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Article type : Original Research Article

Elucidating the Effect of a Brief Drinking Intervention Using Neuroimaging: A Preliminary Study

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Funding Sources: Supported by NIH grants R21AA023669 (LAR and MPK), and training grants T32 DA007272 (AL) and T32DA024635 (EG).

Financial Disclosures: None of the authors have any conflicts of interest or financial disclosures.

This article has been accepted for publication and undergone full peer review but has not been through the copyediting, typesetting, pagination and proofreading process, which may lead to differences between this version and the Version of Record. Please cite this article as doi: 10.1111/acer.13941

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Abstract

Background: Brief interventions have empirical support for acutely reducing alcohol use among non-treatment seeking heavy drinkers. Neuroimaging techniques allow for the examination of the neurobiological effect of behavioral interventions, probing brain systems putatively involved in clinical response to treatment. Few studies have prospectively evaluated whether psychosocial interventions attenuate neural cue-reactivity that in turn reduces drinking in the same population. This study aimed to examine the effect of a brief intervention on drinking outcomes, neural alcohol cue-reactivity, and the ability of neural alcohol cue-reactivity to prospectively predict drinking outcomes.

Methods: Non-treatment-seeking heavy drinking participants were randomized to receive a brief interview intervention (n = 22) or an attention-matched control (n = 24). Immediately following the intervention or control, participants underwent an fMRI scan comprised of the alcohol taste cues paradigm. Four-weeks after the intervention (or control) participants completed a follow-up visit to report on their past-month drinking. Baseline and follow-up percent heavy drinking days (PHDD) were calculated for each participant.

Results: There was no significant effect of the brief intervention on PHDD at follow-up or on modulating neural activation to alcohol relative to water taste cues. There was a significant association between neural response to alcohol taste cues and PHDD across groups (Z>2.3,
\( p<0.05 \), such that individuals who had greater neural reactivity to alcohol taste cues in the precuneus and prefrontal cortex had fewer PHDD at follow-up.

Conclusions: This study did not find an effect of the brief intervention on alcohol use in this sample, and the intervention was not associated with differential neural alcohol cue reactivity. Nevertheless, greater activation of the precuneus and prefrontal cortex during alcohol cue exposure predicted less alcohol use prospectively suggesting that these neural substrates subserve the effects of alcohol cues on drinking behavior.

Key words: brief intervention; mechanisms of behavior change; fmri; alcohol; precuneus

Introduction

Brief interventions have empirical support for acutely reducing alcohol use among non-treatment seeking heavy drinkers. For example, randomized clinical trials of brief interventions have found favorable results among heavy drinkers reached through primary care (Saitz et al., 2003, Fleming et al., 1997), trauma centers (Gentilello et al., 1999) and emergency departments (Bernstein et al., 2007, D’Onofrio and Degutis, 2002). Brief interventions also have shown effectiveness in reducing alcohol use in non-medical settings among a young adult college population (Carey et al., 2006). Given this sizable evidence base, there is considerable interest in understanding the underlying mechanisms toward optimizing this approach.

Neuroimaging techniques allow for the examination of the neurobiological effects underlying behavioral interventions, probing brain systems putatively involved in clinical response to treatment. To date, one study has examined the effect of a motivational
interviewing-based intervention on the neural substrates of alcohol reward (Ewing et al., 2011). In this study, neural response to alcohol cues was evaluated while individuals were exposed to change talk and counterchange talk (i.e., sustain talk), which are thought to underlie motivation changes during psychosocial intervention. The authors report activation in reward processing areas following counterchange talk, which was not present following exposure to change talk (Ewing et al., 2011). Feldstein Ewing and colleagues have also probed the nature of the origin of change talk in order to better understand the neural underpinnings of change language (Feldstein Ewing et al., 2014). In this study, binge drinkers were presented with self-generated and experimenter-selected change and sustain talk. Self-generated change talk and sustain talk resulted in greater activation in regions associated with introspection, including the interior frontal gyrus and insula, compared to experimenter elicited client language (Feldstein Ewing et al., 2014). These studies employed an active ingredient of MI within the structure of the fMRI task, thus allowing for a more proximal test of treatment effects.

Neuroimaging has also been used to explore the effect of psychological interventions on changes in brain activation that are specifically focused on alcohol motivation. For example, cue-exposure extinction training, a treatment designed to prevent return to use by decreasing conditioned responses to alcohol cue stimuli through repeated exposure to cues without paired reward, has also been evaluated using neuroimaging (Vollstadt-Klein et al., 2011). Alcohol dependent patients who underwent cue-exposure extinction training had larger decreases in neural alcohol cue-reactivity in mesocorticolimbic reward circuitry than patients who had standard clinic treatment. Cognitive bias modification training, which similarly trains individuals to reduce attentional bias towards alcohol cues, resulted in decreased neural alcohol cue-reactivity in the amygdala (Wiers et al., 2015b) and reduced medial prefrontal cortex (mPFC) activation when approaching alcohol cues (Wiers et al., 2015a). These studies suggest that fMRI tasks may be sensitive to treatment response.
Further, neurobiological circuits identified using fMRI can be used to predict treatment and drinking outcomes, providing unique information beyond that of self-report and behavior. Individuals with alcohol use disorder who return to use demonstrate increased activation in the mPFC to alcohol cues compared to individuals with AUD who remain abstinent (Beck et al., 2012, Grusser et al., 2004). Moreover, the degree that the mPFC was activated was associated with the amount of subsequent alcohol intake, but not alcohol craving (Grusser et al., 2004). Activation in the dorsolateral PFC to alcohol visual cues has been associated with higher percent heavy drinking days in treatment-seeking alcohol dependent individuals (Schacht et al., 2013b). Increased activation in the mPFC, orbitofrontal cortex, and caudate in response to alcohol cues has also been associated with the escalation of drinking in young adults (Dager et al., 2014). Mixed findings have been reported for the direction of the association between cue-induced striatal activation and return to use. Increases (Grusser et al., 2004, Bach et al., 2015, Reinhard et al., 2015) and decreases (Beck et al., 2012) in ventral and dorsal striatal activation to alcohol cues have been associated with subsequent return to use. Utilizing a different paradigm, Seo and colleagues found that increased mPFC, ventral striatal, and precuneus activation to individually tailored neutral imagery scripts predicted subsequent return to use in treatment-seeking individuals with AUD (Seo et al., 2013). Interestingly, brain activity during individually tailored alcohol and stress imagery scripts was not associated with return to use (Seo et al., 2013).

While initial evidence indicates that psychological interventions are effective at reducing mesocorticolimbic response to alcohol-associated cues, few studies have prospectively evaluated if psychosocial interventions attenuate neural cue-reactivity that in turn reduces drinking in the same population. Furthermore, no previous studies have used neural reactivity to alcohol cues to understand the mechanisms of brief interventions. Therefore, this study aimed to examine the effect of a brief intervention on drinking outcomes, neural alcohol cue-reactivity, and the ability of neural alcohol cue-reactivity to
predict drinking outcomes. Specifically, this study investigated: 1) if the brief intervention would reduce percent heavy drinking days or drinks per week in non-treatment seeking heavy drinkers in the month following the intervention and 2) if the brief intervention would attenuate neural alcohol cue-reactivity. In the first case, we predicted significant effects on drinking based on the existing clinical literature and, in the second case, we predicted decrements in alcohol’s motivational salience based on the feedback about the participant’s drinking levels relative to clinical recommendations and their personal negative consequences of drinking. The effects of neural cue reactivity on subsequent drinking outcomes were tested in order to elucidate patterns of neural cue-reactivity that predict drinking behavior prospectively.

Methods

Participants and Screening Procedures

Participants were recruited between November 2015 and February 2017 from the greater Los Angeles metropolitan area. Study advertisements described a research study investigating the effects of a brief health education session on beliefs about the risks and benefits of alcohol use. Inclusion criteria were as follows: (1) engaged in regular heavy drinking, as indicated by consuming 5 or more drinks per occasion for men or 4 or more drinks per occasion for women at least 4 times in the month prior to enrollment (as indicated on the Timeline Follow-back); (2) a score of ≥8 on the Alcohol Use Disorder Identification Test (AUDIT; (Saunders et al., 1993)). Exclusion criteria included (1) under the age of 21; (2) currently receiving treatment for alcohol problems, history of treatment in the 30 days before enrollment, or currently seeking treatment; (3) a positive urine toxicology screen for any drug other than cannabis; (4) a lifetime history of schizophrenia, bipolar disorder, or other psychotic disorder; (5) serious alcohol withdrawal symptoms as indicated by a score of ≥10 on the Clinical Institute Withdrawal Assessment for Alcohol-Revised (CIWA-AR; (Sullivan et
al., 1989)); (6) history of epilepsy, seizures, or severe head trauma; (7) non-removable ferromagnetic objects in body; (8) claustrophobia; and (9) pregnancy.

Initial assessment of the eligibility criteria was conducted through a telephone interview. Eligible participants were invited to the laboratory for additional screening. Upon arrival, participants read and signed an informed consent form. Participants then completed a series of individual differences measures and interviews, including a demographics questionnaire and the Timeline Follow-back (TLFB; 32) to assess for quantity and frequency of drinking over the past 30 days. All participants were required to test negative on a urine drug test (except for marijuana, which was allowed to be positive). A total of 120 participants were screened in the laboratory, 38 did not meet inclusion criteria and 12 decided not to participate in the trial, leaving 60 participants who enrolled and were randomized. Of the 60 individuals randomized, 46 completed the entire study. See Figure 1 for a CONSORT Diagram for this trial.

**Study Design**

The study was a randomized controlled trial. Participants were assessed at baseline for study eligibility and eligible participants returned for the randomization visit up to two weeks later. During their second visit, participants completed assessments, and then were randomly assigned to receive a 1-session brief intervention or to an attention-matched control condition. Immediately after the conclusion of the session participants completed a functional magnetic resonance imaging (fMRI) scan to assess brain activity during exposure to alcohol cues and completed additional assessments. Participants were followed up 4 weeks later to assess alcohol use since the intervention (or control) through the 30-day Timeline Followback interview. Participants who completed all study measures were compensated $160.
The brief intervention consisted of a 30-45 minute individual face-to-face session based on the principles of motivational interviewing (MI) (Miller and Rose, 2009, Miller, 2002). The intervention adhered to the FRAMES model which includes personalized feedback (F), emphasizing personal responsibility (R), providing brief advice (A), offering a menu (M) of change options, conveying empathy (E), and encouraging self-efficacy (S). In accordance with MI principles the intervention was non-confrontational and emphasized participants’ autonomy. The content of the intervention mirrored brief interventions to reduce alcohol use that have been studied with non-treatment seeking heavy drinkers (e.g. (Borsari and Carey, 2000, Longabaugh et al., 2001, Saitz et al., 2007)). The intervention included the following specific components: 1) giving normative feedback about frequency of drinking and of heavy drinking; 2) Alcohol Use Disorders Identification Test score and associated risk level (Saunders et al., 1993); 3) potential health risks associated with alcohol use; 4) placing the responsibility for change on the individual; 5) discussing the reasons for drinking and downsides of drinking; and 6) setting a goal and change plan if the participant was receptive (see Supplement for Brief Intervention Session Checklist). The aim of the intervention was to help participants understand their level of risk and to help them initiate changes in their alcohol use. Sessions were delivered by master’s-level therapists who received training in MI techniques, including the use of open-ended questions, reflective listening, summarizing, and eliciting change talk, and in the content of the intervention. All sessions were audiotaped and rated by author MPK for fidelity and for quality of MI interventions using the Global Rating of Motivational Interviewing Therapists (GROMIT) (Moyers, 2004). On the 7-point scale, session scores ranged from 5.87 to 6.93 with an average rating of 6.61 ± 0.23, which indicates that the MI techniques used in the intervention were delivered with good quality. Supervision and feedback were provided to therapists by author MPK following each intervention session. The treatment manual is available from the last author upon request.
Participants randomized to the attention-matched control condition viewed a 30-minute video about astronomy. In the control condition there was no mention of alcohol or drug use beyond completion of research assessments. Both the intervention and attention-matched control sessions took place within the UCLA Center for Cognitive Neuroscience in separate rooms from the neuroimaging suite.

**Individual Difference Measures**

The following individual questionnaires and interviews were administered during the study: (1) the 30-day timeline follow-back (TLFB) was administered in interview format to capture daily alcohol and marijuana use over the 30 days prior to the visit by trained research assistants (Sobell et al., 1988); (2) the self-report alcohol use disorders identification test (AUDIT) was administered in order to assess for drinking severity (Saunders et al., 1993); (3) the Penn Alcohol Craving Scale (PACS) to measure alcohol craving over the past week (Flannery et al., 1999). Participants also completed the Fagerstrom Test for Nicotine Dependence (Heatherton et al., 1991). Lastly, participants completed a demographics questionnaire reporting, among other variables, age, sex, and level of education.

**fMRI Paradigm**

The Alcohol Cues Task involves the delivery of oral alcohol or control (water) tastes to elicit physiological reward responses and subjective urges to drink (Filbey et al., 2008a, Filbey et al., 2008b). During the task, each trial began with the presentation of a visual cue (alcohol or water; 2 seconds) such that the words Alcohol Taste or Control Taste were visually presented to participants. This was followed by a fixation cross (jittered for an average of 3 seconds), delivery of the taste (1 mL alcohol or water; 5 seconds), and a
fixation cross (jittered using an exponential distribution with a mean of 3 seconds and a range of 0.5 to 6 seconds). Alcohol and water tastes were delivered through Teflon tubing using a computer-controlled delivery system (Infinity Controller) as described by Filbey and colleagues (Filbey et al., 2008a). Participants were instructed to press a button on a response box placed in their right hand upon swallowing. Alcohol tastes consisted of participants’ preferred alcoholic beverage (wine or liquor). Beer could not be administered due to incompatibility of the alcohol administration device with carbonated liquids. The presentation of visual stimuli and response collection were programmed using MATLAB (Mathworks, Natick, MA) and the Psychtoolbox (www.psychtoolbox.org) on an Apple MacBook running Mac OS X (Apple Computers, Cupertino, CA), and visual stimuli were presented using MRI-compatible goggles (Resonance Technologies, Van Nuys, CA). The Alcohol Cues Task was administered over the course of two runs with 50 trials/run.

fMRI Protocol

At the start of the scanning visit, participants were required to have a BrAC of 0.00 g/dL and a urine toxicology screen negative for all drugs (excluding tetrahydrocannabinol). Additionally, female participants were required to have a negative pregnancy test.

Scanning took place immediately following the brief intervention or attention matched control at the UCLA Center for Cognitive Neuroscience on a 3.0T Siemens Prisma scanner. A T2-weighted, high resolution matched-bandwith (MBW) anatomical scan (Time to Repetition (TR) = 5,000 ms, time to echo (TE) = 34 ms, flip angle = 90 degrees, voxel size: 1.5 mm x 1.5 x 4 mm, field of view (FOV) = 192 mm², 34 slices, ~1.5 minutes) and a T1-weighted magnetization-prepared rapid gradient-echo (MPRAGE) sequence (TR = 2,530 ms, TE = 1.74 ms, Time to Inversion (TI) = 1,260 ms, flip angle = 7 degrees, voxel size: 1mm³, FOV = 256 mm², ~6.2 minutes) were acquired for co-registration to the functional data. A T2*-weighted echo planar imaging scan (TR = 2,000 ms, TE = 30 ms, voxel size: 3
mm x 3 mm x 4 mm, FOV = 192 mm², 325 TRs, ~10.83 minutes/run) was acquired to examine the blood oxygen level-dependent (BOLD) signal during two runs of the Alcohol Cues Task (total time: ~22 minutes).

Preprocessing of data followed conventional procedures implemented in FMRIB’s Software Library (FSL 5.0) (www.fmrib.ox.ac.uk/fsl). This included motion correction [Motion Correction Linear Image Registration Tool (McFLIRT, Version 5.0)], high-pass temporal filtering (100 s cutoff) using FSL’s FMRI Expert Analysis Tool (FEAT, Version 5.63), and smoothing with a 5 mm full width half maximum Gaussian kernel. FSL’s Brain Extract Tool (BET) was used to remove skull and non-brain tissue from both the structural and functional scans. Data were denoised using ICA-AROMA (Pruim et al., 2015) to reduce motion artifacts associated with swallowing. Six subjects (5 in the intervention group and 1 in the control group) were excluded from further analysis due to excessive motion (exceeding 3 mm of translation) or incomplete scan data.

Data Analysis

For the intervention effect on drinking, linear mixed model analyses were conducted to test for the main effect of the intervention on the average number of drinks per week and percent of heavy drinking days in the 4 weeks post intervention. One model was run for each dependent variable. The intercept was a random effect. The models accounted for sex, smoking status and age as covariates. The intervention effect was evaluated by testing the time (baseline and follow-up)-by-condition interaction. Comparative effect size estimates for the effect of intervention on drinking outcomes were calculated based on adjusted models using $d = \frac{B_{\text{condition}} \times \text{time}}{SD_{\text{pooled baseline}}}$. In addition, the effects of neural cue-reactivity on drinking outcomes was also examined.
For the analysis of the cues task, all first-level analyses of imaging data were conducted within the context of the general linear model (FSL's FEAT), modeling the combination of the cue and taste delivery periods convolved with a double-gamma hemodynamic response function (HRF), and accounting for temporal shifts in the HRF by including the temporal derivative. Alcohol and water taste cues were modeled as separate event types. The onset of each event was set at the cue period (visual cue indicating trial type) with a duration of 11 seconds. Six motion regressors representing translational and rotational head movement were also entered as regressors of no interest. Data for each subject were registered to the MBW, followed by the MPRAGE using affine linear transformations, and then normalized to the Montreal Neurologic Institute (MNI avg152) template. Registration was further refined using FSL’s nonlinear registration tool (FNIRT).

The Alcohol Taste > Water Taste contrast was specified in the first level models. Higher-level analyses combined these contrast images within subjects (across the two task runs) and between subjects (within study conditions and across study conditions). Age, sex, cigarette smoking status, and positive urine THC were included as covariates. Additional analyses evaluated if neural response to alcohol taste cues was predictive of drinking outcomes. Two models were run, evaluating percent heavy drinking days and the average number of drinks per week in the 4 weeks following the intervention or matched-control. Both models controlled for age, sex, cigarette smoking status, positive urine THC, and baseline percent heavy drinking days or average drinks per week depending on the drinking outcome model. Z-statistic images were thresholded with cluster-based corrections for multiple comparisons based on the theory of Gaussian Random Fields with a cluster-forming threshold of $Z > 2.3$ and a corrected cluster-probability threshold of $p < 0.05$ (Worsley, 2001).
Results

Demographics Info

Forty-six individuals (intervention group = 22; control group = 24) successfully completed the scan and follow-up visits. The intervention and control groups were well matched on demographic measures including age, sex, years of education, smoking status, and cannabis use. The groups did not differ on baseline alcohol use characteristics including total AUDIT score, alcohol craving (PACS), average number of drinks consumed per week, or percent heavy drinking days (see Table 1).

Effect of Intervention on Drinking Outcomes

Overall, there was no statistically significant effect of the brief intervention on drinking outcomes as measured by the TLFB. Results from the analyses did not support an effect of the intervention relative to the control condition on changes in the frequency of heavy drinking days ($p > .4$) or on the average weekly number of drinks consumed ($p > .3$). Estimated marginal means indicated a pattern that favored the intervention in that there was a 53.3% reduction in heavy drinking days from baseline to follow up among participants in the intervention condition versus a 37.4% reduction among participants in the control condition. In terms of drinks per week the model estimated a mean reduction of 37.7% in the intervention condition versus 26.1% in the control conditions. The comparative effect size estimates for the change in alcohol use over time in the intervention versus control condition were $d = -0.182$ for percent heavy drinking days and $d = -0.203$ for average drinks per week.
Intervention Group: Neural Alcohol Cue Reactivity

The intervention group showed increased activation to alcohol taste cues compared to water taste cues in two large clusters: the first consisting of the thalamus, insula, and the putamen, and the second containing the paracingulate and middle frontal gyrus (see Table 2; Figure 2A).

Control Group: Neural Alcohol Cue Reactivity

The control group also showed increased activation in response to alcohol compared to water taste cues. The control group had increased activation in regions including the superior frontal gyrus, middle frontal gyrus, ventral tegmental area, thalamus, and insula (see Table 2; Figure 2B).

Effect of Intervention on Neural Alcohol Cue Reactivity

Across groups, exposure to alcohol taste resulted in increased activation in frontal and limbic regions, compared to water taste (see Figure 2C, Table 2). There was no significant effect of the brief interview intervention on neural alcohol cue reactivity.

Effect of Neural Cue Reactivity on Drinking Outcomes

Across groups, activation to alcohol tastes in the precuneus and medial frontal gyrus was negatively associated with percent heavy drinking days (see Figure 3, Table 3). In other words, individuals who had lower percent heavy drinking days in the weeks following the fMRI visit had greater neural reactivity to alcohol taste in the precuneus and prefrontal cortex.

Similarly, across groups, activation to alcohol tastes in the precuneus was negatively associated with average drinks per week (see Supplemental Figure S2, Table S1). That is, greater neural activity in the precuneus in response to alcohol cues was associated fewer average drinks per week at follow-up.

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Discussion

This study examined the effect of a brief intervention on drinking outcomes, neural alcohol cue-reactivity, and the ability of neural alcohol cue-reactivity to predict drinking outcomes. Results did not find an effect of the brief intervention on alcohol use in this sample, and the intervention was not associated with differential neural alcohol cue reactivity. Exploratory secondary analyses revealed inverse relationships between differential neural activity in the precuneus and medial frontal gyrus in relation to alcohol-related outcomes, but these relationships were across conditions.

The lack of main effect of intervention on either drinking outcomes or on neural alcohol cue reactivity is contrary to the study hypothesis whereby individuals assigned to the brief intervention condition were expected to show greater reductions in alcohol use compared to a no-intervention control condition (Elzerbi et al., 2015, Samson and Tanner-Smith, 2015, Tanner-Smith and Risser, 2016). In the present study, reductions in alcohol use were observed for both conditions and it appears that simply participating in an alcohol research study at an academic medical center prompted notable behavioral changes. Reductions in drinking following study participation may be attributable to assessment reactivity, in which participants curb drinking after completing alcohol-related assessments and interviews (Epstein et al., 2005). This phenomenon has been well-documented across several assessment modalities (Epstein et al., 2005, Helzer et al., 2002, Kypri et al., 2007), including the AUDIT and TLFB interviews, which were used in the present study. In addition, recent studies have highlighted the fact that single session interventions, while efficacious in relatively large RCTS, have modest effect sizes (Samson and Tanner-Smith, 2015, Huh et al., 2015). As such, the present study may have been underpowered to detect small effects sizes, which may account for the null findings regarding intervention effects on drinking outcomes. Future studies are encouraged to recruit larger samples of non-treatment seeking participants to better detect small effects. Furthermore, this finding should be considered in light of the sample, which was comprised of non-treatment seekers from the community,
which is not the typical sample evaluated in brief intervention research. However, non-treatment seeking individuals with similar alcohol use characteristics are open to participating in brief interventions (Bacio et al., 2014). Also of note the drinking outcomes in this study were evaluated using variables derived from the TLFB as the primary outcome measure. There is some evidence that some individuals under-report substance use when the TLFB is administered by an interviewer rather than a computer (Delker et al., 2016), potentially due to a social desirability bias in which participants wish to appear favorably to the interviewer. In the present study, the TLFB assessment was conducted by a trained research assistant and not the clinician who delivered the brief intervention in order to reduce this bias. However, the TLFB is a retrospective self-report measure and as such is subject to limitations including inaccuracies in participant recall. Alcohol use was also not biologically verified in this study.

In light of the null findings regarding intervention effects on drinking in this study, it is perhaps not surprising that intervention condition was not associated with differences in neural cue reactivity in this sample. While it has been argued that neuroimaging techniques may be sensitive to mechanisms of behavior change (Feldstein Ewing et al., 2011, Feldstein Ewing and Chung, 2013), in the present study, neural processing of alcohol taste cues was no more sensitive to intervention effects than traditional measures of drinking outcomes. It should be noted however, that the alcohol taste cues task used in this study was abbreviated from its original version (Filbey et al., 2008a) in order to increase the number of trials without substantially increasing scan duration. Additionally, the current version of the task used water as a control condition, while the original version (Filbey et al., 2008a) employed an appetitive control condition in the form of litchi juice. While the present version was recently validated in a separate sample (Cservenka et al., 2017), it may not have recruited the reward circuitry in response to alcohol cues as robustly as its previous iteration. Importantly, it should be noted that across both conditions, exposure to alcohol taste resulted in increased activation in frontal and limbic regions, compared to water taste, suggesting the task was
fundamentally internally valid. Nevertheless, the magnitude of the activation may have been more limited due to the combination of the shortened trial duration and use of a non-appetitive control thus hindering efforts to detect intervention effects on neural processing of alcohol cues.

Considered together, both factors likely posed significant challenges to the primary aims of the study, which fundamentally represented an interaction effect between treatment type and cue type. Given this, large magnitude main effects for both experimental factor would be optimal to bring the interaction into sharpest relief. Thus, the relatively modest effect size of the intervention and the sufficient but potentially smaller effects in the neuroimaging paradigm constrained the experimental tests. Future studies using neuroimaging to understanding brief interventions will require at least substantially larger sample sizes for a detectable clinical effect and potentially different neuroimaging paradigms.

Regarding the prediction of drinking outcomes, the most compelling finding in the present study is that activation to alcohol tastes in the precuneus and medial frontal gyrus was negatively associated with percent heavy drinking days. The effect was such that individuals who had greater neural reactivity to alcohol taste in the precuneus and prefrontal cortex had fewer percent heavy drinking days in four weeks following the fMRI scan. Likewise, across groups, activation to alcohol tastes in the precuneus was negatively associated with average drinks per week. This pattern of results suggests that greater activation of the precuneus and frontal cortex during neural processing of alcohol taste cues, compared to control cues, predicts less drinking in the subsequent month.

This effect was found across conditions, control and experimental, and is generally consistent with previous work suggesting that the precuneus is sensitive to changes in cue-reactivity and possibly to changes in addiction severity (Courtney et al., 2014). The precuneus has also been implicated in a meta-analytic review of functional neuroimaging...
studies of alcohol cue reactivity (Schacht et al., 2013a). Thus the implication of precuneus activation as a predictor of subsequent drinking in the real world extends this line of research and suggests that this region may serve as an intervention target, particularly with regard to the salience of alcohol cues. Although the vast majority of neuromodulation studies to address motivation in addiction have focused on the frontal lobes (Naish et al., 2018), and dorsolateral prefrontal cortex in particular, recent investigations have shifted attention to the precuneus (Muller et al., 2018, Koch et al., 2018), with some success.

This prospect is particularly exciting in the context of psychological interventions. The precuneus has been functionally implicated in self-related cognition (Freton et al., 2014, Shad et al., 2012, Cabanis et al., 2013, Ye et al., 2018), which in many cases is essential for behavioral interventions to have an impact. For example, in the context of a brief intervention, a person must encode the factual information provided and square it with their own self perceptions. Furthermore, in the current study’s intervention, participants were specifically asked what they wanted to do next and this necessarily demands meaningful self-related cognitive processing to generate behavior change. To illustrate this by contrast, we would have no expectation that a brief intervention would have a meaningful impact for a hypothetical individual who had no capacity to think abstractly about him or herself (in contrast to a pharmacological intervention). Thus, self-related cognition is a necessary (albeit not sufficient) elementary information processing capacity for this type of intervention to be useful and the current study suggests that the extent to which this was engaged (putatively reflected by precuneus activity) was associated with a more favorable outcome. Of course, this interpretation requires considerable caution because it is inherently conjecture and the precuneus has been implicated in a number of other cognitive functions. A recent review of psychosocial interventions for addiction medicine identified increased recruitment of self-referential processing regions, including the precuneus and medial prefrontal cortex, in response to targeted motivational interventions (Zilverstand et al., 2016). Additionally, in cannabis users, greater precuneus activation during a motivational
interviewing intervention was associated with a reduction in cannabis problems at follow-up (Feldstein Ewing et al., 2013); further indicating that activation of self-referential processing circuitry may be important for treatment response. Other psychological interventions, including cue-extinction and episodic future thinking training, may be successful at increasing self-related cognition through precuneus activation. Precuneus activation has been demonstrated in cigarette smokers who were told to engage in self-focused coping during a cue-exposure task (Wilson et al., 2013), indicating the interventions targeting self-focused coping during exposure to drug cues may effectively activate this brain region. Exposure to episodic future thinking activates the precuneus and mPFC (Hu et al., 2016) and results in alcohol dependent individuals increasing their valuation of future monetary rewards while lowering demand intensity for alcohol rewards (Snider et al., 2016). Frontoparietal circuitry, including the precuneus, is activated when participants make voluntary choices to cognitively reappraise craving responses or freely view craving cues (Cosme et al., 2018). Of note, the precuneus is not neuroanatomically uniform, with distinct functional subregions according to both the anterior-posterior and dorsal-ventral axes, and distinct patterns of functional connectivity by subregion (Zhang and Li, 2012). The current study reveals associations for the precuneus in general, but cannot speak to subregional activation.

In sum, the current study sought to examine whether a brief intervention would reduce both drinking and alcohol motivation as measured by neural reactivity to alcohol cues and neither hypothesis was supported. This conclusion, however, must be tempered by effect size considerations for both the intervention and the paradigm, as well as the apparently substantial reactivity effects present in the control condition. Each of these has important methodological implications for future studies of the neural mechanisms of alcohol-related...
behavior change. In addition, independent of intervention, exploratory analyses revealed differential neural reactivity that predicted more favorable outcomes, particularly in the precuneus, suggesting that is a promising neural substrate warranting further study in this line of inquiry.

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Figure Legends

Figure 1.

CONSORT Diagram for the trial.

Figure 2.

Brain activation to alcohol taste compared to water taste cues. A. The intervention group showed increased activation to alcohol taste cues in limbic and frontal regions. B. The control group also displayed increased activation to alcohol taste cues in frontal, limbic, and insula regions. C. Across groups, there was increased brain activation in frontal, limbic, and insula regions during alcohol taste cues compared to water taste cues. See Table 2 for full list of regions activated in this contrast. Z-statistic maps are whole-brain cluster corrected, $Z > 2.3$, $p = 0.05$. Coordinates are in MNI space. Brain is displayed in radiological convention (L=R).

Figure 3.

Brain activation to alcohol taste cues in the precuneus and prefrontal cortex was significantly associated with decreased percent heavy drinking days in the 4 weeks following the fMRI. Z-statistic maps are whole-brain cluster corrected, $Z > 2.3$, $p = 0.05$. Coordinates are in MNI space. Brain is displayed in radiological convention (L=R).
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<thead>
<tr>
<th>Characteristic</th>
<th>Intervention Group (n = 22)</th>
<th>Control Group (n = 24)</th>
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<td>Sex (m/f)</td>
<td>13/9</td>
<td>15/9</td>
<td>$\chi^2 = 0.06$</td>
<td>0.81</td>
</tr>
<tr>
<td>Smokers (n)</td>
<td>11</td>
<td>12</td>
<td>$\chi^2 = 0.00$</td>
<td>1</td>
</tr>
<tr>
<td>Education (years)</td>
<td>15.45 ± 2.13</td>
<td>15.04 ± 1.78</td>
<td>t = 0.72</td>
<td>0.48</td>
</tr>
<tr>
<td>AUDIT Total Score</td>
<td>17.68 ± 6.49</td>
<td>17.17 ± 7.61</td>
<td>t = 0.25</td>
<td>0.81</td>
</tr>
<tr>
<td>PACS Score</td>
<td>19.32 ± 6.94</td>
<td>18.79 ± 7.15</td>
<td>t = 0.25</td>
<td>0.80</td>
</tr>
<tr>
<td><strong>Baseline</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Average number of drinks/week (TLFB)</td>
<td>24.40 ± 17.62</td>
<td>20.77 ± 11.52</td>
<td>t = 0.83</td>
<td>0.41</td>
</tr>
<tr>
<td>PHDD (TLFB)</td>
<td>37.73 ± 27.15</td>
<td>35.00 ± 22.93</td>
<td>t = 0.37</td>
<td>0.71</td>
</tr>
<tr>
<td>THC Positive (n)</td>
<td>6</td>
<td>6</td>
<td>$\chi^2 = 0.04$</td>
<td>0.86</td>
</tr>
<tr>
<td>THC total number days used (MTLFB)</td>
<td>3.50 ± 7.04</td>
<td>1.79 ± 3.46</td>
<td>t = 1.03</td>
<td>0.31</td>
</tr>
<tr>
<td><strong>Follow-Up</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Average number of drinks/week (ATLFB)</td>
<td>15.48 ± 12.11</td>
<td>14.84 ± 9.83</td>
<td>t = 0.56</td>
<td>0.84</td>
</tr>
<tr>
<td>PHDD (ATLFB)</td>
<td>18.56 ± 19.30</td>
<td>21.61 ± 21.58</td>
<td>t = 0.50</td>
<td>0.62</td>
</tr>
<tr>
<td>THC Positive (n)</td>
<td>1</td>
<td>3</td>
<td>$\chi^2 = 0.92$</td>
<td>0.34</td>
</tr>
<tr>
<td>THC total number days used (MTLFB)</td>
<td>1.32 ± 4.81</td>
<td>2.92 ± 6.44</td>
<td>t = 0.93</td>
<td>0.36</td>
</tr>
</tbody>
</table>

AUDIT – Alcohol Use Disorder Identification Test; PACS – Penn Alcohol Craving Scale; ATLFB – Alcohol Timeline Followback; THC – tetrahydrocannabinol; MTLFB – Marijuana Timeline Followback; PHDD – Percent Heavy Drinking Days
Table 2. Whole-brain activation to alcohol taste cues vs. water taste cues by group.

<table>
<thead>
<tr>
<th>Brain Region</th>
<th>Cluster Voxels</th>
<th>Max. Z</th>
<th>x</th>
<th>y</th>
<th>z</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Intervention Group</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>R Thalamus</td>
<td>1,700</td>
<td>4.18</td>
<td>20</td>
<td>-20</td>
<td>-4</td>
</tr>
<tr>
<td>R Middle Temporal Gyrus</td>
<td>3.27</td>
<td>62</td>
<td>-18</td>
<td>-18</td>
<td></td>
</tr>
<tr>
<td>R Parahippocampal Gyrus</td>
<td>2.80</td>
<td>20</td>
<td>-14</td>
<td>-26</td>
<td></td>
</tr>
<tr>
<td>R Hippocampus</td>
<td>2.71</td>
<td>32</td>
<td>-26</td>
<td>-8</td>
<td></td>
</tr>
<tr>
<td>R Putamen</td>
<td>2.65</td>
<td>34</td>
<td>-6</td>
<td>-10</td>
<td></td>
</tr>
<tr>
<td>R Insula</td>
<td>2.61</td>
<td>42</td>
<td>6</td>
<td>-6</td>
<td></td>
</tr>
<tr>
<td>R/L Paracingulate Gyrus</td>
<td>1,199</td>
<td>3.95</td>
<td>0</td>
<td>36</td>
<td>32</td>
</tr>
<tr>
<td>L Middle Frontal Gyrus</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Control Group</strong></td>
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<td></td>
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<td></td>
<td></td>
</tr>
<tr>
<td>L Superior Frontal Gyrus</td>
<td>3,395</td>
<td>4.17</td>
<td>-14</td>
<td>8</td>
<td>62</td>
</tr>
<tr>
<td>R/L Paracingulate Gyrus</td>
<td>3.18</td>
<td>0</td>
<td>36</td>
<td>32</td>
<td></td>
</tr>
<tr>
<td>L Middle Frontal Gyrus</td>
<td>3.13</td>
<td>-54</td>
<td>14</td>
<td>32</td>
<td></td>
</tr>
<tr>
<td>R/L Ventral Tegmental Area</td>
<td>1,497</td>
<td>3.93</td>
<td>0</td>
<td>-20</td>
<td>-20</td>
</tr>
<tr>
<td>R/L Thalamus</td>
<td>2.97</td>
<td>0</td>
<td>-18</td>
<td>8</td>
<td></td>
</tr>
<tr>
<td>R Parahippocampal Gyrus</td>
<td>2.78</td>
<td>28</td>
<td>-30</td>
<td>-16</td>
<td></td>
</tr>
<tr>
<td>R Insula</td>
<td>1,436</td>
<td>4.74</td>
<td>44</td>
<td>-20</td>
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</tr>
<tr>
<td>R Middle Temporal Gyrus</td>
<td>3.54</td>
<td>60</td>
<td>-4</td>
<td>-16</td>
<td></td>
</tr>
<tr>
<td>R Hippocampus</td>
<td>2.52</td>
<td>28</td>
<td>-16</td>
<td>-14</td>
<td></td>
</tr>
<tr>
<td><strong>Combined Intervention and Control Group</strong></td>
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<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>L Superior Frontal Gyrus</td>
<td>3,691</td>
<td>4.51</td>
<td>-14</td>
<td>10</td>
<td>58</td>
</tr>
<tr>
<td>R/L Paracingulate Gyrus</td>
<td>3.26</td>
<td>-2</td>
<td>36</td>
<td>34</td>
<td></td>
</tr>
<tr>
<td>L Precentral Gyrus</td>
<td>3.02</td>
<td>-42</td>
<td>-2</td>
<td>58</td>
<td></td>
</tr>
<tr>
<td>L Middle Frontal Gyrus</td>
<td>2.91</td>
<td>-48</td>
<td>14</td>
<td>34</td>
<td></td>
</tr>
<tr>
<td>R Thalamus</td>
<td>3,380</td>
<td>4.16</td>
<td>20</td>
<td>-26</td>
<td>-2</td>
</tr>
<tr>
<td>R Middle Temporal Gyrus</td>
<td>3.62</td>
<td>62</td>
<td>-18</td>
<td>-18</td>
<td></td>
</tr>
<tr>
<td>R/L Ventral Tegmental Area</td>
<td>3.28</td>
<td>0</td>
<td>-16</td>
<td>-14</td>
<td></td>
</tr>
<tr>
<td>R Parahippocampal Gyrus</td>
<td>3.08</td>
<td>16</td>
<td>-14</td>
<td>-24</td>
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</tr>
<tr>
<td>R Insula</td>
<td>3.04</td>
<td>40</td>
<td>-16</td>
<td>4</td>
<td></td>
</tr>
<tr>
<td>R Pallidum</td>
<td>2.91</td>
<td>24</td>
<td>-12</td>
<td>-6</td>
<td></td>
</tr>
<tr>
<td>R Hippocampus</td>
<td>2.79</td>
<td>30</td>
<td>-14</td>
<td>-14</td>
<td></td>
</tr>
</tbody>
</table>

**Intervention > Control Group**

N/A

**Control > Intervention Group**

N/A

R – Right; L - Left
Table 3. Whole-brain activation to alcohol taste cues negatively correlated with PHDD across groups.

<table>
<thead>
<tr>
<th>Brain Region</th>
<th>Cluster Voxels</th>
<th>Max. Z</th>
<th>x</th>
<th>y</th>
<th>z</th>
</tr>
</thead>
<tbody>
<tr>
<td>R/L Precuneus</td>
<td>2,281</td>
<td>3.85</td>
<td>14</td>
<td>-56</td>
<td>-26</td>
</tr>
<tr>
<td>L Posterior Cingulate Gyrus</td>
<td>3.05</td>
<td>-2</td>
<td>-48</td>
<td>8</td>
<td></td>
</tr>
<tr>
<td>L Medial Frontal Gyrus</td>
<td>1,417</td>
<td>3.87</td>
<td>-6</td>
<td>52</td>
<td>-2</td>
</tr>
<tr>
<td>R/L Anterior Cingulate Gyrus</td>
<td>3.15</td>
<td>0</td>
<td>42</td>
<td>0</td>
<td></td>
</tr>
<tr>
<td>R Superior Frontal Gyrus</td>
<td>3.03</td>
<td>10</td>
<td>52</td>
<td>22</td>
<td></td>
</tr>
</tbody>
</table>

R – Right; L - Left