**TITLE:**

Disruption of Frontal Lobe Neural Synchrony by Alcohol Intoxication

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**KEYWORDS:**

Neuroscience, brain, cognitive control, magnetoencephalography, theta oscillations, phase-locking, neural synchrony, alcohol, Stroop task, multimodal imaging

**SUMMARY:**

This experiment uses an anatomically-constrained magnetoencephalography (aMEG) method to examine brain oscillatory dynamics and long-range functional synchrony during engagement of cognitive control as a function of acute alcohol intoxication.

**ABSTRACT:**

Decision making relies on dynamic interactions of distributed, primarily frontal brain regions. Extensive evidence from functional magnetic resonance imaging (fMRI) studies indicates that the anterior cingulate (ACC) and the lateral prefrontal cortices (latPFC) are essential nodes subserving cognitive control. However, because of its limited temporal resolution, fMRI cannot accurately reflect the timing and nature of their presumed interplay. The present study combines distributed source modeling of the temporally precise magnetoencephalography (MEG) signal with structural MRI in the form of “brain movies” to: (1) estimate the cortical areas involved in cognitive control (“where”), (2) characterize their temporal sequence (“when”), and (3) quantify the oscillatory dynamics of their neural interactions in real time. Stroop interference was associated with greater event-related theta (4 - 7 Hz) power in the ACC during conflict detection followed by sustained sensitivity to cognitive demands in the ACC and latPFC during integration and response preparation. A phase-locking analysis revealed co-oscillatory interactions between these areas indicating their increased neural synchrony in theta band during conflict-inducing incongruous trials. These results confirm that theta oscillations are fundamental to long-range synchronization needed for integrating top-down influences during cognitive control. MEG reflects neural activity directly, which makes it suitable for pharmacological manipulations in contrast to fMRI that is sensitive to vasoactive confounds. In the present study, healthy social drinkers were given a moderate alcohol dose and placebo in a within-subject design. Acute intoxication attenuated theta power to Stroop conflict and dysregulated co-oscillations between the ACC and latPFC, confirming that alcohol is detrimental to neural synchrony subserving cognitive control. It interferes with goal-directed behavior that may result in deficient self-control, contributing to compulsive drinking. In sum, this method can provide insight into real-time interactions during cognitive processing and can characterize the selective sensitivity to pharmacological challenge across relevant neural networks.

**INTRODUCTION:**

The overall goal of this study is to examine the effects of acute alcohol intoxication on spatio-temporal changes in the brain oscillatory dynamics and long-range functional integration during cognitive control. The employed multimodal imaging approach combines magnetoencephalography (MEG) and structural magnetic resonance imaging (MRI) to provide insight into the neural basis of decision making with high temporal precision and at the level of an interactive system.

Flexible behavior makes it possible to adapt to changing contextual demands and to switch strategically between different tasks and requirements in agreement with one’s intents and goals. The capacity to suppress automatic responses in favor of goal-relevant but non-habitual actions is an essential aspect of cognitive control. Extensive evidence suggests that it is subserved by a predominantly frontal cortical network, with the anterior cingulate cortex (ACC) as a central node in this interactive network1-4. While the abundant anatomical connectivity between the ACC and lateral frontal cortices is well-described5,6, the functional characteristics of communication between these regions during cognitive control, response selection and execution, are poorly understood.

The highly influential conflict monitoring theory7,8 proposes that cognitive control arises from a dynamic interaction between the medial and lateral prefrontal cortices. This account purports that the ACC monitors conflict between competing representations and engages the lateral prefrontal cortex (latPFC) to implement response control and optimize performance. However, this account is primarily based on the functional MRI (fMRI) studies using the blood oxygenation level dependent (BOLD) signal. The fMRI-BOLD signal is an excellent spatial mapping tool, but its temporal resolution is limited because it reflects regional hemodynamic changes mediated by neurovascular coupling. As a result, the BOLD signal changes unfold on a much slower time scale (in seconds) than the underlying neural events (in milliseconds)9. Moreover, the BOLD signal is sensitive to alcohol’s vasoactive effects10 and may not accurately represent the magnitude of neural changes, which makes it less suitable for studies of acute alcohol intoxication. Therefore, the presumed interplay between the medial and lateral prefrontal cortices and its sensitivity to alcohol intoxication need to be examined by methods that record neural events in a temporally precise manner. MEG has an excellent temporal resolution since it directly reflects postsynaptic currents. The anatomically-constrained MEG (aMEG) methodology employed here is a multimodal approach that combines distributed source modeling of the MEG signal with structural MRI. It allows for the estimation of *where* the conflict- and beverage-related brain oscillatory changes are occurring and to understand the temporal sequence *("when")* of the involved neural components.

Decision making relies on the interactions of distributed brain regions that are dynamically engaged to deal with increased demands on cognitive control. One way to estimate event-related changes in long-range synchrony between two cortical regions is to calculate their phase coupling as an index of their co-oscillations11,12. The present study applied a phase-locking analysis to test the basic tenet of the conflict monitoring theory by examining the co-oscillatory interactions between the ACC and latPFC. Neural oscillations in theta range (4 - 7 Hz) are associated with cognitive control and have been proposed as a fundamental mechanism supporting the long-range synchronization needed for top-down cognitive processing13-16. They are generated in prefrontal areas as a function of task difficulty and are significantly attenuated by acute alcohol intoxication17-20.

Long-term excessive alcohol intake is associated with a range of cognitive deficits with prefrontal circuitry being especially affected21,22. Acute alcohol intoxication is detrimental to cognitive control under conditions of increased difficulty, ambiguity, or those that induce response incompatibility17,23,24. By affecting decision making, alcohol may interfere with goal-directed behavior, may result in poor self-control and increased drinking, and may also contribute to traffic- or work-related hazards25-27. The present study uses an aMEG approach to measure the oscillatory activity in theta band and synchrony between the principal executive areas with excellent temporal resolution. The effects of alcohol on theta activity and co-oscillations between the ACC and the latPFC are examined as a function of conflict elicited by the Stroop interference task. We hypothesize that increased cognitive demands are associated with greater functional synchrony and that alcohol-induced dysregulation of synchronous activity of the medial and lateral prefrontal cortices underlies impairments in cognitive control.

**PROTOCOL:**

This experimental protocol has been approved by the Human Subjects Protection Committee at the University of California, San Diego.

# Human Subjects

## Recruit healthy right-handed adult volunteers, obtain their consent, and screen them on the inclusion/exclusion criteria.

## NOTE: In this study, twenty young, healthy individuals (mean ± standard deviation [SD] age = 25.3 ± 4.4 years) including 8 women were recruited who drink in moderation, who have never been in treatment or arrested for drug or alcohol related offenses, who report no alcoholism-related symptoms on the Short Michigan Alcoholism Screening Test28, who do not smoke nor use illegal substances, who do not have a history of neuropsychiatric disorders or any current health problems, and who are medication free and have no internal ferromagnetic objects or implants.

# Experimental Design

## Scan each participant four times, including three MEG sessions (a no-beverage introductory session and two experimental beverage sessions in which alcohol and placebo are administered in a counterbalanced manner), and one structural MRI scan.

NOTE: In this within-subject design, participants serve as their own controls by participating in both alcohol and placebo sessions. This design reduces error variance and increases statistical power by minimizing influence of individual variability in brain anatomy, activity patterns, and alcohol metabolism.

1. **Collecting MEG Scans**
   1. Perform familiarization session.

## During the initial introductory session, administer questionnaires to obtain more information about the participants' medical history, their drinking patterns and severity of alcoholism-related symptoms28,29, family history of alcoholism30, and personality traits including impulsivity31,32.

* + 1. Carry out an initial recording in the MEG scanner following the protocol described below in steps 3.2, 3.3, and 3.5. Do not provide any beverage. Explain the task and run the practice version allowing participants to get familiarized with it beforehand.

NOTE: The acclimation to the experimental situation serves the purpose of minimizing potential effects of situation-induced arousal33, thereby equating subsequent alcohol and placebo sessions on that dimension.

# Perform the alcohol/placebo experimental sessions.

NOTE: Follow the same experimental procedures during both alcohol and placebo sessions with the exception of the administered beverage. Counterbalance beverage order by administering alcohol beverage first to one half of participants and placebo to the other half in a random order.

* + 1. Upon their arrival to the MEG lab, run a brief test scan by putting the participant in the scanner and checking the channels for possible magnetization. Measure their weight. Screen them with an electronic breathalyzer. Query them about compliance with the requirements to abstain from alcohol for 48 h and from food for 3 h prior to the experiment.
    2. Collect urine samples for a multi-drug test panel from all participants and exclude those who test positive for any drug. In addition, check female participants for pregnancy with a urine test and exclude those who test positive or if they suspect that they might be pregnant.
    3. Assess dynamic changes in the subjective effects of alcohol by asking participants to rate their momentary feelings and states on a standardized scale34 prior to drinking and on two additional occasions during the experiment - on the ascending limb (~15 min after consuming beverage) and descending limb of the breath alcohol concentration curve (BrAC), after the MEG recording.
    4. Administer a practice run of the Stroop task on a laptop with stimulus presentation software to ensure that the participants understand the task before recording.

NOTE: This version of the Stroop task combines reading and color naming (**Figure 1**). The congruent condition consists of color words (*i.e.*, red, green, blue, yellow) that are printed in the matching font color (*i.e.*, the word "green" is printed in green). In the incongruous condition, color words are printed in color that does not match their meaning (*i.e.*, the word "green" is printed in yellow). Ask the participants to press one of four buttons corresponding to the font color whenever a word is written in color, or, when a word is written in gray, to press a button corresponding to the meaning of the word18,23.

* 1. Prepare the magnetoencephalography/electroencephalographic (MEG/EEG) recording.

NOTE: Details of MEG data acquisition have been described in previous publications35-37.

* + 1. Position the EEG cap or individual EEG electrodes on the head of the participant and check that all impedances are below 5 kΩ.
    2. Attach the head position indicator (HPI) coils on either side of the forehead and behind each ear.

NOTE: This step is specific to Neuromag systems.

* + 1. Digitize positions of the fiducial points including the nasion and two preauricular points, positions of HPI coils, EEG electrodes, and obtain a large number of additional points (~200) delineating the head shape. Use this information for the co-registration with anatomical MRI images (**Figure 2**).
  1. Administer beverage.
     1. Prepare alcohol beverage by mixing premium quality vodka with chilled orange juice (25% v/v), based on each participant’s gender and weight (0.60 g/kg alcohol for men, 0.55 g/kg alcohol for women), targeting a BrAC of 0.06%38. Serve the same volume of orange juice in glasses with rims swabbed with vodka as a placebo beverage. Ask the participant to consume the beverage in approximately 10 min.
     2. Check the participants' BrAC with the breathalyzer starting at ~15 min after drinking and then every 5 min until they enter the recording chamber. Since electronic devices cannot be used in the shielded room, use a saliva alcohol test, which consists of a cotton swab that is saturated in saliva and is inserted into a receptacle that provides a readout.
  2. Acquire MEG/EEG data.
     1. Position the participant comfortably in the scanner. Since the prefrontal activity is of particular interest, ensure that the participant is positioned so that his/her head is touching the top of the helmet and is aligned along the front.

NOTE: Head position can affect activity estimates in significant ways because the magnetic field gradients decrease with the cube of the distance between the sensors and the brain sources39.

* + 1. Connect HPI coils and all of the electrodes to their respective inputs on the scanner. Position response pads so that the buttons can be pressed comfortably. Ascertain that the font is clearly legible on the projection screen in front of the participant.
    2. Back in the console room, check that the intercom is functioning properly. Remind the participant to minimize blinking and to avoid movements including head motion caused by talking. Instruct the participant to reply to questions by pressing response buttons instead.
    3. Check that all response and stimulus triggers are recorded correctly. Examine all channels for artifacts and measure the head position in the scanner.
    4. Start data acquisition and begin the task. Give breaks every ~2.5 min to rest the eyes. Save the data upon task completion and escort the participant out of the recording chamber.
    5. When the participant has exited the scanner, acquire approximately two minutes of data from the empty room as a measure of instrumental noise.
    6. Ask the participant to rate perceived task difficulty, content of the imbibed beverage, how intoxicated they felt, as well as their momentary moods and feelings34.

1. **Image Acquisition and Cortical Reconstruction of Structural MRI** 
   1. Obtain a high-resolution anatomical MRI scan for each participant, and reconstruct each participant's cortical surface with FreeSurfer software40-42.
   2. Use the inner skull surface derived from the segmented structural MRI images to generate a boundary element model of the volume conductor, which is used to provide a model for the forward solution that is consistent with each individual’s brain anatomy43,44.
2. **MEG Data Analysis**

NOTE: Analyze the data with the anatomically-constrained MEG approach which uses each participant’s reconstructed cortical surface to constrain source estimates to the cortical ribbon40,45,46. The analysis stream relies on custom functions with dependencies on publicly available packages including FieldTrip47, EEGLab48, and MNE49.

* 1. During data preprocessing, use a permissive band-pass filter (*e.g.*,0.1 - 100 Hz) and epoch data into segments that include padding intervals on each end (*e.g.*,-600 to 1100 ms for an interval of interest that spans -300 to 800 ms after stimulus onset).
  2. Remove noisy and flat channels, as well as trials containing artifacts by visual inspection and using threshold-based rejection. Use independent component analysis48 to remove eyeblink and heartbeat artifacts. Eliminate trials with incorrect responses.
  3. Apply Morlet wavelets (**Figure 3**)47 to calculate complex power spectrum for each trial in 1 Hz increments for theta frequency band (4 - 7 Hz). Remove any additional artifacts. Compute the noise covariance from empty room data.
  4. Co-register the MEG data with MRI images using the three-dimensional (3D) head digitization information (**Figure 2**).
     1. Open the MRIlab module.
     2. Select **File| Open| Select subject’s structural MRI**.
     3. Select **File| Import| Isotrak data| select raw data.fif file| Make Points**.
     4. Select **Windows| Landmarks| Adjust fiducial landmarks** until co-registration of MEG data and MRI are acceptable.
     5. Select **File| Save**.
  5. Calculate noise-sensitivity normalized estimates of theta source power and phase with a spectral dynamic statistical mapping approach18,50. Express event-related theta source power as percent signal change relative to baseline.
  6. Create group averages of event-related theta source power by morphing each participant's estimates onto an average cortical representation51.
  7. Visualize the source estimates on an inflated average surface to enhance visibility of sulcal estimates (**Figure 4**).
     1. Open the MNE software.
     2. Select **File|** **Load Surface**| **Load inflated group-average FreeSurfer cortical surface**.
     3. Select **File|** **Manage overlays|** **Load stc| Load group-averaged data| Select loaded file from available overlays**.
     4. Select overlay type as **Other**.
     5. Adjust **Color Scale thresholding| Show**.
     6. View **brain movies** and examine spatio-temporal stages of processing by identifying areas and time windows characterized by highest activation.
  8. Create unbiased regions of interest (ROIs) based on overall group-averaged estimates to incorporate cortical locations with most notable source power. Calculate time courses for each subject, condition, and ROI (**Figure 5**).
  9. Submit the obtained theta source power estimates to the statistical analysis.
     1. Extract time windows of interest from each ROI time course and perform analysis of variance (ANOVA) with beverage (alcohol, placebo) and trial type (congruent, incongruous) as within subject factors. Use a nonparametric cluster-based permutation test52 to examine beverage and condition comparisons of event-related theta power as well as phase-locking value (PLV).
  10. Estimate task-related changes in the long-range synchronization between the main activation foci in the ACC and the latPFC by computing the PLV12. Express PLV as percent change relative to baseline.

NOTE: The PLV is an indicator of consistency of the phase angle between the two ROIs across trials as it measures the extent to which they co-oscillate at a particular frequency and in real time (**Movie 1**).

* 1. Calculate correlations between ROI MEG activity estimates, indices of behavioral performance, and questionnaire scores to inform interpretation of the observed results.

**REPRESENTATIVE RESULTS:**

Behavioral results indicate that the Stroop task successfully manipulated response interference because the accuracy was the lowest and the response times the longest on incongruous trials (**Figure 6**). Alcohol intoxication lowered accuracy but did not affect reaction times18.

The spatio-temporal sequence of activity in theta frequency band revealed with the aMEG approach is overall in agreement with generally accepted models of cognitive functions in this type of task. As illustrated in the brain movies (**Movie 2**), the visual cortex is activated at around 100 ms after stimulus onset, followed by a posterior-to-anterior activation pattern that engages primarily frontal cortices during cognitive integration stages after ~300 ms. The ACC is particularly sensitive to incongruous, high-conflict trials, indicating its engagement during conflict monitoring. The ACC is the principal generator of theta oscillations during tasks probing cognitive control but the latPFC is also active during the integration stage at around 350-600 ms. Activation of the motor cortex is visible after ~600 ms during the response preparation stage (**Movie 2B**). Event-related theta power is greatest on incongruous (INC) trials, which is consistent with its sensitivity to conflict demands (**Figure 5**), especially in the prefrontal cortex13,17,19,20. Theta power is decreased by acute alcohol intoxication overall. However, when compared to congruous (CONG) trials, alcohol decreases theta power on incongruous (high conflict) trials selectively in the ACC and latPFC18.

The present study extends the results from Kovacevic *et al.*18 by focusing on dynamic interactions between these areas during the processing of Stroop interference in light of a prevailing account of the cognitive control network7,8. To better understand the timing, degree, and nature of the interactions between these two principally engaged cortical areas, the PLV was calculated for each beverage and task condition, and for each participant. As shown in a group average in **Figure 7**, co-oscillations between the ACC and latPFC vary across time with an overall early increase in co-oscillations during a stimulus processing stage. Under placebo, this is followed by a sustained increase after ~400 ms on incongruous trials during the integration and response preparation stage. Thus, synchronized co-oscillations between the medial and lateral prefrontal cortices are observed only on the more difficult, incongruous trials evoking response conflict *F*(1,19) = 5.5, *p* < 0.05. This evidence supports the proposal that the ACC and the latPFC functionally interact in real time to subserve cognitive control. In contrast, acute alcohol intoxication significantly dysregulates the co-oscillations, yielding a Condition x Beverage interaction, *F*(1,19) = 5.1, *p* < 0.05, in which incongruous trials specifically were affected by alcohol *F*(1,19) = 8.8, *p* < 0.01 (**Figure 7**). This may underlie alcohol-induced impairments of inhibitory control and indicates the vulnerability of top-down regulative functions of the prefrontal cortex to acute intoxication.

**FIGURE AND TABLE LEGENDS:**

**Figure 1: Stroop task combines color naming and reading.** Trial examples for each of the three conditions along with the correct response color are presented. In the congruent condition (CONG), font color is consistent with the word meaning, while incongruous trials (INC) elicit response conflict due to interference from word meaning. Participants are instructed to press a button corresponding to the font color when words are written in color (CONG, INC) and to respond to the word meaning (Read) when they are written in gray. Trials are presented for 300 ms and then replaced by a fixation screen for 1700 ms. Trial types are presented in a randomized order. In this particular version, the CONG and INC conditions were equiprobable and were presented on 16.7% trials each out of 576 trials total.

**Figure 2: Co-registration of MEG and MRI.** Digitized points across the head collected during the MEG recording are used for co-registration with anatomical MRI images.

**Figure 3:** **Morlet wavelet.** Morlet wavelets are used to calculate complex power spectrum for each trial in 1 Hz frequency increments for the theta band frequency (4 - 7 Hz).

**Figure 4: Cortical reconstruction and inflation.** Individual cortical surfaces are reconstructed and are used to constrain estimated source power. Here shown is an average cortical surface which is inflated to enhance visibility of the sources estimated to cortical sulci.

**Figure 5: Group-average time courses of event-related theta source power estimates in selected regions of interest.** Incongruous stimuli (INC) elicited increased event-related theta power compared to congruent stimuli (CONG) in the anterior cingulate cortex (ACC; *F*(1,19) = 34.1, *p* < 0.0001) as well as the lateral prefrontal cortex (latPFC; *F*(1,19) = 11.0, *p* < 0.01), during 480 - 670 ms. Conflict processing is particularly sensitive to alcohol intoxication as theta power to INC was attenuated by alcohol intoxication (*F*(1,19) = 9.9, *p* < 0.01). The y-axis depicts baseline-corrected noise-normalized event-related theta source power. This figure has been modified from Kovacevic *et al.*18.

**Figure 6: Behavioral results on the Stroop task.** Stroop interference was reflected in decreased accuracy and longer response times to incongruous (INC) trials. Alcohol intoxication (Alc) impaired accuracy compared to placebo (Plac) but did not affect reaction times. Error bars signify standard error of the mean. This figure has been modified from Kovacevic *et al.*18.

**Figure 7: Group-average time courses of phase-locking values (PLVs) in the theta band.** Co-oscillatory synchrony between the anterior cingulate cortex (ACC) and lateral prefrontal cortex (latPFC) in theta band expressed as percent change from baseline for placebo (left) and alcohol (right) conditions. Following an early increase in PLVs during a stimulus processing stage (400 - 600 ms), a sustained increase in co-oscillations is observed on incongruous trials (INC) in response to increased cognitive control compared to congruent trials (CONG) under placebo, *F*(1,19) = 5.5, *p* < 0.05. Acute alcohol intoxication selectively dysregulated the co-oscillations on INC trials, *F*(1,19) = 8.8, *p* < 0.01. Activation maps (inset) show the incongruity effect (INC-CONG), which is prominent in the ACC and latPFC. The color scale denotes baseline-corrected source power estimates at 480 ms after stimulus onset, with red (activity > 0.2) to yellow (activity > 0.3) indicating stronger theta power to INC trials compared to CONG trials.

**Movie 1: Co-oscillations.** Phase-locking values were calculated in the theta frequency range (4 - 7 Hz) between the anterior cingulate cortex (ACC) and lateral prefrontal cortex (latPFC) as a measure of synchronization that is sensitive to the consistency of the phase difference between these two ROIs regardless of their theta power amplitude.

**Movie 2: Brain movies.** Distributed source modeling of the MEG signal combined with structural MRI allows for the estimation of principal cortical areas generating theta power and the temporal sequence of their activation in response to the Stroop interference. (**A**) Following early sensory processing, the anterior cingulate cortex (ACC) is selectively activated by incongruous, high-conflict trials after ~350 ms. (**B**) While the ACC is the principal generator of theta oscillations during tasks probing cognitive control, the lateral prefrontal cortex (latPFC) is also engaged during the integration stage around 350 - 600 ms. Activation of the motor cortex is observed after ~600 ms during response preparation. The color scale denotes differential baseline-corrected source power estimates, with red color indicating activation greater than 0.79 medially (0.57 laterally) and yellow indicates activation greater than 0.9 medially (0.8 laterally). Please note that these two movies should be shown together with unfolding time courses pertaining to the ACC and latPFC, respectively.

**DISCUSSION:**

The multimodal imaging method used in this study comprises distributed source modeling of the temporally precise MEG signal along with spatial constraints of inverse estimates derived from each participant’s structural MRI. The aMEG approach combines the strengths of these techniques to provide insight into the spatio-temporal stages of oscillatory dynamics and the long-range integration subserving cognitive control. This method provides greater temporal precision than other neuroimaging techniques such as fMRI-BOLD whose temporal resolution is on the magnitude of seconds due to its indirect sensitivity to neural changes via neurovascular coupling9. In comparison, the millisecond precision of MEG signal allows for the investigation of neural processing stages, as demonstrated by the present study. The aMEG model assumes distributed sources of the MEG signal along the cortical surface which, when reconstructed from structural MRI images, provides spatial constraints for activity estimates45,53. These spatial estimates can be used to investigate not only local activation but long-range communication at an interactive network level in the form of phase-locking16,20. Moreover, the aMEG approach is well suited for investigating the effects of pharmacological manipulation on neural functions, given that the fMRI-BOLD signal is confounded by the vasoactive effects of pharmacological manipulations such as alcohol and may not accurately reflect the magnitude of neural changes10.

The high sensitivity of this method to minute neural changes means that it is also sensitive to non-neural noise including muscle movements or eye blinks, so the various artifacts need to be detected and carefully removed from the raw signal. Moreover, head position can have significant effects on activity estimates due to sensor sensitivity to magnetic field gradients39. Given the assumptions of the aMEG model, source estimates are constrained to the cortical surface45,46, so the activity elicited from subcortical structures cannot be estimated.

Based on previously published results18, the present study has illustrated changes in event-related theta (4 - 7 Hz) power during Stroop-induced conflict as a function of acute alcohol intoxication in healthy social drinkers. As shown in **Figure 5**, theta power is differentially sensitive to cognitive demands imposed by the Stroop task conditions. Incongruity is especially effective in engaging cognitive control as reflected in greater theta power in the prefrontal cortex compared to prestimulus baseline. The principal estimated generator of theta oscillations is the ACC that is sensitive to response conflict during both early and late processing stages18. These findings support the role of the ACC in monitoring for conflict in concordance with prominent accounts7,8. Thus, the aMEG method has provided a temporally-sensitive insight into the sustained engagement of the ACC during trials imposing higher load on cognitive control. Together with extensive anatomical connections between the ACC and distributed brain regions5,6, this evidence corroborates its multifaceted role in self-regulation. On that view, the ACC is a key hub in the neurofunctional system that subserves cognitive control by aligning goals and intentions with contextual and motivational constraints54,55. Inferolateral prefrontal cortex, especially on the right, is another important area within that system which has been associated with inhibition of prepotent responses, attentional control, and working memory in the service of updating task representations56-58.

It has been established that theta oscillations mediate neural integration necessary for cognitive and affective processing13,16,59,60. Neural communication may thus rely on synchronized excitability of distant neuronal ensembles in theta band with nested fast rhythms mediating local processing61,62. PLVs reflect phase consistency between cortical areas and are commonly used to estimate their oscillatory synchrony as it is assumed that two areas interact when they co-oscillate63. Indeed, transient increases in PLV are observed in those intervals of neural activity that would be expected to necessitate synchronous interactions12,20. The present study confirms previous evidence and adds spatio-temporal refinement to the functional synchronization between the sources estimated to the ACC and the latPFC. Consistent with previous reports64, the present results indicate that PLVs are increased and sustained on incongruous trials in the Stroop task. By quantifying phase synchronization between these two areas with high temporal precision, these findings extend the conflict monitoring account and indicate that their interaction is particularly prominent after ~350 ms on incongruous trials. During this cognitive integration stage, the medial and lateral prefrontal cortices are likely to interact to support behavioral performance during more difficult task conditions imposing demands on attention, response inhibition, and working memory. Extensive evidence from MRI-based functional connectivity studies indicates that these cortical areas form a dynamic, interactive cingulo-opercular network that supports top-down cognitive control65-67. More broadly, the brain optimizes responding to environmental demands in an adaptive and coherent manner *via* flexible and dynamic synchronization of distributed neurofunctional systems68,69.

The anatomically-constrained MEG approach used in the present study relies on a combination of complementary imaging methods. It can characterize the spatio-temporal sequence of neural activity and can provide insight into the dynamics of long-range interactions important for integrating top-down influences during engagement of cognitive control. The MEG signal reflects synaptic currents directly, which allows for testing hypotheses about co-oscillatory interactions within and across neurofunctional systems with high temporal precision. Furthermore, this method is suitable for pharmacological manipulations because it is not susceptible to vasoactive confounds. Research from this lab and others indicates that prefrontally-mediated cognitive control functions are particularly vulnerable to alcohol intoxication17-20,23,24,70-74. The present study shows that acute alcohol intoxication decreases activity in the prefrontal areas subserving response conflict. Furthermore, alcohol disrupts synchronized co-oscillations20,75 that may underlie impaired or maladaptive response suppression. As a result, intoxicated individuals exhibit deficient self-control resulting in disinhibition which may contribute to compulsive drinking and the development of alcohol dependence25,26,76. In sum, estimates of synchronous co-oscillations can illuminate real-time interactions of the neural systems engaged by a particular cognitive demand and can inform a realistic brain-based model. They can characterize the selective sensitivity to alcohol challenge across networks and serve as biomarkers of individual vulnerability to pharmacological effects.

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The authors have nothing to disclose.

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