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## Contributions of prolonged contingent and non-contingent cocaine exposure to escalation of cocaine intake and glutamatergic gene expression

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### Abstract

Similar to the pattern observed in people with substance abuse disorders, laboratory animals will exhibit escalation of cocaine intake when the drug is available over prolonged periods of time. Here, we investigated the contribution of behavioral contingency of cocaine administration on escalation of cocaine intake and gene expression in the dorsal medial prefrontal cortex (dmPFC) in adult male rats. Rats were allowed to self-administer intravenous cocaine (0.25 mg/infusion) under either limited cocaine- (1h/day), prolonged cocaine- (6h/day), or limited cocaine- (1h/day) plus yoked cocaine-access (5h/day); a control group received access to saline (1h/day). One day after the final self-administration session, the rats were euthanized and the dmPFC was removed for quantification of mRNA expression of critical glutamatergic signaling genes, *Homer2*, *Grin1*, and *Dlg4*, in the dmPFC, as these genes and brain region have been previously implicated in addiction, learning, and memory. All groups with cocaine-access showed escalated cocaine intake during the first 10 minutes of each daily session, and within the first 1h of cocaine administration. Additionally, the limited-access + yoked group exhibited more non-reinforced lever responses during self-administration sessions than the other groups tested. Lastly, *Homer2*, *Grin1*, and *Dlg4* mRNA were impacted by both duration and mode of cocaine exposure. Only prolonged-access rats exhibited increases in mRNA expression for *Homer2*, *Grin1*, and *Dlg4* mRNA. Taken together, these findings indicate that both contingent and non-contingent “excessive” cocaine exposure supports escalation behavior, but the behavioral contingency of cocaine-access has distinct effects on the patterning of operant responsiveness and changes in mRNA expression.

## Keywords

Cocaine; self-administration; escalation; contingent access; non-contingent access

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## Introduction

Cocaine addiction is a chronic disorder that persists in spite of negative interpersonal, professional, and physical consequences, with the development of tolerance and increased cocaine intake serving as central diagnostic criteria for stimulant use disorders (APA 2013). It has been argued that one major reason for the escalation of cocaine consumption relates to gradual changes in motivation and learning processes associated with responses to the positive and negative reinforcing properties of cocaine (Koob 2004). In order to model the differences in cocaine use observed in humans, differential access to cocaine self-administration has been employed as an avenue to study the behavioral and neurobiological aspects of drug-taking. Ahmed and Koob (1998) demonstrated that “limited” daily-access (1 h/day) and “prolonged” daily-access conditions (6 h/day) are distinct in their ability to model aspects of drug abuse/addiction in two major ways: 1) rats with prolonged cocaine-access escalate their cocaine consumption across daily sessions, whereas the limited-access rats exhibit stable consumption for several weeks and 2) rats with daily “prolonged” cocaine-access dramatically increased consumption of cocaine for the 1<sup>st</sup> 10 minutes of self-administration while “limited” access rats did not escalate cocaine consumption within the 1<sup>st</sup> 10 minutes of self-administration (Ahmed and Koob 1998). This differential pattern of responding has since been replicated with a number of variations to determine the underlying changes in drug-taking behavior (Ahmed and Koob 1999), neurocircuitry (Ahmed et al. 2002; Ben-Shahar et al. 2012; Robinson and Kolb 2004) and cellular/molecular function in rats (Ben-Shahar et al. 2009; Ben-Shahar et al. 2013), as well as in other species (Kirkland Henry et al. 2009; Nakamura et al. 2011).

Despite substantial study, it remains to be determined whether escalated cocaine intake (and the neurobiological changes associated there with) is dependent upon the amount of cocaine exposure or the act of self-administering the cocaine. Indeed, escalated responding and intake of non-drug reinforcers can be achieved by prolonged-access to liquid food, suggesting that the escalation phenomenon may be mediated by behavioral processes underlying the self-administration of appetitive stimuli (Goeders et al. 2009). One way to dissociate the relative contribution of total drug exposure from the behavioral contingency of that exposure is to employ yoked-access procedures (i.e. the administration of cocaine under the control of a separate self-administering rat). For instance, Hemby et al. (1997) investigated the effects of cocaine under response-dependent and –independent conditions in a yoked-triad. Both self-administering and yoked-access rats exhibited elevated levels of dopamine in the nucleus accumbens during the first hour of self-administration, but the self-administering animals exhibited greater dopamine levels than their yoked- counterparts (Hemby et al. 1997). Additionally, relative to rats self-administering cocaine, yoked rats exhibit differential corticosterone levels both in systemic plasma (Galici et al. 2000) and brain (Palamarchouk et al. 2009), have a higher morbidity rate (Dworkin et al. 1995) and exhibit higher indices of distress (measured by ultrasonic vocalization) (Mutschler and

Miczek 1998). These studies demonstrate that, despite equivalent cocaine dosing, animals given contingent-access to cocaine exhibit distinct behavioral and neurobiological effects compared to animals with non-contingent access.

To expand upon the role of behavioral contingency in the behavioral sequelae of cocaine exposure, Kippin et al. (2006) employed a novel mixed self-administration/yoked cocaine exposure procedure in which rats received 1-h access to cocaine self-administration before receiving non-contingent cocaine infusions via yoking procedures during its last 5 hours of cocaine self-administration. Although escalated cocaine intake was not observed in this earlier study, both the contingent and non-contingent excessive cocaine exposure groups exhibited greater cue-induced reinstatement than rats with a history of limited cocaine-access only. However, only the prolonged contingent-access rats exhibited greater cocaine-primed reinstatement of responding. In the present study, we employ the prolonged-access and limited-access + yoked procedures to determine the impact of contingent and non-contingent “excessive” cocaine exposure on the escalation of cocaine intake and operant responding for cocaine to determine how behavioral contingency of cocaine delivery influences these aspects of cocaine addiction-related behavior.

The dorsomedial prefrontal cortex (dmPFC) is dysregulated in human cocaine addicts (Verdejo-Garcia et al. 2015) and this dysregulation is linked to aberrant learning and plasticity that is attributed to anomalies in glutamate transmission (Kalivas et al. 2005; Pascoli et al. 2014a; Ruan and Yao 2017b). To explore the potential neurobiology underpinning the effects of cocaine on dmPFC as a function of contingency, we measured the levels of mRNA for *Homer2*, *Grin1*, and *Dlg4* within the dorsomedial prefrontal cortex (dmPFC). Homer2 is a glutamate receptor scaffolding protein that is up-regulated within PFC by both non-contingent and contingent cocaine administration (Ary and Szumlinski, 2007; Ary et al., 2013; Ben-Shahar et al., 2009; Gould et al., 2013) and while it remains to be determined whether or not cocaine-induced increases in PFC Homer2 protein expression reflects increased gene transcription, Homer2 expression bi-directionally regulates both basal and cocaine-induced changes in extracellular glutamate levels within PFC (Ary et al., 2013) to influence cocaine-conditioned approach behavior in place-conditioning models (Ary et al., 2013) and cocaine-primed reinstatement of lever-pressing behavior in operant-conditioning models (Gould et al., 2013). Grin1 mRNA encodes the obligatory N-methyl-D-aspartate (NMDA) receptor sub-unit NR1 within the mammalian brain (Bai and Hoffman 2009); this receptor is widely implicated and critical for many forms of plasticity (Bear 1996; Hopf 2017; Sweatt 2016; Thiels et al. 1996), and is up-regulated in the PFC following cocaine exposure (Ary and Szumlinski 2007; Blanco et al. 2014; Hemby et al. 2005). Dlg4 encodes the sequence for postsynaptic density-95 (PSD-95), a scaffold receptor protein that regulates plasticity and learning through its actions on NMDA receptor density and function (Wang and Peng 2016). PSD-95 expression in the PFC is increased after prolonged withdrawal from cocaine (Ghasemzadeh et al. 2009; McIntosh et al. 2013) and following extinction testing during prolonged exposure to cocaine self-administration (Ghasemzadeh et al. 2011).

## Methods

### Subjects

Male Sprague-Dawley rats were pair-housed in a 12-h reverse light-dark cycle room and had *ad libitum* access to food and water (except as noted below). The housing and care of the rats followed the guidelines set forth by the “Guide for the Care and Use of Laboratory Rats, 8<sup>th</sup> Edition” (NIH 2011).

### Surgery

Male Sprague-Dawley rats weighing 300–350g were deeply anesthetized using ketamine (60mg/kg) and xylazine (10mg/kg). Chronic indwelling catheters were constructed using a bent steel cannula with a screw-type connector (Plastics One, Roanoke, VA), SILASTIC tubing (11 cm, i.d. 0.64 mm, o.d. 1.19 mm, Dow Corning, Midland, MI), Prolite polypropylene monofilament mesh (Atrium Medical Corporation, Hudson, NH), a silicon ball 2.5 cm from the end, and methyl methacrylate dental cement. The catheters were implanted and maintained as we have reported previously (Ben-Shahar et al. 2013; Kerstetter et al. 2008). Naïve rats were left in the vivarium and handled daily, but had no access to behavioral training or surgery. They were euthanized at the same age as the rats undergoing behavioral training.

### Behavioral Training

Food-training and cocaine self-administration utilized standard operant chambers (Med Associates Inc., St. Albans, VT, USA) and were conducted during a fixed time in the dark phase of the rat’s circadian cycle each day. Before surgical implantation of the jugular catheters, the rats were restricted to 20 g of food for 1 week, and trained on a fixed ratio 1 (FR1) schedule of food reinforcement for two 16 h training sessions where each right lever-press was associated with a 45 mg food pellet (Ben-Shahar et al. 2012). After recovery from the surgery, the rats were placed on a fixed ratio 1 (FR1) schedule of reinforcement for intravenous (IV) cocaine (0.1mL at 0.25 mg/infusion in 0.9% saline) or saline for 1h/day for 5 days. Each active lever-press was associated with a 4 sec infusion of cocaine or saline and a 20 sec timeout was signaled with a 20 sec light cue above the active lever (Ben-Shahar et al. 2012). On the 6<sup>th</sup> day, the cocaine rats were divided into limited (1 h/day) cocaine-access, prolonged (6 h/day) cocaine-access, and limited-access + yoked-access (1 h access followed by 5 h of “yoked” exposure) treatment groups and continued the FR1 schedule of reinforcement for an additional 15 days. During yoked exposure, rats remained in their chambers for an additional 5 h with the levers retracted to eliminate the opportunity to perform the operant response. During this time, yoked rats received a 4 sec cocaine infusion every time a paired prolonged-access rat self-administered an infusion, but without cue light presentation.

### Tissue Collection and mRNA Quantification

Twenty-four hours after the last self-administration session, the animals were sacrificed via rapid decapitation, and their brains were frozen over ice and dissected into 0.5 mm sections with a metal brain mold (Baintree Scientific, Braintree, MA). The dmPFC was dissected out

at 3.20 to 2.20 mm anterior to Bregma and stored at -80 Celsius. The frozen dmPFC was added to 600  $\mu$ L of buffer RLT (Qiagen DNA/RNA/Protein extraction kit) and homogenized with the Qiagen TissueRuptor for 30 sec. mRNA was then extracted through use of the AllPrep DNA/RNA/protein extraction kit provided by Qiagen in accordance to the protocol provided by the manufacturer. RNA was eluted from the spin column with 50 $\mu$ L of nuclease-free water. RNA (500 ng/sample) was incubated with 2 $\mu$ L of gDNA Wipeout Buffer (Qiagen) at 42°C for 2 minutes then cooled over ice. 1 $\mu$ L of Reverse Transcription Master Mix (Qiagen), 1 $\mu$ L of RT primer mix (Qiagen), and 4 $\mu$ L of Quantiscript RT Buffer (Qiagen) were added to the reaction mixture and incubated in an Eppendorf MasterCycler at 42°C for 18 minutes to amplify the product, then incubated at 95°C for 3 minutes to inactivate the reverse transcriptase. A reverse transcriptase-negative reaction was carried out in parallel with the samples from 500ng of pooled sample RNA.

Levels of mRNA were assessed in triplicate using quantitative real time pcr (qRT-PCR) (Biorad) on the BioRad CFX96 Touch Real-Time system. Negative controls consisted of a DNA-negative sample and a reverse-transcriptase free sample. Standard curves were run on each pcr plate with 3x serial dilutions ranging from 50.0ng/ $\mu$ L to 1.85ng/ $\mu$ L. The data were normalized using three control genes (Gapdh,  $\beta$ Actin, and Tubb5) and dmPFC tissue from naïve age-matched rats according to the equations outlined by Hellemans et al. in 2007.

### Statistical Analyses

The self-administration data for individual sessions during differential access to cocaine were compared to baseline responding (average of days 6, 7, & 8 of differential access) at 10 min (i.e. “loading phase”; e.g. Ahmed & Koob, 1998) and 1 h intervals to compare across all four conditions, as well as across the entire 6 h sessions for the prolonged-access and limited-access + yoked groups. Separate two-way, between-within (group X day), repeated measures ANOVAs, were conducted for the numbers of cocaine infusions followed by Dunnett post-hoc comparisons to deconstruct significant interactions/main effects using the Prism 6 statistical software (Graphpad). Non-reinforced responding (i.e. during the time-out period) was analyzed separately from total responding via a two-way, between-within (group X day), repeated measures ANOVAs followed by Tukey’s post-hoc comparison for the first 10 min and 1 h of self-administration to assess the efficiency of behavioral responding before (days 4 & 5) and after differential access (days 6 to 20). Inactive lever presses were also analyzed but no significant effects or interactions were detected and in all cases, mean inactive lever presses were < 5 (data not shown). Additionally, self-administration data was separated into separate 10 min blocks on day 6 and day 20 to assess loading during initial daily access. A two-way ANOVA (time block X condition) was conducted to assess the differences in cocaine consumption during 10-minute time blocks for the first hour of self-administration in all four experimental groups during the 6<sup>th</sup> day and 20<sup>th</sup> day of self-administration; Tukey’s post-hoc comparison was used to deconstruct significant interactions using the Prism 6 software. Lastly, analyses of normalized quantitative PCR data were performed by one-way MANOVA for *Homer2*, *Grin1*, and *Dlg4* mRNA and decomposed via post-hoc LSD tests using SPSS Statistics 24 (IBM). All graphics were plotted by using the Prism 6 statistical software (Graphpad).

## Results

### Cocaine Intake

A two-way repeated measures ANOVA of cocaine consumption (mg/kg) during the loading phase of self-administration sessions (first 10 minutes) revealed an interaction between time and treatment ( $F_{45, 2055} = 2.88, p < 0.0001$  Figure 1A). Post-hoc comparisons revealed increases in cocaine consumption for day 20 vs baseline in the limited-access group, days 12, 18, 19, and 20 vs baseline in prolonged-access group ( $p < 0.05$ ), and days 11 through 20 vs baseline in limited-access + yoked rats ( $p < 0.05$ ). These results indicate that there is an escalation of cocaine consumption in all cocaine groups for the first 10 minutes of self-administration, where the limited-access + yoked rats show escalation earlier and the limited-access rats show escalation later than the prolonged-access rats.

Two-way repeated measures ANOVA of cocaine consumption (mg/kg) for the first hour of cocaine self-administration revealed a significant interaction between time and treatment ( $F_{45, 1875} = 1.385, p < 0.05$ ; Figure 1B). Dunnett's multiple comparisons *post-hoc* analysis showed increased cocaine consumption in the limited-access rats for day 20 vs baseline ( $p < 0.05$ ), prolonged-access rats for days 18 & 20 vs baseline ( $p < 0.05$ ) and in limited-access + yoked rats for days 14 & 19 vs baseline ( $p < 0.05$ ). These data also indicate that there is an escalation of cocaine consumption in prolonged-access and limited-access + yoked rats that differ in the time of onset.

A two-way ANOVA was run to assess differences in cocaine consumption (mg/kg) during 10 min time blocks for the 1<sup>st</sup> 1 h of self-administration on day 6 and day 20 of the experiment. The results revealed a significant interaction between time block and condition ( $F_{35, 1644} = 1.968, p < 0.001$ ). Tukey's *post-hoc* comparison revealed that rats in the limited-access, prolonged-access, and limited-access + yoked conditions all consumed more cocaine in the first 10 minutes of self-administration than during any other 10-minute block ( $p < 0.01$ ). Additionally, rats in the prolonged-access and limited-access + yoked conditions consumed more cocaine in the first 10 minutes of day 20 than during the first 10 minutes of day 6 of the self-administration procedure ( $p < 0.0001$ ). These results indicate that rats in all cocaine-access conditions consumed more cocaine in the first 10 minutes of self-administration than in the rest of the first hour, and also that rats in the prolonged-access and limited-access + yoked conditions escalated cocaine consumption between day 6 and day 20 of self-administration.

A two-way repeated measures ANOVA was run to assess any differences in total cocaine consumption (mg/kg) over 6 h for the prolonged-access rats versus the limited-access + yoked rats. The results revealed a significant interaction between time and treatment ( $F_{14, 392} = 1.779, p < 0.05$ ), an effect of time ( $F_{14, 392} = 6.874, p < 0.0001$ ), but no effect for treatment ( $F_{1, 28} = 0.0318, p = 0.8597$ ). Dunnett's *post-hoc* comparison revealed that the prolonged-access and limited-access + yoked conditions exhibited escalated cocaine intake from day 12 to day 20. And that there was no difference between prolonged-access and limited-access + yoked conditions. These data indicate that both groups had increased cocaine exposure from baseline and, expectedly as the majority of daily intake in both

groups was controlled by prolonged-access rats, there is no observable difference in cocaine exposure between these two conditions (Figure 1D).

### Non-reinforced Lever Responding

A two-way repeated measures ANOVA of the numbers of non-reinforced active lever-responses (i.e. responses during the time-out period) during the first 10 min of self-administration revealed an interaction between time and treatment ( $F_{45, 2085} = 2.061$ ,  $p < 0.0001$ ) (Figure 2A). Dunnett *Post-hoc* comparisons indicated significant differences in the non-reinforced lever-responding for only the limited-access + yoked-access rats on days 12, 13, 14, 15, 18, and 19 versus baseline (days 6, 7, & 8) ( $p < 0.05$ ) (Figure 2A). These data indicate that for the first 10 min of self-administration, the limited-access + yoked animals exhibited significantly more non-reinforced lever responding than limited-access and prolonged-access conditions well after differential-access to cocaine was initiated.

A two-way repeated measures ANOVA of the non-reinforced lever responding during the first 1 h of self-administration revealed a significant interaction between time and treatment ( $F_{45, 2040} = 2.388$ ,  $p < 0.0001$ ; Figure 2B). Dunnett's multiple comparisons *post-hoc* analysis indicated a significant decrease in the number of non-reinforced active lever responses compared to day 4 of self-administration for days 8–14 & 16–18 for rats with limited-access, days 7–11 & 13–20 for prolonged-access, and days 8 & 9 for limited-access + yoked rats ( $p < 0.05$ ). These data indicate that before differential access, all cocaine-access animals exhibited inefficient behavioral responding at day 4. However, only limited-access + yoked animals continued the pattern of inefficient behavioral responding throughout the experiment.

### Quantitative Real-Time PCR of mRNA levels

A one-way MANOVA of mRNA expression for *Homer2*, *Grin1*, and *Dlg4* resulted in a significant main effect of condition (Hotelling's trace = 0.652,  $F_{12, 110} = 1.991$ ,  $p < 0.05$ ). LSD *post-hoc* pairwise comparisons revealed greater *Homer2* mRNA expression in the prolonged-access group relative to the naïve, saline, and limited-access + yoked groups (figure 3A) LSD *post-hoc* pair-wise comparisons also revealed increased *Grin1* mRNA expression in the prolonged-access group compared to naïve, saline, limited-access, and limited-access + yoked groups (figure 3B). Lastly, LSD *post-hoc* pair-wise comparisons revealed increased levels of *Dlg4* mRNA expression in the prolonged-access group compared to the naïve and limited-access + yoked access groups (figure 3C). These data indicate that prolonged-access rats have a unique molecular phenotype, even though rats in the limited-access + yoked group received equivalent amounts of cocaine and escalated cocaine consumption at about the same rate as the prolonged-access rats.

Additionally, Pearson correlation coefficients were calculated for total cocaine consumption versus mRNA expression. *Homer2* expression was correlated positively with cocaine exposure for both prolonged-access groups ( $R^2 = 0.1975$ ,  $p < 0.05$ ), and limited-access ( $R^2 = 0.4763$ ,  $p < 0.05$ ) groups, whereas mRNA expression correlated negatively for the limited + yoked access group ( $R^2 = 0.4046$ ,  $p < 0.05$ ) (Figure 3D, E, F). Pearson correlation

coefficients failed to reveal significant correlations between total cocaine consumption versus *Grin1* and *Dlg4* mRNA ( $p > 0.05$ ).

## Discussion

The major finding of the present study is that rats self-administering cocaine under all three cocaine-access conditions exhibited escalation of cocaine intake, but with distinct temporal and molecular profiles. Rats in all self-administration conditions escalated their cocaine intake during the loading phase of self-administration (aka the first 10 minutes, Figure 1A,C), during the first 1 h of self-administration, and over entire daily sessions (Figures 1B, D). Additionally, both prolonged-access and limited-access + yoked conditions escalated cocaine intake faster than limited-access animals. However, despite equivalent “excessive” cocaine exposure, the associated patterns of active lever responding are distinct between prolonged-access and limited-access + yoked groups, with the limited-access + yoked condition exhibiting more non-reinforced lever responding on the active lever during the first 10 min of self-administration (i.e. during the time-out period which did not result in cocaine infusion; see Figure 2), suggesting that partially distinctive behavioral mechanisms may underlie the escalation of cocaine intake induced by contingent versus non-contingent cocaine exposure. Furthermore, we observed an overall increase in *Homer2*, *Grin1*, and *Dlg4* mRNA expression only in the prolonged-access rats and total cocaine exposure and *Homer2* mRNA expression are positively correlated in both prolonged- and limited-access conditions but negatively correlated in the limited-access + yoked condition indicating distinct neurobiological consequences of both amount and mode of cocaine exposure. Thus, the present study demonstrates that escalation of cocaine-taking induced by differential cocaine-access, both with respect to session duration and behavioral contingency of cocaine delivery, can produce distinct behavioral and neurobiological consequences.

The finding that all cocaine-taking groups exhibited an escalation of drug intake, but at different rates and to different degrees, is generally consistent with prior findings. Although the capacity of prolonged drug-access to escalate drug-taking, over that observed under limited-access conditions, is a highly replicable finding (e.g. Ahmed and Koob 1998; Ben-Shahar et al. 2004; Ben-Shahar et al. 2013), an escalation of cocaine intake is also reported in rats self-administering cocaine during slightly longer (2 h) daily sessions, albeit to a lesser extent than counterparts with daily 6-h-access (Mandt et al. 2015). Further, some rat strains exhibit escalation under daily 1 h sessions of cocaine-access (Perry et al., 2006), particularly when allowed to self-administer cocaine over a protracted test period (e.g. 75 days) (Belin et al. 2009). The present report extends the literature on escalation by demonstrating that cocaine intake during the initial 10 minutes of the self-administration paradigm (i.e., the loading phase; see Ahmed & Koob, 1998) also escalates with drug experience in all conditions but at somewhat different rates (Figure 1). Thus, it appears that the loading phase (i.e. first 10 min) is more sensitive to escalation of intake than the overall duration of daily access.

The escalation of cocaine consumption observed in the limited-access + yoked condition was additionally associated with a pronounced, but transient, increase in non-reinforced lever-responding during cocaine access (Figures 2A, B). Marked differences in non-



reinforced responding have been reported in the absence of differences in cocaine intake; e.g. in female relative to male rats (Fuchs et al., 2005; Kippin et al., 2006; Kosten & Zhang, 2008); thus, the two measures appear generally dissociable. In the case of the limited-access + yoked rats in the present study, the rats had acquired responding for cocaine, accompanied by low levels of non-reinforced responding which was markedly elevated, particularly during the loading phase (Figure 2A), following exposure to the yoking procedure. This disruption of “efficient” (i.e. lever presses are not tied to the drug reinforcer) operant behavior elicited by the addition of non-contingent cocaine exposure is likely due to discontinuous reinforcement schedules. Further, contextual cues can modulate escalation behavior; when rats are allowed alternating days of 1 h and 6 h access to cocaine with differential cues, they only escalate during the 6 h sessions (Beckmann et al., 2012). An equally-viable explanation for increased non-reinforced lever responding induced by non-contingent drug exposure may pertain to differences in the aversive, stress-inducing, and glucocorticoid-releasing properties of cocaine delivered under yoked procedures (Twining et al. 2009) which have been implicated in the processes underlying escalation (Mantsch et al., 2007, 2008). Thus, the combination of yoked cocaine with subsequent re-introduction to the operant chamber with differential cues (i.e. lever extension and operant light) appears to serve as a potent elicitor of lever-responding.

Given the central role of drug-induced neuroadaptations, particularly alterations in glutamate function, in theories of addiction (see e.g. Kalivas & Volkow, 2005), it is critical to discern the role of behavioral contingency in neurobiological changes associated with escalating drug intake. Here, we identified differences between the prolonged-access and limited-access + yoked conditions; we examined the dmPFC which has been implicated in the escalation of cocaine intake (Smith et al. 2008) for changes in the expression levels of molecular markers implicated in addiction (i.e. *Homer2*, *Grin1*, and *Dlg4* mRNA). Consistent with prior findings (Ben-Shahar et al., 2009), we observed increased levels of *Homer2* mRNA only in the prolonged-access condition, (Figure 3A). Furthermore, the level of intake in the limited- and prolonged-access conditions correlated positively with *Homer2* mRNA expression, whereas cocaine exposure in the limited-access + yoked group correlated negatively with *Homer2* mRNA expression (Figure 3D–F). In addition, we also observed increases in *Grin1* and *Dlg4* mRNA within the dmPFC of only in the prolonged-access rats (Figures 3B–C). Overall, the differences between prolonged-access and limited-access + yoked conditions are consistent with other studies employing yoked procedures (Krawczyk et al. 2013; Ma et al. 2013; McFarland et al. 2003; Radley et al. 2015) but the present finding furthers this literature by demonstrating that contingent and non-contingent cocaine exposure induces distinct neurobiological changes even when both are associated with escalation of cocaine intake, with only the prolonged access condition producing elevation of several genes that are suggestive of enhanced glutamatergic signaling.

The increases in expression of glutamate-related genes observed specifically in following prolonged contingent access to cocaine to current enhanced glutamate neurotransmission and cocaine-specific neuroplasticity in the prefrontal cortex. Repeated contingent access to cocaine results in cocaine-specific synaptic plasticity including: increased dendritic spine density (Frankfurt et al. 2011), increased long-term potentiation (LTP) in the PFC (Huang et al. 2006), and lowered induction threshold for inducing cocaine-specific LTP (Ruan and Yao

2017a). Lastly, optogenetic reversal of cocaine-induced plasticity within the PFC eliminates cocaine seeking behaviors (Pascoli et al. 2014b), indicating that cocaine-specific plasticity of glutamatergic receptors develops with repeated contingent-access to cocaine, and is necessary for cocaine-seeking behavior. We have previously demonstrated that prolonged-access to cocaine enhances baseline glutamate neurotransmission (Shin et al. 2016) after 30 days of withdrawal, and increased expression of the NMDA GluN2b receptor subunit at 3 and 30 days of withdrawal (Szumlinski et al. 2016). Other groups have also shown increases in NMDA as well as AMPA and Kainate receptor subunits after withdrawal from contingent cocaine self-administration (Crespo et al. 2002; Ghasemzadeh et al. 2009; Tang et al. 2004).

Briefly, *Homer2* is a gene encoding for a scaffolding protein that links metabotropic glutamate receptors (mGluRs) and NMDA ionotropic glutamate receptors and has been implicated in addictive behaviors (c.f., Szumlinski et al., 2008). Homer2a/b protein within the mPFC is increased following prolonged-access to cocaine (Ben Shahaar et al., 2009) and repeated cocaine injections (Ary and Szumlinski 2007). Further, viral-mediated *Homer2b* overexpression in the mPFC increases basal glutamate levels and cocaine-preference in mice, whereas *Homer2b* knockdown reduces basal glutamate in this area (Ary et al. 2013). Additionally, *Grin1* encodes for the obligatory NR1 subunit of NMDA receptors, while *Dlg4* encodes for PSD-95, a scaffolding protein that regulates NMDA receptor function (Bai and Hoffman 2009). Increases in NR1 protein within the PFC have previously been observed in response to repeated cocaine injections (Kovacs et al. 2010) and cocaine self-administration (Hemby et al. 2005). Furthermore, NR1 is essential for cocaine-mediated learning; mice expressing a mutant version of the NR1 subunit (which reduces calcium flow through the NMDA receptor) fail to form conditioned place preference and locomotor sensitization in response to repeated cocaine exposure (Heusner and Palmiter 2005). PSD-95 is critical for synaptic plasticity and regulation of NMDA receptor location and function (Wang and Peng 2016). PSD-95 is also implicated in behavioral plasticity associated with chronic cocaine administration (Yao et al. 2004), extinction of cocaine self-administration (Knackstedt et al. 2010; Ghasemzadeh et al. 2011), as well as prolonged withdrawal from cocaine (Ghasemzadeh et al. 2009; McIntosh et al. 2013). Thus, our RNA data is generally consistent with findings examining protein levels of glutamatergic signaling molecules with the increases in RNA observed here coinciding or preceding latent increases in protein.

Escalation of drug consumption is an important diagnostic criterion of addiction in humans and an integral component in various theories of addiction. Therefore, understanding the behavioral and neurobiological underpinnings of escalation is likely to facilitate addiction management programs in humans. In addition to facilitating cocaine intake, prolonged daily access to cocaine is associated with several behavioral changes, such as reduced brain reward function (Ahmed et al. 2002; Ahmed and Koob 2005), increased breakpoints for cocaine reinforcement under progressive ratio schedules (Paterson and Markou 2004; Wee et al. 2009), diminished aversive properties of cocaine (Ben-Shahaar et al. 2008), increased extinction responding during protracted withdrawal (Ferrario et al. 2005), as well as increased responding during cocaine-primed and cue-induced reinstatement of cocaine-seeking (Ahmed and Cador 2006; Kippin et al. 2006; Knackstedt and Kalivas 2007; Mantsch et al. 2004). Similarly, other approaches to modeling “excessive” intake also produce increases in measures of cocaine-taking and -seeking behaviors (Deroche-Gamonet et al.

2004; Roberts et al. 2007). Further study is required to determine the relations between nature of cocaine exposure and induction of addiction-like behavior, as well as between behavioral and molecular outcomes. To this end, the present study demonstrates behavioral contingency plays an important role in the nature of the behavioral and molecular changes induced by cocaine exposure.

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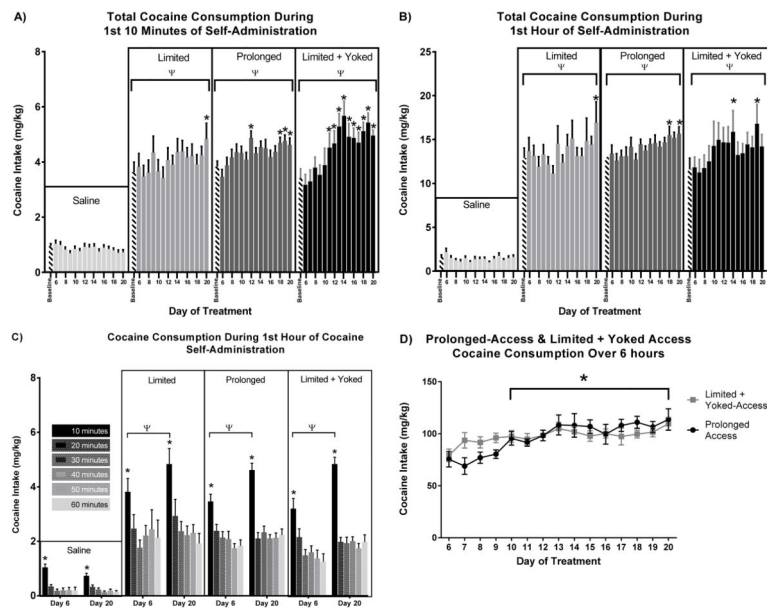
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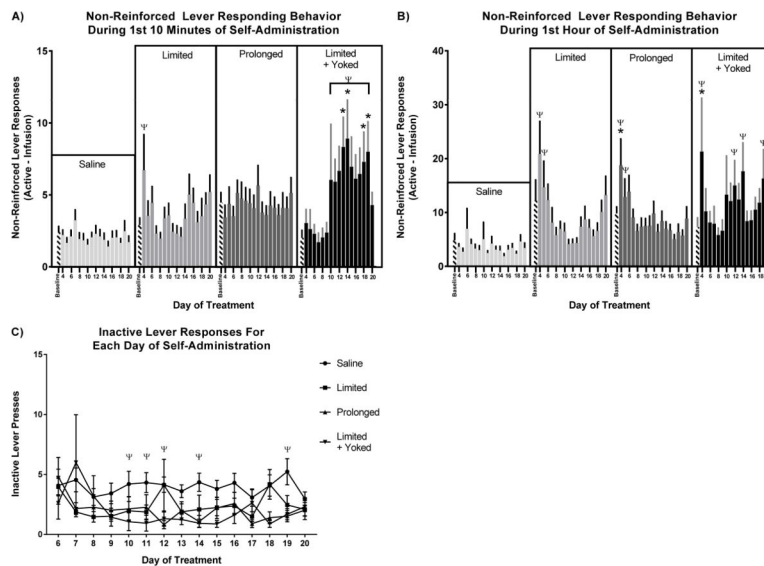
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**Figure 1. Cocaine consumption during self-administration**

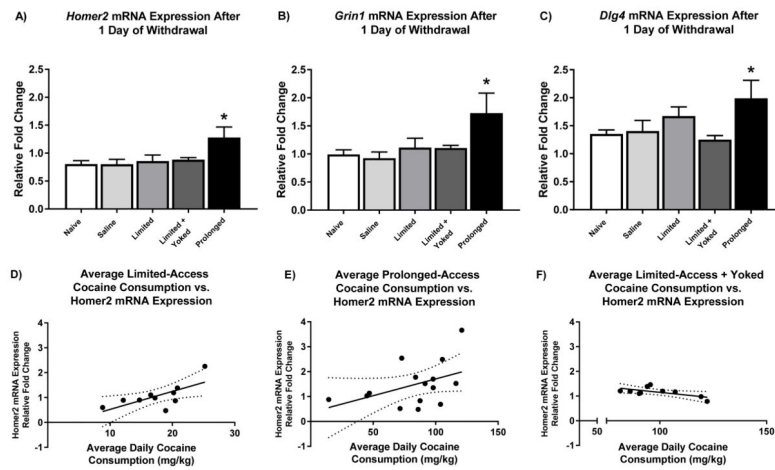
**A) 1<sup>st</sup> 10 min of self-administration.** Limited-access rats escalated cocaine consumption on day 20 of self-administration. Prolonged-access rats exhibited escalated cocaine consumption on days 12, 14, 15, 18, 19, and 20 of self-administration. Limited + yoked rats escalated cocaine consumption in blocks between days 11 and 20 of self-administration. (\*  $p < 0.05$ ,  $\psi p < 0.05$ ) **B) 1<sup>st</sup> 1 hour of self-administration.** Prolonged-access rats exhibited escalated cocaine consumption in time block 5. Limited + yoked rats exhibited escalated cocaine consumption in time blocks 2, 3, 4, and 5. Saline and limited access rats did not escalate cocaine intake in the first hour of cocaine self-administration. (\*  $p < 0.05$ ) **C) Cocaine consumption recorded every 10 minutes during 1<sup>st</sup> hour of self-administration.** Limited-access, prolonged-access, and limited + yoked-access rats all displayed increased cocaine consumption during the first 10 minutes of self-administration sessions on both day 6 and day 20 of self-administration. Additionally, prolonged-access and limited + yoked-access rats exhibited increased cocaine consumption within the 1<sup>st</sup> 10 minutes of self-administration between day 6 and day 20. (\*  $p < 0.05$ ,  $\psi p < 0.05$ ) **D) Total daily cocaine infusions for prolonged and limited + yoked access rats.** Total cocaine consumption did not differ between prolonged and limited + yoked access rats. Cocaine consumption escalated in both groups beginning on the 10<sup>th</sup> day of extended access to cocaine. (\*  $p < 0.05$ )





**Figure 2. Non-reinforced lever responding behavior**

**A) 1<sup>st</sup> 10 min of self-administration.** Only limited + yoked rats exhibited high levels of non-reinforced cocaine responding on days 14, 15, 18, and 19 during the first 10 minutes of cocaine self-administration. **B) 1<sup>st</sup> 1 hour of self-administration.** All cocaine-access groups showed an initial non-reinforced active-lever responding for the first 1 hour of self-administration during cocaine acquisition on day 4. **C) Inactive lever responses over entire day.** There were no differences in the inactive lever responses within or between groups across all days.



**Figure 3. mRNA expression for glutamatergic genes in the dmPFC**

**A)** Prolonged-access to cocaine resulted in increased levels of *Homer2* mRNA within the dmPFC after 1 day of withdrawal. **B)** Prolonged-access to cocaine resulted in increased levels of *Grin1* mRNA within the dmPFC after 1 day of withdrawal. **C)** Prolonged-access to cocaine resulted in increased levels of *Dlg4* mRNA within the dmPFC after 1 day of withdrawal. **D)** *Homer2* mRNA was positively correlated with total cocaine infusions in the prolonged access group ( $R^2 = 0.1975$ ,  $p < 0.05$ ). **E)** *Homer2* mRNA was positively correlated with total cocaine infusions in the limited access group ( $R^2 = 0.4763$ ,  $p < 0.05$ ). **F)** *Homer2* mRNA was negatively correlated with total cocaine infusions in the limited + yoked access group ( $R^2 = 0.4046$ ,  $p < 0.05$ ).