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Effect of Brain CYP2B Inhibition on Brain Nicotine Levels and Nicotine Self-Administration

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The CYP2B enzyme is expressed in human and rat brain, and metabolizes many CNS-acting drugs. The gene that encodes human CYP2B6 is highly polymorphic, where the variation in brain enzyme levels could result in altered brain drug levels. CYP2B can metabolize nicotine, the main psychoactive ingredient in cigarettes; if altered brain CYP2B activity can influence nicotine brain levels, it could influence nicotine-mediated behaviors. To investigate this, a mechanism-based inhibitor selective for CYP2B, C8-xanthate (20 μ g), was administered intracerebroventricularly (ICV) into the brain of rats, and 22 h later, nicotine levels were measured by *in vivo* microdialysis following nicotine (150 μ g/kg intravenous). Brain nicotine levels from 15 to 30 min and the AUC_{0-45min} were both twofold higher (*p* < 0.05) with C8-xanthate vs vehicle pretreatment; there was no difference in peripheral nicotine levels. Rats were then given ICV pretreatment with C8-xanthate/ASCF and underwent intravenous nicotine self-administration with 3.75–30 μ g/kg per infusion dose. C8-xanthate increased responding in progressive ratio (15 μ g/kg per infusion dose, *p* < 0.05). In a separate cohort, C8-xanthate increased the percentage of rats that acquired self-administration (7.5 μ g/kg per infusion dose, *p* < 0.05) from 40% after vehicle pretreatment to 100%, with no difference in peripheral nicotine levels measured at the end of behavior. In a third cohort, C8-xanthate increased the number of sessions required to meet extinction criteria (*p* < 0.05). Together these data demonstrate that the brain CYP2B activity can influence nicotine brain cYP2B brain levels could have different nicotine levels and reinforcement, which might have a role in smoking behaviors and dependence.

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INTRODUCTION

There is increasing evidence to support that extra-hepatic expression of cytochrome P450 (CYP) enzymes, which are involved in the metabolism of numerous clinical drugs, can influence local drug levels and response (Ferguson and Tyndale, 2011). The CYP isoform CYP2B is expressed in various regions of the brain in humans (Gervot et al, 1999; Miksys et al, 2003), monkeys (Lee et al, 2006), and rats (Miksys et al, 2000). Brain CYP2B is active and is able to metabolize its substrates in situ (Miksys and Tyndale, 2009), and many CYP2B substrates act within the central nervous system (CNS), such as bupropion (Hesse et al, 2000), methadone (Gerber et al, 2004), propofol (Ekins et al, 2008), and nicotine (Yamanaka et al, 2005; Yamazaki et al, 1999). Therefore, CYP2B in the brain may contribute to local metabolism of its CNS-acting substrates and to resulting drug response.

CYP2B expression is influenced by both genetic and environmental factors. The human CYP2B6 gene is highly polymorphic (Zanger et al, 2007), and genetic variation in the liver expression of this enzyme can influence the systemic levels of its substrates (Crettol et al, 2005; Dobrinas et al, 2013; Kirchheiner et al, 2003; Tsuchiya et al, 2004). One variant of CYP2B6, CYP2B6*6, results in a reduction of CYP2B protein levels (Lang et al, 2001); this protein reduction is also found in the brains of individuals with a CYP2B6*6 allele (Miksys et al, 2003). CYP2B expression can also be altered by environmental factors, for example, phenobarbital can induce CYP2B expression in both the liver and brain of rats (Anandatheerthavarada et al, 1992; Schilter and Omiecinski, 1993) and monkeys (Lee et al, 2006). Thus, variation in brain CYP2B activity may alter the levels of a substrate locally, possibly altering the resulting drug response. We have previously shown that the selective inhibition of CYP2B activity in the brain by central administration of the mechanism-based inhibitor C8xanthate (C8X), which is selective for CYP2B (Yanev et al, 1999), can modify the brain levels and subsequent behavioral response of the CYP2B substrates propofol (Khokhar and Tyndale, 2011) and chlorpyrifos (Khokhar and Tyndale, 2012, 2014). These studies first established that altering the

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brain CYP2B activity can affect the pharmacological and the toxicological response to CNS-acting compounds.

Nicotine, the main psychoactive ingredient in cigarettes, can be metabolized by CYP2B in humans (Yamanaka *et al*, 2005) and in rats (Hammond *et al*, 1991). Nicotine enters the brain from the periphery where it is thought to exert its reinforcing effect by acting on the brain reward center nicotinic acetylcholine receptors that release dopamine (Di Chiara and Imperato, 1988; Marshall *et al*, 1997; Rapier *et al*, 1988). *In vitro* rat brain membranes are capable of metabolizing nicotine to cotinine (Jacob *et al*, 1997); however, it is unknown whether altering the brain CYP2B activity can significantly influence nicotine brain levels *in vivo* following peripheral administration. To address this question, nicotine brain levels produced by the IV administration of nicotine were measured via *in vivo* microdialysis following intracerebral ventricular (ICV) injection of C8X.

If reducing brain CYP2B activity can increase nicotine brain levels, is this increase sufficient to alter the behavioral response to nicotine? People with the CYP2B6*6 variant are more likely to become smokers (Hoffmann et al, 2006) and to relapse during cessation compared with normal metabolizers (Lee et al, 2007a; Lerman et al, 2007); however, this variant has no effect on peripheral plasma nicotine levels (Lee et al, 2007b), suggesting that the observed differences in acquisition and cessation may be due to genetic variation in brain CYP2B6 levels and resulting higher brain nicotine levels. To examine whether inhibiting brain CYP2B could alter nicotine self-administration (NSA) behavior (Corrigall and Coen, 1989; Donny et al, 1999), rats were given C8X ICV before or during NSA; if inhibition of brain CYP2B increases nicotine levels in brain, then a left shift in the dose-response curve would be anticipated. The effect of brain CYP2B inhibition was first tested on the motivation to obtain nicotine through the progressive-ratio (PR) schedule. On the basis of these findings, we examined the impact of brain CYP2B inhibition on acquisition of NSA, extinction, and reinstatement of nicotine-seeking behavior. In both microdialysis and behavioral studies, peripheral nicotine levels were assessed to confirm that central injections of CYP2B inhibitors had no effect on peripheral nicotine or cotinine levels.

MATERIALS AND METHODS

Adult male Wistar rats (250-300 g, Charles River Laboratories, Quebec, Canada) were housed singly on a 12-h light/ dark cycle, and water was provided ad libitum. Stainless-steel guide cannulae were implanted into the right lateral ventricle for ICV injection and for the insertion of microdialysis probes (anterior-posterior 0.9 mm, lateral 1.4 mm, and dorsoventral 2.6 mm (microdialysis) or 3.6 mm (ICV injection), from Bregma (Paxinos and Watson, 1986)). For brain CYP2B inhibition, C8-xanthate (C8X, Toronto Research Chemicals, Toronto, ON, Canada), a CYP2B-specific inhibitor (Yanev et al, 2000), was injected ICV, 20 µg in 4 µl of artificial cerebrospinal fluid (ACSF) over 1 min; the injector was removed after 1 min. This C8X dose was previously optimized for brain, but not liver, CYP2B inhibition (Khokhar and Tyndale, 2011, 2012, 2014) and retested here with nicotine as a substrate. Sterile nicotine solution

(Sigma-Aldrich, Oakville, ON, Canada) was prepared daily as concentration of base in saline, pH 6.8-7.2.

In Vivo Nicotine Microdialysis

Microdialysis experiments were carried out in the light phase. Twenty-two hours before microdialysis, rats were given a single ICV injection of ACSF vehicle (n = 10) or C8X (n=9) (Figure 1a). To provide an estimate of the increase in nicotine dialysate levels due to brain CYP2B inhibition (ICV C8X) in terms of peripheral nicotine dosing, a dose-response curve for nicotine dialysate levels was generated with a separate cohort of animals (No ICV Tx, n=5 per nicotine dose). Ringer's solution (147 mM Na⁺, 2 mM Ca²⁺, 4 mM K⁺, 155 mM Cl⁻, pH 6.0) was perfused through concentric silicacoated probes with 2 mm membranes (MD-2200; Bioanalytical Systems, West Lafayette, IN) by an infusion pump at 2 µl/min. Baseline dialysate was collected for 15 min before IV nicotine injection (into the tail vein, 150 µg/kg for ACSF and C8X groups, and 150, 225, and 300 µg/kg for No ICV Tx groups in saline, pH 6.8-7.2), after which the dialysate was collected on ice in 15-min bins for 2 h and 15 min. Blood (200-300 µl) was collected from the saphenous vein at 15 and 75 min postinjection. For nicotine and cotinine quantification, 30 µl of dialysate was diluted to 100 µl with deuteriumlabeled internal nicotine and cotinine standards (20 ng/ml in 0.01 M HCl), while 100 µl plasma samples were prepared as previously described (Craig et al, 2014; Vieira-Brock et al, 2011); dialystate and extracted plasma were then analyzed by liquid chromatography-mass spectrometry (LC-MS) (Craig et al, 2014).

Nicotine Self-Administration

Seven days after ICV cannula implant, catheters were implanted into the right jugular vein as previously described (Garcia *et al*, 2014; Shram *et al*, 2008). To maintain patency, catheters were flushed daily with sterile heparin in saline (0.1 ml, 50 U/ml). Thiopental (2–4 mg, IV, 20 mg/ml) was used to test catheter patency after testing, and nonpatent animal data were removed from analysis. NSA was carried out in 16 operant chambers operated by a computer-controlled interface system (Med Associates, St Albans, VT) under previously described conditions (Garcia *et al*, 2014).

Progressive-Ratio Schedule

Before surgery, rats were food trained to learn the selfadministration operant behavior as previously described (Garcia *et al*, 2014). Following recovery from IV surgery, rats (n = 12 per dose) underwent daily 1-h NSA sessions under FR1 and FR2 for 5 days at a single nicotine dose of 7.5, 15, or 30 µg/kg (Figure 1b). Rats then underwent a PR schedule of reinforcement at their same training dose, in 2-h daily sessions for five sessions to create a baseline of responding, as previously described (Garcia *et al*, 2014). At 22 h before the sixth PR session, all animals received one ICV injection of ACSF, then underwent three PR sessions (sessions 6, 7, and 8). At 22 h before the ninth PR session, all animals received one ICV injection of C8X, then underwent three PR sessions (9, 10, and 11).

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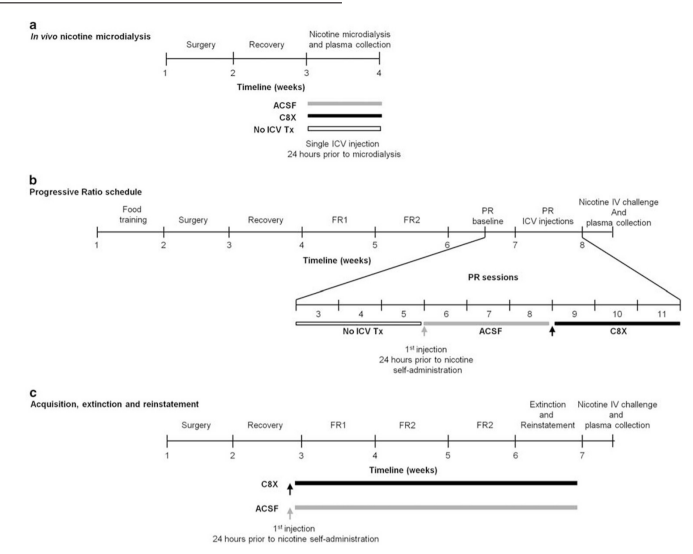


Figure I Experimental study design and timelines for (a) rats that underwent *in vivo* nicotine microdialysis, (b) rats that underwent the progressive-ratio (PR) schedule of reinforcement where they received a single ICV injection of ACSF and then C8X in a within-animal design, and (c) rats that underwent acquisition, maintenance, extinction, and nicotine reinstatement experiments under a fixed-ratio (FR) schedule of reinforcement where they received daily ICV injections of ACSF or C8X.

After PR testing, rats received one ICV injection of ACSF or C8X, 24 h before a bolus injection of nicotine ($200 \mu g/kg/$ IV). Blood was collected from the saphenous vein at 15 and 60 min post-nicotine injection, and trunk blood was collected at 120 min. Plasma was analyzed for nicotine and cotinine by high-performance liquid chromatography (Siu *et al*, 2006).

Acquisition, Extinction, and Reinstatement

Rats were assigned to receive either C8X or ACSF (n=12 per dose) ICV injections daily, 22 h before each behavior session. Seven days after IV surgery, animals underwent NSA without food training (Figure 1c) at three infusion doses (3.75, 7.5, or 15 µg/kg) that were found previously to fall along the ascending limb of nicotine's dose-response curve for spontaneous acquisition (Garcia *et al*, 2014). Rats underwent 2-h daily NSA sessions under a FR1 schedule for 5 days and then a FR2 schedule for another 5 days, which was used to determine acquisition (Garcia *et al*, 2014; Shram *et al*, 2008).

Another cohort of rats (n = 24 each for C8X and ACSFgroups) trained with 7.50 µg/kg nicotine underwent extinction and reinstatement. Following acquisition, where rats were trained to self-administer nicotine and given daily ICV injections as described above, rats continued NSA under the FR2 schedule for 3 extra days and then underwent four 2-h extinction sessions (one per day) during which responding on the active lever had no consequences. The effects of CYP2B inhibition on reinstatement of nicotine seeking were then evaluated. On reinstatement test day, rats were subjected to multiple 1-h extinction sessions until they met extinction criteria (<10 active lever presses in two consecutive sessions). They were then given saline IV (0.1 ml/kg) 5 min before one 1-h session. Following the saline session, reinstatement of nicotine seeking induced by different IV priming doses of nicotine (15, 30, and 60 µg/kg, 0.1 ml/kg) was evaluated in ascending order over 3 different test days. After 1 week, this extinction and reinstatement cycle was repeated with SC saline and nicotine (15, 30, 60, and 150 µg/kg).

Twenty-four hours after the last ICV injection of C8X or ACSF, rats received four nicotine infusions (7.50 μ g/kg, one per min) in the operant chambers. Trunk blood was collected either 1 or 5 min after the last (fourth) infusion and analyzed by LC–MS for nicotine and cotinine as described above.

Statistical Analysis

Nicotine dialysate levels and AUCs after C8X and ACSF treatment were analyzed by Student's *t*-test. Plasma nicotine dose response was analyzed using a one-way ANOVA. The log-transformation of final completed ratios in PR, and active and inactive lever presses and reinforcements in FR were analyzed with repeated-measures ANOVA. Post-hoc analyses for all ANOVAs were conducted using Bonferroni tests. NSA acquisition data were analyzed using a Fisher's Exact test (two sided). Extinction sessions were analyzed using Kaplan–Meier survival analysis followed by a log-rank test. Significance level for all comparisons was p = 0.05.

RESULTS

ICV Injection of C8X Increases Circulating Nicotine Levels in the Rat Brain

Brain levels of nicotine in dialysate collected at 15-min intervals following IV infusion of 150 µg/kg of nicotine in C8X- and ACSF-treated rats are shown in Figure 2a. Mean nicotine dialysate levels were higher with C8X treatment over the first 45 min compared with ACSF treatment, resulting in significant differences at the time of maximum nicotine; t(17) = -2.27, p = 0.03) (Figure 2a) and for AUC₀₋₄₅ (t(17) = 2.23, p = 0.04) (Figure 2b). Mean AUC₄₅₋₁₃₅ was not different (p = 0.79) between ICV treatments (Figure 2b). Plasma levels of nicotine and its major metabolite cotinine, obtained during microdialysis, were not significantly different between C8X and ACSF, confirming that ICV injection of inhibitor did not alter peripheral nicotine metabolism (Figure 2c).

Nicotine dialysate levels increased with increasing nicotine dose, demonstrating that the microdialysis method was capable of detecting higher levels of nicotine (Figure 2d, shown for 15-30 min post-nicotine) consistent with the increasing plasma levels of nicotine observed (Figure 2e, shown for 15 and 75 min post-nicotine). Nicotine dialysate levels after ICV ACSF treatment were similar to nicotine levels without ICV pretreatment (No ICV Tx), as expected, whereas nicotine levels after C8X treatment were higher than both control groups (tested at 150 µg/kg) corresponding to a 25% increase in the apparent dose of nicotine. Plasma nicotine levels in animals pretreated ICV with ACSF and C8X were similar to the levels measured in No ICV animals (150 µg/kg, Figure 2e). Together, this indicates that ICV injections of C8X can increase nicotine levels in the brain, while leaving the systemic levels unchanged.

ICV Injection of C8X Increases Final Ratios Completed under a PR Schedule

To test whether single ICV C8X injections could alter NSA behavior, the effect of C8X on PR breakpoint (final ratio of

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responding completed) was examined. Rats that acquired NSA and underwent five PR baseline sessions were then treated with ACSF ICV and tested under PR. There was no difference between final ratios completed during baseline PR compared with ACSF treatment indicating little to no effect of ICV injection, or time, on the measure of PR (data not shown). All rats were then given C8X ICV and tested under PR. Mean final ratios completed for rats after C8X treatment were higher than ACSF at all infusion doses (Figure 3a) and repeated measures ANOVA within each dose group revealed that this was significantly different at 15 µg/kg (F(1,7) = 7.365, p = 0.03).

There was no difference between C8X and ACSF ICV treatment in the plasma nicotine and cotinine collected after an IV bolus injection of nicotine (200 μ g/kg) (Figure 3b and c). This indicated that central injections of C8X were not inhibiting the hepatic CYP2B and the peripheral nicotine metabolism.

ICV Injection of C8X Shifts Nicotine's Dose-Response Curve for Acquisition of NSA

Training-naive rats given daily pretreatment ACSF ICV injections showed an ascending dose-response curve for the proportion of animals that acquired NSA between 3.75 and $15 \,\mu$ g/kg infusion doses (Figure 4). Around 40% of the rats that self-administered 3.75 and 7.5 µg/kg of nicotine acquired NSA, whereas almost all rats (86%) that self-administered 15 µg/kg met acquisition criteria. Rats given C8X ICV injections also showed dose-dependent acquisition under the same infusion doses where ~20% of rats acquired NSA at the lowest dose tested; however, at 7.5 and 15 µg/kg, essentially all animals acquired NSA (Figure 4) resulting in a significant difference between ICV treatments at the 7.5 µg/ kg acquisition dose (p < 0.001, Fisher's exact). This apparent leftward shift in nicotine's dose-response acquisition curve with C8X ICV is consistent with the higher levels of nicotine observed with C8X ICV during microdialysis (Figure 2a).

There was no difference in active and inactive lever presses by session during FR1 and FR2 between C8X- and ACSFtreated rats trained at 7.5 µg/kg. Mean active lever \pm SEM was 40 ± 15 (ACSF) vs 83 ± 23 (C8X) in FR1 and 101 ± 24 (ACSF) vs 122 ± 24 (C8X) in FR2, and mean inactive lever \pm SEM was 11 ± 8 (ACSF) vs 25 ± 6 (C8X) in FR1 and 33 ± 17 (ACSF) vs 41 ± 10 (C8X) in FR2. In ACSF-treated rats that did not meet acquisition criteria, there was no difference in active vs inactive lever presses (FR1: 14 ± 6 vs 10 ± 6 , FR2: 8 ± 2 vs 9 ± 6) and reinforcements dropped from FR1 to FR2 (14 ± 7 to 9 ± 6); all C8X-treated rats met acquisition criteria.

ICV Injection of C8X Increased the Number of Sessions to Extinction but Did Not Influence Reinstatement of Nicotine-Seeking Behavior

There was no difference in active lever presses during the four extinction sessions between ACSF- and C8X-treated rats (Supplementary Figure). As animals underwent multiple extinction sessions until they met extinction criteria, the number of sessions that animals needed before they met extinction criteria were examined, where C8X-treated animals, on average, required more sessions to meet the criteria compared with ACSF animals (Figure 5a, p = 0.03).

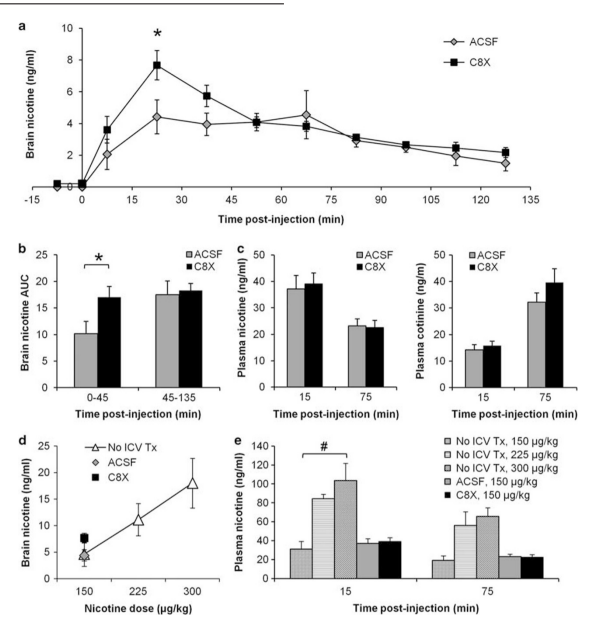


Figure 2 ICV injection of C8X increases circulating nicotine levels in the lateral ventricles measured by *in vivo* microdialysis. (a) Nicotine dialysate levels were higher after C8X treatment (n=9) compared with ACSF (n=10) 15–30 min postinjection (150 µg/kg/IV, which was the lowest dose of nicotine that produced brain dialysate concentrations of nicotine above the limit of detection for our LC–MS assay (1 ng/ml)). (b) Nicotine dialysate AUCs were higher after C8X treatment compared with ACSF from 0 to 45 min. (c) Nicotine and cotinine plasma levels obtained during microdialysis were not different between C8X and ACSF treatment. (d) Nicotine dose–response curve for dialysate collected 15–30 min post-nicotine in No ICV Tx animals (n=5 per dose). Mean nicotine dialysate collected during microdialysis in No ICV Tx animals. Plasma levels increased dose-dependently at 15 min post-nicotine (F(2, 12) = 10.41, p < 0.001). Mean plasma nicotine levels for C8X- and ACSF-treated animals are shown for comparison. Data shown as mean \pm SEM, *p < 0.05, C8X vs ACSF, #p < 0.05, between doses within No ICV Tx.

After extinction, rats were given a single saline injection (IV or SC), and they underwent one session to record responding after an injection of the same volume. There was no difference between active lever presses during extinction and after saline injection (data not shown). After the saline session, animals were tested for reinstatement with increasing IV nicotine priming doses (15, 30, and $60 \,\mu g/kg$). C8X- and ACSF-treated rats did not reinstate their nicotine-seeking behavior at any nicotine priming dose

tested: active lever presses after a saline injection *vs* all nicotine priming doses were not significantly different for both groups (Figure 5b). When active lever presses between C8X and ACSF groups were compared, there was no significant difference at any nicotine priming dose tested. After reinstatement, animals returned to extinction and were tested with SC nicotine priming doses (15, 30, 60, and 150 µg/kg). Within treatment, there was a significant effect of dose on active lever presses for C8X (F(4, 88) = 22.403,

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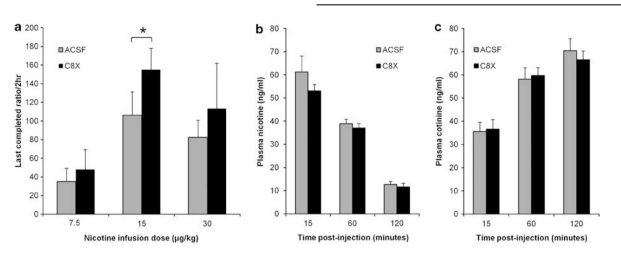


Figure 3 ICV injection of C8X can increase motivation to obtain nicotine in the PR without altering peripheral nicotine metabolism. (a) C8X increased final ratios completed during PR compared with ACSF for animals trained at 7.5 μ g/kg (n = 8), 15 μ g/kg (n = 8), and 30 μ g/kg (n = 7). Mean final ratio completed was calculated using the mean log transformed final ratio from the three sessions following ICV injection of ACSF or C8X, *p < 0.05, C8X vs ACSF. Plasma (b) nicotine and (c) cotinine levels were not different between ACSF- (n = 11) and C8X (n = 12)-treated animals following a bolus IV injection of nicotine (200 μ g/kg) at the end of NSA. Data shown as mean+SEM.

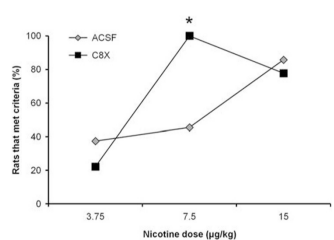


Figure 4 ICV injection of C8X results in a leftward shift of the acquisition dose response for nicotine, but does not alter NSA after behavior is acquired. In rats that self-administered 3.75 µg/kg, 3 out of 8 acquired with ACSF vs 2 out of 9 acquired with C8X; for the 7.5 µg/kg dose, 10 out of 22 acquired with ACSF vs 21 out of 21 acquired with C8X; and for the 15 µg/kg dose, 6 out of 7 acquired with ACSF vs 7 out of 9 with C8X. Data for 7.5 µg/kg dose was combined from two animal cohorts that produced the same results. In the first cohort, 4 out of 10 animals acquired with ACSF vs 10 out of 12 animals acquired ACSF vs 10 out of 10 acquired with C8X. C8X was significantly higher when data was analyzed separately (p<0.05 for each cohort) or together. *p<0.001, C8X vs ACSF.

p < 0.001) and ACSF (F(2.735, 51.985) = 15.669, p < 0.001), where the rats reinstated their nicotine-seeking behavior after injection of the 60 (C8X, p < 0.001; ACSF, p = 0.003) and 150 µg/kg (C8X, p < 0.001; ACSF, p < 0.001) doses, but active lever presses were not different between ICV treatments (Figure 5c).

Plasma nicotine and cotinine levels were not different between ICV treatments after four $7.5 \,\mu$ g/kg infusions of nicotine, tested at the end of NSA (Figure 5d and e). These results confirm that repeated administration of the inhibitor ICV did not alter peripheral metabolism of nicotine.

DISCUSSION

This present work demonstrates that (1) a dose-related increase in nicotine brain levels can be measured by microdialysis after IV nicotine administration, (2) within this dose range, the CYP2B inhibitor C8X given ICV increased nicotine brain levels without altering peripheral levels of nicotine, and (3) C8X given ICV increased acquisition and motivation during NSA. The increases in nicotine concentration measured after C8X treatment provides the first evidence to support the idea that altering brain CYP2B activity can influence nicotine levels in the brain. As nicotine brain levels increase with increasing systemic dose, it suggests that the higher brain nicotine levels after CYP2B inhibition are akin to being given a higher dose of nicotine. Human brain CYP2B levels are highly variable (Miksys et al, 2003) suggesting that individuals who are CYP2B6 slow metabolizers, or who are taking clinically used drugs that can inhibit this enzyme, such as the antiplatelet agent clopidogrel, the antifungal clomitrozole, and the antidepressant sertraline (Richter et al, 2004; Walsky et al, 2006), could have reduced metabolism of nicotine in the brain.

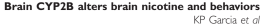
Injecting the CYP2B inhibitor directly in the brain modified the reinforcing effects of nicotine. C8X-treated rats were willing to press more on the active lever to receive one nicotine infusion during PR, and C8X treatment also increased acquisition of NSA, suggesting a leftward shift in the dose-acquisition curve for nicotine. This increase in motivation and in acquisition suggests that inhibiting brain CYP2B activity can increase the reinforcing effect of nicotine in NSA, possibly by increasing the nicotine levels in the brain. The effect of CYP2B inhibition on acquisition of NSA was found at a low infusion dose of 7.5 µg where 100% of the rats treated with C8X acquired NSA, but only 50% of those treated with ASCF acquired NSA. This effect was observed in two separate experiments. Higher nicotine doses result in greater dopamine release in the nucleus accumbens (Di Chiara and Imperato, 1988; Marshall et al, 1997), which is a characteristic response to systemic administration of drugs of abuse (Di Chiara and Imperato, 1988), and is considered

KP Garcia et al a 120 50 b p = 0.03■ACSF 100 40 ■C8X 0 Met extinction criteria (%) Active lever presses/1 hr 80 30 60 20 40 ♦ ACSF C8X 10 20 0 0 0 15 30 60 0 3 5 7 8 1 2 4 6 IV Nicotine priming dose (µg/kg) Extinction sessions С е d 50 30 30 * ACSF ACSF Active lever presses/1 hr 40 Plasma nicotine (ng/ml) ■C8X Plasma cotinine (ng/ml) ■C8X 20 20 ACSF 30 C8X 20 10 10 10 0 0 0 0 15 30 60 150 5 1 5 1 Time post-infusion (min) Time post-infusion (min) SC Nicotine priming dose (µg/kg)

Figure 5 ICV injection of C8X can increase motivation to obtain nicotine in nicotine-seeking behavior, and repeated ICV injections of C8X does not alter peripheral nicotine metabolism. (a) Mean active lever presses (\pm SEM) for the first four daily extinction sessions (C8X: 40 \pm 6, 28 \pm 4, 21 \pm 2, and 23 \pm 3; ACSF: 39 \pm 4, 26 \pm 3, and 25 \pm 2) were not different between C8X (n=24) and ACSF (n=21) groups (Supplementary Figure). Animals then underwent multiple extinction sessions the same day until they reached extinction criteria. Data shown is the proportion of C8X and ACSF rats that met extinction criteria over these extinction sessions where C8X-treated animals needed more sessions to meet criteria. During nicotine reinstatement testing, mean active lever presses between C8X and ACSF were not different after (b) IV (C8X: n=17, ACSF: n=18) or (c) SC (C8X: n=24, ACSF: n=21) nicotine priming injections at all doses tested. Mean active lever presses between C8X and ACSF after 60 µg/kg/IV approached significance (p=0.06). *p<0.05, saline (0 µg/kg nicotine) vs 60 and 150 µg/kg/SC for both C8X and ACSF. There was no difference in plasma (d) nicotine and (e) cotinine between ACSF and C8X treatments after four nicotine infusions were given in the operant SA chamber (7.5 µg/kg/infusion) at the end of NSA in a subset of rats (n=8 for both C8X and ACSF). Data shown as mean \pm SEM.

as one mechanism of creating a drug-reward association that leads to addiction. CYP2B is expressed in multiple brain regions (Miksys et al, 2003) including the nucleus accumbens, and also in other regions associated with nicotine addiction, such as the frontal cortex (Markou, 2008) and cingulate gyrus (McClernon et al, 2007). As our C8X injections were given ICV to globally inhibit CYP2B, it is possible that inhibition in any of these regions or nearby regions that express CYP2B contributed to the microdialysis and behavioral findings. Pilot data in humans suggest that CYP2B6 slow metabolizers show a faster conversion to dependence in adolescence (Hoffmann et al, 2006), which is consistent with potentially higher brain nicotine levels during initial stages of smoking. Higher brain nicotine levels when smoking behavior is first established could result in a greater reinforcing effect of nicotine.

C8X-treated animals also needed more sessions to meet extinction criteria, demonstrating that the animals were more persistent in responding to the active lever. However, there was no effect of C8X on nicotine reinstatement; priming with IV in doses ranging from 15 to 60 µg/kg did not reinstate nicotine seeking in either treatment groups albeit at the 60 µg/kg dose, a trend toward reinstatement of nicotine seeking was observed in the C8X group (p=0.06). It is possible that the training dose $(7.5 \,\mu\text{g/kg})$ was not sufficient to elicit a significant effect of C8X on reinstatement. This is, however, quite unlikely as significant reinstatement of nicotine seeking was observed with SC priming of 0.15 mg/kg of nicotine in both experimental groups. It should be noted that priming with nicotine through the IV route can produce a relatively modest reinstatement of nicotine seeking relative to that induced by SC priming (Chiamulera et al, 1996;



Shaham *et al*, 1997). *CYP2B6* slow metabolizers report greater craving scores during abstinence and are more likely to relapse (Lee *et al*, 2007a; Lerman *et al*, 2007), therefore it is possible that higher brain nicotine levels during smoking resulted in greater reinforcement that may result in greater behavioral response to nicotine withdrawal during abstinence in these individuals.

It is possible that the effects of C8X on NSA could be unrelated to CYP2B-mediated nicotine metabolism. However, brain, but not plasma, nicotine levels were elevated during this treatment. In addition, greater acquisition was not seen at higher and lower nicotine doses, suggesting that C8X did not increase acquisition of NSA behavior in general, but instead resulted in a leftward shift in nicotine's doseresponse curve. Also, active and inactive lever presses were not different after acquisition, suggesting that C8X itself did not increase responding for both levers once rats established NSA. Another possibility is that C8X increased responding for the cues; however, more extinction sessions were required to extinguish NSA behavior without the presence of cues, suggesting that C8X-treated animals still persisted in pressing on the active lever even without cues. Also, it is possible that C8X's effects on brain nicotine levels and NSA behavior could be indirect; however, although other alkyl xanthate compounds have antitumor and antiviral properties (Sauer et al, 1984; Schick et al, 1989), there are no known biological effects of C8X other than its ability to inhibit CYP2B (Yanev et al, 1999, 2000). As C8X has been previously established as a specific CYP2B inhibitor (Khokhar and Tyndale, 2011, 2012, 2014; Miksys and Tyndale, 2009; Yanev et al, 2000), these observations, along with the increase in nicotine levels measured via microdialysis after C8X treatment, support the idea that C8X's effects are due to CYP2B inhibition and reduced nicotine metabolism.

Our findings demonstrate that brain CYP2B can influence nicotine levels independent of liver CYP2B and that reducing brain CYP2B activity can increase brain nicotine levels and resulting nicotine-mediated behaviors. The effect of reducing brain CYP2B activity in this rat model is consistent with observed smoking-related behaviors in human CYP2B6 slow metabolizers, which supports the idea that reduced brain CYP2B6 activity might influence smoking behaviors. Together with data using other CYP2B substrates (Khokhar and Tyndale, 2011, 2012, 2014,), this data suggest that brain CYP2B can influence the drug levels in the brain and their subsequent behavioral effects. These findings also indicate clinical implications for individuals with CYP2B6 genetic variation and those prescribed CYP2B6 inhibitors or inducers, because variation in brain-specific enzyme activity may alter drug response. As many CNS-acting, clinical drugs are substrates and/or inhibitors for CYP2B, brain-specific activity of CYP2B could be another source of variation in the inter-individual response to these drugs.

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