saline; dashed line on Figure 1B) were conducted. The magnitude and persistence of nicotine self-administration behavior was assessed using a 2x6 (Lever × Session) repeated measures ANOVA. The effect of iptakalim on nicotine self-administration behavior was analyzed using a 2x4 (Lever × Dose) ANOVA. To examine the effect of iptakalim on general chamber activity during nicotine self-administration, a one way ANOVA with iptakalim doses as independent and activity as dependent variables was performed. To examine the effect of iptakalim and cocaine on striatal and hippocampal synaptosomal \[ ^3\text{H} \]dopamine and \[ ^3\text{H} \]serotonin uptake, respectively, IC50 values were assessed using a least squares, non-linear regression fit with a minimum of seven data points (determined in triplicate) per curve.

RESULTS

Discriminated goal-tracking task

Figure 1 shows the impact of iptakalim in the DGT task. Iptakalim attenuated nicotine-evoked conditioned responding [main effect of Dose; F(3,27)=20.87, p<0.001; Figure 1A]. Dipper entries following pretreatment with 30 and 60 mg/kg iptakalim were significantly lower than with 0 (saline) or 10 mg/kg iptakalim pretreatment (Tukey HSD tests). Iptakalim

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also attenuated chamber activity in these nicotine test sessions [main effect of Dose; 
\( F(3,27) = 12.53, \ p < 0.01 \); Figure 1B]. Pretreatment with 30 and 60 mg/kg of iptakalim 
decreased activity relative to saline or 10 mg/kg iptakalim (Tukey HSD tests). When this 
chamber activity was compared to a saline control value (i.e., both injections were saline; 
dashed line on Figure 1B) only pretreatment with 60 mg/kg of iptakalim significantly 
reduced activity (LSD comparisons). Notably, in these brief 4-min test sessions, nicotine did 
not have a locomotor stimulant effect; no difference in chamber beam breaks between the 
nicotine and the saline alone condition.

Because iptakalim attenuated activity of nicotine-treated rats, an additional mixed general 
linear model ANCOVA was performed with dipper entries as a dependent measure, 
nptakalim dose as the within-subjects variable, and chamber beam breaks as the covariate. 
This ANCOVA revealed that blockade of nicotine-evoked dipper entries [main effect of 
Dose; \( F(3,23) = 19.73, \ p < 0.001 \)] cannot be explained by the variance in activity during the 
test session [no main effect of Activity and no Activity by Dose interaction]. Furthermore, 
nptakalim did not affect dipper entries or chamber activity (n.s.) in saline-treated rats (Figure 
1C and 1D). Combined, these findings suggest that antagonism of the stimulus effects of 
nicotine by iptakalim cannot be attributed to a non-specific motor effect of iptakalim.

**IV nicotine self-administration task**

Figure 2 shows the impact of iptakalim in a second preclinical model of nicotine abuse 
liability—nicotine self-administration. During the first 6 days of self-administration, 
responding on the active lever was significantly higher than on the inactive lever [main 
effect of Lever; \( F(1,216) = 265.15, \ p < 0.001 \); Figure 2A]. This difference was evident from 
day 1 on [Lever \( \times \) Session interaction; \( F(5,216) = 3.00, \ p < 0.05 \); Tukey HSD tests]. 
Pretreatment with iptakalim significantly attenuated responding on the active lever without 
affecting responding on the inactive lever [main effect of Lever; \( F(1,13) = 120.10, \ p < 0.001 \); 
Lever \( \times \) Dose interaction; \( F(3,39) = 12.61, \ p < 0.001 \); Tukey HSD tests; Figure 2B]. 
Specifically, all iptakalim doses significantly attenuated active lever responding in 
comparison to saline [separate one-way ANOVA; main effect of Dose; \( F(3,39) = 15.59, \ p < 0.001 \); 
Tukey HSD tests; Figure 2B]. Iptakalim also decreased chamber activity [main 
effect of Dose; \( F(3,39) = 5.96, \ p < 0.01 \); Figure 2C]. A follow-up mixed general linear model 
ANCOVA on the active lever responding with the total chamber beam breaks as a covariate 
revealed that the attenuation of nicotine intake by iptakalim [main effect of Dose; 
\( F(3,35) = 16.07, \ p < 0.001 \)] cannot be explained by alterations in activity [no main effect of 
Activity; \( F(1,35) = 0.34, \ p = 0.55 \); and no Activity by Dose interaction; \( F(3,35) = 1.61, \ p = 0.20 \). 
\[^{3}\text{H}]\text{dopamine and }[^{3}\text{H}]\text{serotonin uptake}

Ligands that act, in part, by inhibiting monoamine transporters interfere with nicotine-
evoked goal-tracking and nicotine self-administration (Di Chiara, 2000; Dion et al, 2012; 
Rauhut et al, 2003; Wilkinson et al, 2010). Thus, the impact of iptakalim per se on 
dopamine and serotonin transporter function was evaluated. Findings from the present study 
revealed that iptakalim was without effect, at concentrations ranging from 1 nM – 0.1 mM, 
on either \[^{3}\text{H}]\text{dopamine or }[^{3}\text{H}]\text{serotonin uptake, as assessed in striatal and hippocampal 
synaptosomes, respectively (see Figure 3). As in internal control to verify the integrity of the 
assay, }IC_{50} \text{ values for cocaine (present in the assay tubes at concentrations of 1 nM–0.1} 
mM) \text{ were also evaluated. Consistent with our previous report (Hadlock et al, 2011), cocaine} 
inhibited \[^{3}\text{H}]\text{dopamine and }[^{3}\text{H}]\text{serotonin uptake with }IC_{50} \text{ values of 1212 nM and 922 nM,}

respectively.  

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DISCUSSION

The known neurobiological and behavioral effects of iptakalim suggest that it may be an effective pharmacotherapy to aid in smoking cessation. A goal of the present research was to test this notion by examining the effect of iptakalim on the stimulus and on the reinforcing effects of nicotine in two preclinical rodent models: a) drug discriminated goal-tracking and b) IV nicotine self-administration. We found that iptakalim, in a dose-dependent manner, attenuated conditioned responding controlled by the nicotine stimulus and attenuated nicotine intake by selectively decreasing active lever pressing.

Previous research from our laboratory has implicated the DAT and SERT in the stimulus effects of nicotine using the DGT task. For example, bupropion, a dopamine transporter inhibitor, fully substitutes for the stimulus effects of nicotine. Conversely, the SERT inhibitor, citalopram, blocks the conditioned response controlled by the nicotine stimulus (Dion et al., 2012). However, since iptakalim did not appreciably inhibit SERT or DAT function, another mechanism(s) likely underlies its effects on the stimulus and reinforcing effects of nicotine.

Very little data are available concerning the affinity of iptakalim on various receptor subtypes. In fact, one novel finding from the current study is that iptakalim does not inhibit dopamine or serotonin transporter function. Instead, antagonism of α4β2 receptors likely contributes to the efficacy of iptakalim demonstrated in Figures 1 and 2. In fact, heteromeric nAChRs are established contributors to the stimulus effects of nicotine. Evidence of the importance of these receptors includes the finding that agonists like ABT-418, ABT-594, A85380, TC2559, epibatidine, nornicotine, and 5-IA that bind to α4β2-containing nAChRs fully substitute for the nicotine stimulus (Brioni et al., 1995; Cohen et al., 2003; Reichel et al., 2010; Smith et al., 2007). Partial α4β2 agonists like cytisine and varenicline substitute for nicotine stimulus either partially or fully depending on the study (Chandler and Stolerman, 1997; Jutkiewicz et al., 2011; LeSage et al., 2009; Reichel et al., 2010; Smith et al., 2007). The α4β2 antagonist DHβE fully blocks the stimulus effects of nicotine (Struthers et al., 2009; Zaniewska et al., 2006). In addition, α4β2-containing nAChRs appear necessary for nicotine self-administration (Picciotto et al., 1998). Further, selective activation of α4β2-containing nAChRs is sufficient to establish nicotine place preference demonstrating critical involvement of these receptors in the reinforcing/rewarding effects of nicotine (Tapper et al., 2004). Finally, iptakalim inhibits the function of dopaminergic neurons dissociated from substantia nigra by non-competitively antagonizing α4-containing (α4β2 and α4β4) nAChRs localized on these neurons (Hu et al., 2006).

The brief overview in the previous paragraph suggests a likely role for α4-containing nAChRs in effects of iptakalim in the present experiments. It is worth noting, however, that iptakalim’s actions on KATP channels may also contribute to blockade of the reinforcing and stimulus effect of nicotine. For example, the KATP channels are located in brain areas implicated in both reward and learning processes – ventral tegmental area, substantia nigra, the prefrontal cortex, and hippocampus (Ross et al., 2006). KATP channels located in these areas play an important role in regulating glutamatergic, dopaminergic, and GABAergic neurotransmission (Ross et al., 2006), all of which play a role in the behavioral and neural effects of nicotine (Dwoskin et al., 2009; Wooters et al., 2009).

In the discriminated goal-tracking and the self-administration task, iptakalim reduced chamber activity. At first, these finding might suggests that its antagonism of the stimulus and reinforcing effects of nicotine are the by-product of non-specific motor effects by iptakalim. There are several finding that make this account less tenable. First, an effect on activity was not observed after iptakalim per se, but only when iptakalim was administered...
along with nicotine at the highest test dose (60 mg/kg) in the DGT task. In fact, the stimulus effects of nicotine in the DGT task were also blocked by iptakalim at 30 mg/kg; a dose that did not significantly alter activity. Second, the ANCOVAs that used chamber activity as a covariate revealed that the reduction in locomotion did not significantly contribute to the decrease in nicotine-evoked goal-tracking or the reduction in nicotine-maintained active lever pressing. Finally, in other published experiments, iptakalim increases locomotor activity when administered alone (10 mg/kg IP; Schmidt et al., 2012) or decreases locomotion when administered with stimulants like nicotine or amphetamine (Sun et al., 2010; Volf et al., 2012). We did not observe this increase in activity when iptakalim was given alone (recall Figure 1D). Perhaps the brief 4-min test sessions used here were not sufficiently long to detect this enhanced activity (cf. 60 min in Schmidt et al., 2012). Of course, there are many other potential methodological differences that could account for this discrepancy (e.g., size of apparatus, how activity was indexed, experimental history, rat strain, etc.). Regardless, iptakalim attenuated the stimulus and positive reinforcing effects of nicotine with motor impairment seeming an unlikely account of this blockade.

Nicotine is the primary addictive component of tobacco and purportedly its reinforcing effects are important to the addiction process (Balfour, 2002; Pomerleau and Pomerleau, 1992). However, recent research has highlighted the relative import of its reward or incentive-enhancing effects. For example, low levels of responding can be maintained in rats by a change in visual stimuli as the consequence for completing a schedule of reinforcement. Responding maintained by this visual stimulus is increased significantly when nicotine is administered by the experimenter before the session or by the rat during the session (Caggiula et al., 2001; Chaudhri et al., 2006, 2007; Palmatier et al., 2007). In our self-administration protocol, we used cued timeouts (see Methods) following each nicotine infusion. One might justifiably speculate that the nicotine self-administration seen in the present study reflects the primary reinforcing effects of nicotine plus its reward-enhancing effects on the visual stimulus that signaled the infusion and subsequent timeout. If so, then the decrease in nicotine intake seen with iptakalim may reflect blockade of either of these effects of nicotine, or both.

Although we did not design the present experiments to distinguish between the import of the primary reinforcing effects of nicotine or its reward-enhancing effects on the visual stimulus, the potential implication of the research in this report remains the same. The demonstrated behavioral effects here on the nicotine-evoked goal-tracking and nicotine intake suggest that it should be further investigated as a pharmacotherapy for smoking cessation. The known neural effects of iptakalim on nicotine-induced dopamine release and binding to α4β2-containing nAChRs are consistent with this suggestion. Given the limited research on iptakalim within the context of nicotine dependence and smoking cessation, much more research is needed. For example, will iptakalim blunt reinstatement (relapse) of self-administration behavior triggered by an environmental stressor, drug-associated cues, or priming dose of nicotine? Will iptakalim have an impact on nicotine withdrawal or its associated cognitive deficits? Does the action of iptakalim at ATP-sensitive potassium channels contribute in a unique manner to its potential utility as a smoking cessation aid? Although the present research eliminated inhibition of DAT or SERT function as a potential mechanism, what other central nervous system effects of iptakalim contribute to its potential efficacy and are these localized to particular brain regions? Addressing these questions, and others, in future research will no doubt provide insight into iptakalim as a potential medication for aiding in smoking cessation, as well as suggest potential new targets for medication development.
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References


- Iptakalim blocks responding evoked by the stimulus effects of nicotine
- IV nicotine self-administration is blunted by pretreatment with iptakalim
- Motor impairment cannot account for these effects
- Iptakalim’s behavioral effects not due to SERT and DAT inhibition
Figure 1.
Data representing discriminated goal-tracking tests. Upper panels show mean (±SEM) dipper entries during iptakalim/nicotine interaction tests (A) and iptakalim alone tests (C). Lower panels show mean (±SEM) number of chamber crosses during iptakalim/nicotine interaction tests (B) and iptakalim alone tests (D). Dashed line represents saline control (0 mg/kg iptakalim + saline) activity baseline with dot lines delineating the SEM boundaries. *Significantly different from †, and † significantly different from b (p<0.05).
Figure 2.
Panel A shows mean (±SEM) lever presses during nicotine self-administration phase (* significantly different from inactive lever; inset graph shows total infusions earned during the session). Panel B shows mean (±SEM) lever presses during iptakalim dose-response tests (* significantly different from 0 mg/kg active lever responding). Panel C shows mean (±SEM) chamber crosses during the iptakalim dose-response tests (* significantly different from 0 mg/kg; p<0.05).
Figure 3.