A Pilot Investigation of Acute Inhibitory Control Training in Cocaine Users

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A pilot investigation of acute inhibitory control training in cocaine users


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Abstract

Background—Disrupted response inhibition and presence of drug-cue attentional bias in cocaine-using individuals have predicted poor treatment outcomes. Inhibitory control training could help improve treatment outcomes by strengthening cognitive control. This pilot study assessed the effects of acute inhibitory control training to drug- and non-drug-related cues on response inhibition performance and cocaine-cue attentional bias in cocaine-using individuals.

Methods—Participants who met criteria for a cocaine-use disorder underwent five sessions of inhibitory control training to either non-drug-related cues (i.e., rectangles) or cocaine cues (n=10/condition) in a single day. Response inhibition and attentional bias were assessed prior to and following training using the stop-signal task and visual-probe task with eye tracking, respectively.
Results—Training condition groups did not differ on demographics, inhibitory control training performance, response inhibition, or cocaine-cue attentional bias. Response inhibition performance improved as a function of inhibitory control training in both conditions. Cocaine-cue attentional bias was observed, but did not change as a function of inhibitory control training in either condition.

Conclusions—Response inhibition in cocaine-using individuals was augmented by acute inhibitory control training, which may improve treatment outcomes through better behavioral inhibition. Future studies should investigate longer-term implementation of inhibitory control training, as well as combining inhibitory control training with other treatment modalities.

Keywords
Inhibitory control training; response inhibition; attentional bias; cocaine-use disorders

1. Introduction

Effective inhibition of motivated behaviors triggered by external cues is critical for accomplishing adaptive behavior change (Verbruggen et al., 2014). There has been recent interest in developing interventions that improve inhibitory control in individuals with substance use disorder (Sofuoglu et al., 2013). Cocaine-using individuals display disrupted response inhibition (Fillmore and Rush, 2002; Lane et al., 2007; Verdejo-García et al., 2007; Verdejo-García and Pérez-García, 2007) and cocaine-cue attentional bias (Hester et al., 2006; Lui et al., 2011; Marks et al., 2014a; 2014b; Vadhan et al., 2007). These deficits have been associated with poor treatment retention (Carpenter et al., 2006; Marhe et al., 2013; Streeter et al., 2008; Verdejo-García et al., 2012; Waters et al., 2012).

Poor inhibitory performance is worsened following the presentation of cocaine cues in cocaine users, as demonstrated by inhibitory failures on the Attentional Bias-Behavioral Activation (ABBA) task (Pike et al., 2013, 2015). This cue-dependent response suggests that disproportionate attention towards cocaine cues interacts with cognitive-control mechanisms to exacerbate impaired inhibitory control. Prior research has demonstrated that inhibitory control training involving repeated associative pairings between salient cues and successful inhibition is effective at engendering significant behavior change related to problematic alcohol (Houben et al., 2011, 2012; Jones and Field, 2013) and food (Houben and Jansen, 2011, 2015) consumption. Inhibitory control training to cocaine cues might also be a useful therapeutic adjunct for problematic cocaine use, but this approach has not yet been evaluated.

The purpose of this pilot study was to test the effect of acute (i.e., one day) inhibitory control training to cocaine cues and a non-drug-cue control condition on response inhibition and cocaine-cue attentional bias in individuals with cocaine use disorder. We predicted that cocaine users who completed acute inhibitory control training to cocaine cues would display improved performance on both of these outcomes.
2. Methods

2.1. Participants

Twenty non-treatment seeking participants completed this study. Participants were primarily recruited through word of mouth and postings on community bulletin boards. Participants completed screening questionnaires on their physical and mental health history, measures of current psychological functioning, detailed drug use history questionnaires, and a computerized version of the Structured Clinical Interview for the Diagnostic and Statistical Manual of Mental Disorders-IV (SCID; Sevak et al., 2011). All participants reported using cocaine within the last month, provided a cocaine-positive urine sample during screening procedures, and met criteria for cocaine abuse or dependence on the SCID. Exclusion criteria were: 1) current prescription for a psychiatric medication or 2) presence of a major medical or psychiatric condition (e.g., schizophrenia). Participants were also excluded if they reported dependence on any drug that produce significant withdrawal symptoms during testing (e.g., opioids and benzodiazepines), as they were asked to abstain from drug use for 12 hours prior to testing. All participants provided informed consent prior to beginning the study and were paid for their participation. The University of Kentucky Institutional Review Board approved all procedures, recruitment methods, and the informed consent document.

2.2. Procedure

Participants were randomized to complete the inhibitory-control training task with either cocaine-related images or rectangles as cues. Urn randomization matched participants on sex, cocaine use in the past month (>15 days used or not), and years of education (≥12 years education or not). The study consisted of a practice day (total time approximately 1 hour) to familiarize participants with the computerized measures of response inhibition and cocaine-cue attentional bias, and one day of inhibitory control training (total time approximately 6 hour). Response inhibition and cocaine-cue attentional bias were assessed at two time-points on the day of inhibitory control training: pre-training and post-training (both occurring 45 minutes before and after training, respectively). All computerized tasks were administered using E-prime software (Psychology Software Tools, Pittsburgh, PA) on a PC computer.

2.3. Computerized tasks

2.3.1. Inhibitory control training task—Participants completed five sessions (each separated by 1 hour) of inhibitory control training to either cocaine images or rectangles as cues. In the cocaine condition, go cues were neutral images and no-go cues were cocaine-related images. In the neutral condition, go and no-go cues were vertical and horizontal rectangles, respectively. In both training conditions, the color green served as the go target and color blue served as the no-go target. Participants were instructed not to respond to any image or rectangle, but to only respond once the cue was replaced by a green rectangle (i.e., go target). The inhibitory control-training task used a similar procedure as the ABBA task (described in detail in Pike et al., 2013, 2015), except go and no-go cues always predicted go and no-go targets, respectively. Participants began with $1.65 bonus earnings at the beginning of each training session. Five cents were deducted from this $1.65 bonus for each error that occurred (commission errors on no-go trials and omission errors on go trials) during the training session. The dependent variables were the number of go targets (125
total) to which participants correctly responded and the number of no-go targets (125 total) to which participants correctly withheld a response.

2.3.2. Stop-signal task—Response inhibition performance was measured as the failure to inhibit responses following stop-signals. Briefly, letters (X or O) were presented on a computer screen one at a time. Participants were instructed to identify the letter using a keyboard. The stop-signal (a 500 ms 900 Hz tone) occurred on approximately 30% of trials (50) and participants were instructed to withhold their response in the presence of this tone. The stop-signal was randomly presented after the onset of a letter (X or O) at one of five delays (10, 70, 150, 230, or 300 ms) and was divided evenly between letters. The dependent variables from the stop-signal task were the mean proportion of inhibitory failures following the stop-signal and stop-signal reaction time (SSRT; the estimated time to inhibit responses to the stop-signal; Logan, 1994). This task was used as the primary outcome of inhibitory control to: (1) avoid contamination of acute inhibitory control training with a cued go-no go or ABBA task that is very similar to the training task (Pike et al., 2013, 2015; Weafer and Fillmore, 2012a, 2012b) and (2) examine whether acute inhibitory control training conferred benefits in response inhibition performance beyond impulsive responses to cocaine cues.

2.3.3. Visual-probe task—Attentional bias was measured as fixation time to the cocaine and neutral cues using a Tobii X2-60 eye tracker (Tobii Technology, Sweden; procedure described in detail in Marks et al., 2014a). The dependent variable from the visual-probe task was the mean fixation time (ms). Mean fixation time was calculated by summing the total fixation time for each cue type and then dividing by the total number of critical trials (i.e., 40). Two participants’ data from the visual-probe task were removed because eye-tracking data were insufficient (i.e., greater than 2 SDs below the mean number of fixations recorded). One additional participant’s data from the visual-probe task were removed because the eye-tracker was not able to record the participant’s eye movements.

2.4. Data analysis

Statistical analyses were performed in SPSS Version 22 (IBM Corporation, Armonk, NY) with an alpha level set at \( p \leq 0.05 \) for all outcomes. Independent-samples t-tests were conducted to compare demographics for each training condition (cocaine cues vs. rectangles) for continuous variables. Chi-square analyses were conducted to compare sex and ethnicity/race distribution. Behavioral performance outcomes from all tasks were analyzed using mixed-model ANOVAs with training condition as the between-subjects factor. Two ANOVAs were used for comparisons of accurate responding to go targets and accurate withholding to no-go targets on the training tasks, with trial number (i.e., 1–5) as the within-subjects factor. ANOVAs were conducted on both outcomes from the stop-signal task (proportion of inhibitory failures and SSRT) with time-point (i.e., pre- and post-training) as the within-subject factor in both models. An ANOVA was also conducted on mean fixation time from the visual-probe task with cue type (cocaine vs. neutral) and time-point (i.e., pre- and post-training) as the within-subjects factors. Post-hoc comparisons with Bonferroni’s adjustment were conducted following the mixed-model ANOVAs if significant effects were found. Effect sizes for significant results were calculated as partial eta squared (\( \eta^2_p \)). Observed power for significant results are reported.
3. Results

3.1. Sample characteristics

There were no significant differences between the training groups on any demographic variable (Table 1).

3.2 Computerized tasks

3.2.1 Inhibitory control training task—No significant effects on correctly withheld responses or correctly executed responses were found between groups across all trials (all Fs < 2.25, ps > 0.05).

3.2.2. Stop-signal task—ANOVA revealed significant main effects of time-point on the proportion of inhibitory failures following stop-signals ($F_{(1, 18)} = 4.62, p = 0.05$, $\eta_p^2 = 0.20$, $\text{power} = 0.53$; Figure 1A) and on the SSRT ($F_{(1, 18)} = 4.98, p = 0.04$, $\eta_p^2 = 0.22$, $\text{power} = 0.56$; Figure 1B). Post-hoc comparisons revealed that the proportion of inhibitory failures and SSRT decreased after inhibitory control training for both conditions. No other significant effects were detected (all Fs < 1.00, ps > 0.05).

3.2.3 Visual-probe task—A significant main effect of cue type ($F_{(1, 15)} = 8.36, p = 0.01$, $\eta_p^2 = 0.36$, $\text{power} = 0.77$; Figure 1C) was detected for fixation time. Post-hoc comparisons revealed that the mean fixation time to cocaine cues was greater than neutral cues. No other significant effects were detected (all Fs < 2.25, ps > 0.05).

4. Discussion

This pilot study was the first to examine acute inhibitory control training in cocaine-using individuals. This study also extended the previous literature by assessing changes in response inhibition outside of the training task (i.e., the stop signal task) and using a measure of cue-specific attentional bias (i.e., the visual probe with eye tracking), which have not been included in prior studies that examined the acute effect of inhibitory control training to alcohol cues in heavy drinkers.

Inhibition is a key component of executive control that is often impaired in substance-use disorder (Verbruggen and Logan, 2008, 2009). For example, Fillmore and Rush (2002) found that chronic cocaine users displayed greater proportions of inhibitory failures and longer SSRTs than non-cocaine-using controls on the stop-signal task, suggesting that impaired inhibitory control in cocaine-using individuals might be due to a generalized slowing of the processes that inhibit impulsive responses. Therefore, the reduced proportions of inhibitory failures and shortened SSRTs observed in cocaine-using individuals following acute inhibitory control training in this study are noteworthy and suggest that greater executive control over impulsive behavior resulted from acute training. These novel data are clinically meaningful because disinhibition in cocaine-using individuals has been associated with poor treatment retention (Streeter et al., 2008; Verdejo-García et al., 2012). Inhibitory control training might therefore promote improved treatment outcomes for individuals seeking treatment for their cocaine use, though future studies are needed to assess this possibility.
The magnitude of cocaine-cue attentional bias in this study fell within the large effect size range for partial-eta squared ($\eta^2_p = 0.36$; Cohen, 1988), which is within the same effect size range as the reported Cohen’s d values from previous studies that used this task (Cohen’s ds: 1.2 – 1.9; Marks et al., 2014a, 2014b). Neither type of acute inhibitory control training altered attentional bias. There are two potential reasons for this null finding. First, there is no inhibitory-control component of the visual-probe task. The use of an antisaccade task to assess visual inhibitory control in cocaine-cue attentional bias would be needed to reveal an impact of inhibitory control training on attentional bias tracked by eye movements (Dias et al., 2015; DiGirolamo et al., 2015). Second, a single day of acute inhibitory control training is insufficient to offset the learned associations that contribute to cocaine-cue attentional bias. Attentional bias is theorized to develop from acquired motivational salience that occurs from repeated pairings of substance use with proximal environmental cues (Field and Cox, 2008). Behavioral Stimulus Interaction theory (Veling et al., 2008) posits that motivational stimuli, such as cocaine cues, are devalued through multiple pairings of the cues with successful behavioral inhibition. Cocaine-cue stimulus devaluation through successful inhibition may rely more on automatic processes that are more resistant to change, whereas response inhibition relies on executive processes that are more flexible (Wiers and Stacy, 2006; Wiers et al., 2007). Thus, extensive inhibitory control training sessions would be needed to diminish the associative strength of drug-cue pairings that contribute to cocaine-cue attentional bias.

Although the small sample size of this pilot investigation is a limitation, it is important to note that all statistically significant outcomes achieved appreciable power (range: 0.53 – 0.77). There are two additional study limitations that are worth mentioning. First, one day of five training sessions was delivered, and as noted above, this limited amount of training was insufficient to reduce cocaine-cue attentional bias. Future studies should investigate the efficacy of multiple sessions of inhibitory control training. Second, the present study did not assess cocaine use following training. An aim for future studies is to determine if inhibitory control training reduces cocaine use in the natural ecology and on self-administration procedures under controlled laboratory conditions.

This study contributed to our understanding of the impact of inhibitory control training in stimulant users by demonstrating that both neutral and cocaine-related training procedures augmented a clinically relevant behavior (i.e., response inhibition), but did not change attentional bias in cocaine users. Long-term implementation of inhibitory control training, possibly in conjunction with other behavioral interventions (e.g., cognitive-behavioral therapy or contingency management) or pharmacotherapeutics, could lead to adaptive behavior change and improved treatment outcomes.

References


Highlights

- Acute inhibitory control training tested in cocaine users
- Cocaine users were assigned to cocaine or neutral cue training
- Improved response inhibition performance was observed in both conditions
- Cocaine-cue attentional bias was not altered by acute inhibitory control training
Figure 1.
Panel 1A: Mean proportion of inhibitory failures (+SEM) during stop trials on the stop-signal task as a function of inhibitory control training to cocaine-cues (n = 10) and rectangles (n = 10). Proportion of inhibitory failures means are shown for pre-training (left two bars) and post-training (right two bars). Reduced proportion of inhibitory failures represents improved performance.
Panel 1B: Mean SSRT (+SEM) to the stop-signal on the stop-signal task as a function of inhibitory control training to cocaine-cues (n = 10) and rectangles (n = 10). Mean SSRTs are shown for pre-training (left two bars) and post-training (right two bars). Lower SSRT represents faster executive control of inhibiting erroneous impulsive responses.
Panel 1C: Mean gaze time (ms; +SEM) from the visual-probe task as a function of inhibitory control training to cocaine-cues (n = 9) and rectangles (n = 8). Mean gaze times are shown for pre-training (left four bars) and post-training (right four bars). Attentional bias toward cocaine cues is shown by greater mean gaze time to cocaine cues than to neutral cues. * = significant main effect of time-point (p < 0.05)
Table 1

Participant demographics by training condition and group comparisons.

<table>
<thead>
<tr>
<th>Demographic variable</th>
<th>Cocaine-cues</th>
<th>Rectangles</th>
<th>t or χ² -Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Group size (n):</td>
<td>10</td>
<td>10</td>
<td>-</td>
</tr>
<tr>
<td>UDS+ for cocaine metabolites (n):</td>
<td>10</td>
<td>9</td>
<td>-</td>
</tr>
<tr>
<td>Sex (n male/female):</td>
<td>6/4</td>
<td>6/4</td>
<td>0.00</td>
</tr>
<tr>
<td>Mean age (±SEM):</td>
<td>50.4 (1.0)</td>
<td>45.7 (1.1)</td>
<td>2.02</td>
</tr>
<tr>
<td>Mean years of education (±SEM):</td>
<td>11.7 (0.5)</td>
<td>12.3 (0.5)</td>
<td>-0.90</td>
</tr>
<tr>
<td>Race/Ethnicity (n AA/C/MO):</td>
<td>5/4/1</td>
<td>8/2/0</td>
<td>0.26</td>
</tr>
<tr>
<td>Mean cigarettes/day (±SEM):</td>
<td>9.2 (2.1)</td>
<td>10.1 (2.5)</td>
<td>-0.26</td>
</tr>
<tr>
<td>Mean alcoholic drinks/week (±SEM):</td>
<td>18.8 (8.3)</td>
<td>16.4 (7.9)</td>
<td>0.22</td>
</tr>
<tr>
<td>Mean DAST score (±SEM):</td>
<td>10.2 (1.8)</td>
<td>11.4 (1.9)</td>
<td>-0.47</td>
</tr>
<tr>
<td>Mean days of cocaine use (±SEM)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>-Past week:</td>
<td>2.7 (0.3)</td>
<td>3.5 (0.8)</td>
<td>-0.98</td>
</tr>
<tr>
<td>-Past month:</td>
<td>12.6 (1.5)</td>
<td>13.3 (3.1)</td>
<td>-0.21</td>
</tr>
</tbody>
</table>

Note. UDS+ = positive urine drug screens on the day of inhibitory control training, AA = African-American; C = Caucasian; MO = Mixed/Other; DAST = Drug Abuse Screening Test; SEM = standard error of the mean.